Lawrence Berkeley National Laboratory

LBL Publications

Title

HYDROGENATION AND H, D EXCHANGE STUDIES OF ETHYLIDYNE (CCH3) ON Rh(III) CRYSTAL SURFACES AT 1 ATM PRESSURE USING HIGH RESOLUTION ELECTRON ENERGY LOSS SPECTROSCOPY

Permalink

https://escholarship.org/uc/item/19d4c47h

Authors

Koel, B.E. Bent, B.E. Somorjai, G.A.

Publication Date

1984-04-01



UNIVERSITY OF CALIFORNIA

Materials & Molecular Research Division

LAWRENCE BERKELEY LABORATORY

MAY 30 1984

18-17410

LIBRARY AND DOCUMENTS SECTION

Submitted to Surface Science

HYDROGENATION AND H,D EXCHANGE STUDIES OF ETHYLIDYNE (CCH₃) ON Rh(111) CRYSTAL SURFACES AT 1 ATM PRESSURE USING HIGH RESOLUTION ELECTRON ENERGY LOSS SPECTROSCOPY

B.E. Koel, B.E. Bent, and G.A. Somorjai

April 1984

TWO-WEEK LOAN COPY

This is a Library Circulating Copy which may be borrowed for two weeks. For a personal retention copy, call Tech. Info. Division, Ext. 6782.

Prepared for the U.S. Department of Energy under Contract DE-AC03-76SF00098

DISCLAIMER

This document was prepared as an account of work sponsored by the United States Government. While this document is believed to contain correct information, neither the United States Government nor any agency thereof, nor the Regents of the University of California, nor any of their employees, makes any warranty, express or implied, or assumes any legal responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by its trade name, trademark, manufacturer, or otherwise, does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof, or the Regents of the University of California. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof or the Regents of the University of California.

HYDROGENATION AND H, D EXCHANGE STUDIES OF ETHYLLDYNE (CCH₃) ON Rh(111) CRYSTAL SURFACES AT 1 ATM PRESSURE USING HIGH RESOLUTION ELECTRON ENERGY LOSS SPECTROSCOPY

B.E. Koel*, B.E. Pent, and G.A. Somorjai

Materials and Molecular Research Division Iawrence Berkeley Iaboratory and Department of Chemistry University of California, Berkeley Berkeley, CA 94720, USA

*Miller Institute Postdoctoral Fellow

1

Present Address: Cooperative Institute for Research in Environmental Sciences, and Department of Chemistry, University of Colorado, Boulder, CO 80309

HYDROGENATION AND H,D EXCHANGE STUDIES OF ETHYLIDYNE (CCH₃) ON Rh(111) CRYSTAL SURFACES AT 1 ATM PRESSURE USING HIGH RESOLUTION ELECTRON ENERGY LOSS SPECTROSCOPY

B.E. Koel, B.E. Bent, and G.A. Somorjai

Materials and Molecular Research Division Iawrence Berkeley Laboratory and Department of Chemistry University of California, Berkeley Berkeley, CA 94720, USA

ABSTRACT

We demonstrate that combining high resolution electron energy loss spectroscopy (HREELS) with a high-pressure/low-pressure (HPLP) system is a powerful method for studying the behavior of the monolayer structure of stable hydrocarbon species that form on single-crystal metal surfaces during catalytic reactions at atmospheric pressure. We find that a monolayer of adsorbed ethylidyne (CCH₃) on Rh(111) at 310 K does not hydrogenate to ethylene or ethane in one atmosphere of static D_2 . The methyl group hydrogens exchange with deuterium in a slow process. The amount of exchange depends strongly on the amount of uncovered, bare-metal surface, but little on the hydrogen pressure. A mechanism for H,D exchange involving ethylidene (CHCH₃) as an intermediate is proposed.

1. Introduction

Understanding how heterogeneous catalysts work requires knowing the structure and reactivity of the adsorbed overlayer at the catalyst surface. Evidence for overlayer structure and catalytic reaction intermediates (not necessarily the same) has been obtained indirectly, mostly from kinetic studies, trapping intermediates at low temperatures, and from chemisorption and thermal desorption measurements in ultra-high vacuum (UHV). In order to utilize surface science techniques that require UHV for the determination of monolayer structures that form during high pressure catalytic reactions, we have incorporated high resolution electron energy loss spectroscopy (HREELS) in a high-pressure/low-pressure (HPLP) system.

The HPIP system makes surface analysis of a catalyst (by HREELS as well as low energy electron diffraction (LEED) and Auger electron spectroscopy (AES)) possible in UHV prior to and immediately following (within minutes) a high pressure gas exposure or catalytic reaction. By detecting surface vibrations, HREELS is a sensitive probe of surface species.^{1,2} Surface periodicity is not required, but single crystal surfaces are used as well-defined model catalysts. The vibrational spectrum is a fingerprint of monolayer structure; the vibrational frequencies are a function of atomic configurations and bond strengths. We have begun using HREELS in combination with a high pressure (HP) cell to detect stable surface species after atmospheric pressure reactions and to follow hydrogen (deuterium) transfer and reactivity in hydrocarbon monolayers under catalytically interesting conditions. HREELS is ideal for hydrocarbon monolayer structure determination due to its sensitivity to carbon-hydrogen vibrations.

We have used the HREELS/HPLP method to study the reactivity of an

-2-

ethylidyne (CCH₃) monolayer on the Fh(111) crystal surface with 1 atm. of H₂ and D₂. This monolayer is the stable room temperature structure for chemisorbed ethylene on Rh(111)^{3,4} as well as Pt(111)^{5,6} and Pd(111).⁷ Our results show the stability of this monolyer extends to atmospheric pressures. One atmosphere of hydrogen does not rehydrogenate this ethylidyne to ethylene or ethane on Rh(111), but the methyl group hydrogens do undergo H,D exchange. The rate of H,D exchange depends little on D₂ pressure but strongly on the amount of available bare metal surface.

