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STUDIES OF SUPPORTED METAL CATALYSTS

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PETER C. FLYNN

A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH IN PARTIAL FULFILMENT OF THE REQUIREMENTS FOR THE DEGREE OF DOCTOR OF PHILOSOPHY

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EDMONTON, ALBERTA

FALL, 1974

THE UNIVERSITY OF ALBERTA FACULTY OF GRADUATE STUDIES AND RESEARCH

The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research, for acceptance, a thesis entitled "STUDIES OF SUPPORTED METAL CATALYSTS" submitted by PETER C. FLYNN in partial fulfilment of the requirements for the degree of Doctor of Philosophy.

(External Examiner)

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G. Date Oct. 25, 1974

ABSTRACT

Supported metal catalysts provide an economical wall of utilizing noble metals as commercial catalysts. In this work two methods of characterizing such catalysts, electron microscopy and selective gas adsorption, hall been studied in detail. These two techniques have been employed in following the sintering behavior of a variety of Pt/Al_2O_3 catalysts. A model for sintering of supported metal catalysts , has been developed.

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The accuracy of particle size distributions determined from electron micrographs was examined from both theoretical and experimental points of view. Particle detectability and apparent size were found to be sensitive functions of defocus, and hence of elevation of particles in the specimen. Contrast has been shown to vary with orientation of both particles and support material. Sources of contrast inherent in the support in the sub-nm range have been illustrated. It is concluded that particle size distributions become increasingly subject to error as the fraction of particles with sizes below about 2.5 nm increases.

The nature of adsorbed oxygen on Pt/Al_2O_3 supported catalysts and the reaction on the surface of oxygen with hydrogen have been studied. Gas uptake data show that a repeated (H_2-O_2) titration step leads to subsequent enhancement of the hydrogen adsorbed on the surface, and a retardation of subsequent oxygen uptake. The enhancement is correlated to the ratio of initial hydrogen to initial oxygen uptakes, which in turn depends on the dispersion of the platinum crystal-

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lites.

iR spectra of CO adsorbed on catalysts with preadsorbed oxygen reveal two types of adsorbed oxygen species. Type I forms during room temperature adsorption, while Type II is found after high temperature adsorption and repeated (0_2-CO) treatment. Type II adsorbed oxygen is not observed after repeated (H_2-O_2) treatments, and hence is ruled out as a cause of enhancement. H_2O generated during the titration step is postulated as a possible cause of the enhancement/ retardation effect.

An interparticle transport model for the sintering of supported metal catalysts has been developed. The model postulates escape of atoms from crystallites to the support surface, rapid migration of these atoms along the surface, and their recapture by crystallites upon collision. A reduction in surface energy provides the driving force for transfer of metal from small to large particles. The model has been solved by finite difference methods and applied to several theoretical particle size distributions (PSD). The model predicts an increase in the rate of sintening as the width of the a initial PSD increases. The rate of sintering also increases as the surface velocity and the metal loading increase. Sintering behavior is sensitive to the activation energy and temperature. Under certain conditions substantial redispersion is predicted. The model can account for power-law orders from < 2 to > 13, as observed experimentally. Power-law order increases with PSD width and with increases in mean crystallite size.

Changes in the dispersion of supported Pt/Al₂0₃ catalysts following reduction and a variety of thermal treatments have been monitored by gas uptake and electron microcopy. Evidence of redispersion was

found after sintering of one catalyst in oxygen at 450° to 600°C. Sintering is found to be sensitive to gas atmosphere and metal loading. Addition of a portion of presintered catalyst containing large Pt particles increased the rate of sintering of a catalyst. From electron micrographs of the same catalyst area before reduction and after reduction and various thermal treatments, it was concluded that Pt agglomeration occurs during all these steps. Some Pt crystallites remain in a fixed location during reduction and thermal treatments.

vi

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vii

TABLE OF CONTENTS

\$

`

•

. •

..

	, , , , , , , , , , , , , , , , , , ,	Page
CHAPTER 1	INTRODUCTION	. 1
· · · · · · · · · · · · · · · · · · ·	1.1 Supported Metal Catalysts	1
•	1.2 Measurement of Metal Dispersion and Crystal	lite
	Size	• 3
/	1.2.1 X-Ray Diffraction Line Broadening	4
•	1.2.2 Electron Microscopy	4.
	1.2.3 Gas Chemisorption -	. 5 . ,
(1.3 Sintering of Supported Metal Catalysts	- 7
	1.4 References	8
CHAPTER 2	EXPERIMENTAL PROCEDURES	9
•	2.1 Catalyst Preparation ,	9
•	2.2 Gas Adsorption by Dynamic (Flow) System	9 -
	2.3 Electron Microscopy	13
	2.4 Other Measurements	15
	2.5 References	15
CHAPTER 3	THE LIMITATION OF THE TRANSMISSION ELECTRON	
ی پر می بین	MICROSCOPE FOR CHARACTERIZATION OF SUPPORTED	
•	METAL CATALYSTS	17
	3.1 Survey) 17
	3.2 Specimen Characteristics	18
	3.3 Theory of Image Generation	21
	3.3.1 Phase Contrast	22 /
	3.3.2 Diffraction Contrast	26
	3.4 Microscope Use	26
́.,	3.5 Results and Discussion	29
	viii	
•		

TABLE OF CONTENTS (continued)

.

					Paus	
	CHAPTER 3	cont	inued			
			3.5.1	Calculated Image Profiles	29	
	,		3.5.2	Variation of Contrast and Apparent		
-1				Size with Defocus	29	
			3.5.3	The Effect of Tilt on Contrast	44	
		٠	3.5.4	Contrast Structure in the Support		
				Material	48	
		3.6	Conclu	sions	50	
		3.7	Refere	nces	52	
	CHAPTER 4	THE	TITRATI	ON REACTION ON SUPPORTED Pt CATALYSTS	54	/
•		ą`, 1	Survey	• • • • • •	54	1
		4.2	Experi	mental Results	57 /	
			4.2.1	Isotherms and Isobars of Oxygen and		
	,			Hydrogen	57	
			4.2.2	Hydrogen Titration of Oxygen Adsorbed		
•				at Elevated Temperatures	62	
			4.2.3	IR Spectra of CO Adsorbed on Pt with	-	
	×			0 ₂ Preadsorbed at Various Temperatures	62	
	e.		4.2.4	Gas Uptakes in Titration Sequences .	66	
			4.2.5	Long Titration Sequence	72	
	•		4.2.6	IR Spectra of CO Adsorbed on Pt After		
				Repeated Sequences of $(0_2 - CO)^2$ and $(0_2 - H_2)^2$	_)	
				Treatments	75	
3		4,3	The Or	igin of the Enhancement Effect	77	
		4.4	Conclu	sions	84	

•

١

ix

,

.

			TABLE	OF CONTENTS (CONTINUE)	
	•		e		Page
	CHAPTER 4	cont	inued		
		4.5	Refere	nces	85
	CHAPTER 5	A MO	DEL OF	SUPPORTED METAL CATALYST SINTERING	8 7
		5.1	Survey		87
		5.2	Propos	ed Model	93
	· .		5.2.1	Escape of Metal Atoms from Crystallites	93
			5.2.2	Migration of Metal Atoms Over Support	
	,			Surfaces	99
١			5.2.3	Capture of Atoms by Metal Crystallites	99 .
			5.2.4	A Summary of the Model	100
		5.3	Method	of Solution	101
			5.3.1	Solution for $\ll/5_0 > 10^{12} \text{ m}^{-2}$	102
			5.3.2	Solution of $\ll/5_{0} \sim 10^{12} \text{ m}^{-2}$	104
)	5.4		ption of Cases Investigated	105
			5.4.1	Crystallite Size Distributions	105
			5.4.2	Parameter Values	106
		5.5	Result	s and Discussion	106
	• .		5.5.1	Effect of Initial PSD on Rate of	
	J			Sintering	106
	,		5.5.2	The Effect of Surface Mobility and Metal	
	. · · · ·			Loading on Sintéring Behavior	116
			5.5.3	The Effect of E_a/RT on Sintering	
				Behavior	118
			5.5.4	The Order of Sintering	121

Ņ

				•
TABLE	OF CONTENTS	(continued)	:	

5				~	Page
•	CHAPTER 5	cont	tinued		
			5.5.5	Application of the Model to an	
			,	Experimental Case	125
			5.5.6	The Possibility of Multiple Mechanisms	129
		5.6	Conclu	isions	129
		5.7	Nomenc	lature	131
		5.8	Refere	ences	132
	CIMPLER 6	EXPE	RIMENTA	L STUDIES OF SINTERING OF SUPPORTED	
		ΈLAT	INUM CA	TALYSTS	135
	•	6.1	Survey	,	135
•		6.2	Result	S	138
·			6.2.1	Constant Time/Variable Temperature	
				Treat	138
			6.2.2	Variable me/Constant Temperature	
				Treatments	142
			6.2.3	Catalyst Mixture Experiments	143
		d.	6.2.4	Effect of Reduction and Sintering on	
	۲,			Individual Metal Particles	148
		6.3	Discus	sion	161
		6.4	Conclu	sions	164
		6.5	Refere	nces	165
	APPENDIX A	EXP	ERIMENT	AL CONDITIONS	167
		A-1	Catal	ysts and Preparations	167
		A-2	Gas A	dsorption System	168

xi

· ·

•

¥ -

TABLE OF CONTENTS (continued)

Page

,

· /

APPENDIX A continued

		A-2.1	Calibration, Accuracy, and Sample	
			Calculations	168
		A-2.2	The Effect of Reduction and Outgassing	
			Times on Gas Uptakes	177 .
,		A-2.3	Effect of Rate of Addition of Oxygen	
			on Oxygen Uptake	181
•		A-2.4	Experimental Conditions and Results	186
	APPENDIX B	CALCULATION	OF PHASE CONTRAST IMAGES	208
	APPENDIX C	CALCULATION	OF A SINTERING HISTORY	215

LIST OF FIGURES

.

.

•

.

		Page -	e
Ligure			
2.1	Schematic Diagram of the Dynamic Adsorption System	10 :	
3.1	Schematic Cross Section of Typical Specimen Grid	20 1	
3.2	Schematic Diagram of Optical Fiffraction Micrograph		
	Resolution Test, and Optical Diffraction Pattern		
	from Micrograph of Thin Carbon Film	28	
3.3	Calculated Image Intensity Profile for a Single		
	Atom	30	
3.4	Three Calculated Image Intensity Profiles for a		
	Single Atom	32	
3.5	Calculated Maximum Image Intensity as a Function of		
· · ·	Defocus for the One and Seven Atom Cases	¥ 33 ,	,
3.6	Calculated Maximum Image Intensity as a Function of	2	¢
	Defocus for the Three and Four Atom Cases	34	
3.7	Calculated Apparent Image Size as a Function of		
	Defocus	37	
3.8	Two Calculated Image Intensity Profiles for the		
	Four Atom Case	.38	
3.9	Micrographs of a Pt/Al ₂ 0 ₃ Catalyst Showing the Effect	-	,
	of Defocus Change on Image Contrast	39	
3.10	Variation of Measured Particle Size with Defocus		
•	for Three Through Focus Series of Micrographs of a	÷.,	
	Pt/Al ₂ 0 ₃ Catalyst	41 *	
3.11	Variation of Measured Particle Size with Defocus		
•	for Three Through Focus Series of Micrographs of a	۰ و د	

LIST OF FIGURES (continued)

<u>,</u>

Pt/Al ₂ 0 ₃ Catalyst Variation of Measured Particle Size with Defocus for Three Through Focus Series of Micrographs of a Pt/Al ₂ 0 ₃ Catalyst Micrographs of a Pt/Al ₂ 0 ₃ Catalyst Showing the Effect of Tilt on Image Contrast Micrographs of a Pt/Al ₂ 0 ₃ Catalyst Showing the Effect of Tilt on Contrast	42 43 47 49
Variation of Measured Particle Size with Defocus for Three Through Focus Series of Micrographs of a $Pt/A1_20_3$ Catalyst Micrographs of a $Pt/A1_20_3$ Catalyst Showing the Effect of Tilt on Image Contrast Micrographs of a $Pt/A1_20_3$ Catalyst Showing the Effect of Tilt on Contrast	43 47
for Three Through Focus Series of Micrographs of a Pt/Al ₂ O ₃ Catalyst Micrographs of a Pt/Al ₂ O ₃ Catalyst Showing the Effect of Tilt on Image Contrast Micrographs of a Pt/Al ₂ O ₃ Catalyst Showing the Effect of Tilt on Contrast	47
Pt/Al_2O_3 Catalyst Micrographs of a Pt/Al_2O_3 Catalyst Showing the Effect of Tilt on Image Contrast Micrographs of a Pt/Al_2O_3 Catalyst Showing the Effect of Tilt on Contrast	47
Micrographs of a Pt/Al ₂ O ₃ Catalyst Showing the Effect of Tilt on Image Contrast Micrographs of a Pt/Al ₂ O ₃ Catalyst Showing the Effect of Tilt on Contrast	47
of Tilt on Image Contrast Micrographs of a Pt/Al ₂ O ₃ Catalyst Showing the Effect of Tilt on Contrast	
Micrographs of a Pt/Al ₂ O ₃ Catalyst Showing the Effect of Tilt on Contrast	
of Tilt on Contrast	49
• .	49
licrograph of Pure Aloo, ("Alon") Specimen Showing	
2-3 (
the Contrast Teherent in the Support	49
Dxygen Isomerms for a 2.03% Pt on Alon Catalyst	59
Dxygen and twitted Isobars for Three Different	
Pt/Al ₂ 0 ₃ Cate	60
/ariation of the 2060-80 and 2120 cm ⁻¹ Bands from	
CO Adsorbed on Pt/Al ₂ O ₃ Catalysts with Different	
2 Preadsorption Temperatures	64
influence of Evacuation on the 2060-80 and 2120 ${ m cm}^{-1}$	
ands	67
lydrogen Enhancement Ratio (H3/H1 or H2/H1) as a	
unction of the Ratio of Initial Hydrogen Uptake to	
nitial Oxygen Uptake (H1/O1)	71
ariation of Initial Uptake Ratio (H1/O1) with Catalyst	
ispersion as Measured by Hydrogen Uptake (H1) \langle	73
ong Term Titration	74
	xygen Isomerms or a 2.03% Pt on Alon Catalyst xygen and twice and Isobars for Three Different t/Al_2Q_3 Cate ariation of the 2060-80 and 2120 cm ⁻¹ Bands from 0 Adsorbed on Pt/Al_2Q_3 Catalysts with Different 2 Preadsorption Temperatures nfluence of Evacuation on the 2060-80 and 2120 cm ⁻¹ ands ydrogen Enhancement Ratio (H3/H1 or H2/H1) as a unction of the Ratio of Initial Hydrogen Uptake to nitial Oxygen Uptake (H1/O1) ariation of Initial Uptake Ratio (H1/O1) with Catalyst ispersion as Measured by Hydrogen Uptake (H1)

· xiv —

LIST OF FIGURES (continued) .

.

			Page
_	Figure		
\	5.14	A Power-Law Order Test of the Calculated Dispersion	
•		History for the Initial Distribution of Wynblatt and Gjostein ^{5.3}	128
	6.1	Effect of Treatment Temperature on Metal Particle	
		Size Range, Measured from Electron Micrographs	139
	6.2	Effect of Treatment Temperature on Metal Dispersion	
		for 0.5% Pt on $A1_20_3$, Measured by Gas Uptake	141
	6.3	Particle Size Distribution from Electron Micrographs	
		of a Sintered Engelhard 0.3% Pt on Al ₂ 0 ₃ Catalyst	144
•	6.4	Low Magnification Micrographs from Grid 6	152
	6.5	Micrographs from_Grid 4	154
	6.6	Micrographs from Grid 7	155
	6.7	Micrographs from Grid 2	156
	6.8	Micrographs from Grid 1	157
	6.9	Micrographs from Grid 6	160
	A - 1	Output from Thermal Conductivity Cell from Run 112	175

LIST OF TABLES

·

ć Sig

•

-

~~

Table		Page
1.1	Rapid Decrease in Dispersion as Metal Crystallite	
	Size Increases	2
3.1	Theoretical Defocus Windows for Dark Contrast for	
	Various Atomic Clusters	35
3.2	Calculated Image Intensity and Apparent Size Over	
	a Range of Defocus and Apertures	45
4.1	Description of Catalysts	58
4.2	0 ₂ Adsorption Uptakes as a Function of Temperature	
	and Room Temperature H ₂ Titration and Adsorption	
	Uptakes	61
4.3	H_2 and 0_2 Adsorption and Titration Uptakes	69
4.4	Hydrogen Enhancement and Oxygen Retardation During	
	Titration	81
5.1	Relative Rate of Loss of Atoms per Crystallite as a	
	Function of Crystallite Size as Estimated by the	
	Kelvin Equation	98
5.2	Data for Generation of Particle Size Distributions	107
5.3	Parameter Values Used in Study	108
6.1	Effect of Addition of a Portion of Presintered	
	Catalyst on the Sintering Rate of a 2.03% Pt on Alon	
	Catalyst	147
6.2	Grid Treatments	149
A-1	0 ₂ Calibration of Sample Loop Size	169
A-2	Hydrogen Calibration	172
A-3	Contents of Partial O ₂ Pulses /	173
•	se en la seconda de la seco	

•

xvii

LIST OF TABLES (continued)

[ab]e		Page
A - 4	Sample Calculations, Run 112	176
A-5	, Effect of Reduction Conditions on Gas Uptake	178
A-6	Effect of Outgassing Time on Gas Uptake	180
A-7	Effect of Pulse Dilution on Oxygen Uptake	183
A-8	Oxygen and Hydrogen Uptakes for Stream and Pulse	
	Addition of Initial Oxygen	185
A-9	Results from Dynamic Adsorption System	189
B-1	Atom Coordinate Positions Used in EMCON	209
B-2	Parameters Used in EMCON	210

xviii

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Ĵ

CHAPTER 1

INTRODUCTION

1.1 Supported Metal Catalysts

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The catalytic properties of metals have been known for over 100 years. Early applications of metal catalysts were generally confined to laboratory research studies, particularly in organic synthesis reactions. Since problems of scale economy or catalyst life are generally not critical in synthesis studies, catalysts were generally used in pure metallic form.

In this century, however, and particularly since the Second World War, metal catalysts have found increasing application in large scale industrial processes. Reactions such as hydrocarbon hydrogenation and dehydrogenation, partial oxidation, and petroleum reforming are all examples of standard manufacturing processes utilizing supported metalcatalysts. In plant operation, catalyst expense and durability become extremely critical and this cost/lifetime pressure led to the development of supported metal catalysts.

A supported metal catalyst generally consists of a relatively stable porous base (carbon or a metal oxide such as alumina or silica) on which small crystals of the metal are grown. Group Vlllb metals (Fe, Co, Ni, Ru, Rh, Pd, Os, Ir, and Pt) are the most common metals supported in this fashion. The metal is generally added to the support by immersing the support in a salt solution, evaporating to dryness, and then either reducing the metal salt directly or converting it to the metal oxide in an oxygen atmosphere and then reducing the metal

oxide. The resulting metal crystallites may vary in size from 200 nm or more to smaller than 1 nm.

The economic value of supported metal catalysts may be readily shown: In purchasing metal, a user pays by bulk; in using the metal as a solid catalyst in a gas phase reaction, he utilizes only those metal atoms which "see" the gas, i.e. surface atoms. The ratio of surface metal atoms to total metal atoms, known as the dispersion ratio, thus measures efficiency of utilization of the expensive metal. Table 1.1 shows the rapid decrease in dispersion as metal crystallite size increases.^{1.1}

Table 1.1: Rapid Decrease in Dispersion as Metal Crystallite Size Increases

<pre># of Atoms in edge of crystallite</pre>	Crystallite size, nm (based on platinum)	Dispersion
	0.78	1.000
 ب	1.95 · ``	0.780
10	3.89	0.490
18	7.00	0.300

Thus use of magneticles "wastes" 70% of the total metal atoms, which are una sticipate in the reaction. By way[†]of further illustration, a metal cube that mm edge has a dispersion of about one of the metal atoms do not contact the gas.

Attempts to physically displanemetal catalysts through grinding or precipitation of a metal sponge (black) — ove unsuccessful for two reasons: first, particles even in the 100 nm range are such a fine dust that containment within a reactor is difficult; second, even at moderate

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temperatures the metal particles, even in a sponge, tend to reagglomerate to larger crystallites. Use of a supporting material reduces both of these effects. In general metal crystallites adhere quite firmly to the supporting material, and thus catalyst loss in the product is negligible. In addition, the support holds small crystallites at a distance and retards sintering, the name given to the process of metal crystallite growth. Thus the support stabilizes highly dispersed metals and allows economical use of expensive metal catalysts in large scale industrial processes. Generally, the support is inert and does not influence the reaction, although petroleum reforming is a notable exception to this rule.

1.2 Measurement of Metal Dispersion and Crystallite Size

As outlined above, metal dispersion, a function of the metal crystallite size, is an extremely critical parameter of a supported metal catalyst. In addition to the problem of efficient use of metal atoms, detailed above, a second factor makes dispersion significant. Boudart^{1.2} has identified two types of catalytic reactions, facile and demanding. In the former, the reaction rate varies directly as the surface area of the catalyst; in the latter, the reaction rate per unit *surface area* of metal depends upon the crystallite size. For example, van Hardeveld and van Montfoort^{1.3} demonstrated that the nitrogen adsorption activity per unit surface area of a nickel on silica catalyst was highest when crystallite sizes were 2 to 7 nm.

Thus knowledge of average metal particle size and particle size distribution are important in characterizing a supported metal catalyst. Three techniques have been employed in the determination of the size of the metal crystallites, and each will be briefly described here, along

with advantages and disadvantages.

1.2.1 X-Ray Diffraction Line Broadening

Klug and Alexander^{1.4} explain the theory behind use of x-ray line broadening in determining small crystal sizes. Briefly, an x-ray diffraction peak for a given crystal plane broadens as the crystal size in the sample decreases. The peak width difference between large crystals (width determined by instrumental response) and small crystals (additional width due to crystallite size) may be related to an average crystallite size. Smith^{1.5} indicates that an average size

$$\bar{d} = \frac{\sum n_i d_i}{\sum n_1 d_i^3}$$

is obtained, where \overline{d} is the average size obtained by this method and n_i is the number of particles in the size interval centered on d_i . Fourier analysis of the broadened diffraction curve yields some information on the particle size distribution of the sample.

However, the great drawback of x-ray diffraction line broadening is its inability to detect particles of a very small size. While some authors 1.6, 1.7 reported detection of 3 nm particles and smaller, it is generally accepted 1.8 that particles less than 5 nm will not be detected by x-ray line broadening and thus will not be included in the average particle diameter. Since 5 nm represented a median to upper limit of the metal crystallite sizes of typical interest in this research, x-ray techniques have not been employed.

1.2.2 Electron Microscopy

Adams et al.^{1.7} and Moss^{1.9} gave some of the first descriptions of the use of the electron microscope in determining particle sizes in

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supported metal catalysts. Since then numerous investigators have utilized electron microscopy to determine particle size distributions. The primary advantage of electron micrograph analysis is that the data are in the form of a distribution, rather than an average particle size.

Chapter 3 examines in detail the accuracy and validity of particle size distributions determined by electron micrographs of catalyst specimens. It is concluded that even under ideal conditions the computation of accurate particle counts is questionable when the size of particles counted drops below 2.5 nm.

1.2.3 Gas Chemisorption,

The third means of determining relative particle sizes in a supported metal catalyst is through measurement of the gas chemisorbed on the metal surface. One advantage of the method is immediately evident--it measures how the catalyst will interact with a feed stream in reactor conditions. Since all metal atoms that interact with the gas are detected, this method "sees" particles of all sizes, even atomically dispersed metal.

Three methods are available for determination of chemisorbed uptake, the classical volume c method using a vacuum rack, a dynamic adsorption system and the gravimetric balance. Borling^{1.10} reviews each method. The classical volumetric approach provides greater accuracy and allows ready determination of isotherms (uptake versus pressure at a variety of temperatures). However, it is more time consuming and involves a higher capital investment than the dynamic method. Samples require two determinations, one measurement of gas uptake on the supported metal catalyst and one on a support blank so that physical

adsorption effects may be eliminated.

The dynamic method's primary advantages are atmospheric operation (i.e., it does not require a high vacuum system), and elimination of support correction; these two factors allow more rapid analysis of gas uptake. The system employed in this research will be briefly detailed in Chapter 2.

The gravimetric method actually weighs a small catalyst sample and records weight increases due to gas adsorption. This method also involves high capital cost, and is also difficult to apply for hydrogen adsorption because of the low mass of the adsorbate atom compared to the sample. The gravimetric technique has found widest application with heavier adsorbed gases such as carbon monoxide and hydrocarbons.

Whatever the method of its determination, the uptake is related to an average size through an adsorption stoichiometry and a crystal-shape model. The most generally assumed models for stoichiometry are either one atom of oxygen or one atom of hydrogen per exposed metal surface atom; both have been reported and used by various workers. Crystallites may be modeled in a variety of ways, for example, as spheres, as cubes, as octahedra, as cubo-octahedra, etc. Each of these models predicts a dispersion (fraction of atoms that are at the surface) as a function of particle size. Van Hardeveld and Hartog^{1.11} summarize these models; deviations among the various models are small.

Thus, by assuming an adsorption stoichiometry, the absolute number of surface metal atoms may be determined from the measured adsorbed gas uptake. The absolute number of total metal atoms is known from the catalyst composition and weight, so the dispersion may be readily calculated. From the dispersion, a surface average diameter is determined.

The surface average diameter, d, determined in this Fashion, is related to the particle size distribution by the relation 1, 11

$$\tilde{d} \approx \frac{\sum n_i d_i^3}{\sum n_i d_i^2}$$

Chapter 4 examines in greater detail the uptake of hydrogen and oxygen on support Pt catalysts. Anomalies in the titration reaction (consumption of adsorbed oxygen by gaseous hydrogen, or vice versa) are discussed in detail.

1.3 Sintering of Supported Metal Catalysts

Although a support material helps to stabilize metal dispersion as compared to pure metal catalysts, changes in the dispersion of supported metal catalysts still occur. This effect is known as sintering: loss of dispersion via this process is one of the major factors limiting the life of supported metal catalysts. Where supported metal catalysts are envisioned in high temperature situations (such as automotive exhaust mufflers), problems of dispersion maintenance are especially critical.

The physical mechanisms accounting for sintering have not yet been established. In Chapter 5 an existing model advanced to explain sintering behavior is reviewed, and an alternate model postulating surface diffusion of atoms is advanced. This atomic surface diffusion model is solved and applied to a number of situations of catalytic interest. The effect of the model parameters on the predictions for sintering behaviour is detailed. Finally, in Chapter 6 experimental evidence of sintering is presented and discussed in light of the theore tical models. \geq

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1.4 References

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CHAPTER 2

EXPERIMENTAL PROCEDURES

2.1 Catalyst Preparation

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Commercial and prepared catalysts were used in this work. Engelhard 0.3° and 0.5% Pt on Al_2O_3 catalysts (all catalyst concentrations are morted by weight), and 0.5% Rh on Al_2O_3 were used after overnight soluction at 500°C.

Prepared catalysts were made with two Al_2O_3 supports, Kaiser 201 spheres (8-10 mesh) and $Alon^R$, a fine alumina manufactured by the Cabot Corporation. In each case the support was wetted by a solution of chlorplatinic acid (H_2PtCl_6) , then dried at 110°C. The catalyst was then given a mild reduction at 250°C, which converted the metallic salt to metal. Details of the preparation are given in Appendix A, Section 1. Before gas uptake measurement the catalyst was further reduced in H_2 at 500°C. Catalyst loadings from 0.10% to 4.76% were prepared in this fashion. Bond^{2.1}, Dorling^{2.2}, and Gil'debrand^{2.3} give a broader discussion of various supported metal catalyst preparation techniques.

2.2 Gas Adsorption by Dynamic (Flow) System

Figure 2.1 shows a flow plan of the dynamic system used in this work for adsorption measurements. The catalyst sample was inserted in a Vycor glass tube, allowing heating to 800°C. The catalyst remained under a constant flow of an inert carrier gas (generally helium) during uptake measurements. Gases to be adsorbed (generally H_2 or O_2)



were flowed through a calibrated sample loop; switching of the sample valve (Carle Instrument Co, Fullerton, Cal; catalogue #2014) thus injected a pulse of known content in the carrier gas stream. Provision of two calibrated rotameters allowed dilution of the sample gas by the carrier gas. Thus a range of pulse sizes could be generated. Gas connections were through flexible Swagelok Quick-connects, eliminating the danger of leakage through pressurized valves. Carrier and sample gas flow rates were measured by bubble flow meters.

The catalyst could be inserted in a temperature controlled furnace or in various temperature baths, thus allowing uptake as a function of temperature to be readily determined. The furnace was a Thermolyne Model 86 (Sybron Co, Dubuque, Iowa) modified to allow introduction of the Vycor tube holding the catalyst. Temperature was c olled by a Thermo Electric 400 temperature controller (Thermo Electric Co, Saddle Brook, N.J., Model No. 3242200) attached to an iron-constantan thermocouple in the vicinity of the catalyst. A second thermocouple was inserted into the furnace 'and its output recorded as a check on the internal temperature. In practice control was within 5°C, with little overshoot when a sample was inserted in a hot furnace (<20°C) and no overshoot on cold starts.

Standard adsorption temperatures were: $-98^{\circ}C$ (a methanol ice bath made by stirring liquid N₂ into a Dewar flask containing methanol; $0^{\circ}C$ (ice bath); 96-97°C (boiling distilled H₂O); and 203°C (in the furnace). Room temperature runs were made by inserting the tube in an H₂O filled beaker; room temperatures varied from 22° to 24°C. Reduction was generally effected at 500°C, followed by helium outgassing for one or two hours at 500°C. One requirement of a dynamic system is

a high purity carrier gas, since even traces of adsorbents such as carbon monoxide or oxygen in the carrier gas will accumulate on the sample. In this system the carrier gas is first passed over a mixed reduction-oxidation catalyst (supported Cu/CuO) are ared according to the method of Meyer and Ronge^{2.4}. Contamination of a carrier gas by H_2 or O_2 after passage over this bed of less than 0.4 ppm was determined by the above authors, and confirmed in this laboratory from a long term degassing study described in Appendix A, Section 2. Water is removed after the carrier gas purification step by a molecular sieve column. The by-pass gas line allows bypassing of the purification catalyst during oxidation or reduction of the sample catalyst. Valving was included to allow bypassing of the thermal conductivity cell when desirable.

Hydrogen used in this work was formed in a Matheson H₂ Generator and further purified by passing over an Englehard Deoxo hydrogen purifier. The gas was dried over a molecular sieve column before introduction to the catalyst. Oxygen and nitrogen were Linde prepurified grade, 99.995% and 99.997% pure, respectively. Instrument grade helium supplied by Canadian Liquid Air was further purified, as described above, before use.

The size of the input pulse of oxygen was established by comparison to a known volume of gas injected by syringe; use of the ideal gas law enabled calculation of the moles of oxygen per pulse. In all runs the flow of 0_2 through the sample loop was held constant to ensure identical pulse sizes. Hydrogen content in the sample loop under standard flow conditions was determined by pressure measurements in the sample loop; comparison to the pressure during oxygen flow allowed

the H_2 content to be calculated by the ideal gas law. Details of the calibration are given in Appendix A, Section 2.

The output pulse was measured by thermal conductivity cell (Gow-Mac Instrument Co, Madison, N.J., cell model 10-785, with Rhenium-Tungsten filaments, powered by a Gow-Mac Power supply, model 40-001) and recorded on a chart recorder. Mechanical integration of peaks by a Disc Integrator was used to calculate the size of the output pulse. In standard operation pulses of adsorbate gas were passed over the catalyst until a steady state output size was achieved (presumed to be identical to the input pulse). Comparison of partial output peaks to the area of the final output peak enabled calculation of the total output of adsorbate gas. The amount adsorbed on the catalyst surface was determined by difference between input and output.

Appendix A, Section 2, details studies of the variation of gas uptake in standard operation of the flow system. The influence of reduction time, outgassing time, rate of adsorbent admission and other parameters on gas uptake is also reviewed.

This flow system for chromatographic pulse measurement of gas uptakes on catalysts is similar to that described by Freel and others 2.5-2.8. Chromatographic desorption measurement systems 2.9, 2.10 and chromatographic breakthrough systems utilizing a constant flow of adsorbent gas 2.11 have also been described.

2.3 Electron Microscopy

The electron micrographs used in this work were recorded on a JEM 100 B Microscope in the University of Alberta Department of Physics. The microscope was equipped with a goniometer (tilting) stage, and

was capable of resolving the 0.34 nm snacings in graphitized carbon. All images were recorded in the bright field mode using a 100 KV accelerating voltage ($\lambda = 3.7$ pm). The maximum magnification recorded on the negative was 330,000, although further increases were obtained through photographic enlargement.

For analysis of pretreated catalyst specimens, samples were crushed into a fine powder and suspended in an inert solvent, then dropped onto a copper grid covered with a "holey" carbon film. The latter was prepared by condensing water droplets in a film of dissolved plastic. Specifically, a pre-soaped dry glass slide was dipped into a 0.1% solution of formvar in chloroform (CHCl₃). Water was added to the slide by breath or as a mist from sprayer. As the solvent evaporated and the plastic film hardened, holes were left whose size could be roughly controlled by the preparation conditions 2.12, 2.13. The plastic film was then floated off onto water, and electron microscope grids placed on it. The film was then picked up by adhesion to a piece of paper or plastic sheet. After drying, the film covered grids were coated with a thin layer of evaporated carbon to conduct electrons and strengthen the film. When the suspended catalyst was dropped on the film, portions of the catalyst extended over the holes, so that any effects of the carbon film on the image contrast were avoided. Astigmatism was corrected at maximum magnification, and very low contamination rates were achieved through use of the standard decontamination device.

For in situ reduction and sintering studies, where the catalyst was treated after placement on the electron microscope grid, two

modifications were employed. First, a tungsten grid was used, because of its greater strength at sintering temperatures. Second, in order the improve film stability the plastic film was dissolved after being carbon-coated; as a result, "noloy" carbon films were prepared which were stable for some time at 600°C in an inert atmosphere.

2.4 Other Measurements

In conjunction with our studies of the titration reaction over supported Pt catalysts. Dr. Eichi Kikuchi measured gas uptakes on a static system and IR spectra. These data are presented in Chapter 4.

The static apparatus was a conventional nigh vacuum glass volumetric adsorption system. The catalyst was typically reduced in hydrogen at 500°C for two hours and then evacuated for another two hours at that temperature.

IR measurements were carried out on a 2.03% Pt on Alon catalyst. IR transparent wafers were prepared by pressing a finely crushed bowder in a one inch diameter die at pressures of 12 tons/in². The catalyst wafer was placed in an in site infrared cell with sodium chloride windows. Pretreatment conditions were identical to those for the static measurements described above. Subsequent IR spectra were recorded at room temperature on a Perkin-Elmer Model 621 spectrophotometer, using a similar cell without a catalyst wafer in the reference beam.

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CHAPTER 3

THE LIMITATION OF THE TRANSMISSION ELECTRON_MICROSCOPE FOR CHARACTERIZATION OF SUPPORTED METAL CATALYSTS

3.1 Survey

In the last fifteen years the electron microscope has found increasingly widespread application for the characterization of supported metal catalysts^{3.1-3.7}. A principle aim in such studies is the determination of the metal dispersion (the ratio of surface to total metal atoms) which can be calculated if the metal particle size distribution is known, and if a particle geometry is assumed. Electron microscopy has been used to provide direct determination of the size distribution from the images of the metal particles, as discussed in Chapter 1.

The claimed resolution in images of supported metal catalysts has gradually increased, to the point where the detection of 0.4 nm particles and calculated average particle sizes of less than 2 nm have been reported^{3.3}, ^{3.6}, ^{3.7}. Such particle size distributions obtained by electron microscopy are based on three implicit assumptions, namely:

- a) the size of a metal particle is equal to the size of its image recorded on the micrograph (corrected for magnification);
- b) detection of a particle of a given size implies that all particles of that size and all larger particles are being detected;
- c) image contrast of the metal particles is distinguishable from contrast arising from the support material.

These assumptions are consistent with the use of a simple mass-thickness

interpretation of image contrast in the electron microscope.

It is clear that the correctness of these assumptions is important, since particle size distributions determined from micrographs have often been used to confirm a proposed adsorption stoichiometry for the selective adsorption of gases on supported catalysts^{3.3}, This latter technique, thus calibrated, has been used for routine a termination of metal dispersions.

Recent work on the bright field imaging of atoms and atomic clusters using conventional high resolution instruments^{3.8}, ^{3.9}, ^{3.10} has emphasized the importance of careful image interpretation, using theories which take into account the defects of the imaging lens^{3.11}. On the basis of such results and of the characteristics of the specimens involved, it was suspected that none of the above assumptions was necessarily valid, particularly for smaller metal particles, and that limits should be determined for both the smallest reliable particle size and the smallest reliable difference between the size of two particles.

In this chapter the results of an investigation into the contrast characteristics of images of platinum particles supported on alumina are presented. The qualitative predictions of both the phase contrast and of the diffraction contrast mechanisms of image contrast have been confirmed by experiments in which the variations in image contrast of specimen particles has been studied as the focus or specimen orientation were varied. The results confirm that the three assumptions listed are not correct for the conditions typical in high resolution microscopy of supported metal catalysts.

3.2 Specimen Characteristics

, Crucial to these considerations are the characteristics of the

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catalyst as a specimen for high resolution microscopy. The metal particles vary from atomic clusters (consisting of a small number of atoms, or possibly single atoms), to true microcrystallites with diameters of the order of 10 nm. The fact that the larger metal particles are crystalline, with the same face-centered cubic structure as bulk material, has been established by x-ray diffraction^{3,1} and by analysis of dark-field micrographs^{3,12}. At the other extreme, Prestridge and Yates^{3,13} have presented micrographs in which images of clusters of a few rhodium atoms have been identified. In these same micrographs, however, may be observed the st ong contrast from the silica support.

The metal particles are supported on silica or alumina particles which are usually porous assemblages of irregularly shaped crystals containing defects, of which some are inherent in the crystal structure. Crushing the catalyst to a powder generally results in clumps of support particles of varying thicknesses greater than 30 nm.

The supporting material is in turn placed on microscope grids in various ways; only in the recent work of Free1^{3.3} and Prestridge and Yates^{3.13} have "holey carbon" support films been used, to eliminate any possibility of interference from the granular image detail observed in continuous amorphous support films (see, for example, Thon^{3.14}). From stereoscopic images it is observed that there is typically a range of elevations in the direction of the electron beam of order 100 nm, often between two apparently adjacent alumina particles. This specimen elevation, illustrated in Figure 3.1, imposes a different focus condition upon various particles imaged in the same micrograph. Thus, within any one micrograph there will be a range of values of defocus (deviation from perfect focus), and within two micrographs of similar regions the


range of defocus values will in general be different.

For the imaging of small atomic clusters, considerable efforts have been made to minimize the background contrast produced by the supporting film (see, for example $^{3.8}$). Such films are essentially flat and only a few nm thick. Clearly the typical catalyst specimen is far from ideal in that the support imposes appreciable background contrast and holds the metal particles at different elevations.

Finally it may be noted that the support particles, and hence the metal particles are oriented at random. Therefore any image contrast effects which are sensitive to the orientation of the electron beam may be expected to contribute to variations in contrast between otherwise identical particles.

3.3 Theory of Image Generation

It is clear from the characteristics of the specimens that more 'than one mode of image contrast formation must be considered. For the very small metal particles (clusters of atoms) the *phase contrast* mechanism will operate. However, for larger metal particles in which a definite crystalline structure has developed, the scattering process will be best described in terms of Bragg diffraction giving rise to *diffraction contrast*. This will apply also to the essentially crystal-line support particles.

We have therefore considered the implications of each of these contrast theories in what follows. There will of course be a gradation from one of these extremes to the other, but for the present purpose the intermediate case need not be considered. It was find necessary to perform fairly detailed calculations of phase contrast in order to

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evaluate the variation of image size with defocus. However the qualitative features of diffraction contrast are sufficiently well documented, so that additional calculations were not required.

3.3.1 Phase Contrast

The contrast of an image obtained in the conventional transmission electron microscope is best understood by considering the situation in the back focal plane of the objective lens, where the Fraunhofer diffraction pattern of the electron wave emerging from the specimen is formed. The objective aperture placed in this plane cuts off from the image that part of the wave corresponding to electrons which are scattered through an angle greater than that subtended by the aperture. But in addition, the spherical aberration and the defocus of the lens have the effect of changing the phase of the wave in the back focal plane, through the "contrast transfer function", which depends also on the electron wave length and the scattering angle. Thus, even with a large objective aperture, the microscope will not image faithfully detail smaller than 1 or 2 nm. (For recent reviews of this theory see Thon^{3.14} and Hawkes^{3.15}).

The way in which the phase contrast transfer function removes certain spatial frequencies from the image while changing (including, reversing) the relative phases of others has been established by Thon, using amorphous thin carbon films^{3.14}. order to be able to interpret the images of particles smaller than a few nanometers, the effects of lens defects must be included in suitable theoretical calculations of image contrast. This was first done by Scherzer^{3.11}; more recently Reimer^{3.10} and Hall and Hines^{3.9} have made detailed calculations of the contrast of single atoms and clusters of atoms. (The atom is considered to change the phase, but not the amplitude, of the electron wave.) Their results show that the optimum defocus values, at which maximum contrast is expected, vary with cluster size, and that for exact focus the contrast may be too low to detect. Hall and Hines^{3.9} obtained through focal series of images of gold particles (on an almost structureless graphite support) which confirmed their predictions.

In order to evaluate the effects on apparent particle size of the lens defects, we have computed image profiles of atoms and atomic clusters using the formulation of Eisenhandler and Siegel^{3.16}. Their solution employs a real atomic scattering factor, values of which are readily available (for example, ^{3.17}). This approach assumes a phase shift of $\pi/2$ upon scattering, a less accurate approach than that of Hall and Hines^{3.9}, who used a complex atomic scattering factor with the correct phase shift due to scattering. However, their results show that the discrepancies between the two calculations are not significant for the qualitative results in which we are interested. In these calculations, the intensity at the point (x_i , y_i) in the image plane (scaled to unit magnification) is given by

where

 $\psi(x_i, y_i) = t$ wave function of the scattered electron $\lambda = \text{electron wave length, set by the voltage}$ $(1 - \beta^2)^{-1/2} = \text{relativistic correction; } \beta \text{ is the electron velocity-expressed as a fraction of the speed of light.}$

 $|\Psi(x_{i}, y_{i})|^{2} = 1 + \frac{4\pi}{\lambda} (1 - \beta^{2})^{-\frac{1}{2}} R(x_{i}, y_{i})$

For a single tom, $R(x_i, y_i)$ is given by

$$R(\mathbf{x}_{i}, \mathbf{y}_{i}) = \int_{0}^{\alpha} \int_{0}^{\alpha} f_{0}(\sin(\frac{\alpha}{2})/\lambda) \cos\left[\frac{\pi}{2} - \frac{2\pi}{\lambda}\left(\frac{C_{s}^{\alpha}}{4}\right) + \frac{2\pi}{\lambda}\left(\frac{\Lambda f^{\alpha}}{2}\right)\right]$$

$$J_{0}\left(\frac{2\pi\alpha r_{i}}{\lambda}\right) \quad \alpha d\alpha \qquad \qquad 3.2$$

where

 α = scattering angle (α_{max} is determined by the objective aperture)

 f_0 = atomic scattering factor for electrons

 $\rm C_s^{=}$ spherical aberration constant

∆f= defocus

 $J_0 = zero order Bessel function$

r_i = radial position in image plane from scattering center

$$(r_{i} = (x_{i}^{2} + y_{i}^{2})).$$

For multiple atom cases, the Bessel function term above is re-

all atoms

$$\sum_{n=1}^{J_{o}} J_{o} \left(\frac{2\alpha\pi}{\lambda} \left((x_{i} + x_{n})^{2} + (y_{i} + y_{n})^{2} \right)^{\frac{1}{2}} \right) \quad 3.3$$

where

 $\boldsymbol{x}_n, \, \boldsymbol{y}_n$ are the locations in the object plane of the various scattering atoms.

The term $\cos\left[\frac{\pi}{2} - \frac{2\pi}{\lambda}\left(\frac{c_s\alpha^4}{4}\right) + \frac{2\pi}{\lambda}\left(\frac{\Delta f\alpha^2}{2}\right)\right]$ is the phase contrast transfer function of the lens, and contains the phase changes imposed on the scattered electron wave within the lens by spherical aberration and defocus.

In order to assess the variation in image intensity for particles

of different sizes, four cases were evaluated. These were arbitrarily chosen as a single atom, a three atom planar array, a four atom pyramidal array, and a seven atom planar hexagonal array. For the multiple atom cases, the scattering centers (atoms) were separated by 0.275 nm, and the intensity profiles were computed along a radius which ran through a non-central atom. Test calculations along a radius running between two non-central atoms gave similar intensity profiles with slightly different spacings.

The calculations were performed using relativistic Hartree-Fock atomic scattering factors for gold^{3.17}. Values for C_s (1.6 nm) and $\frac{1}{max}$ (0.0125 rad) were chosen to be typical for a high resolution microscope at 100 KV (λ = 3.7 pm). Since a range of defocus values are expected within an image, the calculations were performed for of values from -200 to +400 nm in 20 nm steps.

In addition, the single atom case was evaluated over the defocus range -20 to +200 nm for four other values of α_{max} . These calculations were designed to test the effect of aperture size on phase contrast effects.

The R integral, equation 3.2, was iteratively evaluated, with a minimum of 200 steps to α_{max} , by the IMSL DRMBIU subroutine. Evaluation of ϵ^2 at intervals of 0.5 nm to a total of 10 nm allowed determination of the phase contrast image profile, *i.e.* the image intensity as a function of spatial location in the image plane. Contrast, defined as the difference in intensity between image point and background divided by background intensity, is given ψ^2 -1. The program for calculation of intensity profiles, along with details of parameters used, is shown in Appendix B.

3.3.2 Diffraction Contrast

For larger specimen structures the phase contrast calculation is invalid, because multiple scattering and diffraction effects become significant. Once the metal particles reach a size of 2-3 nm diameter, Bragg diffraction effects should dominate in the scattering process, and the use of standard multibeam dynamical calculations (see for example Hirsch *et al.*^{3.18}) is appropriate. The diffraction pattern from a single crystal consists of a series of discrete spots corresponding to diffraction from various planes in the crystal. In the back focal plane the objective aperture intercepts all except the directly transmitted beam, giving rise to bright field diffraction contrast: the variation in the intensity of the directly transmitted beam from point to point across the crystal surface. The theory then predicts a fairly sensitive dependence of image contrast on the thickness and orientation of the crystal^{3.18}, but a reduced sensitivity of contrast to defocus as compared with phase contrast images^{3.9}.

3.4 Microscope Use

In order to test these theoretical predictions, two series of experiments were performed. The phase contrast effects were examined by, taking through focal series of each of a number of areas for both Pt/alumina and pure alumina specimens. Diffraction contrast effects were observed by imaging the same field a number of times, tilting the specimen through a known angle between each set of through focus exposures.

The astigmatism correction was checked for selected cases and the transfer characteristics of the lens were determined approximately

from through focal series of images of thin amorphous carbon which were analysed using a simple optical diffractometer^{3.14}. The optical diffraction pattern reveals which spatial frequencies are present in the image, and which have been filtered out by phase cancellation (the contrast transfer function is zero for certain values of α). An example is show in Figure 3.2, along with a schematic of the resolution test apparatus. The distance from the central spot is proportional to the inverse of the spatial separation, and may be calibrated by imaging a known mesh. Accounting for magnification, the spatial scattering passed unfiltered for this particular pattern was calculated as 0.55, 0.67, 0.89 nm and greater than 1.17 nm with filtering effects between these values. Alternate frequencies passed by the lens are reversed in phase relative to their original values. For this particular case, no meaningful detail less than 0.55 nm was transmitted by the lens, thus representing a limit to resolution. Elliptical rings indicate the presence of astigmatism in the objective lens which had not been completely corrected.

The objective apertures available subtended angles of approximately 0.01, 0.006, and 0.003 radians. The first of these, a 60 nm aperture, was used for most of the work reported, since the project was concerned with the imaging of the smallest particles and atomic clusters. This aperture removes from the image information concerning spacings smaller than 0.37 nm, and is therefore large enough for phase contrast. On the other hand, all Bragg diffracted beams (except one) are intercepted, ? giving rise to diffraction contrast from the crystalline particles. The exception in our studies was the lll reflection of Y-Al₂O₃, for which d₁₁₁ is 0.456 nm. Lattice fringes of this spacing were often



MICROGRAPH RESOLUTION TEST



Figure 3.2 Schematic diagram of optical diffraction micrograph resolution test, and optical diffraction pattern from micrograph of thin carbon film, illustrating the elimination of certain spatial frequencies from the image.

observed and were used as an internal magnification standard. The smaller apertures were used to verify the predicted effects of lower u_{max} values on image resolution.

3.5 Results and Discussion

3.5.1 <u>Calculated Image Profiles</u>

Solution of the phase contrast equation generates an image intensity profile; the result for a single atom and a defocus of 180 nm is shown in Figure 3.3. In order to compute detectability and size as a function of defocus, we assumed limits for distinguishable contrast, and took the width of the central peak (d_0 in Figure 3.3) as the apparent size. Following Eisenhandler and Siegel^{3.16}, 5% light or dark maximum contrast was assumed to be necessary for detection: if the maximum intensity was between 0.95 and 1.05 the size was set for zero.

The arbitrary nature of these assumptions is evident. Other size definitions could have been used, such as the diameter at 5% contrast $(d_{0.95}$ in Figure 3.3). A different detectability limit would alter the sizes determined; under some conditions images can have bright and dark rings surrounding the central peak, with obvious complications in defining size. These and other factors would be critical if one attempted to deduce the actual size of a cluster (number of atoms), for which one would require to know the defocus value for each particle in the image. It is important to emphasize, however, that the assumption of different detectability limits or size definitions in no way affects the qualitative results of the present studies, which apply in general irrespective of the optical constants of the particular microscope being used.

3.5.2 Variation of Contrast and Apparent Size with Defocus



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Calculated image intensity profile for a single atom at a defocus of 180 nm. Dashed lines are detectability limits; two size parameters are shown.

The character of an image profile is a sensitive function of focus, and can shift from light, through undetectable to dark, as shown in Figure 3.4. The three profiles are for a single atom with defocus values of 140 nm, 160 nm and 180 nm.

Figures 3.5 and 3.6 show the calculated maximum image intensity of the four clusters of atoms as a function of defocus. It is evident that all four clusters will be detectable, as light or dark regions depending upon the defocus. In addition, it is evident that the range of defocus over which the arrays will appear as a region of dark contrast, the "window size", increases as the number of atoms in the cluster increases. Table 3.1 shows the calculated window sizes for the four cases.

In order to relate these results to micrographs of supported metal catalysts, the geometry of the specimen must be considered. Referring to Figure 3.1, the specimen in the region of a hole in the grid consists of an irregular stacking of catalyst fragments, with a typical depth (measured from the image shift after tilting the specimen through a known angle) of greater than 50 nm. This depth imposes an immediate restraint on any micrograph, for the defocus over the specimen will have dark contrast regardless of its spatial position and defocus, because of the diffraction contrast mechanism. Small metal clusters, however, have a contrast which is extremely sensitive to the defocus, and thus only some regions of the specimen may be so situated as to provide dark phase contrast for these small particles.

Calculated apparent size of metal clusters is also a sensitive function of the defocus, as shown for the three and seven atom cases in











Calculated maximum image intensity as a function of defocus for t: three and four atom cases. All atom clusters will give detectable dark contrast only over a selected range of defocus values. Figure 3.6

Table 3.1: Theoretical Defocus Windows for Dark Contrast for Various

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Case	•	llindo	de			
0000		Windows				
1 atom	60-120;	150-195;	235-250			
3 atoms	45-215;	240-275				
4 atoms	40-280					
7 atoms	10-390					

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Atomic clusters ($C_5 \approx 1.6 \text{ mm}$, $\lambda \approx 3.7 \text{ pm}$)

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Figure 3.7. Even in the region where clusters give dark contrast, the sharpness and diameter of their images varies strongly with defocus. Thus, again returning to an actual catalyst specimen, identical small metal particles at different elevations in the specimen would be expected to have varying image sizes, because the variation in defocus for these particles is equal to their differences in elevation in the specimen. Similarly, identical particles imaged in different micrographs would in general appear of different sizes because of differences in defocus.

An extreme case of the deviation in calculated image appearance with defocus for the four atom case is shown in Figure 3.8. A 60 nm change in defocus changes the image from a dark region of 0.8 nm apparent diameter, to a ringed dark-light pattern, with an apparent size for the inner dark region of 0.4 nm.

- Analysis of a ser of micrographs of a 4.76% Pt on Alon catalyst confirms that for small contrast regions detectability and apparent size are a sensitive function of the defocus. Figure 3.9 shows micrographs of identical areas of catalyst at various values of defocus. (Reproduction may have reduced the quality of the micrograph; prints are available on request from the author.) Between the two micrographs shown in Figure 3.9 (a) there is a difference in defocus of 80 nm. Regions "a" and "d" show the distortion of apparent shape effected by the defocus change. The dark region "a" appears to resolve into two crystallites in the right hand micrograph, while the shape of particle "d" is substantially altered. The region of contrast above "c" is enhanced in the right hand pic-ure. Regions "b", "e", "f" and "g" show the fine contrast detail which alters with



Figure 3.7

.7 Calculated apparent image size as a function of defocus. The size of the image is not a simple function of the size of the specimen particle.









Change in Defocus ≠ 80 bh





.0 ni

÷.

10 nm

Change in defocus = 40 pm

Figure 3.9 (a), (b). Micrographs of a Pt/Al_2O_3 catalyst showing the effect of defocus change on image contrast.

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defocus; in each region particular sub-nanometer structure is apparent or highlighted in one micrograph, but not in the other. Similar effects are observable near regions "a", "b" and "c" in Figure 3.9 (b), for which the change in focus is 40 nm.

The fact that it is not possible to decide from such micrographs whether this fine contrast arises from platinum particles or from the substrate is discussed later.

Measurements of apparent image size were made for a number of particles in each micrograph of a through focal series. The results are plotted in Figures 3.10, 3.11 and 3.12 for three separate fields of view, and it is evident that "window sizes" are smaller and relative fluctuations in apparent sizes greater, for the smaller contrast regions. These results confirm the qualitative predictions made on the basis of the phase contrast calculations (see Figures 3.5 and 3.6 and Table 3.1).

These results make it quite clear that the number of particles which will be detected in a standard size analysis will depend upon the average defocus value of the micrograph as a whole, and on the particular defocus range existing within the micrograph. Identical small particles at different elevations in the specimen will have different apparent sizes, or may not be detectable simultaneously. The observed variation in apparent size of up to 1 nm imposes an uncertainty on measured sizes which implies that for analysis of size distribution, division of sizes into classes which differ by less than 1 nm is not warranted.

As aperture size is reduced, calculations indicate that the variation in image intensity and apparent size decreases, but at the expense 40

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Figure 3.10 Variation of measured particle size with defocus for three through focus series of micrographs of a Pt/Al₂O₃ catalyst. Zero values mean that the particle was not detected.





A.2



Figure 3.12 Variation of measured particle size with defocus for three through focus series of micrographs of a Pt/Al₂O₃ catalyst. Zero values mean that the particle was not detected.

of contrast and size information^{3.9}. This is shown in Table 3.2, where for the single atom case the maximum image contrast and apparent size are shown over a defocus range for varying apertures, and hence varying values of α_{max} . For the α_{max} value of 0.005 radians, no contrast above 5%, light or dark is realized in this defocus range, while for a very small aperture giving an α_{max} of 0.00125 radians, even 0.1% contrast is not achieved. As the aperture is decreased, the image size increases, until for the smallest aperture, the atom, if detected, would appear to be greater than 2 nm in size. Thus variations in image appearance with defocus are reduced by smaller apertures, but at the cost of image intensity and size definition of the image.

Through focal series of images were recorded of the same specimen area for apertures subtending angles of 0.01, 0.006, and 0.67 radians. The predicted loss of resolution was observed with some very small regions of dark contrast, clearly resolved in the image recorded with an m_{max} of 0.01, but being progressively washed out through lower contrast and increased size as the aperture size was reduced.

It may be noted that if one is not concerned with detection of particles less than about 2 nm, the use of smaller apertures is probably advantageous. The variation of intensity with defocus is reduced for smaller apertures and the diffraction contrast of particles is greater, although we still observed some variation of apparent size with defocus.

3.5.3 The Effect of Tilt on Contrast

The sensitivity of the contrast of both metal microcrystallites and support particles to the orientation of the electron beam was investigated using the high resolution tilting holder. Micrographs of

Table 3.2: Calculated Image Intensity and

Apparent Size Over a Range of

Defocus and Apertures

aperture, um a _{max} , radians	40 0-01	26	32 0.01	100	16 0.00)50	8 0.00)25	4 0.00	125
defocus, nm	1 1 9 9	do b	1 Max	đõ	Imax	do	Imax	do	I max	do
-20	1.039	0,56	1.051	0.56	1.017	0.70	1.001	1.44	1,000	2.0
· 0	1.053	0.50	1.036	0,52	1.007	0.66	1.001	1.30	1.000	2.0
20	1.037	.0.40	1.035	0,40	0.996	0.86	0,999	1.46	1.000	2.0
40	1.005	0,14	1.023	0,25	0.986	0.76	0,999	1.45	1.000	2.0
60	0.950	0.56	0.943	0,56	0.976	0.75	0.998	1.45	1.000	2.0
- 80	0.860	0.43	0.847	0.45	0.969	0.75	0.997	1.45	1.000	2.0
100	0.868	0.46	0.867	Q.48	0.964	0.76	0.997	1.45	1.000	2.0
150	0.954	0.80	0.993	0.74	0.961	0.78	0.996	1.45	1.000	2.0
140	1.105	0.26	1.060	0,26	0.962	0.80	0.995	1.45	1.000	2.0
160	1.014	0.18	1.003	0.12	0.965	0.86	0.995	1.44	1.000	2.0
180	0.865	0.30	0,944	0.40	0.970	0.92	0.994	1.44	1.000	2.0
200	0.049	0.20	0.972	1.20	× 7	1.04	0.994	1.45	1.000	2.0

a - calculated so that background intensity is 1.0

b - in nn

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a specimen being tilted differ not only in angle of orientation to the electron beam, but also differ slightly in the defocus condition. This latter occurs because during tilting the specimen shifts physically in the object plane, and an exact restoration of the previous defocus value is impossible. However variations in regions of contrast greater than 2 to 3 nm can be attributed to orientation effects since phase contrast effects become minimal at these sizes.

The micrographs in Figure 3.13 show identical specimen areas with a variation in tilt angle. In Figure 3.13 (a), a tilt change of 15° significantly alters the appearance of the metal particles. The particle in region "a" disappears in the right hand micrograph, while a third crystallite in region "c" is more clearly evident in the right hand micrograph. The two particles in region "b" appear as one in the right hand micrograph possibly due to superposition. The sensitivity of support contrast to drientation is evident to the right of region "c". Similarly, in Figure 3.13 (b) several contrast alterations are evident with a tilt change of 7. The relative contrast of the two particles in region "a" changes, revealing the sensitivity of contrast to prientation. A contrast region to the right of region "c" is not detected in the right hand micrograph. In regions "b" and "d", the background contrast is reduced substantially as a result of tilt; in region "b" this results in highlighting of a particle virtually obscured in the left hand photograph.

The simultaneous variations in support and metal particle image contrast again raises questions about the reliability of single micrograph particle size distribution analysis. The variation in orientation leads to a change in apparent contrast of a particle. For larger





Figure 3.13 (a), (b). Micrographs of a Pt/Al_2O_3 catalyst showing the effect of tilt on image contrast.

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particles, this contrast change would not be sufficient to prevent detection, but for smaller particles the orientation clearly can prevent their being included in a particle count. Since a typical supported metal catalyst contains particles at all orientations, the

3.5.4 Contrast Stand Material

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An additional concern in assessing the accuracy of particle size distribution analysis is the distinguishability of small crystallite images from the contract inherent in the support. Typical support materials, such as alumina or silica, have a highly irregular structure leading to high surface areas. This irregularity is a desired feature for catalysts, allowing a large gas-solid interface and a high dispersion of metal. Diffraction evidence from alumina support materials confirms a fine polycrystalline structure.

The polycrystalline irregular structure leads to considerable contrast variation in the micrograph of alumina itself. Such contrast structure is on a small (down to 0.5 nm) scale, and is particularly evident in regions of crystal overlap.

Figure 3.14 demonstrates this effect. The left hand micrograph shows two regions of catalyst at different elevation. In the right hand micrograph, the grid has been tilted through 25° so that these regions now overlap. Considerable contrast structure is evident along the line of overlap which does not arise from the presence of metal crystallites detectable in the left hand micrograph. This contrast structure is indistinguishable in a standard micrograph from that which is generated by small crystallites. Examination of micrographs of pure







Figure 3.15 Micrograph of pure $A1_20_3$ ("Alon") specimen showing the contrast inherent in the support.

Non, with no platinum added, show this fine contrast structure, again particularly in overlap and boundary regions; Figure 3.15 illustrates this.

The inherent support contrast structure limits the lower size limit to which particle size analysis may be extended. Even where microscope resolution is better than 1 nm, distinction of metal particle from support contrast in this size range is virtually impossible in a single bright field micrograph.

3.6 Conclusions

A number of conclusions concerning the determination of particle size distribution of supported metal catalysts from electron micrographs emerge from the work reported in this chapter.

1. For metal particles below 2 nm in diameter:

a) Because of the phase contrast mechanism, detection is a sensitive function of the defocus and hence of the spatial elevation in the specimen. As metal clusters get smaller, the defocus range over which they are detectable as a region of dark contrast decreases;

b) Apparent size is a sensitive function of defocus, since the microscope "filters" certain spatial frequencies and thus distorts the images. Thus identical clusters will have apparent images which vary with their spatial elevation in the specimen, within a single field of view. Identical particles in separate micrographs could appear to have differing sizes because of different settings of the objective focus. Further, variation in apparent particle size of up to 1 nm suggests this value as the lowest meaningful division of diameters in a particle size distribution;

c) An inherent contrast is generated by the use of irregular polycrystalline supports. Particularly for metal particle images below 1 nm, distinction of particle images from background contrast is virtually impossible within a single bright field micrograph.

- For larger metal particles, identical crystallites can mave markedly different contrast due to different orientation to the electron beam.
- 3. For all sizes of metal particle, the effects of overlap, and the orientation sensitivity of the contrast of the support material, can seriously affect size analysis. Certain comb nations of particle/support orientation make the particle undetectable.

It is evident that these conclusions are incompatible with the assumptions implicit in some applications of particle size distribution analysis of supported metal catalysts. Clear evidence of the detection of a crystallite of a given size does not imply that all crystallites of that size and larger are being detected, because of contrast window and orientation factors. Small image sizes cannot in general be directly correlated to the particle size in the specimen, preventing computation of a meaningful average size for catalysts containing small metal particles. Analysis of such catalysts is further complicated by the difficulty of distinguishing metal contrast from inherent support contrast.