2. Experimental

The design considerations and specifications of this⁸ and other^{8,9} HPLP systems have been described. A schematic diagram of the two-tier HPLP system with HREELS used here is shown in Figure 1. The HREEL spectrometer design and operation are described elsewhere.¹⁰ We describe here the procedure for the use and performance of the HREEL spectrometer with the high pressure cell.

Both the upper and lower portions of the HP cell translate with welded bellows seals. The crystal translates on-axis with the upper portion of the HP cell to the HREEL spectrometer by an extended-travel manipulator. To seal the HP cell, the upper bellows are contracted so that the upper half of the HP cell locks against the top of the chamber. A hydraulic piston behind the lower half-cell drives and pressurizes a knife-edge seal between the two halves to 2100 psi. The volume of the HP cell together with the external circulation loop is about 0.5 L.

The Rh(111) single crystal was a 0.5 cm² disk, .030" thick, polished on both sides by standard methods.³ The crystal was mounted with 0.020" Pt wire spotwelded between the crystal edges and 1/8" Ta rods. Crystal temperatures

-3-

between 300 and 1500 K were obtained by resistive heating, monitored by a chromel-alumel thermocouple spot-welded to one face of the crystal. Crystal cleanliness was checked by HREELS and AES. The cleaning procedure has been described elsewhere;¹⁰ after heating in O_2 , it was found necessary to Ar⁺ sputter in order to remove trace amounts of boron oxides detectable by HREELS (740 cm⁻¹). Annealing to 1400 K in vacuum dissolved trace amounts of carbon into the bulk.

Gases used as supplied were: Matheson H₂, 99.9%; D₂, > 99.5 atom %; C₂H₄, 99.5%: Merck, Sharp and Dohme C₂D₄, 99 atom %. Gas dosing was performed with a microcapillary array doser in UHV or by pressurizing the HP cell. Exposures in UHV are given uncorrected for ion guage sensitivity, but corrected for a five times enhancement of the doser. Gas exposures (pressures) in the HP cell were measured with a Heise guage (absolute) and were generally 1 atm of static gas.

Gases are pumped from the HP cell through a rapid-pump value at the manipulator top with a liquid nitrogen-trapped, 2" diffusion pump. Pump-down is typically ten minutes or until the ion guage pressure above the liquid nitrogen trap is 1×10^{-6} torr. Upon opening the HP cell, outgassing of the cell causes the chamber pressure to rise to about 4 x 10^{-8} torr, water being the major contaminant. A pressure of about 5 x 10^{-9} torr is reached after 15 minutes, but usually it is several hours before the chamber returns to the base pressure (2 x 10^{-10}).

After retraction of the HP cell, the crystal is lowered into the HREEL spectrometer and precisely positioned using travel guages to previous settings. In these H,D exchange reactions where surface structure is largely unchanged, a minimum of spectrometer retuning was required and a spectrum begun within

-4-

ten minutes of lowering the cell. Changing the crystal bias to compensate for work function changes was the only tuning usually required. Elastic scattering count rates in this work varied from 0.3 to 1.5 x 10^5 counts per second depending largely on the degree of surface order. Resolution (full width at half maximum of the elastically scattered beam) was 45 - 65 cm⁻¹. No major changes in the work function of the graphite-coated lenses after repeated HP work was found. Over 40 high pressure exposures of H₂, CO, and C₂H₄ have been made without spectrometer bake-out.

3. Results and Discussion

3.1 Ethylidyne

The adsorption of ethylene on Rh(111) at room temperature to produce surface ethylidyne (CCH₂) species (Figure 2) has been well-studied previously.^{3,4} Figure 3 shows the HREEL spectra for saturation coverages (saturation coverage = 1 $CCH_3/4$ Rh atoms or 1 C/2 Rh atoms) of CCH₃ and CCD₃ on Rh(111) produced by a 15 L exposure at 310 K to C2H4 or C2D4. These spectra are consistent with those found in previous HREEL studies, 4 but offer improved resolution. The assignments of the loss peaks to vibrational modes of surface ethylidyne are given in Table 1, in complete agreement with frequencies, intensities, and H/D shifts observed in infrared spectra (and normal mode analysis) of $(CH_3C)Co(CO)_9$.¹¹ Except for the degenerate methyl group rock (pCH₂), the antisymmetric methyl hydrogen bending $(\delta_{as}CH_3)$, and the antisymmetric methyl stretching (v_{as} CH₃) modes, the vibrational modes are dipole active (A₁ type), establishing C_{3v} symmetry for the surface complex. Angular-dependent HREELS studies reported elsewhere¹² establish the dipole contribution to the loss peaks and the surface symmetry.

-5-

3.2 Ethylidyne + 1 atm. H₂

These experiments show the remarkable stability of an ethylidyne monolayer. The HREEL spectra of a saturation coverage of ethylidyne on Rh(111) before and after exposure to 1 atm H₂ for 5 min. at 310 K are shown in Figure 4. The spectra are nearly identical. There is no evidence for decreased ethylidyne coverage by rehydrogenation to ethylene or ethane. Ethylidyne is even more stable towards rehydrogenation on Rh(111) than Pt(111) where 1^{4} C radiotracer studies¹³ have shown some hydrogenation and removal of surface ethylidyne (turnover rate < 10^{-4} / Pt atom \cdot sec) under these conditions. Preliminary HREELS studies on Pt(111) support this conclusion.