These conclusions draw into question the extension of particle size analysis into the sub-nm size range; they further imply that

analysis of micrographs cannot currently provide a definitive test of a adsorption stoichiometries for small crystallites^{3.6}. As a general rule we would argue that particle size distributions become increasingly unreliable as the size of particles counted extends below 2.5 nm. While micrographs can give evidence of smaller particles, detection and accurate identification of all particles of a size below 2.5 nm is extremely unlikely.

It should be emphasized, however, that the resent studies have been concerned only with the use of conventional high resolution transmission electron microscopes, using stand of brack field imaging. There are a number of other imaging technic which may prove wore shore after similar detailed evaluation. The use of dark-field which have proved valuable for the detection of single heavy. atoms^{3.21}, and for relatively large supported metal particles^{3.22} may show promise for high resolution work. An even more promising possibility in the long term may well be the use of scanning transmission instruments of the type developed by Crewe and his colleagues^{3.23} which permit a range of new contrast mechanisms to be applied.

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CHAPILI 4

THE TITRATION REACTION ON SUPPORTED Pt CATALYSTS

4.1 Survey

As discussed in Chapter 1, gas adsorption on metal surfaces has become a fairly standard technique for the determination of the dispersion of supported metal catalysts. Conversion of a corrected volumetric gas uptake to a metal surface area or dispersion requires the assumption of a stoichiometry of adsorption. In light of the current controversy over an appropriate stoichiometry, detailed below, experiments were designed to explore the adsorption of oxygen and hydrogen, and their interaction on the surface of a supported to catal. The IR spectra and static system gas uptake results reported in this chapter were recorded by Dr. E. Kikuchi, and then combined with the flow system gas uptake results for interpretation.

In their studies of low concentration supported Pt catalysts, Benson and Boudart^{4.1} proposed that H_2 titration of adsorbed O_2 would increase the sensitivity of measurement. In interpreting their results they used the stoichiometry

 $Pt_{\overline{surf}} 0 + \frac{n}{2} H_2 \rightarrow H_2 0 + Pt_{\overline{surf}} H_{(n-2)}$

4.1

with an n value of 3, but Mears and Hansford^{4.2} found n values closer to 4. Wilson and Hall^{4.3} proposed to resolve this discrepancy by postulating that n is 3 for large crystallites, but incomplete surface coverage by 0_2 on small crystallites leads to apparent $\frac{1}{2}$ values of 4 for highly dispersed catalysts. Dalla Betta and Boudart^{4.4} similarly

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found that small platinum clusters on Y zeolite took up less oxygen per surface atom than larger platinum crystals. Like Wilson and Hall, they attributed this phenomenon to the relative electron deficiency of small particles as compared to large ones. Basset $P t at_{+}^{4.5}$ studied heats of adsorption and reaction for both the titration of oxygen, equation 4.1, and the titration of adsorption hydrogen

 $Pt_{surf} H + \frac{m}{2} 0_2 \rightarrow H_2 0 + Pt_{surf} (m-1)$

They also found an n value of 3 for large metal particles, and calculated low heat of adsorption values (about 18 kcal/mole) for the capture by the support of the H_2O generated during titration.

A preliminary survey of titration data in this laboratory showed considerable deviation from the simple stoichiometric picture presented by equations 4.1 and 4.2. Initial hydrogen uptakes generally exceeded initial oxygen uptakes. By "initial" we mean the gas adsorbed on the "degassed" catalyst surface, as compared to that calculated as remaining on the surface after a titration step, which we refer to as secondary, tertiary, etc. uptakes. In addition, the secondary or tertiary uptakes of gases differed from the initial uptake of the same 'gas on a "degassed" surface. We frequently observed that

 $H_{A}^{+} > H_{A}^{-}$

and $0_{\rm A}^{\prime} \leq 0_{\rm A}^{\prime}$

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where H_A' and O_A' are the uptakes of hydrogen and oxygen after titration, corrected for the amount of gas consumed to form water, and H_A and i are initial gas uptakes. Thus titration seems to enhance the adso tion of hydrogeneric to retard the adsorption of oxygen on Pt-on fluminal
catalysts.

In this chapter the deviations from a simple constant stolchiomethy are documented by static and flow system gas uptake data. The data of other authors are also summarized. Three possible explanations of the deviations are discussed: a modification of the surface area, as proposed by Darensbourg and Eischens^{4.6}, the formation of more than one type of adsorbed oxygen species, and a promotion/retardation effect stemming from the water generated during the titration. IR spectroscopy was used to examine the nature of the surface oxygen species during adsorption and titration.

Infrared spectroscopy has been applied extensively to study adsorbed species on satalysts. Heyne and Tompkins^{4,7} have recorded IR spectra of CO adsorbed on silica-supported Pt pretreated with oxygen at 300°C. They observed two different kinds of CO adsorbed on platinum sites: one on platinum metal (2080 cm⁻¹), the other on platinum ion (2120 cm⁻¹). They also found that oxygen adsorbed on platinum metal could be removed by carbon monoxide (as carbon dioxide), while that on platinum ion could not. Primet at 4.8 concluded from their IR study that a part of the inface platinum adsorbs both oxygen and carbon monoxide on the same site; and gives an IR absorption band of C=0 stretcning shifted to 2120 cm⁻¹. They showed that such species $Pt<_{CO}^{O}$ could be obtained even it noom temperature by repeated (0₂-CO) treatment.

These investigations with IP spectroscopy indicate that the adscrption of oxygen on Pt is not simple: namely there are at least two different kinds of adsorbed species of oxygen. In this work, volu-. metric uptake data has been supplemented by an IR study of the nature of the surface oxygen species after repeated O_2 -CO, and O_2 -H₂ treatments, \$6

and after Ω_p adsorption at temperatures from 25°C to 500°C.

4.2 Experimental Results

Table 4.1 shows the details of the catalysts used in this work. For both the static and flow system, gas uptakes were measured at room temperatures except as noted in the text.

4.2.) Isotherms and Isobars of Oxygen and Hydrogen

Figure 4.1 illustrates typical isotherms of oxygen uptake obtained with the static system at temperatures from 25°C to 500°C for catalyst 1. The adsorption isotherm was extrapolated to zero pressure to obtain the adsorbed amount of oxygen on platinum.

Figure 4.2 shows oxygen and hydrogen uptakes as a function of tem-Perature measured in the static system for catalysts E and I, and in the flow system for catalyst A. Hydrogen uptakes were remeasured at room temperature after reducing the sample used for the measurement of each static oxygen adsorption isotherm (Table 4.2). For catalyst I, reduction at 500°C was enough to restore the original hydrogen uptake value. On catalyst E, howeyer, subsequent hydrogen uptakes were found to be depressed by 24.4% and 35.4% after oxygen adsorption measurements at 400°C and 500°C, respectively. Assuming that these decreases in hydrogen uptake resulted from a decrease in metal surface area, oxygen uptakes at these temperatures were corrected and the corrected values are snown in Figure 4.2 by the dotted line.

Oxygen uptakes were found to increase with increasing uptake temperature, while hydrogen uptakes decreased. In the room temperature region hydrogen uptake is higher than oxygen uptake for both catalysts A and E, as measured by both systems.

Table 4.1: Description of Catalysts

Catalys		Content t%)		Type of Support
Â		0.5		Engelhard (Lot #18-381)
В		0.5		(1)
С		0.5	• •	(2)
Dur		0.5		(3)
E	•	4.76		Alon (4)
F	1)	2.46		Alon
G		0.10		Alon
H		1.00		Alon
Ι	· .	2.03		Alon
ال و ل	х	1.61		Kaiser 201 alumina
· K		2.03		Kaiser 201 alumina

(1) same as catalyst A except sintered for 16 hours in O_2 at 450°C (2) same as catalyst A except sintered for 16 hours in O_2 at 600°C (3) same as catalyst except sintered for 16 hours in O_2 at 700°C (4) Registered trademark of Cabot Corp. \clubsuit 58

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Figure 4.1 Oxygen isotherms for a 2.03% Pt on Alon catalyst



Figure 4.2 Oxygen and hydrogen isobars for three different Pt/Al₂O₃ catalysts.

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Table 4.2:	O_2 Adsorption Uptakes as a Function of Temperature
and Room	Temperature H ₂ Titration and Adsorption Uptakes

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Catalyst	Temp°C	<i>cicomption</i> Uptake atoms/atom)	Room temper <i>hydrogen tit</i> Uptake (atoms/atom)	mation HT/OA	Subsequent room temperature hydrogen adsorp- tion uptakes (atoms/atom)
I	RT	0,31	1.39	4.48	0.73
	100°C	0.36	1.49	4.14	0.72
	197	0.42	1.69	4.12	0.73
	., 290	0.49	1.72	3.44	
	387	0.63	1.62	2.57	0.73
	485	0.75	0.96	1.30	0.74 ⁻
Ē	RT	0.30	1.32	4.40	0.61
	100	0.32	1.13	3.53	
	200	0.40	1.30	3.25	
	300	0.46	י' ן	2.85	0.60
	395	0.40	1.04	2.60	0.45
	500	0.39	0.72	1.85	0.39

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4.2.2 Hydrogen Titration of Oxygen Adsorbed at Elevated Temperatures

Experiments were designed to test the reactivity of oxygen adsorbed at various temperatures with gaseous hydrogen. Oxygen was added at 100 torr to catalysts E and I at temperatures from 25°C to 500°C, following which the catalyst was evacuated for one-half hour and cooled to room temperature. Hydrogen titration uptakes were then measured. Oxygen and hydrogen titer uptakes are shown in Table 4.2.

These data suggest that oxygen adsorbed at higher temperatures is less reducible by hydrogen at room temperature in the oxygen adsorbed at 25°C. Oxygen adsorbed at elevated temperatures may form a norther type of surface species or may oxidize metal below the surface layer. This phenomenon was further studied by IR spectroscopy.

4.2.3 IR Spectra of CO Adsorbed on Pt with O2 Preadsorbed at Various Temperatures

IR transparent discs of catalyst I were contacted with 0_2 at temperatures from 25 to 500°C in the in dita IR cell. Following 0_2 adsorption (100 torr 0_2 pressure) the sample was cooled to room temperature, the cell was evacuated and the catalyst was contacted with CO, at 100 torr, for two hours. IR spectra, following these treatments, showed absorption bands with maxima near 2120, 2060-80, 1800, 1640, 1480 and, 3600 cm⁻¹. The last four bands correspond to those observed by Parkyns^{4.11} when CO₂ was adsorbed on alumina. Eischens and Pliskin 4.12, in their study of CO adsorption on supported Pt, observed bands at 2060-80 and 1800 cm⁻¹, which they attributed to the linear and bridged forms of adsorbed CO. Heyne and Tompkins^{4.7} and Primet at al.

to the nature of the bond responsible for this band differs; the former workers attribute it to CO adsorbed on Pt ions while the latter assign it to CO adsorbed on a Pt atom which also has an adsorbed oxygen atom.

In our work the bands at 1480, 1640 and 3600 cm^{-1} , and part of the band at 1800 cm^{-1} , are probably due to the interaction of CO . formed by the oxidation of CO with adsorbed 0_2 , with the alumina support. CD adsorbed on Pt, probably in the bridged strate, als con tributes to the band at 1800 ${\rm cm}^{-1}$ since it is present when LU is adsorbed on a freshly reduced sample. When oxygen was introduced at room temperatu — to a freshly reduced sample onto which CO has been adsorbed, the absorption bands at 2060-80 $\rm cm^{-1}$ and near 1800 $\rm cm^{-1}$ decreased in intensity, and a new band developed at 2120 cm⁻¹. This . new band is the one also observed by Heyne and Tompkins^{4.7} and Primet et at, 4.8. The intensity of the 2060-80 cm⁻¹ band initially decreased rapidly, but it did not disappear completely over a period of several hours unless the catalyst was heated to 100°C. Heyne and Tompkins^{4.7} reported that this band rapidly disappeared completely at room temperature. This difference is possibly due to the different support, as postulated by Eischens and Pliskin^{4,12}, or it may be caused by differences in Pt crystallite sizes.

The band at 2120 cm⁻¹ was also observed when CO was introduced to the catalyst which had preadsorbed oxygen at elevated temperatures. This is illustrated in Figure 4.3. The intensity of this band increased with increasing temperature of oxygen adsorption up to 400°C, then decreased when 0_2 was preadsorbed at 500°C. The 2060-80 cm⁻¹ band, however, continually decreased with increasing oxygen adsorption



= 4.3 Variation of the 2060-80 and 2120 cm⁻¹ bands from C0 attronted on $Pt/A1_0^{-0}$ catalysts with different Ω_2 preadsorption temperatures. a) freshly reduced sample b) Ω_2 preadsorbed at 200°C c) at 400°C d) at 500°C

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temperature. These results suggest that oxygen is taken up by supported platinum on at least two different sites. One site which gives the 2060-80 cm⁻¹ hand when CO is admitted, is generally associated with adsorption of oxygen on a platinum atom. This species of oxygen is readily removed by CO in a reaction to form CO_2 , and the band thus arises from CO adsorbed on the "clean" site. The second site for adsorbed oxygen, which gives rise to the 2120 cm⁻¹ band when CO is introduced, has been associated with platinum ion, or a platinum atom which simultaneously adsorbs CO and oxygen. This second species of oxygen is thus less reactive with CO, and is formed more extensively as the temperature of oxygen adsorption increases up to 400°C. For convenience we refer to this second species of adsorbed oxygen, and to the previously described site as Type I. However, it should be emphasized that the exact character of the sites is not definitively established.

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The 2120 cm⁻¹ Type II band decreases when the samp is treated with hydrogen at room temperature and is recontacted with CO. At the same time the 2060-80 cm⁻¹ Type I band increases; suggesting that hydrogen is capable of reducing some Type II sites to metallic platinum which subsequently adsorbs CO. However, the higher the temperature of oxygen preadsorption, the less reducible the Type II oxygen becomes.

A series of experiments was designed to further investigate the nature of the two oxygen sites. Oxygen was adsorbed on two freshly reduced samples of catalyst I, at room temperature for one sample and 400° C for the other. Subsequently both samples were evacuated at 400° C for 16 hours. The sample that had O_2 adsorbed at room temperature

did not show any further 0_2 uptake when oxygen was readmitted at room temperature. However, the other sample took up 21.5° of the original 0_2 uptake when oxygen was readmitted at 400°C. These results suggest that oxygen may be more easily removed from Type II sites, formed at 400°C, than from Type I sites, which predominate in 0_2 adsorption at 25°C. The removal could be either resonant or migration into the metal to form a bulk oxide.

In other experiments CO spectra were recorded of wo samples of catalyst I after adsorption of 0_2 at 400°C, one af alf hour evacuation, and the other after a 16 hour evacuation. Spectra, shown in Figure 4.4, again indicate a sizeable decline in Type II sites. But almost no change in Type I sites. Similarly, a spectrum recorded after 0_2 adsorption at room temperature followed by 16 hour evacuation at 400°C shows on 2° a slight decreases in the 2075 cm⁻¹ peak from the Type I site, and no evidence of a peak at 2120 cm⁻¹.

These results not only suggest that oxygen is re-vable from Type II sites far more readily than from Type I sites, but also indicate that Type II species in out formed from adsorbed Type I species. Heating in vacuum a sample on which 0_2 had been adsorbed at room temperatures to temperatures (400°C) where Type II sites are formed does not generate Type II oxygen. Thus Type II sites may be formed only by contact with gaseous oxygen or, as will be subsequently shown, during cyclic treatment at room temperature with CC and 0_2 .

4.2.4 Gas Uptakes in Titration Sequences

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Table 4.3 shows the results from a number of titra is sequences in both directions (i. 0_2 -H_Z- 0_2 and H₂- 0_2 -H₂). While the trend is



not universal, it is evident that secondary and tertiary hydrogen uptakes (H2' and H3') are generally higher than the initial hydrogen uptake, while secondary and tertiary oxygen uptakes (O2' and O3') are generally less than or equal to the initial oxygen uptake. (See Table 4.3 for nomenclature.) A survey of other reported titration data 4.2, 4.3, 4.5, 4.13, 4.14 owed that only-vilson and Hall report instances where secondary hydrogen (See the initial hydrogen uptake. 68

Figure 6.5 shows a plot of the post-titration hydrogen enhancement (12'/H1. or H3'/H1) as a function of the hydrogen uptake to oxygen uptake (H1/O1) ratio. This plot includes the data of Hilson and Hall^{4.3, 4.13} and Basset *et al.*^{4.5} as well as our own. The data of Gruber^{4.14} was excluded because oxygen uptake was measured at a different temperature (350°C) than the hydrogen uptake, and samples were evacuated at 500°C between titrations.

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Two in sting observations emerge from inspection of Figure 4.5. First, all of the points from these experiments which show a hydrogen enhancement factor less than unity were recorded at 0°C of were sintered in 0₂ before measurement. For all room temperature measurements of non-sintered specimens, secondary and tertiary hydrogen adsorption is higher than the initial hydrogen uptakes. One 'titration sequence (Run 83) was run in a methanol ice bath (-98°C). The tertiary hydrogen adsorption is only 10% of the initial uptake, which we postulate arises from the fact that generated H₂O freezes and blocks access to the metal surface. The O°C points similarly may experience some surface blockage, leading to lower enhancement ratios than occur in titrations at room temperature or higher.

69 35 والمدقوه H1/01 H2 Adsorption 02 Adsorption Corrents actios 1:10.1 0.93:1 ි 0.97:1-1:66 0 0.58:1 0.63:0.71:1 0.32:1 0.93:1 0.86:0.99:1 1.21.1 1.51:1.52:1 0.86:0.72:3 1.01:1.07:1 0.77:5.53:1 Ad section and Titration Uptakes 43' : H2' : H1 03' ٢x 1.49 1.25:1.40:1 ۱:36.92 1.77.01.32 1. -0.37:1 -1.36:1.d5:1• 1.06 7-17:1 1.10:1.04:1 0.53:1 1.25:1.05:1 1.35:1.07:1 1:00.1 1:72.1 و .0.95.i 1:01:----:10 1.19:----1 0.09:---:1 1.62:---:1 -⁶7.28 2.36 ري ج . 84 1.59 1.42 05.1. -..... 3.00 1.26 1.37 1.33 1.71 • 0.48 0.64 0.42 0.6. 99. 65 1 0.54 03 0.48 -0.46 0.96 01.1.525.5 ʻó.25 0.91 د 0.58 0.57 0.60 0.70 3.39 1.86 1.34 42 0,12 0.12 0.20 0.20 0.03 0.18 0.26 0.25 0.18 X) o 0.35 0.36 0.31 • •• 1.32 0.46 0.97 0.77 0.50 0.96 0.77 0.86 0.44 i 42 1.36 0.36 0.81 3 Ĥ 0.46 ø 0.29 0.42 0.43 0.22 0.83 0.25 0.29 0.36 0.390.34 0.)2 1.34 0.45 0.26 J. 53 0.55 0.32 0.40 - 0.61 0.22 8 0.28 0.22 0.27 0.23 0.09 0.16 0.380.39 0.73 0.32 0.37 0.300.60 0.22 0.22 0.14 **8**.36 8.1 0.22 Catalyst Temperature able 4 j∘75 ວംດ້ 0°0 206°C -98°С 0°C 0.0 0,0 0,0 л. Ж RI ₽ ₽ 5 E 5 ĥ 5 Ľ, <u>م</u> 74 & 74a 104 & 105 90 1 50a 17 8 02 60 & 61 62 5 63 64 2 65 79 2 80 31 1, 32 106 58 1 59 56 1 57 11 75 Ē 9 3 87 8





"Second, Wilson and Hall's data reveals cases where low enhancement factors are observed. All of these points occur where the initial hydrogen uptake is Jess than about 1.3 times the initial oxygen uptake. Finally, a check of Wilson and Hall's pretreatment conditions indicates that all their points showing enhancement ratios less than Sunity had been exposed to air or hydrogen at temperatures of 545°C or higher, with most treated at temperatures greater than 600°C. Thus from Figure 4.5 the trend is evident that catalysts which have initial i up takes greater than about 1.3 times the initial oxygen uptend to show enhancement of hydrogen adsorption after one or two titrations, particularly at rodm comperatures and highen. Figure 4.6 shows a plot of the ratio of initial hydrogen uptake to initial oxygen uptake (H1/01) versus the catalyst dispersion as measured by hydrogen uptake (H1). Again, the data of Milson and Hall 4.3, 4.13 and Basset of al. 4.5 are included. White considerable scat ter is evident in the data, a trend to a higher H/O ratio with increasing dispersion (smaller particle size) is evident. Thus, as Wilson and Hall suggest, chemisorption data supports a variable stoichiometry. with hydrogen and cxygen uptakes being about the same only of larger platinum crystallites.

4.2.5 Long Titration Sequence

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One experiment (Run 75) on catalyst J was extended for a total of 14 (O_2-H_2) titration sequences. Figure 4.7 shows the resulting uptakes of titered oxygen and hydrogen, as well as the ratio of hydrogen uptake to oxygen uptake for each step. The final HT/OT ratio is does not approach 2.0 as predicted by stoichiometric considerations,





but rather is asymptotic to about 2.02. This 15 factor could well arise from the failure of the flow system to detect a small portion of the hydrogen which is reversibly adsorbed and "bleeds" off. However, it is evident that hydrogen adsorption enhancement continues through the first 8 titration sequences, and thus this effect is not restricted only to the first titrations.

4.2.6 IR Spectra of CO Adsorbed on Pt After Repeated Sequences of (0_2-CO) and (0_2-H_2) Treatments

a) $\sqrt{(0_{2}, 0)}$ cycles

Oxygen (100 torr) and carbor monoxide (1 torr) were cyclically introduced, at room temperature with enacuation for one-half hour between cycles, to catalyst I, and IR spectra were recorded after each addition. Figure 4.8 shows some of the recorded pectra.

After the second CO adsorption the 2070-75 cm⁻¹ band reached a maximum, and subsequent (0_2-CO) sequences caused this band to decrease in intensity. On the other hand, the field cm⁻¹ band increased with cyclic treatment. Thus repeated $(CO-O_2)$ treatments generate increasing amounts of Type II surface adsorbed oxygen, which is not reduced by we next CO titer.

When freshly reduced catalyst I was contacted for 17 hours with oxygen at room temperature, subsequent CO treatment did not generate the 2120 cm⁻¹ Type II oxygen band. Thus it is the cyclic treatment with CO which promotes the formation of the Type II site, rather than the time of contacting with Ω_2 . When hydrogen was introduced at room temperature to a cyclically treated catalyst, the 2120 cm⁻¹ band decreased but was not totally reduced. The 2070-75 cm⁻¹ band shifted

5 с U o treatment ani)i 2000 %0 6th cycle %0 ic. treatment: 9 2000 5 NUMB CVC 5 -- 70 % 0 489 CO adsorptión Figure 4.8 The effect of 2200 lst cych 2) σ \$000 g . n__ 2200 displaced °/₀ NOISSIMSNAAT (ordinates 5.0

to lower frequency ($\Delta v = 20 \text{ cm}^{-1}$).

5) $(0_2 - H_2)$ cycles

In these experiments fresh catalyst was contacted with oxygen, then hydrogen, and then CO was introduced to the cell and the spectra recorded. Typical results are shown in Figure 4.9. As found by Darensbourg and Eischens^{4,6}, the 2070-75 cm⁻¹ band is enhanced by subsequent treatments. In addition, we observed the band to shift to 2060 cm⁻¹ (auxi15 cm⁻¹). This shift is almost identical to the shift observed by Primet at at.^{4,8} when water was added to their CO adsorption system. We thus tentatively ascribe the shift to a modification of adsorbed CO electronic properties arising from the H₂O generated during titration.

Evidence of H-O-H bending, in the 1630 cm⁻¹ band shown in Figure 4.9, further demonstrates the presence of water. Also, absorption in the 3500-3800 cm⁻¹ region, due to alumina OH groups, increased with increasing number of 0_2 -H₂ cycles, indicating that some of the generated H₂0 was being taken up by the alumina.

The 1835 cm⁻¹ band was enhanced by repeated Ω_2 -H₂ cycles, as shown in Figure 4.9. This was not observed in the cyclic Ω_2 -CO treatment. The most outstanding difference; however, between the two treatments is that the 2120 cm⁻¹ band, attributed to Type II oxygen, was not detected in the cyclic $(\Omega_2$ -H₂) treatment, even after six cycles.

4.3 The Origin of the Enhancement Effect

Enhancement of hydrogen uptake following titration is demonstrated by the results in Table 4.3 and Figure 4.5. It may be argued that this apparent increase is due to incomplete H_2 desorption during the

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relatively short degassing periods after reduction (1 to 2 hours at 500°C). However, an absolute measure of enhancement or retardation, independent of initial surface hydrogen coverage, can be calculated, as shown in equations 4.5 to 4.10.

 $Pt-H_x \xrightarrow{H_1}{2} H_2$, $Pt-H_a \xrightarrow{O_2}{2} 0_2$, $Pt-0_b \xrightarrow{H_3}{2} H_c$, $Pt-H_c$. +b H20 w ta H20 $\frac{01}{2} \xrightarrow{0}_{2} \xrightarrow{\text{H2}} \text{Pt-H2} \xrightarrow{\text{Pt-H2}} \text{Pt-H2} \xrightarrow{03} \xrightarrow{02}_{2} \xrightarrow{02}_{1} \xrightarrow{\text{Pt-O}_{f}} \text{Pt-O}_{f}$ +€ H20 A +X.H.0 +d .H20

where 'x' is the number of H, atoms remaining on the platinum per platinum atom after evacuation, and [a',]b', c', d', e'; and \mathcal{T}' are the number of hydrogen or oxygen atoms adsorbed per platinum ... atom at various stages in the titration sequences. All staichiometric coefficients in these equations are expressed in terms of the metal atoms, hence they are not an absolute surface stoichiometry.) The absolute enhancement in Ho adsorption is given by (c-a) and (e-a). According to Equation 4.5

(c-a) ≈ H3 ~ 2(92).

and from Equations 4.5 and 4.6

(e-a) = H2 - 2(01) - H1

4.3

4.10

Similarly, the retardation of O_2 adsorption is given by $(d-f) = \frac{H2}{2} - 03$ 4.9

and

 $(d-b) = 01 + \frac{H1}{2} - 02$

The values of (c-a), (e-a), (d-f) and (d-b) obtained from our data are tabulated in Table 4.4. The results indicate a general enhancement of H_2 uptakes during titration and a retardation of Ω_2 uptakes. Thus the variation in adsorption stoichiometry which we observed during titration is not attributable to insufficient degassing.

Darensbourg and Eischens^{4.6} noted H_2 enhancement in their IR studies of supported Pt catalysts, and attribute it to an increase in metal surface area arising from the room temperature reduction of the adsorbed oxygen. In examining our results, we have considered this hypothesis and others: the creation of different species of surface oxygen; as occurs during repeated (CO-O₂) treatments, and an influence from the water generated during titration.

The primary difficulty with an interpretation of increased surface area following titration is that it fails to explain the retardation or lack of enhancement of oxygen uptake noted during titration. In some cases the reduction in O_2 uptake exceeded 35% on the second titer (O3') and 30% on the first titer (O2'). It is difficult to conceive of n alternate increase in surface area during a hydrogen titration, followed by a decrease in surface during an oxygen titer. For this reason we conclude that a simple physical fize explanation for hydrogen enhancement is not satisfactory.

Run	Temperature	H ₂ en H ator (c-a)	ns / Pt a	t 0_2 tom 0 atoms e-a) (d-f)	taration 2 / n° com (d≖b	
	R.T.	0,26	0.0	1 0.07	0.10	
	R.T.	0.21	0.04	0.13	0.10	
	R.T.	0.26	u 0.06	0.05	0.10	
83	~98°C	-0.34			:	·
56 & 57	0°C		0.03		0.005	
58 & 59	0°C		0.02	1	0.001	
6	+97°C		0.04			
87	+206°C		0.02			
60 & 61	- 0°C		-0.02		0.02	• .
62 & 63	0°C	٢	-0.06	•	0.02	
64 & 65	· O°C		-0.01		0.01	
79 & 80	R.T.	0.06	, 0.02	0.04	0.03	
81 & 32 -	R.T.	0.03	0.07	-0.002	-0.02	
104 & 105 ·	R.T.	0.09				
106	R.T.	0.005				
90 & 90a	R.T.	0.10	0.16	-0.09	-0.14	
77	R.T.	0.05				.)
70 & 71	0°C		-0.04			
74 & 74a	R.T.	0.02	0.01		. o.oi	1. A.
75	R.T.	0.01	·	•		

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Table 4.4: Hydrogen Enhancement and Oxygen

Retardation During Titration

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at 150°C). However, the Type II sites decrease substantially during high temperature evacuation, while Type I sites as not.

Given these results for (n_2-Cn) treatment, we speculated that cyclical (n_2-H_2) treatment at room temperature might similarly generate the second type of oxygen site. If these sites could still adsorb hydrogen, then enhanced hydrogen and retarded oxygen uptakes could be explained. However, CO adsorption after repeated (n_2-H_2) treatments gave no-evidence of the 2120 cm⁻¹ band. We therefore do conclude that Type II sites are not generated during titration sequences. This does not, of course, imply that other surface complexes may not be formed which account for the H₂ enhancement/0₂ retardation effect.

Another factor to be considered in titration anomalies is the effect of the water generated. The results presented in Tables 4.3 and 4.4 indicate that at temperatures of 0°C and below both H₂ and 0₂ adsorption may be retarded. This may be due to incomplete reaction during titration and/or blockage of the Pt surface by H₂0. At room temperature and above H₂ adsorption is enhanced and 0₂ adsorption is retarded. This may be caused by an alteration of the Al₂0₃/Pt interaction due to the H₂0/Al₂0₃ interaction in the vicinity of Pt crystallites. From IR evidence we concluded that some of the water is taken up by the support. However, the water affects CO adsorbed on Pt sufficiently to generate a shift in the 2076 cm⁻¹ band, also observed by Primet at αl . ^{4.8}. This indicates a change in the electronic properties of the platinum.

The changes in H_2 and O_2 adsorption uptakes are much more marked for the static results than for the dynamic ones. The presence of the carrier gas in the dynamic measurements may facilitate the movement of

water from the Pt surfaces to areas of the support sufficiently removed from the Pt crystallites so as to reduce the effect of water on subsequent adsorptions.

It is impossible to duplicate post-titration we ing conditions by addition of H_20 to a fresh catalyst, because during titration water is generated only at the platinum crystallite locations. However, one experiment (' ns 104 and 105) was made on catalyst F in which a 3.1 g Catalyst charge was wetted by one-half cc of degassed distilled H_20 . While this major dose of water caused both initial hydrogen and initial oxygen uptakes to decrease, the oxygen decrease (37%) was greater than the hydrogen decrease (20%). Thus for the wetted catalyst, the H1/01 ratio was 1.90, compared to 1.53 for the dry sample, again indicating that wetting more greatly retarded oxygen uptake. While this single' result is by no means conclusive, it suggests that investigation of the effect of wetting on gas uptakes warrants further investigation.

It is easy to postulate an influence of crystallite size on the effect of H_20 on gas uptakes. Primet *et al.*^{4.8} attributed the shift of the 2075 cm⁻¹ band in CO adsorption on a wetted catalyst to the electronic donation from the water to the platinum crystallite. Both Wilson and Hall^{4.3} and Boudart and Dalla Betta^{4.4} have cited the relative electron deficiency of small platinum crystallites in explaining oxygen uptake differences among catalysts of varying dispersion. If the water does act as an electron donor, it could affect the smaller crystallites more significantly than the larger ones. Thus the mechanism of uptake enhancement/retardation may be electronic, rather than physical blocking of the surface sites as cited for the -98°C results.

The other interesting point to emerge from this study is a

dependence of the relative uptakes of hydrogen and oxygen, as measured by the ratio H1/01, on the dispersion. Wilson and Hall^{4.3} first noted this in accounting for the discrepancy between Benson and Boudart's^{4.1} and Mears and Hansford's^{4.2} titration stoichiometries. In Figure 4.5, it is evident that Wilson and Hall's data predict the start of a deviation between hydrogen and oxygen uptakes as the dispersion measured by H₂ adsorption exceeds about 0.3. Our data and that of Basset *et al.*^{4.5} indicate that hydrogen and oxygen uptakes differ over the entire dispersion range. This discrepancy may well be attributable to Wilson and Hall's generally more severe pretreatment conditions.

Wilson and Hall used particle size distribution analysis of electron micrographs to attribute the deviation of hydrogen and oxygen uptakes in small particles to Pt₂O_{surf} stoichiometry on small platinum particles. They argued that evidence points to an H/Pt_{surf} ratio of unity regardless of metal particle size. Dalla Betta and Boudart^{4.4} also conclude that oxygen uptake on small particles is less than 1 oxygen per surface metal atom, based on calculations of the size of particles in the cages in a Y zeolite. In our experiments we did not attempt to independently determine hydrogen and oxygen adsorption stoichiometries through separate measurement of crystallite size. Rather, we only note the influence of dispersion, and hence particle size, on the relative uptakes of the two gases.

4.4 Conclusions

Titration of adsorbed hydrogen and oxygen on supported platinum ~ catalyst generally results in enhanced post-titration adsorbed hydrogen values, and reduced post-titration adsorbed oxygen values. The - 84

enhancement occurs over a variety of catalyst loadings and temperatures, and is correlated with the ratio of initial hydrogen and oxygen uptakes, decreasing as H1/01 approaches unity.

Laplanation of the enhancement through modified surface areas, according to Darensbourg and Eischens^{4,6}, does not satisfactorily account for a reduced oxygen uptake. Explanation through the formation of Type II oxygen sites, as occurs in the (O_2-CO) reaction is also unsatisfactory since there is no evidence of Type II sites in O_2 -H₂ titration sequences. The addition of H₂O to catalysts appears to retard subsequent oxygen uptake more than hydrogen uptake. More experimentation is necessary to confirm whether the water generated during titration is the key factor in enhancement.

Type II sites associated with the 2120 cm⁻¹ band are formed during O_2 adsorption at elevated temperatures, and during repeated (N_2 -CO) treatments at room temperature. They are not formed by room temperature O_2 adsorption or by repeated (H_2 - N_2) treatments. Once formed, the sites are not completely reducible by hydrogen at room temperature but do disappear in H_2 at 150°C. The sites are not formed from Type I adsorbed oxygen, even after *in vacuo* heating of Type I adsorbed oxygen.

The ratio of initial hydrogen uptake to initial oxygen uptake on supported platinum catalysts depends upon the dispersion of the catalyst. Relative to oxygen, hydrogen uptake increases with the metal dispersion.

4.5 References

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CHAPTER -5

A MODEL OF SUPPORTED METAL CATALYST SINTERING

5.1 Survey

The overall process resulting in a change in metal dispersion of supported metal catalysts during use or treatment at elevated temperatures is called sintering. Since, in general, sintering results in a loss of catalytic activity, an understanding of the processes occurring during sintering is of importance in the design of catalysts with improved stability. Unfortunately, there is insufficient direct evidence to establish the mechanism of supported metal catalyst sintering.

Ruckenstein and Pulvermacher^{5.1, 5.2} have carried out a detailed analysis of the sintering process by means of a model which envisages the sintering process as a migration of metal particles over the support surface, followed by the fusion of metal crystallites upon collision. Their results showed that the rate of metal surface area change is given by an equation of the form

$$\frac{dS}{dt} = -KS^n \qquad 5.1$$

where the value of n varies between 2 and 8, depending on whether the rate of fusion (particle sintering) or the rate of surface diffusion of metal crystallites is rate controlling. Recently, Wynblatt and Gjo-stein^{5.3} in their sintering studies of supported Pt found the value of n to be z13. They proposed an equation, based on the concept of a nucleation barrier, to explain this large value of n. In their formulation, the resulting equation being somewhat similar to that of

Somorjai^{5.4}, the constant K in equation 5.1 decreases exponentially with increasing average particle radius, r, *i.e.*

$$K = \beta \exp(-Ar/RT)$$
 5.2

Mynblatt and Gjostein did not compare the predictions with experiment, but planned to pursue this in the future.

The model of Ruckenstein and Pulvermacher^{5.1, 5.2}, besides being unable to account for the above mentioned large value of n, encounters other difficulties¹ in being unable to explain some of the experimental observations made during sintering studies. These include:

- Under certain conditions, generally in an oxidizing atmosphere, metal dispersion increases during high temperature treatment^{5.5-5.8}. A simple crystallite diffusion model of sintering process cannot account for increases in metal surface area, although Ruckenstein and Pulvermacher propose to discuss crystallite separation in a future work.
- 2. The high activation energies observed for the sintering process, up to 70 kcal/gmole for Pt on $Al_20_3^{5.9}$, are difficult to account for on the basis of the fusion of two adjacent metal crystallites. (Ruckenstein and Pulvermacher do not quantitatively discuss the activation energies of crystallite surface diffusion and particle sintering.)
- 3. The continued growth of metal particles to the extent that the metal particle size is of the order of the support particle size is difficult to explain on the basis of metal crystallite migration. Figure 5.1 illustrates this from electron micrographs of catalyst treated in this laboratory. The specimen is an impregnated 4.76% Pt on Alon alumina catalyst. The support consists

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Figure S.1: Electron micrographs of 4.76" Pt on Alon alumina after various pretreatments, showing formation of large Pt . Crystallites.

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of 10 to 30 nm discrete alumina particles which agglomerate upon wetting. The various micrographs show platinum particle sizes after impregnation, after reduction in hydrogen in mild conditions (250°C), and after 16 hr treatment in an 0_2 atmosphere with temperatures ranging from 500 to 700°C.

It is evident that the platinum continues to grow in size on the Alon support even when it reaches 30 nm particle sizes, exceeding the average size of the particles in the support. The occurrence of this growth by crystallite transport over long distances of the irregular support, by diffusion from one Alon particle to the next, seems highly unlikely. A transfer of atomic or molecular metal species, either across the bridge where the support particles contact or through desorption, vapor phase transport, and readsorption, seems a far more plausible hypothesis for the Formation of these large particles.

4. Despite the frequent citing of crystallite migration as a sintering mechanism (e.g. 5.1, 5.2, 5.10, 5.11 5.12), there is little concrete evidence of motion over the appreciable distances needed to account for sintering of supported metal catalysts. For example, in *in situ* film growth studies by transmission electron ficroscopy Pashley *et al.* 5.13 observed small reorientations, such as rotations of the order of 1°, when particles merged. However, they did not report significant crystallite motion. Phillips *et al.* 5.12 cite crystallite diffusion in interpreting their electron micrographs. However, the authors were not able to describe their observations by their proposed model. In particular, Figures 1 and 2 of reference 5.12 are odd in that larger particles (~100 nm)
appear to move more rapidly than smaller particles ($\infty 5$ nm).

Wynniatt and Gjøstein^{5.14} observed minimal migration, less than 10 nm in 16 hours at 1000°C in a hydrogen atmosphere, for Pt particles of an average size of 7.5 nm on an AI_2O_3 substrate. They conclude that particle migration is too slow a process for particle sizes greater than 5 nm to account for observed sintering rates. Gous' review^{5.10} makes it evident that most evidence of substantial crystallite motion is inferential.

It should be noted that the conditions in which crystallite diffusion is cited as a mechanism in film growth are drastically different from those in typical supported metal catalyst sintering studies: first, temperatures are generally 200 to 500°C lowe. In film growth studies; second, coverage of the support is typically about 0.1 to 0.5 in film growth studies, while typical catalysts nave about 10^{-3} times that coverage of support by metal; and finally. typical film growth supports are smooth, often cleaved planes, while typical catalyst supports are polycrystalline and highly irregular. Thus even if definitive evidence of crystallite diffusion at film growth conditions is obtained, the translation of this mechanism to subported metal catalyst sintering would not necessarily follow.

In Tight of these difficulties in interpreting experimental evidence of sintering via the model of diffusing crystallites, we have looked to alternate mechanisms to account for the sintering phenomenon. In this chapter a model for the sintering of supported metal catalysts is developed based on individual metal atoms leaving the metal crystallites, migrating over the support and being captured by metal crystallites

upon collision. This approach is by no means devoid of conceptual difficulties, but it can_account for a variety of experimental observations of sintering.

5.2 Proposed Model

The mechanism of supported metal catalyst sintering in the proposed model is postulated to consist of three steps: one, individual metal atoms (or molecules such as PtO in an oxygen atmosphere) move from the metal crystallite to the surface of the support; two, the metal atoms migrate over the support surface; and three, the migrating atoms are either captured by collision with a metal crystallite or are immobilized by a drop in temperature or by encountering an energy well on the support surface. Each of these three steps will be discussed in detail.

5.2.1 Escape of Metal Atoms from Crystallites

Previous workers considering metal migration and particle agglomeration have generally discoulted the possibility of loss of individual atoms, due to the linge activation energy, E_a , that this deciwould require. The heat of dimension of platinum, for example, is 135 kcal/gmole^{5.15}. Arguing by extension from measurements on other metals, Geus^{5.10} estimated the dissociation of a pair of Pt atoms to be about 65 kcal/gmole. Thus most investigators of nucleation effects (e.g. ref.^{5.10-5.13}, 5.16-5.19) have concluded that for metals such as Bi, Au, Ag, Pt, etc. the dissociation of atoms from clusters of larger than a few atoms is neglectably small. Similarly, the forces between particle and support are presumed to be van der Waals type only, so that heat of adsorption, H_a, of single metal atoms is taken to be less than 20 kcal/gmole.

Supported metal catalyst sintering is generally observed at temperatures of 800 - 1000°K, which is higher than typical nucleation study temperatures of 500 - 600°K. At these temperatures, an activation energy of less than 60 to 75 kcal/gmole is necessary if a process is to occur with a significant rate. Thus the large sublimation energy and presumed low interaction of metal with support appear on first glance to rule out escape of metal atoms to the support surface from metal crystallites as a plausible mechanism.

However, Geus' extensive review 5.10 of data observed in metal film formation studies indicates that several factors may substantially increase the interaction between metal and support surface. In advancing this model we suggest that localized metal-support interactions are sufficiently large to reduce the net energy difference between an atom in the crystallite and an atom on the surface to the required value less than 75 kcal/gmole.

The presence of oxygen has been reported by many authors to significantly increase the interaction of metal crystals and oxide supports^{5.10}. Formation of a metal oxide layer at the support surface is believed to result in a strong chemical interaction, thought to involve metal incorporation in the support structure. For nickel, iron, chromium and titanium ageing in air at room temperature has been observed to substantially increase the adherence of metal particles on films. This type of strong interaction between metal and support, not necessarily reversible by reduction, could readily enhance removal of a metal atom from the particle to the surface. (In certain atmospheres the escaping metal species would presumably be molecular rather

than atomic.)

A second factor explored by $Geus^{5.10}$ which increases localized metal-support interaction is the presence of defects in the support structure. The well known decorating effect, where crystals are seen to nucleate in the region around defects, is attributed to the favorable energetics of adatom adsorption in these regions. Thus again metal escape from particle to support surface can be enhanced in localized regions by the presence of defects.

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Finally, $\text{Geus}^{5.10}$ reviews evidence that impurities on the support surface increase the interaction of metal and support. The presence of cracking products of hydrocarbons or of carbon has been found to increase the localized adatom population. The role of such impurities in serving as a bridge for the "spillover" of adsorbed gases from metal to surface has been discussed^{5.20}. By increasing metal-support interaction, such impurities may also help to bridge the spillover of metal atoms to the support.

These arguments for loss of metal atoms to the support surface are admittedly tenuous. However, they point to an interaction far more complex than the van der Waals forces cited in ruling out the possibility of particle dissociation. Because of the large heat of condensation of metals, and their ready mobility on support surfaces, no direct measurement of the heat of adsorption of Pt on typical supports is possible^{5.10}. Illips et al. ^{5.12} found that an H_a of 35 kcal/gmole for Au on SiO was required to explain their results in terms of their particle migration model. Tabatadze et al. ^{5.21} measured the conductance of metal oxides during the adsorption of noble metals. They observed large changes in conductance as long as metal atoms were supplied to the oxide surface. Their results indicate a strong electronic interaction of single metal atoms with the oxide surfaces which disappears once these atoms agaregate to form metal crystallites. This indirect evidence supports conclusions from nucleation and film growth studies that forces other than van der Waals are involved in the adsorption of metal atoms on oxide surfaces.

The reduction in surface energy is the driving force for the transfer of metal from smaller to larger crystallites. We are interested in obtaining an approximate relationship for the rate of loss of metal atoms from a crystallite to the subport as a function of crystallite size. The Kelvin equation relates the spreading pressure, \$\$, to the crystallite radius, r, by

$$\phi = \phi_0 \exp(\beta/\gamma)$$

5.3

where the value of B depends on the Shape of the crystallite, the metalsupport contact angle, the metal-support and metal-vapor interfacial energies, the metal molar volume, and the temperature. For a specific support/metal system, at a constant temperature, the value of B should be relatively constant.

At equilibrium the rate at which a crystallite cabtures atoms is proportional to ro, and therefore the rate of atom loss, $\frac{dl_1}{dt}$, is also proportional to ro. The rate of loss is independent of whether ar not the system is in equilibrium if energy transfer is not controlling. Hence the ratio of the rates of loss for two crystallites of different size is given by

$$\left(\frac{dL_1}{dt}/\frac{dL_2}{dt}\right) = \frac{r_1\phi_1}{r_2\phi_2} = \frac{r_1}{r_2} \exp\left[8\left(\frac{1}{r_1}-\frac{1}{r_2}\right)\right] = 5.4$$

The Pt/Al_2O_3 contact angle is $.90^\circ$, and therefore the crystallites were taken to be hemispheres. For this case the value of B is defined by

$$\beta = \frac{(\gamma_1 + 2\gamma_2)v}{RT}$$
5.5

where y_1 and y_2 are the Pt - Al₂O₃ and Pt - atmosphere interfacial energies, V is the molar volume of Pt, and T was taken as 1000°K. In order to assess the range in reported values of y(e.g. 5.22, 5.23) and . the variation of Pt species (such as metallic Pt, PtO, etc.), three values of β , ranging from 4 to 6 nm, were employed in calculating relative rates of loss by equation 5.4. In Table 5.1 the rates of atom loss relative to a crystallite with a diameter of 25 nm are reported for the three cases over the size range 2 to 50 nm.

From the values reported in Table 5.1, it is evident that in the size range of interest in sintering studies the rate of loss of atoms from a crystallite may reasonably be approximated as independent of the crystallite size. The variation in loss is less than 30% up to 30 nm, except for very small particles where equation 5.3 predicts questionably high values of ϕ/ϕ_0 . Accordingly, we have modeled the rate of loss of atoms from the ith particle as

$$\frac{dL}{dt} = A e^{-E_{d}/RT}$$

where $\frac{dL_i}{dt}$ is the rate of transfer of atoms to the surface, A is an arbitrary constant, and E_a is the activation energy required to move from a particle to the surface. For particle sizes above 30 nm the surface energy and size variance of the rate of loss do not compensate

5.6

Table 5.1: Relative Rate of Loss of Atoms per Crystallite as a Function of Crystallite Size as Estimated by the Kelvin Equation (Normalized to 25 nm

Diameter Crystallites)

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Crystallite diameter nm	Case I () = 4.0 nm	Case II ß = 5.0 nm	Case III ß = 6.0 nm
2	3.17	7.96	19.97
4	0.86	1.31	1.99
6	0.66	0.85	1.10
8	0.63	0.75	0.89
10	0.65	0.73	0.82
12	0.68	0.74	0.81
15 .	0.74	0.78	0.83
18	0.82	0.84	0.87
20	0.87	0.88	0.90
25	1.00	1.00	1.00
30	1.14	1.12	1.11
40	1.42	1.38	1.34
50	1.70	1.64 •	1.57
100	3.15	2.96 .	2.79
/			· _

for each other, and the assumption of a constant loss rate, equation 5.6, becomes increasingly subject to error.

5.2.2 Migration of Metal Atoms Over Support Surfaces

The extensive work on nucleation and film growth leaves little doubt about the mobility of atoms on supports, even at temperatures much lower than those encountered during supported metal catalyst sintering 5.10-5.13, 5.16-5.19. The atoms may be considered as a twodimensional gas, in which case their speed is given by

$$v = \left(\frac{\pi k_B T}{2\pi}\right)^{\frac{1}{2}}$$
 5.7

or the motion may be described by the jumping From one surface site to net which is described by

$$v = a v \exp(-E_s/RT)$$
 5.8

It should be noted that E_s , the activation energy for surface diffusion, is not necessarily equal to E_a , but is generally considerably smaller than E_a .

In the present of it is immaterial whether equation 5.7 or 5.8 is used, but equation distrates that a drop in temperature could readily immobilize and ones if certain sites on the support surface have large values of the presume a surface valocity rapid enough, at sintering temp is the support surof free surface metal atoms on such of sub-tree.

5.2.3 Capture of Atoms by Veta: C., Sallies,

Upon collision of a migrating a set with metal crystallite, the atoms may become incorporated in the crystallite. The mate at which a crystallite gains metal atoms by this process depends on the concentration of metal atoms on the support surface, the velocity of these atoms, and the effective diameter of the crystallite, D_i , *i.e.*,

$$\frac{dG_{i}}{dt} = \alpha \frac{v}{N_{t}} \frac{F_{s}}{S_{o}} D_{i}$$
5.9

where " = sticking probability of an atom colliding with a crystal-

lite (assumed to be independent of D)

 $N_{+} = total metal atoms$

 $S_0 =$ support area per metal atom

 F_s = number of atoms migrating on support with an area $N_t S_o$ (*i.e.* $\frac{F_s}{N_t S_o}$ is the concentration of atoms on support surface) Once the atom has become attached to the periphery of the particle it can migrate over the metal surface and become part of the crystal. The surface self diffusion rates of metals are generally quite large, the activation energy for Pt only being 26 kcal/gmole^{5.24}, and the resulting crystals should be three-dimensional rather than metal islands on the support.

5.2.4 A Summary of the Model

The net rate of change of the number of atoms in the ith particle is given by

$$\frac{dN_{i}}{dt} = \frac{dG_{i}}{dt} - \frac{dL_{i}}{dt}$$
5.10

which, according to equation 5.3 and 5.6 may be written as

$$\frac{dN_{i}}{dt} = \alpha v \frac{F_{s}}{N_{t}S_{o}} D_{i} - A e^{-E_{a}/RT}$$
5.11

The rate of change of the number of migrating surface atoms is given by the material balance

$$\frac{dF}{dt} = \sum_{i=1}^{M} \left(\frac{dL_i}{dt} - \frac{dG_i}{dt} \right) = -MAe^{-E_g/RT} + \alpha v \frac{F_s}{N_t S_o} \sum_{i=1}^{M} D_i = 5.12$$

where M is the number of metal crystallites on the support of area $N_{\pm}S_{0}$.

This model of supported metal catalyst sintering may be likened to an evaporation condensation process among multisized droplets. In that case the variation in vapor pressure among drops of different size leads to a transfer of liquid from smaller to large: droplets. By our sintering model a single particle on a support surface would establish an equilibrium concentration of metal atoms migrating along the surface. With several particles of different size, however, the surface concentration is not equilibrated with each particle, and thus growth or decay in individual particle sizes occurs. Because smaller crystallites would equilibrate with higher concentrations of migrating surface atoms than larger crystallites, the larger crystallites grow at the expense of the smaller.

5.3 Method of Solution

Equations 5.11 and 5.12 were written in the following finitedifference form for solution

$$\Delta N_{i} \approx \left(\alpha v \frac{\overline{F}}{N_{t} S_{o}} D_{i} - A e^{-E_{a}/RT} \right) \Delta t \qquad 5.13$$

$$\Delta F_{s} = - \sum_{i=1}^{M} \Delta N_{i} \qquad 5.14$$

where \overline{F}_{S} = arithmetic average of F_{S} during time increment Δt and v is given by equation 5.7. In the above formulation the constant α includes

the reduction in the velocity of migrating atoms, v, due to an activation energy for surface diffusion. The support surface area per metal atom, S_0 , is related to the metal loading of the catalyst. For example, $S_0 \approx 10 \text{ nm}^2/\text{atom}$ corresponds to a catalyst containing $\sim 1 \text{ wt} \approx 2 \text{ Pt}$ on a support having an area of $300 \text{ m}^2/\text{g}$.

Two methods of solution for equations 5.13 and 5.14 were employed, depending on the value of $\frac{\alpha}{S_o}$ (v being given by equation 5.7 and F_s/N_t , the fraction of atoms migrating on the support, being determined in part by the value of $\alpha v/S_o$).

5.3.1 Solution for $\alpha/s_0 > 10^{12} \text{ m}^{-2}$

For values of $\alpha/S_0 \ge 10^{12} \text{ m}^{-2}$ it was found that F_s is always small, $f_s = 0$, and the approximation $\sum_{i=1}^{M} \Delta N_i \approx 0$ could be made. This condition corresponds to rapid rates of surface atom migration and capture compared to the rate at which atoms leave crystallites. For this situation, the rate at which a crystallite captures atoms, assuming all crystallites are equally accessible to migrating atoms, is proportional to diameter, D_i . Equations 5.13 and 5.14, for this case, reduce to

$$AN_{i} = (- - - 1) A e^{-E_{a}/RT}$$

$$\Delta N_{i} = (- - - 1) A e^{-E_{a}/RT}$$

$$\Delta t = 5.15$$

$$5.15$$

The values of N_i , for all particles, as a function of time can readily be computed by a single-step method (Euler's formula) once a relationship between D_i and N_i is chosen. In this work we described the crystallites as fee cubes and used the equations of van Hardeveld and Hartog^{5.25} to determine the total number of atoms, N_i , and the number of atoms, $N_{s,i}$, at the surface of each crystallite, *i.e.*

5.16

$$N_1 = 0.5 + 0.5 [1 + 2q]^3$$

and

$$N_{s,i} = 12q^2 + 2$$
 5.17

for $q \geq 1$

where

 $q = \frac{(l_{i} - a_{i})}{\sqrt{2} a_{i}}$ (*i.e.* the number of atoms along the edge of cube minus one)

 ℓ_i = the length of the edge of cube

 $a_0 = \text{atomic diameter (0.277 nm for Pt)}$

The effective diameter, D_i, of a crystallite for the capture of migrating atoms was taken as

 $D_{i} = 4(l_{i} + a_{0})$

By starting with a given particle size distribution, $i.e. l_i$ for all particles at time = 0, equation 5.15 was used to obtain N_i 's as a function of time. The corresponding dispersion, D, was calculated by

 $D = \frac{\underbrace{\substack{i=1 \\ j=1 \\ M}}^{M} N_{j}}_{i-1} N_{j}$

5.19

5.18

Any crystallites for which N_i became <14 (*i.e.* q < 1) during a time interval were considered to have disappeared, and the remaining atoms in these crystallites were distributed among the remaining crystallites in the following time increment in proportion to their size.

5.3.2 Solution for $a/S_0 < 10^{12} m^{-2}$

Equations 5.13 and 5.14 were solved by a trial and error for these cases. The number of atoms, F_s , was taken as zero at time equal to zero. In order to avoid oscillatory values of F_s in the initial time increments a heating period was incorporated in the calculations. The temperature was increased from 300 K to the desired final temperature over a period of one hour in 0.01 hour increments. The trial and error procedure used to calculate N_i and F_s at time t' = t + Δt was as follows:

1. The value of F_s at t' was guessed (*i.e.* ΔF_s for Δt). 2. N_i was computed for each particle using equation 5.13

$$(\overline{F}_{s} = F_{s}|_{t} + \frac{\Delta F_{s}}{2}).$$

3. ΔF_s was computed using equation 5.14.

4. If ΔF_s from step 1 differed from ΔF_s by step 3 by less than 0.1 atoms and if $F_s|_t + \Delta F_s$ was ≥ 0 calculation proceeded to the next time increment; if the difference was > 0.1 steps 1 to 3 were repeated, with the new ΔF for step 1 being

estimated using the magnitude of the difference as a guide. If only a negative value of F_s (physically impossible) at t' could satisfy the above conditions, the size of Δt was decreased to one-tenth of its value and steps 1 to 4 were repeated. If any N_i became < 14, these N_i were added to F_s in the next time increment.

The dispersion for this case was calculated in a manner analogous to equation 5.19, except that F_s was included in the number of metal surface atoms, *i.e.*

5.20

A value of 0.1 hour for Δt was used for the computations. Calculations using a Δt of 0.017 hour yielded essentially the same results. The complete programs used for each case are attached in Appendix C.

 $D = \frac{F_{s} + \sum_{i=1}^{m} N_{s,i}}{\sum_{i=1}^{M} N_{i}}$

5.4 Description of Cases Investigated

5.4.1 Crystallite Size Distributions

The initial (zero-time) particle size distributions (PSD) were generated in the following manner:

- 1. the maximum and minimum size of particles in the distribution was chosen $(\ell_{max} \text{ and } \ell_{min})$,
- 2. the size range was split into K equal size increments of size Δl ,
- the number of particles, P_k, was specified for each size increment,
- 4. the size of each particle was calculated by

$$\lambda_{i} = \ell_{min} + (k-1)\Delta \ell + (\frac{j-1}{P_{k}})\Delta \ell$$
 5.21

for k = 1 to K + 1 and j = 1 to P_k for each k. Where $i = j + \sum_{m=0}^{k-1} P_m$ with $(P_0 = 0)$, $P_{K+1} = 1$ was included to obtain a particle of size k_{max} . This procedure results in $\sum_{k=1}^{K+1} P_k$ particles all having a different size.

Two general types, I and II, of PSD were generated by this procedure. The Type I distribution is approximately Gaussian in nature, and Type II is skewed with a large fraction of small particles. The values

of min, max, k and P_k used to generate the specific PSD are given in Table 5.2.

The PSD listed in Table 5.2 contain 121 particles, except 8 and 9 which contain 122. In these two distributions one large particle (10 nm) was added to PSD 1 and 7. To check the sensitivity of the calculations to the number of particles present, calculations were carried out with PSD for which the values of P_k for k = 1 to K were dou! (P_{K+1} was kept equal to one). These calculations showed that as long as the remaining number of particles was ≥ 7 the dispersion as a function of time remained unaltered.

5.4.2 Parameter Values

The values of parameters used in solving equations 5.13 and 5.14 (or equation 5.15) for the specific cases reported in the next section are listed in Table 5.3. Although many other combinations of parameters were examined the cases listed illustrate the general behaviour of the model. It should be mentioned that, although all the cases listed used a value of $A = 8 \times 10^{13} \text{ sec}^{-1}$, the results are readily extended to lower values of A (significantly higher values of A are chysically unlikely). Thus, for a specific PSD and fixed values of T and α/S_0 , the dispersion as a function of time is determined by the value of A exp(-E_a/RT), i.e., $A = 1.65 \times 10^7 \text{ sec}^{-1}$ and $E_a/RT = 15.0$ is equivalent to $A = 8 \times 10^{13}$

5.5 <u>Results</u> and Discussion

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5.5.1 Effect of Initial PSD on Rate of Sintering

From equation 5.11 it is evident that for a certain value of D_i , and the corresponding crystallite size, 2_i , according to equation 5.18,

Distribution Number	² min (nm)	دmax (nıır)	Туре	Dispersion
1	2.70	3.30	I	0.35
2	2.10	3.90	I	0.34
3	1.20	4.80	I	0.31
4	2.950	3.065	I	0.35
5	2.00	12.50	I	0.13
6	5.00	15.50	I	0.10
7 .	1.50	4.29	II	0.35
8*	1.50	4.29	II	0.28
9*	2.70	3.30	I	0.29

Table 5.2: Data for Generation of Particle Size Distributions

Type I

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k for k = 1 to 10 10 for k = 11 and 12 P_k = 22 - k for K = 13-21 1 for k = 22

Туре 11

k 1 2 3 4 5 6 7 8 9 10 P_k 20 20 18 16 14 12 10 7 3 1

* one 10 nm particle was added to these distributions to simulate a bimodal distribution.

Table 5.3: Parameter Values Used in Study	Table	5.3:	Parameter	Values	Used	in	Study	
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Case Number	E _a /RT	T(K)	$\frac{\alpha}{S_0}$ (m ⁻²)
,			. 12
1	35.0	1000	1×10^{12}
2	35.0	1000	2×10^7
3	33.3	900	1×10^{12}
4	33.0	900	2×10^{7}
5	31.6	950	1×10^{12}
· 6	31.6	950	2 x 10 ⁹
7	31.6	950	2×10^{7}
8	31.6	950	4×10^{6}
9	31.6	950	2 x 10 ⁶
10	30.0	1000	1 x 10 ¹²
11	30.0	1000	2×10^7
12	27.5	1000	1 x 10 ⁷²

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 $A = 10^{13} \text{ sec}^{-1}$ for all cases.

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the net rate of growth is equal to zero. Particles with sizes larger than this 'critical' size will grow while smaller particles will shrink. This critical size is given by

$$\ell_{CRIT} = \frac{A N_t S_o}{4\alpha v F_s} e^{-E_a/RT} - a_o \qquad 5.22$$

J

or according to equation 5.15

$${}^{k}_{CRIT} = \frac{\frac{\sum_{i=1}^{D} D_{i}}{\sum_{i=1}^{M} - a_{o}} = \frac{\sum_{i=1}^{M} \ell_{i}}{M}$$
 5.23

As sintering progresses the number of particles, M, decreases and the number of free surface atoms decreases as well. This results in an increase in *e*_{CRIT} as sintering progresses, and crystallites which initially grow in size will eventually decay.

Figure 5.2 and 5.3 shows the particle size history for individual particles of the same initial size for PSD 1 and 9, Case 3. PSD 1 and 9 are the same except one 10 nm particle has been added to PSD 1 to get PSD 9. The effect of the added 10 nm particle is evident when comparing Figures 5.2 and 5.3. This large particle captures many of the migrating atoms, resulting in an increased rate of sintering.

The effect of large particles or broad initial PSD on the rate of sintering is further illustrated in Figure 5.4 where the normalized dispersion D/D_0 is shown as a function of time for various initial PSD. The addition of one large particle (10 nm) to PSD 7 (i.e. to get PSD 8) significantly increases the rate of sintering. PSD 1 to 4 are all Type I distributions with a mean crystallice size of 3.0 nm, but the size range ($\ell_{max} - \ell_{min}$) varying from 0.105 nm (PSD 4) to 3.50 nm (PSD 3).







TIME (hours)

Figure 5.3: Particle size history for selected individual particles, PSD9, Case 3. (Number is i, the particle number is the PSD.)