Figure 4 also shows no evidence for contamination by other molecules that may coadsorb during high pressure gas exposure. Contamination from hydrocarbons and CO in the ambient was expected. On clean Rh(111) during 1 atm H₂ exposures, hydrocarbon contaminants from the HP cell loop adsorb dissociatively on the surface, giving fingerprint HREELS peaks at about 800 and 1400 cm⁻¹. Similar features were found in a recent study of surface structure following Fischer-Tropsch synthesis on Fe(110) and assigned to CH and C_xH fragments.¹⁴ But Figure 4 shows no evidence for formation of such surface species in this case. Figure 4 also shows no change in the relative intensity of the contaminant CO (< 5% of a monolayer) peaks at 1938 and 1745 cm⁻¹, further evidence that the ethylidyne coverage doesn't change.

The stability of an ordered c(4x2) ethylidyne monolayer towards hydrogenation on Rh(lll) was unexpected for such an open overlayer. Even exposure of a saturation coverage of CCD₃ to 10 torr of C₂H₄ failed to produce any detectable CCH₃ by coadsorption or replacement. We conclude that C₂H₄ cannot

-6-

reach the surface and dissociate. The methyl group packing is shown in Figure 5. There does indeed appear to be room for H₂ but not C_2H_4 to reach the surface and dissociate. TPD studies show that about 1/4 monolayer of H atoms can be coadsorbed in a similar CCH₃ saturated monolayer on Pt(111) by exposure to 1 atm of H₂.¹⁵ Apparently the strong metal-carbon bonds prevent complete hydrogenation of the carbyne carbon. Partial hydrogenation of the carbyne carbon is, however, likely based on the LEED studies reported next and the H,D exchange reported in section 3.3.

Previous LEED studies³ showed that careful annealing of the adsorbed monolayer produced by exposure of Rh(111) to ethylene at 240 K could produce a well-ordered c(4x2) CCH₃ structure. Exposure of Rh(111) to 15 L C₂H₄ at 310 K does not produce this structure, but instead the one shown in Figure 6(A). The diffuse intensity of the (1/3,1/3) positions is sometimes resolved to a triangle of spots, but the unit cell for this partially-ordered phase remains unexplained. Exposure of this surface to 1 atm of H₂ causes a two-dimensional phase transition from the structure characterized by the LEED pattern in Fig. 6(A) to a structure with a sharp c(4x2) LEED pattern as shown in Fig. 6(E). This transition occurs without significant change in the coverage of ethylidyne or contamination by other species, since the HREEL spectra are identical for the two phases.

One plausible explanation for the influence of 1 atm H_2 on the ordering of the ethylidyne monolayer is the partial hydrogenation of the carbyne carbon atom to form an ethylidene species (CHCH₃) that is stable under high pressures of H_2 . Breaking one of the carbon-metal bonds would greatly decrease the fragment-metal bond energy allowing the bridge-site ethylidene species greater mobility on the Rh(111) surface. Removal of the high pressure of H_2 would

-7-

result in dehydrogenation to give ethylidyne, in the most stable, well-ordered two-dimensional phase.

Precedence for the formation and fluxionality of ethylidene-like species comes from organometallic chemistry. H,D exchange in a tricobalt biscarbyne¹⁶ (<u>1</u>) (cyclopentadiene ligands not shown here) has been proposed to proceed by protonation at the carbyne carbon to give the fluxional intermediate

2:

followed by deprotonation to give the exchanged species.

Unfortunately, we have not been able to detect adsorbed hydrogen on Rh(111) by HREELS. HREEL spectra of a CCH₃ monolayer after treatment with 1 atm H₂ or of clean Rh(111) in UHV with 10^{-7} torr background H₂ or D₂ show no peaks obviously assignable to M-H or M-D vibrations. Previous temperature programmed desorption (TPD), however, suggests that H atoms are present on the surface in both cases.^{17,15} The absence of a detectable Rh-H stretch by HREELS is partially due to the covalent nature of the Rh-H bond. Previous study of H₂ adsorption on Pt(111) has shown the difficulty of detecting the Pt-H vibration.¹⁸

3.3 Ethylidyne + 1 atm. D_2

3.3a HREELS Peak Assignment

In addition to the stability of ethylidyne towards rehydrogenation or contamination by replacement or coadsorption, we have studied ethylidyne H,D exchange. HREELS is a sensitive probe for identifying the partially deuterated ethylidyne species at the surface. The symmetry of ethylidyne is lowered from C_{3V} to C_S on partial deuteration. This lower symmetry means more vibrational modes are dipole active and expected to have large scattering amplitudes in the specular direction in HREELS. Besides this symmetry reduction, partial deuteration requires a new approximate normal mode description because of the large percentage mass change. The methyl group modes are now best described as a combination of $CH_2 + CD$ or $CD_2 + CH$ vibrations.

Exposure of a saturation CCH₃ monolayer to 30 L D₂ at 310 K gave the HREEL spectrum shown in Figure 7(A). Indeed two new peaks characteristic of methyl group H,D exchange are seen at 1246 and 1407 cm⁻¹. (The 806 cm⁻¹ peak is due to contamination in the original CCH₃ monolayer.) Neither peak is present in the spectrum of CCD₃ and therefore must be the result of partial exchange. However, less than 20% completely exchanged methyl groups would be hard to detect in this experiment, because of the weak carbon-deuterium dynamic dipole (see Fig. 3(B)). Treatment of a similar ethylidyne monolayer with 1 atm D₂ for 5 min. at 320 K gave the HREEL spectrum shown in Figure 7(B), which differs little from the spectrum in (A). The amount of H,D exchange of a saturation coverage of ethylidyne in 1 atm D₂ is surprisingly insensitive to D₂ pressure.

We also followed H incorporation in deuterated ethylidyne. Treatment of CCD_3 with 1 atm H₂ for 2 min. at 310 K, as shown in Figure 7(C), results

-9-

in new peaks at 1246, 1300 - 1450, 2920, and a small increase in the peak at 780 cm⁻¹. The absence of a peak at 1340 cm⁻¹ for the intense symmetric methyl deformation (δ_8 CH₃) mode means there are few completely exchanged methyl groups.