The rate of sintering, as shown in Figure 5.4, is larger for the broader distribution in an initial period of about 10 hours. The reduced rate of sintering for narrow distributions occurs because in the initial stages of sintering these distributions are becoming gradually broader without the disappearance of particles. The narrow size range does not generate a substantial driving force for transfer of metal from the smaller to the larger particles,

Figures 5.5 and 5.6 show the cumulative distribution function F as a function of time for Type I and II initial distributions. This illustrates the spreading of the distributions with time. In each case, as sintering progresses the model predicts the presences of particles smaller than the initial minimum size. During early stages of sintering the broadening, and specifically the decay of a substantial fraction of the particles to smaller sizes, is evident. This prediction of the presence of small particles at all stages of sintering is one significant difference between the proposed interparticle transport model and the crystallite diffusion model^{5.1; 5.2}. The latter model does not allow for formation of particles smaller than the initial minimum size, nor does it predict the type of induction period for narrow PSD shown in Figure 5.4. Further, it predicts that sintering would occur with a unisized distribution (practically unattainable), whereas the model developed here predicts that such a distribution would not sinter because of an absence of a driving force for interparticle transport. Some experimental evidence exists that even in heavily sintered samples small crystallites are still present 5.26. Further experiments that follow the nature of the particle-size-distribution broadening would be useful in elucidating the mechanism of sintering.



Figure 5.4: Effect of initial particle size distribution on the rate of sintering (Case 5, the numbers identify the PSD).









5.5.2 The Effect of Surface Mobility and Metal Loading on Sintering Behavior

If no initial free surface atoms are postulated, then during the early stages of sintering the value of F_s grows as particles lose atoms to the surface and relatively few are captured by crystallites. The parameter α/S_0 determines the extent to which surface atoms are captured, and hence the size of F_s . The increases in dispersion (defined by equation 5.20) shown in Figure 5.7 are due to this increase in F_s .

For large values of α/S_0 , corresponding to high metal loadings (*i.e.* small values of S_0) or high velocity of migration, the number of migrating surface species approaches zero and the rate of sintering is controlled by the loss from particles only. In these cases, the rise in dispersion due to the surface species is negligible, and this model does not predict any redispersion of metal.

Lower values of α/S_0 , however, can account for significant redispersion. The lower values could arise from lower metal loadings or a reduced velocity of migrating species (larger E_S). In such cases a significant amount of free surface atoms builds up, leading to higher dispersions and reduced overall rates of sintering. Redispersion could occur through cooling of a sintering sample during this period of growth in number of free surface atoms. If the migrating atoms were frozen as single atoms or small particles during cooling, the increased dispersion would remain. Experimental evidence^{5.27} indicates that rapid cooling does give higher dispersions after sintering treatments than slow cooling. Figure 5.7 shows that for α/S_0 values less than about 2 x 10⁶ m⁻² the capacity of the surface to accomodate free surface atoms exceeds the number of atoms present, and thus all the particles disappear.



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Figure 5.7: Effect of surface mobility of metal atoms on support, and metal loading, on sintering behavior. (PSD1, Cases 5 to 9.)

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The predictions of the model as α/S_0 becomes smaller may become unreliable because the assumption that the surface concentration of migrating atoms is position independent is no longer valid at low migration rates. The extent of the error introduced by this approximation has not been investigated.

5.5.3 The Effect of E_a/RT on Sintering Behavior

Figure 5.8 shows, for the case where α/S_0 is large and the free surface atoms are rapidly captured, the influence on sintering behavior of variations in E_a/RT . The plots correspond to either a change in temperature or in the activation energy. For instance, if E_a is taken as 60 kcal/gmole, the curves in Figure 5.8 would represent temperatures ranging from about 850°K ($E_a/RT = 35$) to 1100°K ($E_a/RT = 27.5$). If the temperature is postulated as 1000°K, the curves represent activation energies from 55 to 70 kcal/gmole.

As would be expected for such large activation energies, the sintering behavior is extremely sensitive to variations in temperature. Sintering is extremely slow for $E_a/RT = 35.0$, with less than 10% loss of dispersion in 40 hours, while for $E_a/RT = 27.5$, sintering is extremely rapid, resulting in the transfer of all the metal in the initial 121 particles to a single large particle within 5 hours.

Similar trends are evident, as shown in Figure 5.9, when values of α/S_0 are lower and a significant surface concentration of migrating atoms is present. In this case, however, a lower value of E_a/RT , corresponding to higher temperatures or a lower activation energy, affects not only the rate of sintering but also the extent of initial redispection. While lower values of E_a/RT lead eventually to more rapid sintering, the



Figure 5.8: Effect of E_a/RT on sintering behavior, PSD1, high value of \ll/S_0 , Cases 1, 3, 5, 10 and 12.



Figure 5.9: Effect of E_a/RT on sintering behavior, PSD1, low value of $</\overline{S}_0$, Cases 2, 4, 7 and 11.

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initial rise means that for a period of time the dispersion is lower for higher E_a/RT values.

Changes in atmosphere would affect the activation energy for metal movement from the particle to the support surface. The actual species leaving would probably differ in oxidizing and reducing atmospheres. In the former case a metal oxide molecule would probably be the migrating species. Oxidizing atmospheres are found to enhance interactions between metal and support^{5,10}, and thus could lower the activation energy of escape. The pre-exponential factor, A, could also be affected. It is possible that in the case of oxidizing atmospheres the rate determining step in the sintering process is the formation of the surface ox: species since oxygen adsorption on metals such as Pt is an activated process. Different atmospheres could also affect the velocity of migration along the surface, *i.e.* changes in the activation energy for surface migration, a factor included in the parameter a discussed above.

5.5.4 The Order of Sintering

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Several authors $(e.g.^{5.3}, 5.9)$ have fit experimental data of sintering to a power-law model of the form

 $\frac{dD}{dt} = - KD^n$

5.24

where D is the dispersion and K and n are constants. Values of n from 2 to 13 have been reported.

The rate of change of dispersion has been fitted to equation 5.24 . to determine the range of n values predicted by the proposed model. One immediate difficulty with the power-law model is its inability to account for an initial period of redispersion as the surface atoms concentration builds up. Accordingly, we have fitted only those data for the regions where the dispersion is decreasing. If equation 5.24 is valid, then a plot of $(D_0/D)^{n-1}$ versus time will result in a straight line with an intercept of unity ($n \neq 1$). Figure 5.10 shows these plots for PSD 5 and 6. These distributions are identical, except the mean size is displaced by 3.0 nm. The behavior of PSD 5 (mean size = 7.25 nm) is well fitted by an n value of 6, while PSD 6 (mean size = 10.25 nm) is well fitted by an n value of 3. This tendency of decreasing values of n with increasing mean particle sizes for symmetrical PSD was observed for several other PSD.

Figure 5.11 illustrates the effect of PSD width on the value of n for PSD with the same mean size (PSD 1 and 2). Neither case is well described by the power-law model, but the narrow PSD 1 shows very low values of n (< 2) at times < 15 hours. This is the result of the 'induction' period for the sintering behavior of narrow PSD that has been discussed previously. As PSD 1 broadens with time (see Figure 5.5) the value of n increases and at even larger values of time n decreases. A similar, but less marked, trend is observed with PSD 2. Not only does n vary with the extent of sintering, but the value of n also depends on the width of the initially symmetric PSD.

The inability of the power-law model to describe the progress of sintering over wide variations of D/D_0 is well recognized^{5.3, 5.4}. It is evident that the sintering process should depend on the nature of the PSD and not only on the dispersion. The proposed model predicts the nature of the dependence of n on PSD, but experimental determinations of n as a function of PSD are not presently available.



Figure 5.10: Effect of mean crystallite size on power-law order (Case 5; ● PSD5, ⊅ PSD6).

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Figure 5.11: Effect of particle size distribution width on powerlaw order (Case 3; ● PSD1, o PSD2).

5.5.5 Application of the Model to an Experimental Case

To test the ability of the proposed model to describe some experimetal sintering data we chose the data of Wynblatt and Gjostein^{5.3}. An approximate initial PSD was obtained by measuring the size of approximately 200 particles in Figure 1a of reference 5.3. The distribution was adjusted to yield an average initial dispersion of 6.7% corresponding to the average particle radius of \sim 8.0 nm reported by Wynblatt and Gjostein. The resulting bimodal PSD is shown in Figure 5.12 (line at t = 0)

This result agrees well with the experimentally observed n value of $13^{5.3}$. The initially broad and bimodal PSD is responsible for high value of n at low time. As sintering progresses the bimodal nature of the distribution becomes less pronounced and at t \approx 20 hours the PSD is approximately symmetric. As the PSD becomes more symmetric the value of n starts to decrease rapidly and changes from 13 to 4. To further test the validity of the model initial PSD and PSD as sintering progresses should be determined experimentally and compared to predictions. 125

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Predicted dispersion as a function of time for the initial distribution of Wynblatt and Gjostein^{5.3}


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5.5.6 The Possibility of Multiple Mechanisms

Sintering behavior occurs over at least two orders of magnitude of particle size, from 0.5 to 50 nm, and over a range of atmospheres, temperatures, supports, and metal loadings. While this work has postulated interparticle atomic or molecular migration as an alternate to particle motion, the two mechanisms are not exclusive. Wynblatt and Gjostein^{5.14} suggest particle migration may be prevalent for particles under 5 nm, but that some form of interparticle transport must take place at larger sizes to account for observed rates of sintering.

5.6 Conclusions

A model for the sintering of supported metal catalysts, based on the dissociation of individual atoms from the metal crystallites, has been developed. The model postulates that large interactions between the support and metal atoms may potentiate the escape of metal to the surface. Evidence of high interaction between metal and a support is found for oxygen atmospheres (on oxide supports), in defect regions, and in the presence of contaminants. The model has been applied to a variety of simulated catalysts. The model can account for a wide variety of experimental observations of sintering. These include a strong influence of atmosphere, a high apparent activation energy, possible redispersion, and a variation in order from < 2 to \geq 13 when sintering behavior is fitted to a simple power-law equation.

The model predicts that the rate of sintering increases with increasing width of the initial PSD. Metal is transported from small to large particles, and transport is more rapid as the difference in size of particles is increased. This model predicts the presence of particles

smaller than the initial minimum particle size as sintering progresses.

During the early stages of sintering the model predicts a buildup of a concentration of migrating metal atoms or molecules. If this buildup is substantial then redispersion of the catalyst will be significant. Factors affecting the extent of surface species buildup are migration velocity, collision accommodation, activation energy for escape from the particle to the support surface, temperature and metal loading. The first three factors would be influenced by the atmosphere in which sintering occurs.

This model predicts a considerable variation in order of fit to à power-law equation of sintering. Variation of the order is predicted within a given sintering experiment. Distribution width and initial particle size are especially critical parameters with respect to the power-law order.

A variety of improvements to the presented model can be made, such as: 1. using the Kelvin equation to calculate the rate of loss of atoms from larger crystallites rather than approximating this by a constant rate of loss per crystallite, 2. including variations in the concentration of migrating atoms with position, and 3. modelling the support surface as inhomogeneous. It is felt that at the present time improvements in the model along the above lines are unwarranted because insufficient information is available to a prior: estimate the extra parameters in these extended models. These new parameters, such as interfacial tensions, energy distribution of inhomogeneous is more than face, etc., in the refined models would be determined by filling them to experimental data and any improvement in the fit over the simple model would be due to an increase in the number of adjustable parameters. One 130

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of the strengths of the presented model is that it contains only three parameters, *i.e.* α , A, and E_a, (in the case of equation 5.15 A and E_a are the only adjustable parameters) and it is easier to assign physical significance to these parameters and hence use their magnitude in designing catalysts with greater stability.

5.7 Nomenclature

- A' and B constants in equation 2.
 - A pre-exponential factor in rate of loss equation (equation 3).
 - a_0 diameter of metal atom (0.277 nm for Pt).
 - D dispersion.

 D_{i} - effective collision size of crystallite i.

- $\cdot E_a$ activation energy for atom moving from crystallite to support.
 - E_s activation energy for metal atom migration over support surface.
 - F cumulative distribution function.
- F_s number of metal atoms migrating on support surface.
 - \overline{F}_{s} average number of free surface atoms in time increment.
 - ${\rm G}^{}_i$ atoms gained by crystallite i.
 - ${\rm H}_{\rm a}$ heat of adsorption of single metal atoms on support.
 - K rate constant in equation 1.
 - k number of size increments used when generating intial PSD.
 - k_R Boltzmann's constant.
 - L_i atoms lost by crystallite i.
 - κ_{i} particle size, the length of the cube edge.

CRIT - the particle size above which increase and below which decrease in size occurs (function of time). min - the size of the smallest particle in the initial PSD.

- M total number of crystallites (function of time).
- m mass of metal atom.
- N_i number of atoms in crystallite.
- N_t total number of metal atoms in all crystallites.
- ${\rm N}_{\rm S,i}$ number of surface metal atoms in crystallite it.
 - n order in the power-law model (equation 1).
 - P_k number particles assigned to size increments when generating initial PSD.
 - q the number of metal atom along cube edge minus one.
 - r particle radius (equation 2).
 - S metal surface area (equation 1).
 - S_0 support surface area per metal atom (N_tS_0 = total support surface area).
 - T temperature.
 - t time.
 - V molar volume of metal.
 - v velocity of atoms on support surface.
 - a sticking probability of atoms colliding with crystallite.
 - β defined by equation 3.
 - v frequency factor (10^{13} sec⁻¹).
 - 4 spreading pressure of metal on support surface.

5.8 References

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CHAPTER 6 EXPERIMENTAL STUDIES OF SINTERING OF / SUPPORTED PLATINUM CATALYSTS

6.1 Survey.

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As discussed in Chapter 5, supported metal catalysts show changes in metal surface area during use or treatment at high temperatures, a process known as sintering. Sintering is of concern industrially because a loss of catalytic activity generally occurs, associated with an increase in the particle size of the metal. A number of patents i have been issued for processes designed to redisperse sintered catalysts (v.g. references^{6.1}, ^{6.2}).

A number of investigators have studied catalyst systems under treatment conditions leading to sintering. In this chapter their work is briefly reviewed, and a series of experiments performed in this laboratory are described. From this data the sintering process appears to be extremely complex, and further experiments will be necessary to establish the critical parameters in this process.

Many authors 6.3 - 6.10, in trying to describe the kinetics of sintering, have fitted data to the simple power law equation of the form

$$\frac{ds}{dt} \approx Ks^n$$
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where s is the metal surface, n is generally reported as an integer, and an activation energy is lumped in the constant K. Both Herrmann $\frac{6}{2t} al.^{6.3}$ and Matt and Moscou^{6.4} report n values of 2 for Pt/Al₂0₃

catalysts in oxidizing environments. The latter authors examined intersemples under the electron microscope before and after sintering. Initially the Pt in their catalysts consisted of particles in the 1 nm range. After treatment in air, they reported the presence of 1 to 50 nm particles. Somorjai^{6.5} studied supported Pt catalysts sintered in an oxidizing atmosphere by small angle x-ray scattering. Based on data obtained by this technique, he proposed a modified growth law similar to that advanced by Wynblatt and Gjostein^{6.6}, of the same form as Equation 6.1 but with a K value which decreases exponentially with increasing average particle radius (see Equation 5.2). Somorjai^{6.5} also estimated the activation energy to be 52 kcal for oxidizing atmospheres. However, small angle x-ray diffraction is not equally sensitive to all sizes of Pt particles.

Sintering kinetics change in a reducing environment, in general associated with a higher value of n. Somorjai^{6.5} found a lower activation energy for an H₂ atmosphere, about 20 kcal, but a much lower rate pre-exponential factor, and hence a lower overall rate of sintering as compared to an oxidizing atmosphere. Gruber^{6.7} and Hughes et al.^{6.8} found n values in the 6 to 8 range for sintering at 500°C in the H₂ atmospheres.

Finally, two special studies have been made of Pt/Al_2O_3 systems. Wynblatt and Gjostein^{6.6}, 6.9 used electron microscopy to study Pt particles and specially thin (10 nm) alumina substrates. They found the surface for a sample where Pt particles were larger than 5 nm, to terparticle transport for a mechanism in these and larger F particles. High values of n (28) are attributed to facet inhibited growth; 2., expansion of a Pt particle must await

nucleation of a ledge on a smooth face. Experimentally they observed n values from about 3 to 13 for sintering in air. Huang and $Li^{6.10}$ studied Ostwald ripening for large (>100 nm) Pt particles on $a - Al_2O_3$. Their n value of the order of 5 led them to conclude that surface diffusion is the mechanism of the growth process. Gregg and Howlett^{6.11} similarly concluded that surface diffusion was the agglomeration process for the sintering of metal spheres observed under the optical microscope. Both Wynblatt and Gjostein^{6.9} and Huang and $Li^{6.10}$ observed that the sintering rate increased with increases in oxygen concentration in an O_2/N_2 mixture.

As detailed in Chapter 5, two models have been developed to account for particle size changes occurring during sintering. Ruckenstein and Pulvermacher^{6.12}, 6.13 postulated crystallite migration and collision as a sintering mechanism and developed and solved a model for homogeneous and partially heterogeneous surfaces. This model can account for power law orders from 2 (rate controlled by fusion of particles) to about 8 (rate controlled by diffusion of particles). It also predicts that after a certain time a universal dimensionless curve will fit the particle size distribution of a catalyst. In Chapter 5 a model was developed which postulated the loss from particles of atomic or molecular species, followed by surface diffusion and recapture, as the sintering mechanism (see also references 6.14 and 6.15). This model can account for power law orders less than zero (i.e. redispersion) up to 13 and higher. The order is found to depend heavily on the nature of the particle size distribution. While under either model the decrease in surface energy provides the driving force for metal agglomeration, the difference between transport as a crystallite

or as an atomic species leads to significant differences in predicted growth kinetics. This point is discussed in further detail below in conjuction with some experimental results.

6.2 Results

Four series of experiments were conducted to explore the sintering behavior of supported Pt catalysts. In one series catalysts were sintered at varying temperatures for a fixed time period, and changes in the catalyst were monitored by electron microscopy and/or gas uptake. In a second series, the temperature was fixed and the time of sintering varied. In a third set of runs the sintering behavior of a catalyst batch into which a portion of pre-sintered catalyst containing large Pt particles was mixed was compared to that of an unmixed catalyst. Finally, electron micrographs were recorded of the same catalyst specimen area before and after reduction and sintering in an inert atmosphere.

6.2.1 Constant Time/Variable Temperature Treatments

Both commercial catalyst samples (Engelhard 0.5% Pt on $Al_{2}O_{3}$) and prepared specimens (4.76% Pt on Alon $Al_{2}O_{3}$) were heated 15 nours in oxygen at varying temperatures, and the dispersion or particle size were measured. In the case of the prepared catalyst, small samples (<0.5 g) were used, and the progress of sintering was checked by electron micrographs. For the commercial catalyst, batches of sufficient size (8 - 10 g) were used to allow gas uptakes to be measured after sintering.

Figure 6.1 shows the size range of particles observed in the micrographs of several specimen areas of the Alon supported catalyst. The



Figure 6.1: Effect of treatment temperature on metal particle size range, measured from electron micrographs. Batches of 4.76% Pt on Alon catalyst were heated 16 hr. in flowing oxygen at the indicated temperature.

lower limit of 1 nm shown in Figure 6.1 is considered the minimum size where metal size contrast can accurately be distinguished from the contrast inherent in the support (Chapter 3, reference 6,16). No change was observed in the metal particle sizes following 16 hour heating in oxygen at 500°C. Significant particle growth was observed for 16 hour treatment at 550°C and for all higher temperatures. For the specimens sintered by 600° and 700°C, particle growth was sufficiently great to significantly lower the number of particles in a given field of view, hence the size ranges shown in Figure 6.1 for these two cases, based on about five fields of view, must be considered approximate. (Micrographs from this ceries of experiments are shown in Figure 5.1.) It is clear that the platinum particles grow to the same order of magnitude as the particles in the support, 10 to 30 nm, a point which will be discussed further below.

Figure 6.2 shows the variation of metal dispersion with treatment temperature for the 0.5% Pt commercial catalyst as measured by hydrogen and oxygen uptake at 0°G. The open triangles in Figure 6.2 show oxygen uptakes for the same catalyst sample, which was successively sintered overnight in flowing oxygen at 500°, 600° and 700°C. The circles in Figure 6.2 show oxygen uptakes (open circles) and hydrogen uptakes (solid circles) for separate batches of catalyst sintered at the indicated temperatures. Prior to gas adsorption measurements the catalysts were reduced by cooling or heating in flowing hydrogen from the indicated sintering temperature to 500°C, the sole exception being the 700°C successively sintered case where the catalyst was cooled to 600°C in oxygen and reduction was started at 600°C. Catalysts were outgassed in flowing helium for two hours (successively sintered cases) 140



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or one hour (separately sintered cases) at 500°C before being cooled for gas uptake measurement. Use of the catalysts for other experiments prior to the sintering study may account for the 15% difference in initial oxygen uptake observed for the untreated catalyst samples.

It is immediately evident from Figure 6.2 that at moderate sintering temperatures a redispersion of metal occurs for both catalysts. The increase in hydrogen uptake for the sample sintered at 450° C is far greater than the increase in oxygen uptake of the same sample. However, several authors (Chapter 4, references 6.17-6.19) have noted that relative to oxygen, hydrogen uptakes increase as dispersion increases: Hence, for redispersion, particularly to very small Pt particles, larger increases in hydrogen uptake as compared to oxygen would be expected.

It appears that changes in dispersion are strongly related to the atmosphere in which the catalyst is treated. For example, a portion of the 0.5% Pt on Al_2O_3 catalyst for which the oxygen uptake had been determined was used in another series of experiments, during which time it was exposed to flowing hydrogen at 500°C for a total of 10 hr, and to flowing helium at 500°C for 22 hr. The subsequent oxygen uptake increased by only 0.004 atoms/Pt atom (2% of the total uptake) from the initial oxygen uptakes, a value of the order of the accuracy of the uptake measurement. Comparing this with the results in Figure 6.2 indicates that dispersion changes are far more extensive for shorter treatment in oxygen, for which 16 hr treatment at 500°C gave a 19% increase in oxygen uptake.

6.2.2 Variable Time/Constant Temperature Treatments

Several samples of a second commercial catalyst (Engelhard 0.3%

Pt on $A1_20_3$) were treated in air at 575°C for varying time periods, from 6 hr to over one month. Based on the results from overnight sintering reported above, it was felt that sintering at 575°C would provide a rate of change of dispersion sufficiently slow to allow monitoring of the kinetics. However, sintering was unexpectedly rapid; after 6 hr the uptake of oxygen had dropped from 0.350 atoms/atom Pt (for the unsintered specimen) to 0.019. At this low an uptake, gas adsorption measurements are subject to large relative error.

The unsintered catalyst was examined by electron microscopy, and metal particles were observed between 2 and 4 nm. Figure 6.3 shows the observed particle sizes for four of the sintered catalyst samples (6, 12, 120 and 360 hr of sintering). The rapid growth of large metal particles even after a six hour sintering period is evident from the observed metal particle sizes.

Comparison with the results of the earlier experiments involving the 0.5% Engelhard catalyst is difficult, however. Both catalysts are of the surface coated type, where the metal particles are in a thin layer on the outside of the alumina pellet. Thus while 0.3% and 0.5% are nominal metal loadings, the actual concentration of metal in the outer region is unknown and could vary from one catalyst lot to another.

6.2.3 Catalyst Mixture Experiments

A series of experiments were designed to check the effect of addition of a small portion of heavily pre-sintered catalysts on sintering kinetics. A portion of the 2.03% Pt on Alon alumina catalyst was heavily sintered (16 hr at 700°C in flowing oxygen) and had an oxygen uptake of 0.0313 atoms/Pt atom; examination of electron



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3: Particle size distribution from electron micrographs of a sintered Engelhard 0.3% Pt on AJ $_{90}$ cataJyst. Batches were heated at 575°C in air for the indicated time.

micrographs confirmed the presence of large metal particles (range observed was 2.5 to 27 nm). The fresh catalyst had an initial oxygen uptake of 0.256 atoms oxygen/Pt atom.

Four batches of catalyst were then prepared; two of the fresh catalyst, and two of a mixt wt'% of the heavily sintered catalyst and 76.9 wt % of the TaTyst. The batches of mixed catalyst were crushed, were redried an 110°C to promote a thorough mixing of the sintered and unsintered portions. One mixed batch and one fresh batch were sintered 16 hr in flowing oxygen at 575°C and their oxygen uptakes were measured. These samples were subsequently sintered at 610°C for 16 hr, while the other two batches were sintered at 630°C for 16 hr. Samples were sintered, reduced (by cooling in flowing hydrogen from the sintering temperature to 500°C) and outgassed 1 hr (at 500°C in helium) under identical conditions, the exception being the mixed sample sintered at 610°C which was reduced and outgassed at 610°C.

Table 5.1 shows the initial oxygen uptakes for the four cases. The treatment at 575°C appeared to cause a slight redispersion in the samples. However, for the other two temperatures the mixed catalyst sinters significantly more extensively than the unmixed catalyst: after the 610°C treatment the unmixed catalyst had 66% of its original area compared to 43% for the mixed, after the 630°C treatment the unmixed catalyst had 29% of its original area compared to 3% for the mixed. From this data it appears that the presence of a portion of heavily sintered catalyst. The implications of this conclusion on proposed sintering mechanisms is discussed below.

Table 6.1: Effect of Addition of a Portion of Presintered Catalyst on Sintering Rate of a 2.03% Pt on Alon Catalyst

, K	Axygen Uptake,	atoms O/Pt atom
Treatment	Unmixed	Mixed
Unsintered	0.256	0.204*
, Sintered at 575°C	0.258	0.218
Sintered at 610°C	0.169	0.087
Sintered at 630°C	0.073	0.006

*By calculation from the dispersions of the unsintered and presintered samples.

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comparison of the results in Figure 6.1 and Table 6.1 indicates a dependence of sintering behavior on metal loading for the prepared catalysts. The two Alon supported catalysts (4.76% and 2.03% Pt) were similarly prepared, but the more heavily loaded catalyst shows sintering at 550°C (growth of Pt particles up to 80 nm), while the less heavily loaded catalyst shows no significant change in dispersion after similar treatment at 575°C. Pt particles from 1 to 3 nm were observed in the reduced 2.03% catalyst, compared to 1 to 4 nm for the reduced 4.76% catalyst.

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6.2.4 Effect of Reduction and Sintering on Individual Metal Particles

Unreduced 2.03% Pt on Alon catalyst was deposited onto a specially prepared 'holey' film described in Chapter 2. Benchmarks on the grid made it possible to record electron micrographs of the same area before reduction, after reduction (1/2 hr in hydrogen at 300°C), and after various thermal treatments in flowing helium. Table 6.2 lists the details of treatment for the grids.

While the intention of these experiments was to follow the effect of various treatments on representative catalyst areas, caution must be used before conclusions based on these studies are drawn about catalyst sintering. In several important ways a catalyst supported on a holey carbon film mounted on a tungsten grid differs from a pure catalyst specimen. Other factors limit the accuracy of size and spatial position determinations of supported metal particles. For example:

a) Evidence appeared in about half the micrographs of small shifts (10 nm) in portions of the Alon substrate. In

Table 6.2: Grid Treatments

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Grid	Areas Examined	Treatment History*
1	9	The grid was reduced at 300°C for 1/2 hr and some contamination was noted. After sintering 3 hr at 500°C (inserted in hot furnace), heavy contamination was noted. The film ruptured after a further 3 hr at 500°C.
2	9	The grid was reduced at 300°C for 1/2 hr and some contamination was noted. The film ruptured after 5 hr sintering at 500°C.
3	4	The grid was reduced at 300°C for 1/2 hr and some contamination was noted. The film ruptured after 4 hr sintering at 500°C.
4	4	The grid was reduced at 300°C for 1/2 hr and heavy contamination was noted; sin- tered 2.5 hr at 500°C, contamination re- duced; film ruptured after 72 hr sintering at 550°C.
5	4	The grid was reduced at 300°C for 1/2 hr and deposits on grid noted. Subsequently the grid was discorded.
6	4	The grid was reduced at 300°C for i/2 hr and some contamination was noted. After sintering 4 hr at 500°C, heavy contamina- tion was noted. The grid was sintered an additional 4 hr at 500°C and a very heavy contamination was noted. After sintering, 4 hr at 550°C, very heavy contamination.
	tering treatments w nace unless otherwi rogen.	vere in flowing helium and started in cold se noted; all reductions were in flowing

Table 6.2: (continued)

Grid Areas Examined	Treatment History.
7 3	The grid was reduced at 300°C for 1/2 hr and heavy contamination was noted. After sintering 4 hr at 500°C, some contamina- tion was noted. The film ruptured after 4 hr sintering at 600°C.
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addition, a few micrographs showed evidence of major movements (10 nm) of the substrate. Thus the position of metal particles from micrograph to micrograph may in part be affected by movement in the substrate, arising from 285of H₂O during the reduction step (the first time the substrate is heated above 110°C) or the high temperatures realized in thermal treatment.

b)

c)

Crushing the catalyst and dropping it on the grid gave catalyst clumps of various size. In selecting an area to be examined, a balance had to be struck between very thin clumps of catalyst, where electron microscope imaging clarity was good but the amount of catalyst was low, and thick clumps, where imaging clarity was reduced but the catalyst more closely approximated the bulk typical in a sintered catalyst. The result was a compromise which often reduced the quality of the image over we could be expected in ideal conditions. Where possible, thin regions at the edge of thick clumps were chosen, but this frequently led to a considerable range of elevation in the specimen, a factor which casues resolution and detection problems^{6.16}.

Contamination by carbon was far greater than that observed in conventional micrographs, both after reduction and after thermal treatment. There are two sources of carbon to contaminate the catalyst. One is from the cracking by the electron beam of residual hydrocarbon vapors within the microscope from the oil diffusion pump. However, under normal operation of the Licroscope, with a liquid nitrogen 150

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filled "cold finger" in the area of the sample, such contamination is quite low (see, for example, the micrographs in Chapter 3 and 5). The exone source is the carbon in "e film itself, which du action and sintering treatments may have migrated onto the catalyst from the film. Given the high contamination evident in many of the micrographs as compared to conventional micrographs, we suspect this latter source to be significant; if so, then catalysts treated on grids differ from ordinary catalysts due to the presence of mobile or vaporised carbon.

Evidence from micrographs also suggests that platinum can d) magrate onto the Amrbon, and may accumulate there in preference to remaining on the alumina. Figure 6.4 a shows two micrographs of the same specimen area of grid 6 at relatively low magnification, before reduction and after the second thermal treatment. The large dark region in over left hand corner of Figure 6.4 a appears to be a the Pt crystal on the carbon. Similarly, Figure 6.4 b shows three micrographs of another region of grid 6 before reduction and after the second and third thermal treatments. No only are the development of metal deposits (in the areas marked "a", "b", "c", and "d" in Figure 6.4 b, for example) evident after sintering, but their growth or decay with increasing sintering is also demonstrated. Access by the migrating platinum to the carbon film, and possibly to the metal grid, is a second major difference between grid treated catalysts and ordinary samples.

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With these cautions in mind, several observations about the growth of Pt particles on supported metal catalysts may be made from micrographs of the grid treated catalysts. (Photographic reproduction may have reduced the quality of the micrographs, especially for fine detail; prints of the original micrographs are available from the author on request.)

In the unreduced catalyst the Pt is extremely disperse. While typically Pt particles with images corresponding to 2 nm were noted, comparison of a sintered and unreduced catalyse (for example, Figure 6.5) shows that much of the Pt evident after sintering does not show up in the unreduced state, presumably because it is atomically dispersed and has not agglomerated into crystals. (In Chapter 3 the difficulty of relating image size to object size, particularly for detail smaller than 2.5 nm, was discussed. All size measurements reported here are image sizes corrected for magnification, with a probable variance of up to \pm 1 nm from true object sizes.) In Figure 6.6 larger Pt crystallites than normal appear in the unreduced catalyst; this will be discussed below.