Peak assignments are made by comparison with variously deuterated methyl chlorides^{19,20} whose peak frequencies and approximate mode descriptions are shown in Figure 8. The v_{C-C1} modes have been replaced with expected C-C stretching frequencies of ethylidyne for comparison with HREEL spectra.

Definitive peak assignments in the partially deuterated ethylidyne spectra are hampered by the presence of mixtures of variously deuterated species on the surface; at a resolution of 50 cm⁻¹, peaks due to the various CCH_xD_y species overlap (as can be seen from the shaded modes in Figure 8). However, by comparing many HREEL spectra of variously deuterated ethylidyne mixtures with those of the partially deuterated methyl halides, 19,20 we are able to make in Table 2 tentative peak assignments of partially-deuterated ethylidyne.

In the case of CD_2H , comparison is also made with a normal coordinate calculation by Skinner et al.¹¹ on $(CCD_2H)Co(CO)_9$. Similar calculations with the same force constants and geometry on $(CCH_3)Co(CO)_9$ and $(CCD_3)Co(CO)_9$ gave good agreement with adsorbed ethylidyne frequencies. The agreement here is best with the δ_sCH , the least overlapping peak. The most distinctive feature of both partially-exchanged species is a band at about 1250 cm⁻¹. This frequency corresponds to a CH_2 wag for the monodeuterated species and a CH in-plane bend for the dideuterated species.

-10--

3.3b H,D Exchange Rate

We have also studied the H,D exchange rate of ethylidyne. Some results from these studies are shown in Figure 9. Fig. 9(A) (like Fig. 7(E)) shows deuterium exchange in CCH₃ at room temperature for 5 min. at a saturation coverage of ethylidyne. Figure 9(E) shows the vibrational spectrum after the same monolayer is additionally treated with 1 atm D₂ for 2 min. at 350 K. Raising the temperature 40 K still does not cause any significant change in the total ethylidyne coverage (as also monitored indirectly by CO adsorption), but increases the amount of exchange, indicated by the growth of the 1246 and 1432 cm⁻¹ peaks. Exposing this D and CCH₃ covered surface at 300K to UHV for 36 hours results in additional exchange between the remaining adsorbed deuterium and partially exchanged ethylidyne, as shown in Figure 9(C).

It is difficult to quantitatively determine the amount of H,D exchange from the HREEL spectra even with the peaks assigned. The major problem is quantifying the dynamic dipoles perpendicular to the surface (that should be proportional to dipole scattering peak intensity) for the different molecules in the isotopic mixture CCH_xD_y . Because of different dynamic dipole strengths, and overlapping peaks, the peak heights don't change linearly with the concentration of the various molecules in a particular CCH_xD_y in a mixture. However, using the CH_2 wag dynamic dipole equal to the C-C stretch dynamic dipole²¹ and comparing the intensity of the 1120 and 1246 cm⁻¹ peaks (assumed to be all CH_2 wag for small amounts of exchange), we estimate using Figure 9(A) that five minutes of 1 atm D₂ over CCH_3 gives 20% exchange (#D atoms/3 x # methyl groups).²² This value corresponds to methyl group H replacement at a turnover rate of < 10⁻³ / Rh atom \cdot sec. This rate is faster than ethylidyne rehydrogenation. We are currently using temperature programmed desorption to help confirm this estimation.

-11-

Subsaturation coverages of CCH₃ exchange much more rapidly. Reaction of < 50% saturation coverage (1.5 L C₂H₄ exposure) CCH₃ with 30 L D₂ is shown in Figure 10. The presence of the δ_{s} CD₃ peak at 984 cm⁻¹ and the nearly complete loss of the δ_{s} CH₃ at 1340 cm⁻¹ indicate complete exchange in some methyl groups. Some partially exchanged ethylidyne is still observed as evidenced by the 1218 cm⁻¹ peak.

3.3c H,D Exchange Mechanism

The spectra in Figures 7 and 9 with peaks assigned as in Table 2 can be used to show that deuterium atoms are incorporated in the CCH₃ methyl group one at a time. Compare the intensities of the 1246 and 1430 cm⁻¹ peaks. The 1430 cm⁻¹ peak is characteristic of the CH₂ scissor in CH₂D. It is the same intensity as the 1246 cm⁻¹ CH₂ wag when deuterium exchange first begins, as seen in Figures 8(A),(B) and 9(A),(B). With continued exchange, the 1246 cm⁻¹ peak grows in relative intensity (Figure 9(C)). Since neither the 1246 nor 1430 cm⁻¹ peak is present in CCH₃ or CCD₃, the change in their relative intensities implies two partially-deuterated species. As CD₂H forms the 1430 cm⁻¹ CH₂ scissor has a smaller intensity relative to the 1246 cm⁻¹ CH₂ wag that gains a contribution from the δ_8 CH.

The presence of both CCH_2D and $CCHD_2$ and the gradual conversion of one to the other rules out the exchange mechanism whereby one molecule of D_2 always replaces two H atoms in the methyl group by a concerted exchange. The mechanism for the process that we monitor is single atom H,D exchange. Furthermore, there was no evidence from the HREEL spectrum of complete methyl group deuteration by some rapid exchange process at selected sites after 5 - 10 min. at 310 K under 1 atm D_2 .

-12-

These data are consistent with the exchange mechanism proposed on $Pt(111).^{23}$ In this mechanism, the role of the metal surface is both to atomize H₂ and to help break the methyl group CH bond. The rate-limiting step is probably tipping of the CH₃ group for CH bond breaking. The insensitivity of the amount of exchange to the D₂ pressure despite more hydrogen being adsorbed with increased pressure^{17,22} and the sensitivity to CCH₃ coverage support this hypothesis.

However, methyl group rocking at its zero point energy $[1/2(988 \text{ cm}^{-1})]$ is about 7° from the surface normal²⁴ — not enough for much surface-aided CH bond scission. We can explain the surface assisted CH bond scission in the methyl group by again postulating ethylidene as an intermediate. Deuterium incorporation at the carbyne carbon might force the lower carbon into a bridge site with the C-C bond in CDCH₃ tipped about 50° from the surface normal. This moves a methyl group hydrogen nucleus to within 1.8 Å of a Rh atomic center (Rh-H bond about 1.7 Å). Carbon-hydrogen bond scission is possibly concurrent with D migration from the carbyne carbon into the methyl group (back-side attack) followed by reorientation of CCDH₂:

Since the ethylidyne H,D exchange rate is inversely dependent on CCH_3 coverage and D₂ pressure-independent, the rate limiting step is hydrogenation of the carbyne carbon and tipping of the ethylidene.

-13-

5. Conclusions

We have shown the applicability of HREELS within a HPLP system for studying monolayer structure after HP gas exposure. The HREELS/HP cell combination is a feasible and powerful way of studying surface reactions. We can, with isotopic labelling, study the composition and structure changes in a stable adsorbed monolayer — a relatively unstudied part of catalysis. HREELS, with its sensitivity to carbon-hydrogen bonds, is ideal tool for unraveling hydrogen transfer in adsorbed hydrocarbon monolayers.

Hydrogen transfer in an ethylidyne monolayer on Rh(111) has been examined in this H,D exchange study. Using the HREELS/HP cell combination, the following conclusions have been reached:

- 1) Saturation coverage of ethylidyne on Rh(lll) is stable towards irreversible hydrogenation under 1 atm H₂ at 310 K.
- 2) The relatively open c(4x2) CCH₃ monolayer structure is not contaminated by coadsorption or replacement during gas exposures at atmospheric pressure at 300-350 K on Rh(111).
- 3) H,D exchange in ethylidyne is a slow process for a saturation coverage CCH_3 overlayer on Rh(lll) at 310 K in 1 atm D₂. Exchange is rapid and extensive at subsaturation ethylidyne coverages of the Rh(lll) crystal face.
- 4) The H,D exchange rate of saturation coverage of CCH_3 on Rh(111) at room temperature is essentially independent of D_2 pressure. We postulate that this is due to a rate-limiting step of H incorporation at the carbyne carbon to form ethylidene.

ACKNOWLEDGEMENT

This work was supported by the Director, Office of Energy Research, Office of Basic Energy Sciences, Materials Sciences Division of the U.S. Department of Energy under contract Number DE-AC03-76SF00098. One of us, B.E.B., would like to acknowledge the support of an NSF graduate fellowship. We want to thank Dr. P. Davies for design of the HPLP apparatus and HREELS power supply and F. Zaera for his critical comments and suggestions.

ve

REFERENCES

ł

63

U

U.

47

1.	H. Ibach, H. Hopster, and B. Sexton, Appl. Surf. Sci. 1 (1977) 1.
2.	B.E. Koel, M.A. Van Hove, and G.A. Somorjai, Advances in Catalytic
	Chemistry II Symposium, Salt Lake City, Utah (1982).
3.	R.J. Koestner, M.A. Van Hove, and G.A. Somorjai, Surf. Sci. 121 (1982) 321.
4.	L.H. Dubois, D.G. Castner, and G.A. Somorjai, J. Chem. Phys. 72 (1980)
	5234.
5.	R.J. Koestner, M.A. Van Hove, and G.A. Somorjai, J. Phys. Chem. 87 (1983) 203.
6.	H. Steiniger, H. Ibach, and S. Lehwald, Surf. Sci. 117 (1982) 685.
7.	J.A. Gates and L.L. Kesmodel, Surf. Sci. 124 (1983) 68.
8.	A.L. Cabrera, N.D. Spencer, E. Kozak, P.W. Davies, and G.A. Somorjai, Rev.
	Sci. Instrum. 53 (1982) 1888.
9.	D.W. Blakely, E.I. Kozak, B.A. Sexton, and G.A. Somorjai, J. Vac. Sci.
	Technol. 13 (1976) 1091.
10.	E.E. Koel, J.E. Crowell, C.M. Mate, and G.A. Somorjai, J. Phys. Chem.,
	in press.
11.	P. Skinner, M.L. Howard, I.A. Oxton, S.F.A. Kettle, D.B. Powell, and N.
	Sheppard, J. Chem. Soc. Faraday Trans. 2, 77 (1981) 1203.
12.	B.E. Koel, J.E. Crowell, C.M. Mate, B.E. Bent, and G.A. Somorjai, to be
	submitted.
13.	S.M. Davis, F. Zaera, M. Salmeron, B.E. Gordon, and G.A. Somorjai, J. Am.
	Chem. Soc., July (1982).
14.	W. Erley, P.H. M ^C Breen, and H. Ibach, J. Cat., 84 (1983) 229.
15.	F. Zaera and G.A. Somorjai, J. Amer. Chem. Soc., July (1983).
16.	J.R. Fritch and K.P.C. Vollhardt, Angew. Chem., 92 (1980) 570; Angew. Chem.
	Int. Ed. Engl., 19 (1980) 559.

17. J.T. Yates, Jr., P.A. Thiel, and W.H. Weinberg, Surf. Sci. 84 (1979) 427. 18. A.M. Baro, H. Ibach, H.D. Bruchmann, Surf. Sci., 88 (1979) 384.