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Some agglomeration of Pt takes place during reduction of the catalyst; examination of Figures 6.6, 6.7, and 6.8 illustrate this. In Figure 6.6, despite heavy carbon contamination in the micrograph of the reduced catafyst, growth of the Pt clusters as compared to the unreduced catafyst can be noted. In both Figures 6.7 and 6.8 numerous larger particles up to 4 nm appear; and a general agglomeration of Pt is evident. This agglomeration is somewhat remarkable considering the mild temperatures (300°C) used in the reduction step. Close inspection of Figures 6.7 and 6.8 show that some of the particles noted 153

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6.5 Micrographs from grid 4 showing the same area before reduction and after one thermal treatment at 500°C.



Figure 6.6

Micrographs from grid 7 showing the same area before and after reduction and after one thermal treatment at 500°C. AL CONTRACT



Figure 6.7 Micrographs from grid 2 showing the same area before and after reduction.





 ${\cal C}^{K}_{n}$

in the unreduced catalyst remain fixed in Tocation during reduction. In Figure 6.7, for example, five particles are illustrated (one each at "a", "b", and "c" and two at "d") which appear in both micrographs. Similarly, two particles each by regions "a" and "b" in Figure 6.8 appear before and after reduction. Not all particles in the reduced catalyst correspond to a particle in the unreduced catalyst; however, the small size of particles involved and the thickness of the specimen make simultaneous resolution of all small Pt particles unlikely^{6.16}.

Thermal treatment at 500°C leads to further agglomeration of Pt, as shown in Figure 6.6 and 6.8. In Figure 6.8, a consolidation of Pt particles is generally evident. In the region "c", a number of small particles evident in the reduced catalyst disappear and are replaced by one large (/4 nm) particle in the sintered catalyst. A similar agglomeration is strikingly evident in Figure 6.6, where particles of the order of 20 nm in size appear after thermal treatment at 500°C. Once again, there is evidence of some swrticles remaining stationary during treatment. In Figure 6.5, particles observed after thermal treatment at regions "a", "b", and "c" cc spond to particles observed in the unreduced state. This is not true of all the particles in Figure 6.5, however, such as those by "d" and "e". In these cases no particles are evident in the unreduced state, although the problem of detectability again exists for the very small unreduced particles. In Figure 6.8, the two particles each in regions "a" and "b" again remain stationary during remain treatment at 500°C, although two apparently new particles appear by "d".

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a.)

Both Figures 6.6 and 6.8 suggest that some localized effects are significant in the changes in metal particles associated with reduction

and thermal treatments. For example, in Figure 6.6 an unusually high density of Pt exists in the unreduced catalyst in the catalyst area. While the reasons for this concentration of Pt salt are unknown, it results in Pt particles 5 to 10 times larger than normal after sintering. In light of the apparent influence of larger Pt crystallites on the over i sintering rate (see Section 6.2.3 above), such localized effects may be significant in the sintering of a catalyst. In Figure 6.8, the apparent agglomeration of the particles in region "c" to the large particle evident after thermal treatment again suggest a localized effect due to the close proximity of so many Pt particles is a section of the solution.

In only one case, grive 6, did the holey carbon film in the areas micrographed survive more than one sintering treatment. Figure 6.9 shows the same specimen area after three thermal treatments (the middle micrograph is the mirror image of the other two because the arid was inverted). The extremely high carbon contamination, evident in all three micrographs, makes the distinction of individual Alon particles impossible, and severely impedes the clarity of the Pt crystallites. Despite this problem, the continued agglomeration of Pt can be noted for increasing thermal treatments: typical particle sizes change from 2.5 to 5 nm. While the poor clarity limits particle detection, we were able to note that some particles did not shift with the thermal treatment. Six particles noted in region "a" after the first sinter were identified in the micrograph taken after the second sinter. Similarly, five particles evident after a thermal treatment correspond to an existing particle in the same location prior to the treatment. The caution must also be re-emphasized that the catalyston grid 6, contaminated by carbon (presumably from the holey film),

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and losing metal to the film surface (shown in Figure 6.4), deviates substantially from a typical Pt/Al_2O_3 catalyst system.

6.3 Discussion

The purpose of these experiments was in part to try to establish the mechanism of sintering, specifically to distinguish between crystallite migration^{6.12, 6.13} and atomic or molecular interparticle transport^{6.14, 6.15} as the sintering process. This pursuit is by no means of academic interest only, since knowledge of the mechanism of Pt agglomeration would suggest appropriate schemes to control or reverse to loss or metal dispersion. Unfortunately, we do not consider the evidence gathered in this work to be sufficiently definitive to allow elimination of one or the other mechanism.

Sintering of the 4.76% Pt on Alon catalyst suggests strongly, however, that crystallite diffusion is not the sole mechanism of catalyst sintering, as previously noted (Chapter 5). The Aloh substrate consists of 10 to 30 nm alumina particles; upon drying these apped to form irregular porous stacking arrangements. As Figure 6.1 shows, growth of Pt particles continues even when the metal crystallite size exceeds the size of a typical support particle; it is difficult to conceive of crystallite motion along the support surface continuing to occur in such a case.

Sintering of the commercial 0.5% Pt catalyst confirms the ability of supported metal catalysts to show an increase in dispersion (mea-.ured by gas uptake) with certain thermal treatments. The atomic surface diffusion model developed in Chapter $5^{6.14}$, 6.15 accounts for redispersion as a transitory effect arising from the buildup of a high

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surface concentration of mobile metal species: cooling of a catalyst could "freeze" this metal as small crystallites or atomically dispersed atoms. Emelianova and Hassau^{6.20} compared the dispersions of supported Pt catalysts which were rapidly and slowly cooled following otherwise identical thermal treatments. Their finding that the rapidly cooled catalyst always had a higher dispersion than the slowly cooled catalyst does suggest a "freezing" of the agglomeration process. Ruckenstein and Pulvermacher^{6.12}, 6.13 do not account for redispersion in their model, but propose to do this in a future work, attributing it to particle breakup caused by a spreading surface pressure. Attempts to determine whether redispersion is a transitory phenomenon followed by particle growth were frustrated when the 0.3% Pt commercial catalyst showed an unexpectedly high rate of sintering at 575°C.

The higher rate of sintering observed when presintered catalyst containing large Pt particles is added to a catalyst batch sug that the large Pt particles speed agglomeration by acting as i "sink" for the transported metal. This in turn strongly suggests \pm highly mobile rapidly transported Pt species under the sintering conditions' employed, since if the process were controlled by slowly diffusing Pt crystallites the effect of large metal barticles on sintering rate would be minimal. The atomic diffusion model^{6.14}, 6.15 assumes a very rapid migration of metal atoms or molecules along the substrate such as is noted in film growth studies^{6.21}. Ruckenstein and Pulvermacher 6.12, 6.13 describe a coalescence controlled condition, in which the rate of merger of metal particles is slower than their migration álong is surface. However, Wynblatt and Giostein^{6.9} consider a sintering rate being coalescence controlled as impossible. If so, this suggests 165

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interparticle transport rather than crystallite migration as a sintering mechanism that can account for the influence of the added large Pt particles on the overall sintering rate.

An ideal way of distinguishing between crystallite migration . and atomic diffusion mechanisms would be to sinter a catalyst while observing it in the electron microscope, using a heated grid holder. However, our attempts at this were frustrated because drift of the specimen, presumably due to thermal gradients in the grid, prevented resolution and recording of the metal particles of the order of 10 nm and less. The selection of an area of catalyst, to be examined after various treatments, was intended to approximate as closely as possible the in site experiment. However, two detectabilit ms affected the electron micrograph studies of catalysts treated .ds. For small particles, the problem of simultaneous detection of all Pt crystallites^{6.16} arises. For larger particles, carbon contamination, which we think arises from evaporation from the holey carbon film during thermal treatment, reduces the clarity of the micrographs. WED conclude that at least some Pt particles remain in a fixed location during reduction and continued thermal Areatments, and that age omeration of Pt occurs during all of these steps. For a perfectly nomogeneous surface, the crystallite mustion model of sintering 6.12, 6.13 predicts all Pt particles will move, while the atomic surface diffusion model predicts none will (although some will disappear). On a heterogeneous surface, however, which probably more accurately describes a typical catalyst support, the former model can account for fixed fanticles and the latter can account for the appearance of particles. at new locations. Hence these em studies do not allow either model

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definitively selected.

Contamination of the catalyst by material from the supporting grid would probably be reduced and disappear if a high boiling metal (such as tungsten) holey support film were used; Fukami of aZ 6.22, 6.23 describe the preparation of the support film, shown in Figure 6.4.

6.4 Conclusions

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Sintering of supported metal catalysts was studied in a variety of experiments. For a 4.76° Pt on Alon catalyst, metal particle growth continued to a point where it exceeded the particle size of the support. Sintering of Engelhard 0.5° Pt on alumina catalysts in oxygen gave evidence of redispersion at lower temperatures (450° to 600°C) and loss of metal surface at higher temperatures (600°C). Ji rate of sintering was found to be sensitive to the gas atmosphere and to the metal loading.

Including a portion of presintered catalyst the catalyst batch to be sintered appears to substantially increase the overall rate of sintering. This is attributed to the large Pt particles found in the presintered specimen acting as a "sink" for transported metal.

Electron micrographs were recorded of the same catalyst area before reduction and after reduction and thermal treatments on the grid. Problems were noted with metal transport to the grid carbon support film and carbon transport to the catalyst. Agglomeration of Pt was noted after reduction in hydrogen at 300°C and after thermal treatments in helium at 500° and 550°C. Some metal particles kept a fixed location on the support during reduction and thermal treatment

The data can be interpreted in terms of an interparticle transport sintering mechanism. While some of the data cast doubt upon crystallite migration as a sintering mechanism, we do not consider the results sufficiently conclusive to rule out this model as a possible mechanism of sintering.

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APPENDIX A

EXPERIMENTAL CONDITIONS

A-1 Catalysts and Preparations

Both commercial and prepared catalysts were used in this work. Commercial catalysts were purchased from Engelhard Industries of Canada, Ltd. (512 King Street East, Toronto, "Ontario). The catalysts were 48" alumina pellets with a surface coating of metal. 0.3% Pt (Lot 12, 514), 0.5% Pt (Lot 18, 381) and 0.5% Rhodium (Lot 17, 941) catalysts were employed. Specific pretreatments are noted with the experimental disults.

Catalysts prepared in this laboratory were made with two different supports: Kaiser 201 Alumina spheres (Kaiser Chemicals, Baton Rouge, Louisiana, U.S.A.), and Alon, fumed alumina, (Cabot Corporation, Boston, Mass., U.S.A.). The former was screened in this laboratory, and the 8-10 mesh range was used in all preparations. The latter was used as received.

Catalyst K1 (1.62% Pt on Kaiser 201 alumina) was prepared in the following fashion: to 24.8129 grams of Kaiser 201 alumina was added 30.ml of an H_2PtCl_6 solution containing 0.401 grams of Pt metal. This solution was prepared by dissolving Engelhard Platinum Chloride (40%) (Engelhard Industries, Toronto, Ontario) in distilled H_2O . After allowing to stand for 1/2 hour, the mixture was dryed at 110°C overnight, and then reduced in flowing hydrogen at 250°C for 1 hour. During crushing it was noted that the catalyst had a dark exterior ring, presumably Pt, that extended about 1/3 of the radius. The center of each

alumina pellet was relatively free of Pt. Catalyst K2 (2.035 Pt on Kaiser 201 alumina) was prepared in a similar fashion except that the alumina was first wetted with pure H_20 , then the Pt solution was orded. This mixture was allowed to stand 24 hours before being dried at 110 C. The dark ring of metal penetrated a shorter distance into the pellet for this catalyst, about 0.25 of the radius.

Catalysts Al (0.10° Pt on Alon), A2 (0.51° Pt on Alon), A3 (1.00° Pt on Alon), A4 (2.03° Pt on Alon), A5 (2.46° Pt on Alon) and A6 (4.76° Pt on Alon) were prepared in a similar fashion. The alumina was wetted for form a thin paster and then an appropriate amount of the platinum salt solution was added. This paste was dryed at 110°C, then reduced at 250°C in flowing hydrogen.

-2 Gas. Adsorption System

4-2:1. Calibration, Accuracy, and Sample Calculations

During normal operation of the dynamic system adsorbate gases were added to the continuously flowing carrier gas stream. This was achieved through the Carle sample valve described in Chapter 2, which has two loops that alternately hold a continuously flowing stream of sample gas. Switching of the valve passes the carrier gas through the loop and sweeps a pulse of the adsorbate gas past the catalys. The contents of the loops were calibrated by comparison to known volume cted by syringe. Table A-1 shows the results of an Ω_2 calibration in (Run A), sfor various oxygen flow results of an Ω_2 calibration is a calibrated rotameter) the average peak size from a pulse passing through an empty sample tube is shown. The maximum variation observed among peak areas from the same loop and flow rate was



less than 0.75%. The maximum variation observed among peak areas from the two different loops was less than 2.1%; nowever, this is not necessarily due to differences in the size of the two loops. The carrier gas passes through small ports in the sample valve, and the alignment of the valve (set by hand) was slightly different for the two loops, leading to small variations in the carrier gas flow rate through the two loops. Thus as the valve was operated over a long period of time the relative areas of the peaks from the two loops varied. In the calculations the two loops were assumed to hold the same amount of adsorbate gas.

-437

Table A-1 also shows the area as a function of size of n_2 pulse injected by syringe, with the maximum rigition in peak areas indicated. Syringe contents were calculated from the ideal gas laws. From this data and the peak area as a function of flow through the loop, the size of an O_2 pulse was determined to be 20.3 ± 0.5 u moles at a flow rate of 60 cc (STP)/min through the loop. The variation in sample loop contents are from a variation in atmospheric pressure; which was 700 ± 10 mm Hg. In all standard runs O_2 flow through the sample valve was maintained at the rate of 60 cc (STP)/min, and the 20.3 u mole/pulse figure was used in uptak calculations.

Hydrogen, when used as a sample gas, was also flowed through the loop at 60 cc (SPT)/min, but the lower viscosity of hydrogen relative to oxygen lowered the absolute pressure, and hence the contents, in the sample loop. To calibrate the hydrogen contents, the pressure directly above and below one of the sample loops was measured by an H_2O , U-tube manometer read by a cathatometer. The upper, lower, and average

pressures are shown in Table A-2. Using 13.55 as the specific gravity of Hg, the results were converted to mm_Hg. Using 700 mm Hg as the average barometric pressure (variation from 690 mm to 710 mm was actually observed), the hydrogen pulses are found to contain 0.92 as many moles as the 0_2 pulses. In all standard runs where H_2 was to be adsorbed flow was maintained at 60 cc (STP)/min, and the 0.92 x 20.3 u moles/ pulse figure was used in uptake calculations.

In some runs, the effect of rate of addition of Ω_2 on total uptake was checked by adding smaller Ω_2 pulses. This was achieved by mixing the Ω_2 with helium (the carrier gas) prior to flowing it through the sample loop. Most runs were made using a Type A or Type B dilution, and Table A-3 shows the calibration of the contents of each pulse in these two cases, calculated from the area of the two peaks through an empty sample tube. Two runs (5 and 15) were made at mixing rates for which final areas were not recorded. In these cases the contents of a 'pulse was estimated from the nominal composition of the sample gas and a total content of 20.3 u moles of gas in the sample pulse. This procedure is less acr rate than the calibration by peak area used for the Type A and Type B dilute pulses.

During gas adsorption studies pulses were passed over the catalyst, and the output peak was recorded, taking two to five minutes per pulse. Pulses were communic until a steady output peak for each sample loop was achieved. The area, of these final peaks were assumed to represent 20.3 u moles c C or 3.7 u moles of H₂, and the fractional output peaks were computed by comparison to these output areas, assuming a linear relationship between sample gas amount and peak area. Thus the **V** variation of carrier gas flow rate did not affect the results, since

172 Table A-2: Hydrogen Calibration Gas Flow cc (STP)/min Pupper, cm H_20^1 Plower, cm H_20^2 Pavg, cm H_20 H_2 60 35.6 34.8 35.2 . 117.5 0, 60 114.7 116.1 Based on an average atmospheric pressure of 700 mm Hg and a specific gravity of 13.55 for Hg: $P_{H_2} = 700 + \frac{35.2 \times 10}{13.55} = 726 \text{ mm Hg}$ $P_{0_2} = 700 + \frac{116.1 \times 10}{13.55} = 786 \text{ mm Hg}$ U 1415 $P_{H_2}/P_{0_2} = 726/786 = 0.3$ -3 at upstream end of sample loop ² at downstream end of sample loop 33

	0 ₂ Flow ¹ cc (STP)/	He Flow ¹ cc (STP)/ min	Nominal 0 ₂ Conc	Peak Area Disc Integrator Counts (correct- ed to the same • attenuation)	0 ₂ ,u Moles in Pulse,by Area
Pure	60	3 0	100.0%.	12,740	20.32
Type A	12	49	19.7%	2,850	4.54 •
Туре В	<u>, 5</u>	190	_2.56 ²⁰	493	a (), 79
5	rotameter	tion*(see Ta	· · · · · · · · · · · · · · · · · · ·		

an internal area standard was available. Figure A-1 shows a typical chart output for an oxygen run; the data are from Run 112. Table A-4 shows a typical calculation of oxygen uptake in atoms of 0 per atom Pt for this run.

Helium was always used as a carrier gas for oxygen pulses; both helium and nitrogen were employed for hydrogen pulses. Nitrogen has a much greater difference in thermal conductivity relative to hydrogen than helium. When used as a carrier gas for hydrogen pulses, a higher attenuation is used on the thermal conductivity cell output than when helium was used. This higher attenuation damped out noise and baseline wift due to temperature fluctuations and other funcontrol able factors; hence hitrogen carrier gas gave more accurate and drift free hydrogen uptake results. However, in titration runs, switching from helium to nitrogen as a carrier gas required about a one hour sweep period before a stable baseline was observed. For this reason, later runs employed helium only as a carrier gas. Hydrogen peak areas were checked and found to be linear with pulse size up to a full pulse. An attenuation of unity for H₂ output pulse meant that baseline drift reduced the accuracy of the H₂ peak areas relative to 0₂ peak areas.

A second factor intreasing the relative error in H_2 results was 'the presence of substantial tailing of the hydrogen peaks, attributed to reversible adsorption on the Pt catalyst of a portion of the H_2 pulse. Comparison of O_2 and blank tube peak areas showed virtually no retention of O_2 once the surface was saturated with O_2 . The reversible hydrogen tail could be readily confused with baseline drift. For this reason O_2 uptakes in pulses are generally reported to two decimals (to one-one hundredth of a pulse) in Section A II 4, while



176 Table A-4: Sample Calculations, Run 112 -3 Catalyst Charge: a) 1.901 grams of catalyst A4 (2.03% Pt on Alon) 1.901 g x $\frac{0.0203 \text{ g metal}}{9 \text{ catalyst}} \times \frac{1 \text{ mole}}{195.1 \text{ g Pt}} \times 10^6 \frac{\text{u moles}}{\text{mole}}$ = 197 u moles of Pt atoms 劉xygen Uptake Area, Discillitegratos Counts ۲. ج Fraction Out 7,280 Peak J 7280/12,250 = 0.59412,470/12,640 = 0.985Peak 2 12,470 Peak 3 12,250 1.000 Feak 4 1.000 12,640 Ũ Gross pulses in 4.000 Total pulses out 3,579 Net pulses in 0.421 20.3 u moles pulse 2 atoms molecules 02 X 0.421 pulses χ = 17.1 u atoms 0 c) Uptake: uratoms () 0.087 atoms C/atom Pt a-atoms Pt.

 H_2 results for an He carrier gas are reported to one decimal (one tenth of a pulse) unless manual integration of peaks was employed. Similar tailing was observed in adsorption of O_2 on supported Rhodium catalysts, indicating some reversible adsorption of O_2 in this case.

Exact repeatability of dynamic gas uptake can not be assessed, for two reasons. First, catalysts show some variation from sample to sample in terms of catalyst loading and dispersion, making exact paralel uptake determinations difficult. Second, each successive reduction and outgassing treatment can alter the surface of the metal catalyst. This latter effect seemed far more significant for the catalyst prepared on Alon than for the commercial catalysts or those prepared on Kaiser 201. For commercial catalysts oxygen uptakes were generally repeatable within 0.02 atoms 0/atom Pt if the samples were run at about same time. Shelf ageing does appear to change gas adsorption amounts. Hydrogen uptakes were somewhat less repeatable than oxygen uptakes due to the lower measurement accuracy discussed above.

A-2.2 The Effect of Reduction and Outgassing Times on Gas Uplaces

Whether reduction time affects gas uptake appears to depend in part on the prior history of the catalyst sample. The Englehard 0.3% and 0.5% Pt on alumina cated ats were checked for gas uptake after various reduction treatments. Results of these experiments are shown in Table A-5. None of the 0.3% Pt samples had any prior reduction before the indicated treatment, which the 0.5% sample had been reduced overnight in flowing hydrogen at 500°C before the treatments shown in Table A-5. For the 0.3% catalysts, overnight reduction at 500°C

exertic .	Comments	fresh catalyst	frest catalyst		catalyst was sintered 3	days in air at 575°C.		catalyst had previously	been reduced overnight at		•	-
nditions on Ge.	H2, atoms/atom	1	8 1 1 1		1 1 1	8 1 2	1 1 1	0.591	0.587	0.587	+ + + + +	
Reduction Co	01, atoms/atom	0.078	0.352		0.038	0.043	0.081	0.178	9.178	0.180	0.180	
Table A-5: Effect of Reduction Consitions on S2, Jotake	Reduction Temperature, °C	500	500	4	. 500	500	600	500	500	500	500	
Table .	Reduction Time, hr	~	overnight			overnight	overnight	1.5	4.0	14.5	2.0	,
	Run Catalyst ⁷	. E2	. Е2		E2	E2	E2	Ē	El		El	
	Run	67	98		95	00 L	101	46	<u>47</u>	48	55	

E1 = Englehard 0.5% Pt on alumina

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E2 = Englehard D.3% Pt on alumina

raises the gas uptake. Since the catalyst is treated in air by Engle and temperature unknown), the long initial reduction may be necessary to convert bulk platinum oxides. The increase in uptake observed in run 101 for reduction at 600°C may be due to redispersion of the catalyst; as noted in Chapter 6, such increases in dispersion have been noted during sintering of catalysts in oxygen.

The results for the 0.5% Pt catalyst indicate that once the initial reduction is achieved, subsequent long term rereduction is not necessary. For the various reduction times, the deviation in initial oxygen⁶ uptake corresponds to about 0.01 pulses, the reliability of the peak measurements. Similarly, the titered hydrogen uptakes, measured for three of the runs, show a variance within the precision of the peak area measurements.

Changes in outgassing time cause a more substantial variation in initial oxygen uptake as shown in Table A-6. Again, the Englehard 0.5% Pt on alumina catalyst sample used in the study of reduction times was employed in these runs. Oxygen uptake decreases with increasing outgassing times, probably due to the desorption of surface hydrogen which otherwise consumes oxygen in a reaction to form H_20 . Table A-6 also shows the surface coverage by hydrogen, both in absolute amounts and as a fraction of the oxygen uptake. These were calculated assuming that the surface coverage by hydrogen is zero after 16 hours of outgassing; that the higher oxygen uptakes reacted partially with the hydrogen to form water, leaving the same surface coverage by oxygen, in all cases; and that the carrier gas did not contain any oxygen to build up on the surface. The ability of the catalyst to retain hydrogen even after two hours outgassing is evident from Table A-6.

Tab	ole A	A-6:	Effect	0Ť	()ų	tgassing	Line	on	Gas	Uptake'	
-----	-------	------	--------	----	-----	----------	------	----	-----	---------	--

	•	· ·	Calculated Residual . Hydrogen						
Run	Outgassing Time, mrs	0] atom/atom	atom/atom	fraction of surface					
52	0.25	0.183	0.42	0.26					
49	0.5	0.176	0.28	0.17					
54	0.5	0.180	0.36	0.22					
51	1.0	0.178	0.32	0.20					
55	2.0	0.172	0.20	0.12					
50	4.()	0.168	0.12	0.07					
53	16.0	0.162	0	0					

¹Reduction time two hours in Flowing hydrogen at 500°C

²Assuming: a) no 0_2 in carrier gas :

- b) all hydrogen consumed via $2H + \frac{1}{2} O_2 \rightarrow H_2 O_2$
- c) 16 hr outgassing gives "clean", *i.e.* hydrogen free, surface

The results from Runs 50 and 53 may also serve, with a different set of assumptions, as a test of the maximum oxygen content of the carrier gas. Thus one may assume that the metal surface is "clean" after four hours and that subsequent decreases in oxygen adsorption are due to the continual adsorption from the carrier gas of traces of oxygen during the degassing period. At the carrier gas flow rate of 53.5 cc (STP)/min, 1.72 moles of helium flowed over the catalyst in the 12 hours difference in outgassing time between the two runs. The decrease in oxygen uptake for Run 53 corresponds to 0.65 micro moles of oxygen, so the maximum oxygen in the carrier gas, assuming that all oxygen present would be adsorbed on the catalyst, would be 0.38 ppm. This value compares favorably to an upper limit of 0.4 ppm. found by Meyer and Ronge (Reference 2.4 Chapter 2) in first testing out the Cu/CuO catalyst used in this work. A decrease of oxygen uptake with increasing outgassing time due solely to oxygen in the carrier gas would be linear with time. Since the results shown in Table A-6 do not correspond to a linear decline in oxygen uptake, at least part of the higher uptakes associated with shorter degassing times are attributed to residual surface hydrogen.

A-2.3 Effect of Rate of Addition of Oxygen on Oxygen Uptake

Supported platinum and rhodium catalysts were used to determine the sensitivity of oxygen uptake to the rate of addition of the oxygen to the catalyst. Since the adsorption of oxygen is exothermic and activated, it was anticipated that significant differences in localized heating might arise during the addition of adsorbate gas at different rates, leading to differences in uptake of the adsorbate gas.

In the first set of experiments the oxygen in the sample loop was diluted by mixing with the carrier gas (helium). The range of dilution in the sample loop was from pure 0_2 to 4% 0_2 . The results of uptales, for various stages of dilution for 0.5% platinum (E1) and rhodium (E3) catalysts are shown in Table A-7.

For the platinum catalyst runs, the uptakes at various dilutions are generally within 10° of each other, and show no consistent trend towards higher or lower uptakes for the diluted cases. For the one series where a rhodium catalyst was employed (Runs 13a to 19) significantly higher uptakes were observed for the very dilute pulses. However, as noted above, oxygen adsorption on supported rhodium catalysts includes a reversible slowly eluted adsorption. The high uptake for Run 13a is attributed to the failure to detect extremely broadened peaks. This run was terminated by adding a pure O_2 pulse after 330 dilute type B pulses were added. Total uptake included the 330 pulses and a small fraction of the final pure O_2 pulse. A portion of the first 330 pulses probably bled off without detection.

From these data it was concluded that dilution of a 0.5 cc oxygen pulse did not significantly affect the uptake of oxygen on the catalyst surface. However, even when a pure oxygen pulse is used, the pulse gets diluted by the carrier gas as it is swept into the specimen tube and over the catalyst. A second experiment was designed to see if uptake of oxygen varied when a pure oxygen stream was passed over the catalyst, as compared to an oxygen pulse.

The data, from Runs 93a, b, c, and d are shown in Table A-8; 2.03% Pt on Kaiser 201 catalyst was used. In Runs a and c, the catalyst was contacted with a stream of pure oxygen at about 1.2 atmospheres, then

Run	Catalyst	Temperature, °C	a per Pulsa?,	Dygen Uptake. atoms/atom
5	E1 ³	0	9.2	0.350
· 6		(A.54	0.370
. 7		2	20.3	0.381
8	1		0.29	0.350
9		97	0.29	0.365
10 .			20.3	0.330
11	-		4,54	0.323
13a	E3_	. 0	0.79	1.000
15			1.85	1 Q.519
16			4.54	0.492
18	•	· ·	20.3	0.539
يمر 19 م	•		· A.54	0.499
21	E1	- 96	0.79	0.299
22			A.54	. 0.254
23	`		20.3.	0.256
24	E٦	-98	0.79	0.197
25			4.54	0.186
26			20.3	0.198
27 _	E1	0	0.79	0.214

Table A-7: Effect of Pulse bildtion on Oxygon Uptake

lEl is Englehard 0.5% Pt on alumina E3.is Englehard 0.5% Rh on alumina

²For details on calibration see Section A-2.1

³This sample of El had a varied thermal history in the early trials of the adsorption system and appears to have a significantly higher uptake than subsequent samples of El.

		Table A-7 (ontinued)	
Run	Catalyst Ten	nperature, "C	O ₂ pe pulse, u molias	Oxygen Uptak e , atoms/atom
28	/	,	4.54	. 0.196
29	•		20.3	0.206
30a	El	203 (0.7 <i>9</i>	0.339
31	in the		4.54	0.310
32	() · · ·		20.3	0.319
<u>.</u> 33 .	· ·		20.3	0.301
34	E1	.97	0.79	0.270
35		<i>t</i> ;	4.54	0.277
36	•	J	20.3	0.278
37	E1 sintered at 500°C	0	0.79	0.268
38		·	: [*] 20,3	, 0.254
39	Él sintered at 600°C	0	20.3	0.197
40			0.79	0.195
4]	El sintered at 700°C	0	0.79	0.082
42 '	αι 700 ζ		20.3	0.075

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Table A-7 (continued)

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Table A-8: Oxygen and Hydrogen Uptakes for Stream

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and Pulse Addition of Initial Oxygen

in the second	Run	Initial Oxygen Added by	Ol, Initial Oxygen Uptake, atoms/atom	H2 Hydrogén Titer Uptake, atoms/atom	03, Oxygen Titer Hptake, alums/atom
	93a	~stream	0.276 ¹	0.992	0.438
	93b	pulse	0.226	0.891	0.434
	93c	stream	0.2512	0.926	0.442
	93d	'pulse	0.225	0.874	0.431

¹By calculation, assuming H2' (run a) = H2' (run b)

²By calculation, assuming H2' (run c) = H2' (run d)

hydronan and oxygen titer steps were carried out. Runs bland diwere ridentical to Runs a and c except that the initial oxygen was added by pulses for Runs b and d. If one assumes that the Bydrogen left on the surface after titration (H2') is the same in Run a as in Run b, and similarly the same in Run c as in Run d, then the initial oxygen uptake (O1) for Runs a and c may be calculated. Accounting for the water formed during titration, H2' (Run b) = 0.439 atoms/atom, so that O1 (Run a) = 0.276 atoms/atom. This represents about a 200 increase in initial oxygen uptake in Run a as compared to Run b. Similar calculations for Runs c and d give H2' (Run d) = 0.424 atoms/atom, O1 (Run c) = 0.251 atoms/atom, or about a 10% increase in initial oxygen uptake in Run c as compared to Run d.

Before concluding that oxygen uptake is significantly enhanced, a comment must be made on the assumption that H2 (Run a, c) is equal to H2 (Runs h, d). If the nydrogen uptake following titration is higher for a higher initial oxygen uptake, Ol, then the calculated increase in Ol for Runs a and c will be correspondingly lower. One factor raising Ol for cases a and c is the higher partial pressure of oxygen when a pure 0_2 stream is used. Whether localized heating effects are also involved cannot be determined from these four runs.

In conclusion, further dilution of the 0.5 cc pulses of oxygen used in this work appear to have no significant or consistent affect on oxygen uptake on support Pt catalysts. Use of a pure oxygen stream appears to increase oxygen uptake slightly as compared to addition of oxygen by pulses.

A-2.4 Experimental Conditions and Results

Table A-9 gives in summary form the experimental conditions and

results for the dynamic gas adsorption system. Gus uptake results are in pulses of 0_2 or H_2 ; 01 is an initial oxygen uptake, H_2 is the subsequent uptake of hydrogen uptake, etc. Thus the series $0_2-H_2-0_2-H_2$ leads to uptakes 01, H2, 03, and H4, while the series $H_2-0_2-H_2-0_2$ leads to uptakes H1, 02, H3, 04. The pulse contents are 20.3 u moles 0_2 and 18.7 u moles H_0 , as discussed in Section A-2.1.