- 19. T. Shimanouchi, "Tables of Molecular Vibration Frequencies," Consolidated Volume 1, NSRDS-NBS 39 (1972) 51.
- 20. J.R. Ritter and D.F. Eggers, J.Chem. Phys., 44 (1966) 745.
- 21. These modes are both reported to be of medium intensity in reference 11.
- 22. The only complication is that the δ_{s} CH₃, whose intensity changes with deuteration, is not symmetrically placed between the 1246 and 1440 cm⁻¹ peaks. Its decrease in intensity will contribute some to changes in the 1246 and 1440 cm⁻¹ relative intensities.
- 23. M. Salmeron and G.A. Somorjai, J. Phys. Chem., 86 (1982) 341.
- 24. Force constant for the methyl group rocking was taken from reference 11: $H_Y = 59 \text{ N/m}$

FIGURE CAPTIONS

- Fig. 1. (A) Schematic diagram of a two-tier HPLP system containing a spectrometer for HREELS, with the high-pressure cell open. (B) Detail with high-pressure cell closed.
- Fig. 2. Specular HREEL spectra obtained following saturation exposure of ethylene to Rh(111) at 310 K to form surface ethylidyne.

e 7

- Fig. 3. Ethylidyne on Rh(111): the stable, room temperature, chemisorbed structure of ethylene.
- Fig. 4. Surface ethylidyne at saturation coverage on Rh(111) is extremely stable at 310 K to rehydrogenation. HREEL spectra of a Rh(111) surface: (A) saturated with ethylidyne under UHV conditions, and (B) following exposure of this monolayer to 1 atm. H₂ for 5 min. at 310 K.
- Fig. 5. Structure of the c(4x2) CCH₃ monolayer on Rh(111). Approximate Van der Waals radii of ethylidyne and H₂ are shown.
- Fig. 6. IEED photographs corresponding to the same conditions as in Fig. 4: (A) 15 L C₂H₄ at 310 K, and (B) following exposure of this monolayer to 1 atm. H₂ for 5 min. at 310 K. A phase transition and improved ordering of the ethylidyne monolayer is observed, even though no new species stable in UHV are formed.
- Fig. 7. Methyl group vibrational frequencies and approximate mode descriptions in variously deuterated methyl chlorides. Shaded peaks are modes of A₁ or A' symmetry and are dipole allowed, while unshaded peaks are modes of E or A" symmetry and are not dipole allowed. C-Cl modes are replaced by approximate C-C stretching frequencies for direct comparison with ethylidyne. (A) CH₃Cl, (B) CDH₂Cl, (C) CD₂HCl, (D) CD₃Cl.
- Fig. 8. H,D exchange in the saturated ethylidyne monolayer on Rh(111) at 320 K.
- Fig. 9. Increasing temperature and time influence the extent of H,D exchange in ethylidyne. Under these conditions, almost no ethylidyne species have undergone complete exchange, and some species remain that have undergone no exchange.
- Fig. 10. HREEL spectra taken after a partial monolayer of ethylidyne was exposed to 30 L D₂ under UHV conditions at 310 K. All of the ethylidyne undergoes some deuterium exchange, with a large fraction undergoing complete exchange in the ethylidyne methyl groups.

Table I.	Comparison of the vibrational frequencies (cm^{-1}) of the ethylidyne surface species formed on Rh(111) with those of the ethylidyne cluster compound. ¹¹

Assignment	CH3CC03(CO)9	CH3C-Rh(111)				
$v_{as}(CH_3)/v_{as}(CD_3)$	2930(m)/2192(w) e	2920(vw)/2178(vw) e				
$v_{s}(CH_{3})/v_{s}(CD_{3})$	2888(m)/ a ₁	2880(w)/2065(vw) a ₁				
$\delta_{as}(CH_3)/\delta_{as}(CD_3)$	1420(m)/1031(w) e	1420(vw)/ e				
$\delta_{s}(CH_{3})/\delta_{s}(CD_{3})$	1356(m)/1002(vw) a ₁	1337(s)/988(w) a _l				
ν(CC)	1163(m)/1882(ms) a ₁	1121(m)/1145(m) a _l				
ρ(CH3)/ρ(CD3)	1004(s)/828(s) e	972(vw)/769(vw) e				
ν _S (MC)	401(m)/393(m) a ₁	435(w)/419(w) a ₁				

.'

•

and a second second second second

Table II.	Dipole-allowed vibrational frequencies (cm ⁻¹) for a partially deuterated methyl group: assignment for ethylidyne on Rh(111) and
	comparison with similar species. Rh(111) assignments were made using the average peak frequencies from many partially deuterated ethylidyne spectra.

δCD2 wag	δCD2 scissor	δ _s H	vs ^{CD} 2	νCH	Source
 880	1019	1256	2117	2919	calculation ¹¹
784	1000	1239	2166	2952	ethylidyne Rh(lll)
813	1034	1209	2195	3023	methyl halides ²⁰

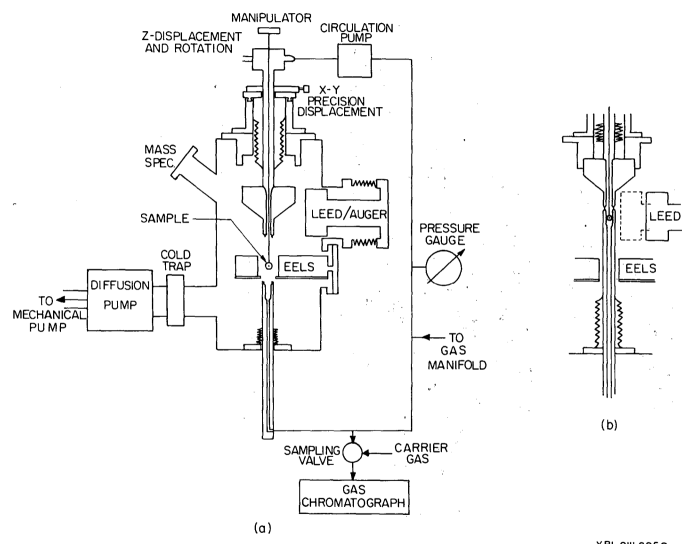
				· · · · · · · · · · · · · · · · · · ·		
	δ _S CD	δCH2 scissor	δCH2 wag	νCD	v _s CH ₂	Source
H D, I H	810	1430	1248	2169	2918	ethylidyne Rh(111)
Ċ	769	1425	1222	2236	2998	methyl halides ²⁰

-20-

Q,

¥

.