Specific comments on each run follow Table A-9. The following codes are employed in the table:

Catalyst Code: (all concentrations are weight %)

El Englehard 0.5% Pt on alumina (commercial)

E2 Englehard 0.3% Pt on alumina (communcial)

E3 Englehard 0.5% Rh on alumina (commercial)

K1 1.62% Pt on Kaiser 201 alumina (prepared)
K2 2.03% Pt on Kaiser 201 alumina (prepared)

A1 0.10% Pt on Ale ale na (prepared)
A2 0.51% Pt on Alon alumina (prepared)
A3 1.00% Pt on Alon alumina (prepared)
A4 2.03% Pt on Alon alumina (prepared)
A5 2.46% Pt on Alon alumina (prepared)

A6 4.76% Pt on Alon alumina (prepared)

Reduction Code:

1 - Overnight (14 hr) in flowing H_2 at 500°C.

2 - 1 hour in flowing H₂ at 500°C.

3 - Other, see comments.

A - Reduction started in room temperature furnace, heatup rate was ${\sim}500^{\circ}\text{C}$ per hour.

B - Reduction started at 500 °C,

C - Reduction started at some other remperature. Unless otherwise noted, in these cases the remperature controller set point was 500°C, and the furnace was either heating or cooling to that temperature.

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188

Outgassing Code:

1 - Two hours in flowing He at SODOC.

2 - One hour in flowing He at 500° C.

3 - Other, see comments.

Sample Gas Code:

A starred Ω_2 (Ω_2^*) means that the oxygen pulses were diluted for this run; details of the dilution are in the comments. From Section A-2.1, a Type A dilute pulse contains A.54 u moles Ω_2 per pulse, a Type B dilute pulse contains 0.79 u moles Ω_2 per pulse.

	System
	Adsorption
	ı Dynamic
	from
	Results
	A-9:
	lable
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	Pulses					4.93	2,01	2.46	0.62	0.336	58.9	0.5	<u> </u>	9.32	08.1	•	
	Uptakes, P			1		6 = 10	2 = (0	3 = 5	0 = ,10	$\mathcal{D} = \dots \mathcal{D}$	5 = 5	0J = 60	01 = 2.	6 = <i>\U</i>	{ = }∂) 	
iystem	Temperature °C	1 1 1		Q		0	6	₹ 7	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~		0.	97	97			с.	
Results from Dynamic Adsorption System	Sample Gas	02	· 0,	,,	0 ⁵	*~0	* ~				*~~ ~	02*.	62	о х + С	* ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	***	•
n Dynamic A	Carrier Gas	He	Не	Не	Не	Не	He	, ak	v	122	lle	He	Не	Не	÷t y	He	
ts fron	Out Gas	1 1 1	1 . 	-	1 1 1	_	م ر ب	~			~		~	2	·.	~	•
:6-	Reduçtion	 	- - - - 	10	8 8 1 1	JC	3C	35	,		31	10	2A .	łC	ЧC	SC	-
Table A	Catalyst Weight, Grams,	1 1 1	1 1 1 1	10.277	1 2 5	10.227	10.227	30.223	, X		10.227	10.227	10.227	19.227	10.223	10.305	•
1	Catalyst) 	י ג נ	El .	 	El	٤٦	£		•	Ē1 .	٤٦	Ε)	٤٦	5	83	• • •
	Run	-	2	د ۲	4	5	è. P	N [°] ≥	• .	. ,	8	6	10	~~	, 12	e C	

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Uptakes, Pulses	01 = 319.6	1	01 = 70.5	01 = 27.6		01 = 6.67	01 = 27.2		01 = 47.7	01 = 7.03	D) = 7.59	D) = 32.3	01 = 5.28	01 = 1.26	01 = 36.3	01 = 5.77
Temperature °C	0	0,	0	0	0	0	0	0 76	201	96	96	-98	- 98	- 93	0	0
Sample Gas	02*	02	۰°* 0°*	* ^د 0.	0,00	- 02	- 0	02	, 0°*	• * • 0	0	· 0°* .	- ~ ~ 0	2 6	- 00	0 ⁵ *
Carrier Gas	Не	He	Не	Не	. He	Не	Не	He	Не	Не	He	He	Не	Не	Не	Не
Qut Gas		1 L J S	~	—		1	~	 1	1	2	٢	Σ	-	(·	
Reduction	2A	1 1 1 1	ZA	Vc .		<u>.</u>	2A	 	10	2A	, SA	SC ,	74	2Þ	JC	2A
Catalyst Weight,Grams	10.305		10.305	10.305	10.305	10.305	10.305	10.305	9.836	9.836	9.836	10.125	10.125	10.125	10.468	10.468
Catalyst	E3	1 1 1 1	E3	E3	E3	E 3	E3	Ē3	٤١	E)	٤٦	ET.	El	E1	El	Ē
Run	13a	14	15	16	17	18	19	20	21	22	23	54	25		27	58

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Uptakes, Pulses		+			11	н	п	н	, 1 ¹	IJ	, H	і П	D) = 32.5	ti	ų	1		
Jemperature °C		203	203	203	203	- 203	67	67	28	D	Ö	D .	0	C	D	1 2 1		
SamJe Gas	0	2 °C	√ [×] 0	2 × °0	- 0°	°0	° ° ° ° °	ر 0, *	5 G	* °D	× °0	× °0	0°*.	0,* 0,*	ν ₂	0, 20	J	
Carrier Sas	He	He	Че	He	Не	Не	Не	Не	ЭН	Не	Не	Не	Ke	He	Не	Не	n Antonio	
nut Sas	~	1	, , , , , , , , , , , , , ,	~	, (-	-	—	-	∽,	5	(, ,	~	~	1	1		
Reductian	38	1C	ZA	2A	2A	2A	1C	3A	24	38	ZA	3C	2C	4C	ZÅ	1		
Catalyst Weight, Grams	10.468	8,85	8.85	8.85	8.85	8.85	8.065	8.065	8.065	8.065	8.065	10.468	10.468	8.85	8,85	. 1 	•	
Catalyst	Ē	. E]	El	EI	E]	El	, Ц	E	٤٦	٤٢	E۱	El	Ε1	Ε)	Εl	1 1 1		
Run	29	30	30a	31'	32	- 33	34	, 35	36	. 37	38	39	40	ţ1]	42	43		

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Uptakes, Pulses	1 1 1	6 	1 1 1	01 = 0.95	H2 = 3.43	01 = 0.95	H2 = 3.37	0] = 0.96	H2 = 3.37	01 = 0.94	01 = 0.90	01 = 0.95	01 = 0.98	01 = 0.87	. 01 = 0.96	01 = 0.91	
Temperature	Q	0	0	0		Ċ				0	0	0	0	, 0		0	
ⁱ Sample Gas	D_2	02	02	0 ²		02		02		02	02.	02	² 0 .	02	02	020	
Carrier G a s	Нe	Не	Не	Не		Не		Не		Не	Не	Не	He,	Не	Не	Не	
Out Gas.	\sim	۲	т		-	-		-		m	m	2	ε	ω [´]	ŝ	-	
Reduction	75	JC	3B	38		3B		38		38	38	3B	38	38	38	38	
Catalyst Weight, Grams	1 1 1 1	1 1 1 1		8.459	-	8.459		8.459		8.459	8.459	8.459	8.459	8.459	8.459	8.459	
Run. Catalyst	E3	El	,	Ē		E		E]		EJ	Ц	El	EI		EJ		-
Run .	<i>なな</i>	45		46		47		48		49	50	51	.52	/53	54	55	

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192

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	Uptakes, Pulses -	H1 = 1.30	п	0] = 0.95 u2 - 2 24		H2 = 3.Q2	HI = 1.18	02 = 1.35	01 = 0,88	H2 = 3.35	H1 = 1.52	02 = 1.51	H1 = 1.24	02 = 1.32	01 = 0.83	H2 = 2.59		
	Temperature °C	0	,	0	G	·	D		0		0		0		0			
	Sample Gas	H ₂ , 0 ₂	-	⁰ 2, ^п 2	02, H ₂		H2, 02		02, N2		H ₂ , 0 ₂		H ₂ , 0 ₂		0 ₂ , H ₂		•	
	<i>Larrier</i> Gas	иг, Не		, su	He, N ₂		N2, He		He, N ₂		N2, He	•	N2, He		He, N ₂	·		
	Out Gas	2	~	L	2	·	\sim		2	·	2		2		∼,			
	Reduction	28	. ЯС	7	3C	-	2C	-	, 2C		ZA		30	,	2C	••• •		
	. Catalyst Weight, Grams	8.459	8.459) - -	7.305		7.305		6.952		. 6.952		6.600		6.600			
- ,	Run Catalyst Catalyst Weight, Grau	Ē	 ניבו		Ē٦		٤	1	Ē	ŕ	El		E]	• • •	EJ			
•	Run	56	27		58		56		60		61		62		63		·	

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Run	Catalyst	Run Catalyst Catalyst Weight, Grams	Reduction	Out Gas	Carrier Gas	Sample Gas	Temperature °C	Uptakes, Pulses
64	E I	6.265	ЭС	2	не, И ₂	02, H2	0	01 = 0.36
		~						H2 = 1.06
65	EJ	6.265	2C	5	N ₂ , He	H ₂ , N ₂	0	H1 = 0.39
- (02 = 0.47
6 6		5.913	ЗА	\sim	He, N ₂	H ₂ , 0 ₂	0.4	01 - 0.24
			-					H2 = 0.96
6/	ш	5.913	3C	2	N ₂ , Не	^{,4} 2, 0 ₂	0	H1 = 0.032
				، •				02 = 0.135
68	E	5.913	2C	2	He, 1,2	02; H ₂	0	~ 01 = 0.087
								H2 = 0.241
69	<u></u>	8.5	2A	\sim	He, N ₂	02, H ₂₀	Q v	- 0) = 2.94
						,		H2 = 8.5
20	۲ کر	8.5	2A	\sim	N ₂ , He	H ₂ , 0 ₂	0	H1 = 5.2
			C		١	-		02 = 4.50
71	К К	8.5	2A	2	He, N ₂	0 ₂ , H ₂	0	01 = 2.10
					-			H2 = 9.1

1																	
Uptakes, Pulses	Q1 = 2.76	K2 = 9.0	no uptake of S ₂ or H ₂ observed ²	01,= 2.02	H2 = 8.6	H1 = 4.0	D2 = 3.74	H3 = 8.5	04 = 3.84	H5 = 8.8	06 = 3.99	4.0 02 = 3.67	8.2 04 = 3.72	·8.4 06 = 3.82	8.6 08 = 3.89	8.8 010 = 3.22	
Temperature V; °C	98		IJ	RT	•					· u		= \ <i>\</i>	H3 =	. H5 = '	= 7H	= 6H	
Ten					÷										,		
Sample Gas	02, H2		G2, K2	02, H ²		H ₂ , 0 ₂						H2, 02					
Carrier Gas	не, N ₂		tte, tt	He		Не						Не	·				
Out Gas	\sim		~	° ∾		2						\sim		r			
Reduction	. 2C		34	3A		ЗА						ЗА					
Catalyst ight, Grams	8.5		X	8.32		8.32		~		•		8.32			i.		
Run Catalyst Catalyst Weight, Gran	\$	/	pure Kaiser 201 alumina	27		12						ſX					
Run	72		23	74		74a				4		75					

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re Urtakes, Pulses	= 8.9 012 = = 9.0 014 = = 9.0 016 =	= 9.0 018 = = 9.0 020 = = 9.0 022 =	123 = 9.0 1224 = 2.02 125 = 9.1 026 = 4.08 115 = 9.0 028 = 4.18	kl = Q.9 Q2 = 2.31 H3 = 5.3 04 = 2.52))5 = 5.6 D6 = 2.53	
Temperature • °C		- \ \ \			Ĺä	
Sample Gas	H ₂ , 0 ₂			² , ² , ² ² , ²	¢ 5	•
Garren Gas	Не		,	भू इस	, , , , ,	-
Out Gas				$\sim \sim$		
Reduction	36			4C .		
t Catalyst Weight, Grams	8.32			8.32	1 . I . I . I	- Conder
Catalyst Wei	Ϋ́			×3 ×3		
Run	75 con.			38	78	

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с н е			
		58	28
Ч	(78 7	
He	0	28 2	
)			
He	2	28 2	
• 、 、			
Не	\sim	2B 2	
He	\sim	28 28	
ЯЮ	5	28 2	
Не	ς	2B 3	
			•
Не	5	28 28	-
	•	•	•
			$\langle \rangle$

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		re Uptakes, Pulses	01 = 2.15		H1 = 0.137 02 = 0.192	H3 = 0.445	01 = 0.84 H2 = 0.377	03 = 0.200	negligible uptake		H2 = 11.6 03 = 4.72	0] = 2.43 H2 = 10.4	03 = 4.67	H2 = 10.8 03 = 4.75	Q1 = 2.42 H2 = 10.2	Q3 = ·4 .64 -		0) = 0.061		•
	ý)Témperature °C	<u>я</u> т	-196	RT		RT		RT	RT	RT ·	RŢ		RŢ	RT	~	RT	AT		
\sim		Sample		H ₂ , D ₃	H2, 02	,	02, H ₂	·	#2 .	H2, 02		0 ₂ , H ₂		H ₂ , 0 ₂	² , ² 2		02	02		·
		Carrier Gas	He	He	Не		He .		Не	He	Не	Не		H,e);e		Не	He		
•		Out Gas	′∾	, 0	\sim		5	1	1 1	2	2	5		5	2	•	2	2		
		Reduction	, 28	28	28		2B ·		1 t 1 1	28	, 2A	2A		2A	, 2,4		2А	28		
,		Weight, Grams	7.595	8.889	2.49	•	2.49		, , , ,	1 8 8 8 8	4.198	4.198	-	4.198	4,198		4:960	4.263		
	+ - 1 - 1	varaiyst W	A6	٤١	AI .		٩١		. 46	A2	K2	K2		K2	, K2		A4	E2	×	
	Pun	1	88	, 89 ,	06-		90a		-16	92	93a	93b		93c	93d		94	95		

-1	e Temperature Uptakes, Pulses		RT DJ =	RT 01 = 0.626	RT 01 = 0.028		1	RT 01 = 0.138		1 25 2	L	H3 = 8.0 RT H1 = 3.75 02 = 4.1	H3 = 9.4	. RT 01 = 1.45	RT 01 = 1.12	RT. $QI = 0.407$
	Sample Bas	6	°0°	с О, Н,		0, 0	ີ ດ	0, H,	2 °D	N C	42, 07	H ₂ , 0 ₂		02	02	6
	Carrier Bas	Э,Р	He	Не	Не	He	, He	Нe	+ He	Не	He	Не		ט בי	. He	He
	Out Gas	2	· ~	2	N	N	3	Ċ	\sim	∼∪	ŝ	Ś	с С	JC	N	2
•	Reduction	28	28	lВ	1B	18	3A .	30	٩ŀ.	2B	, 2A	ZA	۲ ۲		с С	30
	catalyst Weight, Grams	4.731	9.232	4.705	3.972	4.263	4	1.118	8.473	3.066	3.066	3.066	2.189	2 006	1	2.125
	Kun Catalyst ₩∈	E2	£2 +	E2	. E2	E2	E2	> 44	A4	A5	45	A5	A4	A4		44
C	Kun	. 96	32	, 98	66	100	101	102	103	104	105	106	107	108	002	501

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199

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Uptakes, Pulses 01 = 0.031 01 = 1.131 01 = 0.421					
Temperature °C RT RT RT					
Sample Gas 0_2 0_2 0_2 0_2					
Carrier Gas He He He		``````````````````````````````````````	·	· · · · · · · · · · · · · · · · · · ·	
Out Gas 2 3				•	
Keduction 3C 3C 3C 3C	• •				
ud ta 1955 leight, Grams 2.62 1.901					
A4 - A4 - A4			. ·		
110 111 112			·		547

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Comments on Dynamic System Runs

Run	
1.	Calibration, injected 0.5 cc (STP)/min pulse 02.
2.	Calibration, injected 0.3, 0.5, 0.7 cc (STP)/min pulse 02.
3.	Recorder test, uptake not measured, reduction started at 250°C.
4.	Calibration, injected 0.4, 0.5, 0.5, 0.7 cc (STP)/min pulse 0_2 .
5.	Dilute n_2 pulse (~9.2 × 10 ⁻⁶ moles n_2 per pulse), reduction
	started at 250°C.
· 6.	Type A dilute O2 pulse, reduction started at 250°C.
7.	Pure O_2 pulse, reduction started at 250°C, Ol' is after 2 hrs
	in He at 250°C, 01" is after subsequent overnight purge in He
	at 250°C, peak areas computed for four different flow rates.
8.	Type B dilute O2 pulse, reduction started at 250°C.
9.	Type B dilute N2 pulse, reduction started at 250°C.
10.	Pure O ₂ pulse.
11.	Type A dilute O2 pulse, reduction started at 250°C.
12.	Reduced 6 hr in flowing H _z at 500°C.
13.	Type B dilute O2 pulse, run aborted by plugged line, reduction
	started at 250°C.
13a.	Type B dilute O_2 pulses (330); then pure O_2 pulse added.
14.	Rhodium catalyst causes tailing in oxygen peak.
15.	Dilute Ω_2 pulse (~1.85 x 10 ⁻⁶ moles Ω_2 per pulse).
16.	Type A dilute O ₂ pulse.
17.	Some desorption of n_2 observed at $200°C$.
18.	Pure 0 ₂ pulse.
19.	Type A dilute O ₂ pulse.
20.	Attempt to determine AH_a for reversible Ω_2 on RH by getting holdup

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Type B dilute n_2^{\prime} pulse, reductive started af 250°C. 22. Type A dilute O₂ pulse. Pure 0, pulse. 23. Type B dilute 0_2 pulse, reduction started at 250°C. 24. 25. Type A dilute 0, pulse. 26. Pure 0₂ pulse. Type B dilute $\hat{0}_2$ pulse, reduction started at 250°C. 27. 28. Type A dilute 0, pulse. Pure 0_2 pulse, reduction time 1.75 hr. 29. 30. Run aborted, reduction started at 250°C. 30a. Type B dilute 0₂ pulse. 31. Type A dilute 0, pulse. Pure 0, pulse. 32. Pure 0, pulse. 33. Type B dilute 0_2 pulse, reduction started at 250°C. 34. 35. Type A dilute 0_2 pulse, reduction time 0.6 hr. Pure 02 pulse. 36. 37. Catalyst from Runs 34-36 was sintered in 0_2 at 500°C overnight: Type B dilute 0_2 pulse, reduction time 2 hr. Same catalyst as Run 37, pure 0_2 pulse. 38. 39. Same catalyst from Runs 27-29 was sintered in 0_2 at 600°C overnight; pure O2 pulse; reduction started at 600°C, lasted 2 hr. Same catalyst as Run 39, Type B dilute 0_2 pulse reduction 40. started 150°C. 41. Same catalyst from Runs 30-33 was sintered in 0_2 at 700°C

for various temperatures; excessive mixing prevented accurate

assessment of holdup.

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overnight; Type B dilue 0_2 pulse; reduction started at 600° C, lasted 2 hr.

- 42. Same catalyst as Run 41; pure O_2 pulse.
- 43. Correlation of dilute peak sizes.
- 44. Check on tailing, reduction started at 250°C.
- 45. Reduction started at 250°C; run aborted due to open valve. Reduction time was 0.5 hr., outgassing time was 4 hrs; tube burst after this run, catalyst lost.
- 46 \therefore eduction time 1.5 hr.
- 47. Reduction time 4.0 hr.
- 48. Reduction time 14.5 hr.
- 49. Outgas time 0.5 hr., reduction time 2 hr.
- 50. Outgas time 4.0 hr., reduction time 2 hr.
- 51. Outgas time 1.0 hr., reduction time 2 hr.
- 52. Outgas time 0.25 hr., reduction time 2 hr.
- 53. Outgas time 16.0 hr., reduction time 2 hr.
- 54. Outgas time 0.5 hr., reduction time 2 hr. A recheck of Run 49, no significant change in dispersion.
- 55. Outgas time 2.0 hr, reduction time 2 hr.
- 56. Titration, H₂ first.
- 57. Same catalyst as Run 56, titration, O_2 first.
- 58. Reduction started at 250°C, lasted 3 hr., titration, 0_2 first.
- 59. Same catalyst Run 58, titration, H_2 first, reduction started at 250°C.
- 60. A portion of the catalyst used in Runs 58 and 59 is sintered overnight in flowing 0_2 at 450°C. Titration, 0_2 first. Reduction started at 450°C.

- 61. Same catalyst as Run 60, titration, H₂ first.
- 62. A portion of the catalyst used in Runs 60 and 61 is sintered overnight at 600°C. Titration, H₂ first. Reduction started at 600°C, lasted for 1 hr.
- 63. Same catalyst as Run 62, titration, 0_2 first. Reduction started at 310°C.
- 64. A portion of the catalyst used in Runs 62 and 63 is sintered in O_2 overnight at 700°C. Titration, O_2 first. Reduction started at 700°C, lasted 1.5 hr.
- 65. Same catalyst as Run 64, titration, H_g first. Reduction started at 300°C.

66. Same catalyst as Runs 64 and 65, Duplication of Run 64.

- 67. A portion of the Catalyst used in Runs 64 and 65 is sintered in O_2 overnight at 800°C. Titration, H_2 first. Reduction started at 800°C, lasted 1.4 hr.
- 68. Same catalyst as Run 67, titration, O_2 first. Reduction started at 200°C.
- 69. Titration, 0₂ first.
- 76. Catalyst exposed to H_2 at 600°C prior to this run due to furnace malfunction, Same catalyst as Run 69, titration, H_2 first.
- 71. Same catalyst as Run 69, a repeat of that run.
- 72. Same sample as Run 69, titration, O_2 first, Reduction started at 100°C.
- 73. A test of adsorption on an alumina blank. Heated to 500° C in H₂. 74. A portion of the Catalyst used in Runs 69-72 is used in this run

and run 75. Reduction 0.5 hr. Long term titration.

75. Carrier gas was passed through liquid nitrogen trap for this

long term titration. Reduction 0.5 hr., same catalyst as Run 74. 76. Same catalyst as Run 75. Reduction 0.15 yr., 0.2 cc of CCl_4 injected in carrier gas line prior to first adsorption, no uptake observed. Overnight degassing at room temperature does not lead to uptake, but reduction in H₂ at 500°C does.

- 77. Titration, H₂ first.
- 78. Test of linearity of peak size of H_2 to amount of H_2 , using a blank tube and varying H_2 pulse sizes. Response was linear over region observed.
- 79. Titration, H₂ first.
- 80. Same sample as Run 79, titration, n_2 first.
- 81. Titration, 0₂ first.
- 82. Same sample as Run 81, titration, H₂ first.

83. Titration, H₂ first, at -98°C.

84. Same catalyst as Run 83.

85. Same catalyst as Run 83. Peaks were too diffuse to be recorded.

86. The catalyst as Run 83. Outgassed at 400°C for 1 hr., 500°C for 0.5 hr. Titration at 97°C.

87. Same catalyst as Run 83. Titration at 206°C. Reuptake of H₂ after 1 hr was 0.23 pulses.

88. A portion of the catalyst used in Runs 79 and 80 is finely crushed.

- 89. A portion of the catalyst used in Runs 83-87 was used in a duplication of Run 85. Again both H_2 and O_2 peaks were too diffuse to be measured.
- 90. Titration, both ways.
- 91. A portion of the catalyst from Runs 79 and 80 was sintered in 0_2 overnight at 700°C, then cooled to room temperature without

reduction. No further uptake of hydrogen at room temperature was observed; however, the sample was small and extensively sintered (which EM pictures show).

92. Titration--run was aborted by malfunction in alignment of sample valve.

- 93a. Catalyst exposed to pure oxygen stream after cooling but before first hydrogen adsorption, hence first hydrogen adsorption is H₂.
 93b. Same.catalyst as 93a.
- 93c. Same catalyst and treatment as 93a.
- 93d. Same catalyst as 93a, a repeat of Run 93b.
- 94. Run aborted due to water in carrier gas line.
- 95. Catalyst was sintered three days in air at 575°C.
- 96. Catalyst was sintered 12 hr. in air at 575°C.
- 97. Unsintered catalyst.
- 98. Unsintered catalyst.
- 99. Catalyst was sintered 6 hr. in air at 575°C.
- 100. Same catalyst as Run 95.
- 101. Same catalyst as Runs 95 and 100, reduced overnight at 600°C and outgassed at 600°C.
- 102. Catalyst was sintered overnight at 700°C in 02; reduction started at 700°C, lasted 1.5 hr., outgassing started at 520°C, lasted 1 hr.
- 103. Unsintered catalyst.
- 104. Catalyst wetted with 0.5 cc H_2^0 prior to adsorption.
- 105. Same catalyst as Run 104, catalyst wetted with 0.5 cc H_2^0 prior to adsorption.
- 106. Same catalyst as Runs 104 and 105.

- 107. A portion of the catalyst used in Run 103 is sintered at 575° C overnight in O₂. Reduction started at 575° C, lasted 1 hr.
- 108. A mixture of 23.1% of catalyst from Run 102 and 76.9% of catalyst from Run 103 is prepared and treated identically to the catalyst in Run 107.
- 109. Catalyst from Run 107 is sintered overnight in n_2 at 630°C. Reduction started at 630°C, lasted 1 hr.
- 110. Catalyst from Run 108 is treated identically to the catalyst in Run 109.
- 111. A portion of the catalyst used in Run 103 is sintered at $610^{\circ}C$ overnight in O₂. Reduction started at $610^{\circ}C$, lasted 1 hr.
- 112. A mixture of 23.1% of catalyst from Run 102 and 76.9% of catalyst from Run 103 is sintered overnight in O₂ at 610°C. Reduction carried out at 610°C, lasted 1 hr., outgassing started at 610°C, lasted 1 hr.

APPENDIX B

CALCULATION OF PHASE CONTRAST IMAGES

Chapter 3 described the theory whereby phase contrast profiles in the image plane may be calculated for a variety of assemblies of scattering centers. The EMCON program was developed to perform this calculation.

Table B-1 gives the spatial coordinates of the scattering centers, which are based on a close packed spacing of 0.275 nm. These values were read into EMCON as XN(J) and YN(J), in Angstrom units. Table B-2 lists the parameters used in the EMCON program. The Doyle-Turner scattering equation is detailed in reference 3.17 (Chapter 3); the parameters for gold were us

The EMCON program is listed. It uses the IMSL DRMBIU integration routine (Library 1, International Mathematical and Statistical Libraries, Inc., Houston, Yexas, U.S.A.) and the SSP BESJ program for the zero order Bessel function value (IBM System 360 Scientific Subroutine Package Programmer's Manual (H20-0205), IBM, New York, N.Y., U.S.A.). The program determines the square of the wave function, ψ^2 , as a function of the spatial position in the image plane. Thus a contrast profile of the atom or atoms scattering the electrons may be determined. Directly following the program a sample output from EMCON is also shown.

Case	Atom	number	Coordinates of atom, A
(1)		1	(0, 0)
		1	(58771, 0)
$\begin{pmatrix} 2\\ -\infty \end{pmatrix}$		Ś	(-0.79386, 1.375)
		3	(-7,79386, -1.375)
	to 3 atom	1	(1,68771, 0)
case with in center	4th atom	2 ·	(-0.79386, 1.375)
•		3	(-0.79386, -1.375)
	-	4	.(0, 0)
$(\overline{4})(\overline{3})$]	(0, 0) .
5 1 2		2	(2,75,0)
.67		3	(1.375, 2.38157)
У	· · ·	4	(~1.375, 2.38157)
↓ ×		5	(-2,75,0)
		6	(~1.375, -2.38157)
		7 ·	(1.375, -2.38157)

Table B-1: Atom Coordinate Positions Used in EMCON

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All reported profiles were stepped off in the +x direction.

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Table B-2: Parameters Used in EMCON

All four cases were evaluated for the following input data:

· •'			
	WAVEL	(wavelength of incident electron)	0.037 Å
	APER	(aperture size)	40 /microns
	HMIN	(minimum integral step size)	5 x 10 ⁻⁷ radians
	FOCLEN	(focal length of objective lens)	1.6 mm
	TOL `	(tolerance of integral in DRMBIU program)	0.005
	DELRAD	(radial spatial increment)	0.5 Å
	СЅРН	(spherical aberration factor)	1.6 mm
	DELFOC	(defocus in Angstroms)	-200. to +400. nm, in 20 nm increments
	LI	(number of radial increments)	21

In addition, the single atom case was evaluated over the DELFOC range -20. to +200 nm, in 20 nm increments, for the above conditions with APER values of 32, 16, 8 and 4 microns.