Ģ

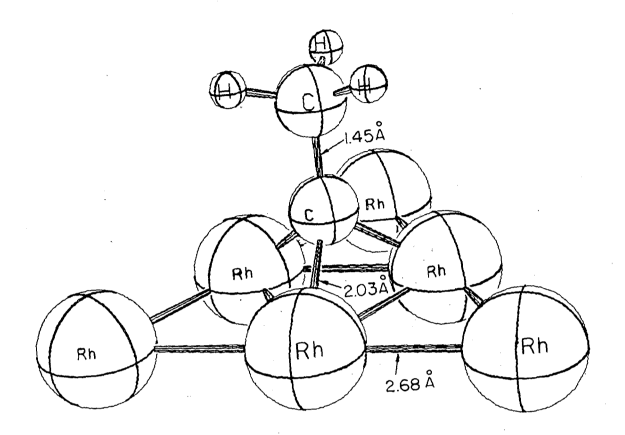
¢

XBL 8111-6950

Fig. 1

e c

-21-



-22-

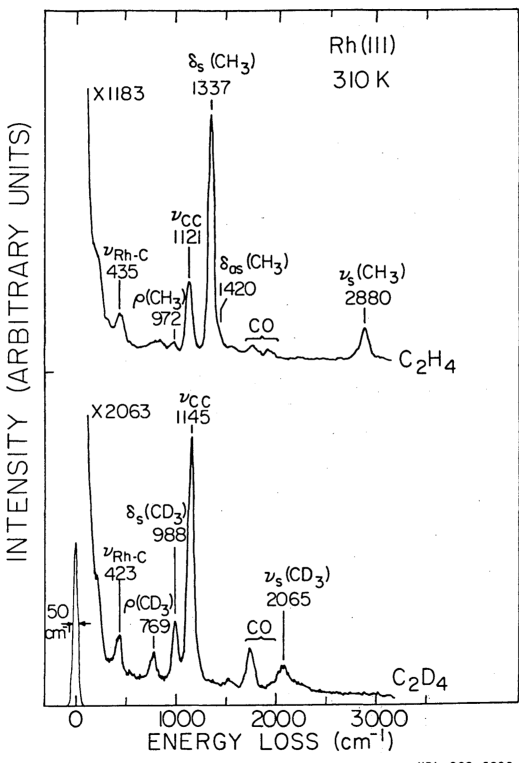
Rh(111) + ethylidyne

XBL794-6167A

Q.