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COMMON XN, YN, XI, YI, WAVEL, CSPH, DELFOC, AL, A2, A3, A4, RI, 82, 83, A4, LUMP ATOM COURDINATES IN ANGSTROMS FROM CFNTER EMCON DETERMINES THE CONTRAST BY CALCULATING THE WAVE FOT READ ILUMR, 101 WAVEL, APER, HMIN, FOCLEN, TOL, DELRAD, CSPH, DFLFOC WAVELENGTH DF INCIDENT FLECTRON IN ANGSTROMS AL THRU R4-ARE DOYLE/TURNER SCATTER FACTOR COMSTANTS MIMIMUM INTEGRAL STEP SIZE IN RADIANS FUCAL LENGTH DF DBJECTIVE LENS IN ME SPHERICAL ABERATION CONSTANT IN MM I, L2, PI, C1, C2, C3, R1ADJ, B2ADJ, B3ADJ, B4ADJ, CD RADIAL INCREMENT IN ANGSTROMS NUMBER OF RADIAL INCREMENTS APFRATURE SIZE IN MICRONS READ (LUNR, 20) A1, B1, A2, B2, A3, B3, A4, B4 TULFRANCE OF JNTEGRAL DEFOCUS IN ANGSTROMS DIMENSION WK(300), XN(50), YN(50) 1,22,1 1, 22, ATNMS = ((() NX) IJ LYNLJIJ YN(J) ARE IF (WAVEL) 1000, 1000,40 NUMBER DF RFAD (LUNR, 301 L1, L2 FURMAT (8F10.0) FORMAT (RFID.)) 2.*PI/WAVFI READ (LUNR, 35) READ ILIMR, 35) FORMAT (RFIG.3) FORMAT 1 21207 ULU AND DATA ARF PI=3.141592 DELRAD = P1/2. DELFNC FOCLEN EXTERNAL NAVFL LUNR = 59 11 SPH **CONTINUE** NIWH APFR TUL L2 LUNP IJ 5 35 2 40 20 30 ں

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3, F7,1,5X, ORJECTIVE LEWS FOCAL LENGIN =1, F6,21, 2PERTURE IN KICY α 29/1 SPHERICAL ABFRATION IN MM = 1, F5.2, 5K, 1 GEACUS IN ANGSTROWS =1 IAXIMUM ANGLE IN RANIANS=", F8.5,5%, " TOLERANCE IN INTEGRAL =, F1. ч У , FOCLEN, APER, L2 =1,F6.2 FORMAT (////' RASIC DATA'/' WAVELENGTH DF FLEGTROM = ', F7 PSI SOUARED = 1, F8, 5// FURMAT (PUSITION IN ANGSTROMS, XI = , F6.2,2X;, YI = DRMRIU(F,0., ANGMAX, HMIN, HMAX, T0L, WK, JER) WRITE (LUNP, 50) WAVEL, ANGMAX, TOL, CSPH, DELEAC 40WS = . , FS. 1, 5%, , NUMBER DE ATOMS = . , [3][] I. + 4.*PI*I.196*PI/MATEL) WINP, ICH, XI, YI, RI, PSI, PSISOD 2•*P1*CSPH*10000000./(4•*WAVEL =',F8.5,5X,' = • 00 I* APFR / (2.*FnCLEN) 200.*(K-1) C3 = 2.*P1*DELFOC/(2.*WAVFL) 10 1) XI + PSI SO D CALL UFRISIIER, DRMBIND COSTAR (OSTAN MALENS) AND SAND + DFLRAD*(I-1) BI/IWAVFL**2) 821(\\\\\\\\\\ = B3/1HAVFL**21 = R4/1WAVFL **2) .>2 * DNGMAX IN = ', FIO. 7, 5X, ' PSI 101 FURMAT (F10.3, F20.6 I PSI SONI + = 1,12 (ARC) = -20). 1,[1 FUNCTION FLXI SORT 11 WRITE (7, ABC = ABS7 999 K PSISOD =11 CONTINUE Ð CONTINUE 1 55 UU • C II • C H 60 TO 1 ANGMAX = XVWH DFLFDC BIADJ RZANJ 84105 BBADJ WRIJE = 1SdH STOP END 23 Ι× 'Å ۲ ر ۲ 52 100 50 666 1000

XN,YN,XI,YI,WAVEL,GSPH,NELFOC,AI,A2,A3,A4,RI,R2,83,R4, UUMP FURMAT (* SSP FRRUR MESSAGF FUR BESJ IS*, I2, 5X * NEW RESJ PRFCISIO 1A2*FXPI-B2ADJ%ISIN(X/2.1**211 + A3*EXP(-R3ANJ%(SIM(XJ2.)**2)) + 24 F = GOS (GO -C2*X**4. +C3*X**2) * 123*X**21 * 143*FXD1-B22D3*(S1*(X72. **21) 1,42,P1,C1,C2,C3,B1ANJ,B2ADJ,B3ADJ,B4ADJ,G0 2**((X)NX+ IX) + 2*EXP(-84403*(SIM(X/2.)**2))) * BJSDM * X CALL RFSJ (C1*C*X , 0, RJ, PRECIS, JKL) IF (ICOUNT ⁴- 4) 910,910,920 WRITE(LUMP,901) JKL, PRECIS IF (C1*C*Y) 187,180,190 $C = S^{nR}T ((XI + X^{N}(K)) **2$ 5. * PRECIS IF (JKL)200,200,000 ICOUNT = ICOUNT + 1 BJSUM = RJSUM + RJ $D\Gamma 250 K = 1, L2$ PPFCIS = .001 IN IS', FIJ. R), BJSUM = 0.ICOUNT = 0 GN T0 190 GN TO 201 PRECIS = CONTINUE CONT I NUE COMMON RFTURN STOP END ВJ 90 I 180 0 I 0 250 190 006 200 9 20

EASIC DATA EASIC DATA EMAYELENGTH OF ELECTEON = 0.0370 MAXINUM ANI SPHEFICAL ADEFATION IM M = 1.60 _ D2FOLUS II ADEFHIPP IM MICPONS = 40 0 UNMAREN OF 22008	AXINGM ANGLE IN PAJIANS= 0.01250 D2P02US IN ANSTROMS = 200.6 OP 22048 = N	TOLEBANCE IN INTEGRAL =0. OBJECTIVE LENG FOCAL LENGTH		
IY DIAL		14846.0 = ISG	18(18.0 = C35,4000 IST	· · · · · · · · · · · · · · · · · · ·
	С UIGHEU С UIGHEU 7(UI000.0-= КН	0.5079.0 = 129	3-2+5 = 3-2+5 IEd	
••• I N S LINEATSI) ••• NON-NEFINED POSITION IN ANGSTRUMS, XI = 1.50 YI = 0.0	0 014411 28 =-0.03(1102	48666 d = 15d	45579.0 = CZEKUQC IEA	-
••• I N S LIVEFICT) ••• KON-DEFINED POSITIOS IN ARGUTEDES, XI = 1.50 YI = 0.0	0 012490 640300-0-= K4	154 - 154 - 154	(זנועיט = מפראטיט ופע	
10 - 1 - 7. 10.11.57.37) KON-DEFINED 7.1104 14 ANUSTRONS, 11 - 1.00 11 - 0.0	0 010280 84 5.000020%	TT222.0 = 109	גנזאינים - ניפאאניעפ דצ	
RISCICION IN RESERVED 504-0611410	0444414 0 ,54 = 0.0000151	11100.1 - 129	451 SUNATUR = 1.23615	
ניינ ני . נוֹזבנרגינו ייי אטע-טבוראונס פטבנרנמא נא אאנקינטאיי, אנו = זיט וני טיט	6110010 - 3.8 5110010 - 3.8		י פינ גזמאזנט - וייזנטי	
402 CL COR (KRESSE) +++ KOK-355 CREO +++ C K 2 C(RRASSE) +++ KOK-355 CREO	115555 = 11 0148950 0	22 - גיפשעאג	גיער מלמונגבים = גיפעלאב	- - - - - - - - - - - - - - - - - - -
POSITION IN ANGSTRONS, XI = 0,00 POSITION IN ANGSTRONS, XI = 0,000 POSITION	034010 0 0000143 2 0 0000143	53860°L = ISA	PST SCHARED = 1.5:002	
●●● I M S L(UERTSI) ●●● 40K-DEFINED FOSIFION IN ANGSTROAS, XI = 4.50 XI = 0.0	ржжаіс о Элсодне 0 вкжа	154 = 0.4420	PJT 303420 = 0.4463	
••• I # 5 L(VEATST) ••• HON-DEFISES POSITION IN AAGSTRCMS, XI = 5.00 YI = 0.0	DAMBIN 0 PAMEN 0 PAMEN 0	7uu9.0 = 124		

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APPENDIX C

CALCULATION OF A SINTERING HISTORY

Chapter 5 outlines the equations which are postulated to describe the change in individual particle sizes during sintering. The SINTR programs were used to solve these equations, for the cases described in Chapter 5.

SINTR 1 was used for α/S_0 values $<10^{12} \text{ m}^{-2}$, where a trial and error solution is needed because F_s , the number of free surface atoms, is significant. In the case of $\alpha/S_0 > 10^{12} \text{ m}^{-2}$, F_s could be approximated as zero, a trial and error solution was no longer necessary, and SINTR 2 was used.

Both programs are reproduced here, along with a sample output from SINTR 2 which directly follows the listing of SINTR 2. Particle sizes for every tenth time increment were printed. The sample output shows only one of these complete particle size outputs after 1 hr. Computations were continued for 50 hrs (500 time increments) or until only one particle remained.

- 215

RE TO BE ADDED TO THE DRIGINAL DISTRIBUTION OF 121 PARTICLES 3 GIVES ANNTHER RUN WITH - USUALLY THIS CARD IS BLANK UNLES SOME LARGE PARTICLES REPEAT DF CARD A BLANK A NEW PSD. NTHFRWISE NY(I) = NUMBER OF PARTICLES'IN FACH SIZE INCREMENT SIZE DISTRIBUTION CUBE J STARTS A NEW RUN. = TOTAL NUMBER OF TIME INCREMENTS (1110) (THIS IS THE SIZE ALONG THE EDGE OF SIZE OF PARTICLE SIZE JNEREMENT (A) 4 SURFACE AREA PER PT ATOM (SO.A) F BLANK THEN THE NEXT CARD SOULD BE SIZE OF SMALLEST PARTICLE (A) PRE-EXPONENTIAL FACTOR (1/SFC) TIME INCREMENT (SEC) = ACTIVATION ENFRGY (CAL/MOLE) INFORMATION ON INITIAL PARTICLE CARD CAUSES THE PROGRAM TO EXIT. AMOTHER CARD FORMATED LIKE CARD = ACCOMMODATION COFFFICEINT THE CARD'I AND 2 COMDITIONS BUT DIMENSION NY(25),P(300),SS(300),DN(300) WIMBER GIVES PRIZZI, PIIZ3), ETC. NI = NUMRFR DF SIZE INCREMENTS (4F10.1,2F10.2) A CARD FORMATED LIKE CARD 1 FORMAT FOR CARD 4 - (2F10.4) FORMAT FOR CARD 3 - (2713) INSERT A BLANK CARD. TEMPFRATURE (K) FORMAT FOR CARD 2 SIZE NF OR BE A RLANK * * * SINTR I PROGRAM TUDATA INPUT н 11 NIIN n NEWS 11 ACC " ∀ ΔS 10 AC ŝ C3=14.32 C4=7.84 I I I t ŝ \sim 4 ŝ CARN 6 \sim CARD CARD CARN CARD CARD CARN 公子 C C $\cup \cup \cup$ S \circ

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۵ GENERATION OF INITIAL PARTICLE SIZE DISTRIBUTION (TO 2). FACH INLIM = MAYMUM ENWVERGENCE ERRAR IN MUMBER OF DIMIS HERRE PRAGRAM PRINEFFINS IN WEXT TIME INCREMENT. READIS, 56801 20, (P(1) , (=122, CP REARCES IIIS SMIN, DS , DISPI, NPSD READ(5,110) MI, INY(1), 1=1,N1) PARTICLE HAS & DIFFERENT SIZE. 1F(DT-0.0011999,999,115 FORMAT(4 FIG. 1, 2 FIG. 2) 1717 56801 FURMAT(3FIG.4, IIIG) FURMAT(11110, 7F13.0) IF (NI 1 114, 114, 113 RFAD15 5551 NTTN DON 2 KK=1, NI NP NY (KK) FURMATI2713) READ IN DATA CCINT 5 11155 CONTINUE CONTINUE DTC=DTC AC=-AC TUTN=D. D1=01C X Y Z = J 24 X=1 0=V 555 112 511 113 110 222 5680 4 . ω \mathcal{O} \checkmark ω υÓ $\omega \, \underline{\mathcal{O}} \, \omega \, \overline{\mathcal{O}}$

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217

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LOOR ACTIVATED IF LARGE PARTICLES ARE ADDED TO DISTRIBUTION (TO 3681). SAS = TÖTAL SURFACF ARFA DF SUPPORT, SO. CM. P(I)=0.5*(1.+(1.+(SI-2.77)/1.96)**3) TOTN=TOTN+P(I) SS(I] = SITE OF PARTICLE 2. ANGSTROMS. P(I) = NUMBER OF ATOMS IN PARTICLE 2. S(=Sw(w+ ([]-1.+SUS+ []+J-]) >) 3/*US ·S=0 •5*(1•+({2•*p(1)-1•)**0•3333)) FORMAT('1', 5X, 'SINTERING MODEL'; ġ PRINT OUTPUT HEADINGS (TD 40) ê r IF(LP-1))3681,3681,3682 SS(I)=(S-1.)*3.92+2.77 SAS=TUTN*SA*1.0F-16 DD 3683 I=122, LP ÝY=(T-300.)/100. TOTN= TOTN+ P(I)00 2 KP=1,NP WRITE(6,103) IS-=(I)SS CONTINUE CONTINUE l + l = l 7 Y = NP 7X=X5 Z Z = K K L=LP T1=T L =] 3681 3683 3682 103 \sim ပပပ္ပ \mathcal{O} S C S ပ ပ ں ن ں

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FORMAT(/,5X, OISTRIBUTION NUMBER, 115,10X, 'INITIAL DISPERSION, 1F FORMATI5X, PRF-FXPOMENTIAL FACTOR 11/SEC) -. JE9.2.11.5X, ACTIVAT IION ENERGY IEJR) KELVIN - 1 LEIS.Q. 15K, SURFACE GREA PER DIDIN ISD D FORMAT(5X, 'NUMBER OF ATOMS LEAVING PER PARTICLE IN', IF6.1, ' SEC -FORMAT(/,5X,'TOTAL ATOMS =',1F10.0,5X,'TOTAL ARFA (SQ CM) =', -', IF10.1, /, 5X, 'TIME INCREMENT (SFC) 21 -1.1 1F9. 1.1.5 5K, 4CCOMMONATION COEFICIENT -1.169.21 FORMAT(/, 5X, INITIAL NO. OF PARTICLFS -', 118, /, 5X, FORMAT(5X, INITIAL PARTICLE SIZES (AMGSTROMS)") START OF THE TIME INCREMENT LOOP (TO 60). WRITE(6,403) (SS(I), J=1,L) WRITE(6, 105) A, AC, SA, ACC WRITF(6,109) NPSD, DISPI WRITE16, 401) JNTN, SAS 1'TEMPERATURE (KFLVIN) WRITE(6, 104)L, T, NT FORMAT(5X, 1) FI1.21 AL=DT*A*EXP(AC/T) WRITE(6,40) NT,AL DD 60 J=1, NTIN WRITE(6,402) $l = 1, l \in 10, 1$ TIME =0. 1169.3,/) FN 1=0 . FN=0. DFN=0. DT = DTCDTT = DT21F7.0) 0=00 0=11 110.2) Y Δ= J 109 104 105 403 40 I 152 40 $\cup \cup \cup$

FN.IS THE MUMBER DE FREE SURFACE ATOMS AT THE START DE TIME INTERVAL FNI IS THE AVERAGE MUMBER DE FREE SURFACE ATOMS DURING THE TIME INTERVAL FN2 IS THE NUMBER DE FREE SURFACE ATOMS AT END DE TIME INTERVAL DEN AND DENI ARE THE CHAMGE IN NUMBER DE ERFE SURFACE ATOMS IN TIME PRAVISIAN OF HEATING AND CODLING PERIDDS (TD 227). الح C2=1./SAS*RIR.3*SORT(T)*ACC*1.0E-08 INCREMENT IN ITERATION I AND I+1 ζ IF(J+103-NTIN)227,227,228 IF(300-NNCN)479,476,476 IF(J-101)225,225,226 IF(FN2)330,330,331,331 T=300 •+YY*!YA-1•) FN1=(FN+FN2)/2. Cl=A*EXP(AC/T) TIME=TIME+DT FN2=FN + NFN NOCO=NOCO+ 1 **P**T=0 • **I** * **∩**T T TTO☆I. (=TO Gn Tn 227 CONT I NUE TIMETIME CONTINUE DFNI=DFN CONTINUE CON TINUIF $T_{19} = D_{T}$ FN2=0. NDCD=0 C = 0225 226 228-227 330 331 22 $\cup \cup \cup$ O \mathcal{O} 000000

PARTICLES NUMBERED LESS THAN LL HAVE OISAPPEAREN. CORRECTION TO FN2 FOR NEXT ITERATION IF (ULIM WAS EXCEEDED ITO 605+1) THE METHOD OF OBTAINING THE NEXT GUESS FOR FW2 VARIES OFPENDING OM THE VALUF OF ERROR . CALCULATION OF CHANGE OF NUMBER OF ATOMS FOR FACH PARTICLE, ON(I), CHECK TO SEE IF MATERIAL BALANCE ON TOTAL NUMBER OF ATOMS IS MET. 0N(I)= (C2*FN1*(C3+C4*(2•*P(I)-1•)**0.3333)-C1)*nT FORMAT(5X, ******** ODES NOT CONVERGE ******* SIGN = NET CHANGE NE ATOMS IN ALL PARTICLES. IF(DN(I)+P(I)) 470,470,471 IF (ABERR-ULIM) 301, 301, 302 IF(ERRAR)411,411,412 IF(ICN)800,601,602 LL IS A COUNTER, ALL ABERR=ABS (ERRUR) SIGD = SIGD + DN(I)ERRDR = SIGD+DFNERR I = SIGD + DFNWRITF(6,478) (I) = -b(I)DD 31=LL+L DFN=FN2-FN ERR 1=EAR DR 60 TO 115 60°T0 728 CONTINUE CONTINUE SIGD=0I = 0.0 = 1(TO 3). 479 37 2 478 476 470 471 601 60 2 7 28 m υU 00000 ပပ Q $\cup \cup \cup \cup \cup$

EACH PARTICLE CONTAINING LESS THAN 24 ATONS IS WADE TO BISAPPEAR AND THE THE NUMBER OF ATOMS (M EACH PARTICLE AT THE END OF THE TIME INCREMENT IS IF ENZ NF LESS THAM ZERN (PHYSICALLY IMPUSSIBLE) WAS RFOUIRFN, THE SIZE DF THE TIME INCREMENT WAS NECREASED BY A FACTOR OF 10. FN2=(FN3*ABS(ERRDR)+FN4*ABS(FRR1))/(ABS(ERRDR)+ABS(FRR1)) l LF (P (I) + MW (I) - 14 . 1 30R, 307, 310 CALCULATED IN THIS LODP. IF(ERR1)425,425,414 ATOMS ARE ADDED TO FN. IF(FN2)603,603,604 FN=FN+P11) + MM11) FN1=(FN+FN2)/2. FN1=(FN+FN2)/2. FW L= < FM+ FN2 > 52. FN2=FN2-FRROR I = h(1) + h(1) = (1) d200, 307 1=22,42 DT=07/11. FMZ=FWS2. FN=FN+DFN GN TN 22 GN TN 22 5N 7D' 22 GN TN 317 BN 71 22 FN3=FN2 FN4=FN2 . ()= ([.] NU P111=1. FNZ=0. 425 117 603 604 412 414 405 3.3 1 30.8 310 $\cup \cup \cup \cup$ ~~~~ 0000

CHECK WHETHER SJZF DF TIME INCREMENT HAN BEEN DECPEASED DURING FM2 ITER-THIS LOOP COUNTS ALL PARTICLES THAT HAVE DISAPPEARED , I.E., INSTRUCTIONS TO PRINT IN INCREMENTS OF IN TIME INTERVALS. \$ IF(0T1-1.7F-78)311,312,312 AA 201 1= 44,4 151 011) 1800,211,210 IF(FN2)451,451,452 FN2=FN+DFN]*DT/DT9 IF (P(L-1)) 800,71,60 IF! JJ-10)78,71,78 FN1=(FN+FN2)/2. DTI = TIMF - TIMDETERMINES LL. TIM=TIM+UT CDNT I NULF GO TO 22 CONTINUE CONTINUE 5D 70 15 CONTINUE CONTINUE DT=DT1 FN2=0. 11=11+1 TDTN=0. SUR T=0. I C 0 =0 5=77 *0=CC* . 307 312 6.51 452 311 210 201 27 78 15 \cup \cup U \mathcal{L} $\cup \cup \cup \cup$

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CALCULATION OF PARTICLE SIZES AND DISPERSIONSIONLY DONE FOR THE PRINTING DISI IS THE DISPERSION OF PARTICLES ONLY, NOT TAKING FREE SURFACE ATOMS CORNER LOSS PER PART. ", IF 7.0,5%, SAIN PER 20 & PART." FORMAT(//,5X,'INTAL ATOMS',1F10.0,5X,'SURFACE ATOMS',1F10.0, 15X, FREE ATOMS AT END DE TIME IMC. (, 1610.0.1, 5X, 1015PERSIANS) =', 1F 8.1.5X, TIME INC. =', 1F8.1, SURFACE ATOMS INTO ACCOUNT 2FIG.3, (WITH AND WITHDUT FREE ATOMSI',/I FORMAT(5X, ·******************************** S=0•5*(1•+({2•*P(I)-1•)**0•3333) WRITE(6,32) TOTN, SURT, FN, DIS, DISI GGLL= I • / SAS#0T#818. 3# SDR7 | 7) #AEE WRITE(6,698) T, NT, ACLD, AVGG DIS IS ISPERSION TAKING FREE AV66=CDLL*80 ** 1.97E-98*FM2 DI S= (FW+ SURT | / TOTN* 100. FORMAT(5X, TEMPFRATURE SS(1)=(S-1.)*3,92+2.77 ACLD=A*DT*8.*EXP(AC/T1 PL=12,*S**2-36.*S+30. DI S1=SURT/T0TN*100. ·IF(P(I))800,79,73 SUR T = SUR T+ 8. + E+ PL TDTN=TOTN+P(I) TOTN= TOTN+FN TIME INTERVAL). DD 70 1=1,L F=12.*S-24. WRITE(6,91) SS(I)=0. G0 T0 7) CONTINUE 21F7.0.1 5X , ~ .16 ۶ ۲ 698 73 32 $\cup \cup \cup \cup$ ں $\cup \cup \cup \cup$

TIMES=TIMF/3600. WRITF(6,33) J, TIMES FORMAT(5X, 100. WUMMER, 117,5X, TOTAL FLAPSED FIME (HRS1), IIF10.3,)
WRITE(6,403) (SS(1),1=1,L) 33

INSTRUCTIONS TO GO TO NEXT PSO WHEN ONLY OME PARTICLE REMAINS. ມບບ

IF(P(L-1))R00,115,60 CONTINUE . 60

GU TO 115

WRITE(6,871] 800

FURMATI 5X, 1:*** FRRDR IN PROGRAM #*** 1 666 801

CALL EXIT

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-Alt

DATA INPUT AND INITIAL PSD GENERATION ARE THE SAME AS IN SINTR THIS PROGRAM IS FOR THE LIMITING CASE OF LOSS CONTROLLING THE CONCENTRATION OF FREE SURFACE ATOMS IS ALWAYS SMALL INPUT AND DUTPUT FORMATS ARE PARALLEL TO SIMTR 1. SINTR 2 HAS NO HEATING OR CODLING PERIOD DIMENSION NY (30), SS(300), P(300) READIS, 5690) 2 P. (P(11, [=122, 10 · NISPI, MPSD READ(5,110) NI, (NY(I), I=1,NI) READ(5,112) DT, SA, AC, T, ACC; A IF(DT-0.001)999,999,115 FORMAT(4F10.1,2F10.2) FDRMAT(3FI0,4,1110) FORMAT(1110,7F13.0) RFA0(5,111)SMJN, DS IF(NI)114,114,113 CONTINUJE RFAD(5,555) MTIN SINTR 2 PRIGRAM FORMAT(1110) IF (NP) 2, 2, 183 CONTINUE FORMAT(2713) DD 2 KK=1, MI CONTINUE CONTINUJE NP=NY (KK) SIGS=0. TOTN=0. DTC=DT AC = -ACT=XYZ X Y Z = T $\gamma = NP$ Ŷ 114 555 112 115 113 110 11 5680 183 C S ں ر 0000

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FORMATIJ, SY, NJSTRIANTIAN NUMBER, SIGS, SOSTAL DISPERSION, 25
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         110N ENERGY (E/R) KELVIN -' 1F15.0,/5X,'SURFACF AKEA PER ATOM (SO A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         FURMATISX, PRE-EXPONENTIAL FACTOR (ILSEC) -., 159.2, 11, 5X, 14CTIVAT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  11. PMPERATURE (KFLVIN) -1, IFI0.1, /, SX, 'TIMF INGREMENT (SEC) -1,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ', 1F9.1, /, 5X, 'ACCOMMODATIOM CDEFFICIENT - VERY LARGE')
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 RART(1,5X, "INITIAL ND. DF PARTICLES -", II8, 1,5K,
                                                                                                                                                                                                                                                                       -1.1**0.3333))
                                                                                                              P(I) =0.5*(1.+(1.+(<I-2.77)/1.96)**3)
                                                                         SI = SMJ N+ (ZZ - 1 \cdot ) * NS+ (ZX - 1 \cdot ) / ZY * NS
                                                                                                                                                                                                                                                                                                                                                                                                                                      FORMATS ... 53, . 53 MJ FR 3 MR MUREL . 2
                                                                                                                                                                                                                                                                 S =0.5*(1.+((2.*P(1)
                                                                                                                                                                                                        IF(LP-I0.)3681,3681,3682
                                                                                                                                                                                                                                                                                                                                                                                                                                                        WRITELA, LAGI NPSA, ALSA,
                                                                                                                                                                                                                                                                                  SS([)=(S-],)*3,92+2,77
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               WRITE(6,401) TOTN, SAS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       WRITE(6, 105). A, AC, SA
                                                                                                                                                                                                                                                                                                    5165=5165+ SS<1+2.7
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              79,75,16,20472,7,07
                                                                                                                                                  SIGS=SIGS+SI+2.77
                                                                                                                                                                                                                                                                                                                                                              245=21774#544 245
                                                                                                                                                                                                                                              NG 3683 I=122,LP
                                                                                                                                                                                                                                                                                                                                                                                44=1J-300. 1200.
                                                                                                                                (I) = TUTN+P(I)
                                                                                                                                                                                                                                                                                                                       507W=707W+ P13
                  DO 2 KP=1,NP
                                                                                                                                                                                                                                                                                                                                                                                                                  WR17F16, 1731
                                                                                           iS = (I)SS
                                                                                                                                                                                       CONTINUE
                                                                                                                                                                                                                                                                                                                                          CUM 75 MUS
                                                       7 X = K P
                                      [ + ] = [
77 = KK
                                                                                                                                                                                                                           (=[p
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (0.7.1.
                                                                                                                                                                                                                                                                                                                                                                                                   2 = 72
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               12.021
                                                                                                                                                                      109
                                                                                                                                                                                                                                                                                                                                        3682
                                                                                                                                                                                                                                                                                                                       3683
                                                                                                                                                                                                                          3682
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                                                                                                                                                                                                                                                                                                                                                                                                                                      KQ 3
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2<mark>,2</mark>7

FORMAT(5X, INUMARE OF ATOMS LEAVING PEP PARTICLE IN', IF6.1,' SEC ATOMS =', JF10.0,5%, TOTAL AREA (SO CW) START DF TIME INCREMENT LOOP (TO 60). NO TRIAL AMD FRRDR CALCULATION NEEDED FOR THIS LOOP. FORMAT(5X, 'INITIAL PARTICLE SIZES (AMGSTROMS)'I WRITE(6,403) (SS(I),I=1,L) AG=SIGL*(SS(I)+2.77)/SIGS SIGS= SIGS- SS(I)-2,77 FORMAT(5X, 12FI0.2) IF(PA- (.)]),11,11 AL=D (* A*FXP(AC/T) WRITE(6,4) DT,AL FORMAT(/, 5X, CT) AL=DT*A*EXPIAC/T) SIGL = SIGL = SIGL = AL + PIIIF(P(I))800,3,4 DD 6D J=1,NTIN PA=P(I)-AL+AG WRITF(6,402) TIME=TIME+NT SIGL=TUTP*AL DD 3 I=1,L I = 1, IFIO, 1) SIGS1=0. 71NF=D. CONTINUE 10110-01 · SS(])=0. llF9.3,/) דח דף= ננ SURT=n. NTGT=0. • U=(1)d ノーノ I C = 0しか 40 2 403 40 <u>г</u>2 S 4 0000

: 1

S=0.5*(].+((2.*PA

CONTINUS

11

60 JN 3

r

IF(IC)12,12,13

60 TO 14 SS(I)=SI

3

<1

P[])=PA

4

(14%0,3333))

SIGSI=SIGSI+SS[1)+2.77 SI=(S-1.)*3.92+2.77 SS(I)={SS(I)+S1)/2.

CONTINUE

ÍF(IC)16,16,17 IF (P(1))R00, 20 DD 20 1=1,L SAGS = SIGS160 TO 15 I C= 1 2

CONTINUE 20 22

IF(LL-1)800,71,23 11=00+1

[F(JJ-1))60,71,60 CONTINUE 23

T01N=0. SURT=0. JJ=0

 $DD \cdot 70 I = 1 \cdot 1$

S=0.5*(1.+((2.*P(I)-1.)**0.3333)) ,79,73 IF(P(I)80)

73

SS(1)=(S-1.)*3.92+2.7

E=12.*S-24.

PL=12.*S**2-36.*S+30. -SURT=SURT+8.+E+PL

TOTN=TOTN+P(I)GO TO 70

FARMAT(//, 5K, 7 TATAL ATAMS', 1F10.0, 5%, 'SWRFALE ATAMS', 1F10.0, "DTAL ELAPSED TIME (HRS)", FURMAT(5X, TEMPERATURE = , IF 8.1,5X, 'JIME INC. =', IFR.11 15%, "REMAINING PARTICLES, LIS, L, 85K, "UISPERSION", 1F10.3> FORMAT(5X, **** FRRDR IN PRDGRAM ***) FORMAT(5X, 'INC. NUMBER', 117,5 WRS 7516, 32) JUN, SURT, 22, D752 WRITE(6,403) (SS(1),I=1,L) WRITE16, 33) J, TIMES IF(P(L-1))800,115,60 5) 52=SURTJTRTN# 200. 10,1 TIMES=TIME/3600. WRITF(6,698) WRITE(6,801) GO TO 115 CALL EXIT 11F10.3,// SS([)=0. CAN TINKIF CONTINUE END 864 666 8() I 800 202 $\frac{3}{2}$ 6 60 33

REMAINING PARTICLES 106 29.62 30.17 30.52 30.52 31.42 27.85 28.57 29.07 880.9 35.00 15.71 74.75 71.44 71.75 71.75 33.93 33.93 33.93 33.93 33.93 33.93 33.93 33.93 33.93 33.93 00.0 28.52 29.03 29.19 29.19 30.14 30.P9 31.38 32.07 1.000 THTA (ARFA 150 CM1 =0.201E-08 INITIAL DISPERSION 0.00 78.51 78.51 78.51 75.52 75.52 99.05 81.61 8 29.41 29.41 30.11 30.85 31.33 31.33 27.66 28.47 28.99 65954 . ATTMS LEAVING PER PARTICLE IN 360.0 SEC 41HE INC. = 360.0 TOTAL FLAPSED TIME (HRS) -15000. Ç. [wittal wo. of particles - 12] tevetatios (kelun) - 1003.0 twee (velun) - 350. pre-exponential factor (1/560) - 0.80F 07 27:57 28.42 28.95 29.38 29.74 30.08 30.42 30.82 31.28 31.92 URFACE ATOMS ACTIVATION EMFRGY (E/P) KELVIM -SURFACF AREA PER ATOM (SO A) - 100.0 ACCOMMONATION COEFFICIENT - VERY LARGE INITIAL PARTICLE SIZES LANGSTRONS 27. 12 28. 32 19. 82 19. 82 17. 92 17. 92 17. 92 29. 06 30. 06 31. 25 31. 85 *************** TDIAL ATOMS = 201105. 2011105 1000.00 0.00 0.00 22,55 28,59 28,89 332,51 332,51 332,58 332,58 332,58 332,58 DISTRIBUTION NUMAFR 1 27.28 28.81 29.81 29.83 29.68 30.02 30.74 31.79 31.79 9 SINTERING MUDEL JEMPERATURE = TOTAL ATONS * * * * * * * * * * * 0.0 69.15 69.15 60.45 61.45 61.45 61.45 61.45 61.45 61.45 61.45 61.45 61.45 61.45 61.45 27.00 28.25 29.83 29.65 29.99 37.34 30.71 31.16 31.74 32.99 NUMBER OF

30.58 30.99 31.52 32.33

29.53 29.88 30.22

29.10 29.50 30.19 30.55 30.55 30.55 31.47

0.00 75.75 75.65 78.55 78.55 31. 83 34. 72 34. 66 36. 77 38. 77 24.92 29.92 14.05 14.05 84.15 84.15 84.15 84.15 29.59 29.94 30.78 35.12 30.65 31.08 31.62 32.57 25.79 12.02 30.21 33.11 0.00 70.06. 75.49 78.49 31.60 31.60 31.60 31.60 31.60 31.49 35.97 35.13 30.61 29.47 29.57 29.91 30.25 32.42 32.795

DISPERSION

28.79 29.24

28.14 28.75 29.21

7 U. A Z

27.99 8.66 9.14

29.72 28.61

28.19

0.00 11.81 11.81 11.81 11.85 11.85 11.85 11.85 11.85 11.85 11.85 11.85 11.85 11.85 11.85 11.81 1

0.00 25.19 25.19 25.19 25.19 25.19 33.48 33.48 33.48 33.48 33.48 33.90

38.45

0.00 24.55 24.55 21.65 21.65 31.65 337.66 332.66 332.65 35.57 37.45