്

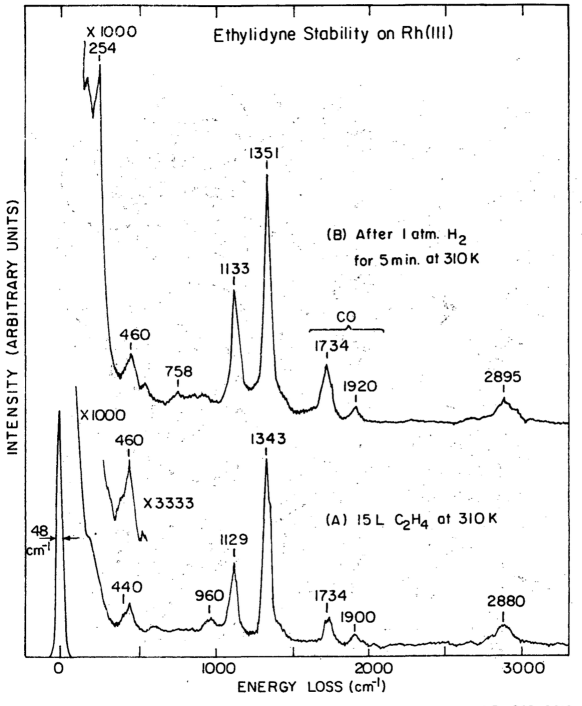




-23-

XBL 838-6209

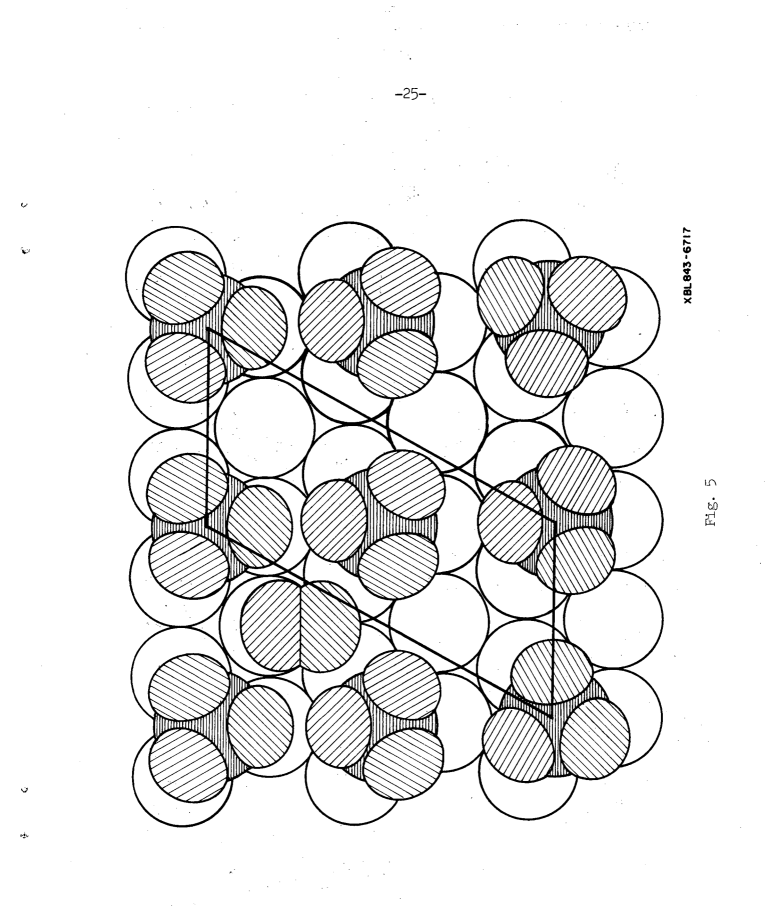
д,

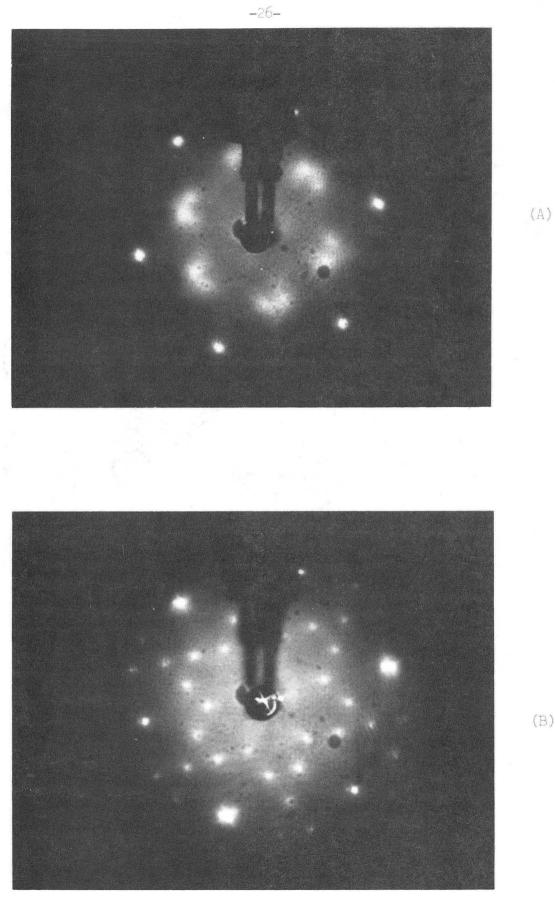


XBL 842-6616

ŧ,

Fig. 4

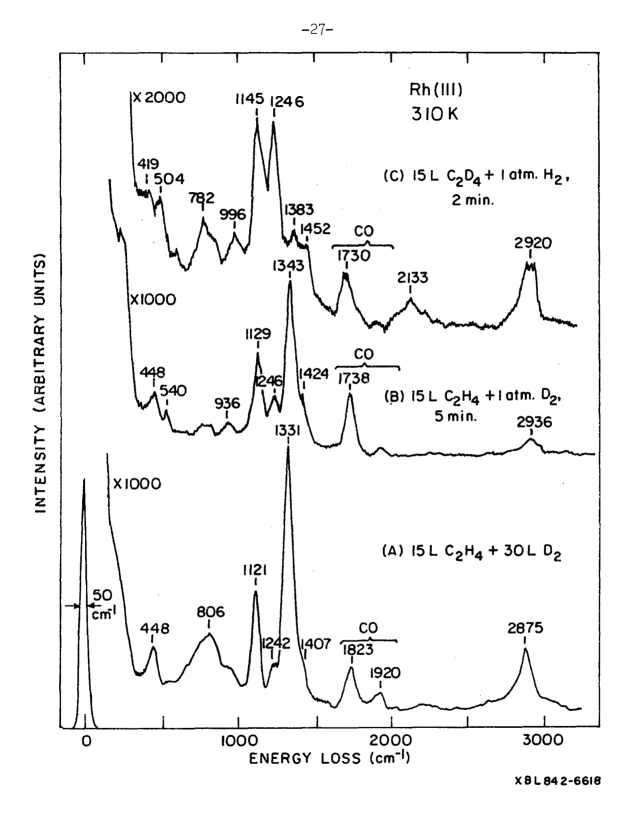




(B)

Fig. 6

XBB 841-626



ų

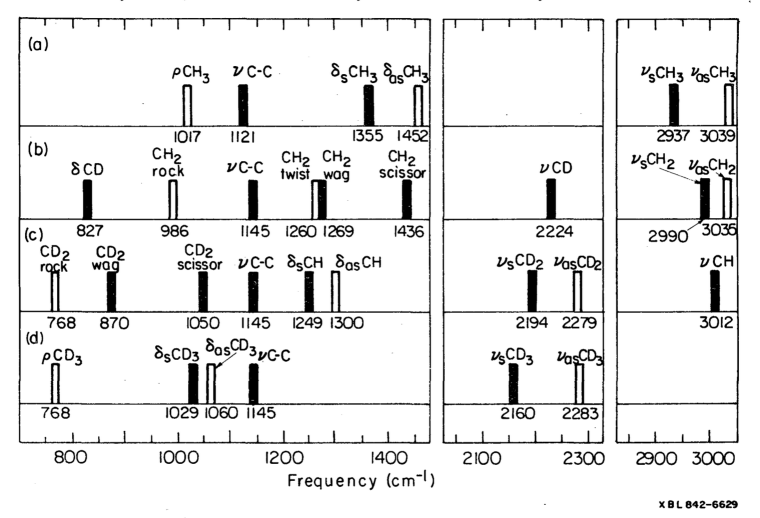
Ø

Ų

ۍ

4) |

Fig. 7



Methyl Group Modes in Variously Deuterated Methyl Chlorides

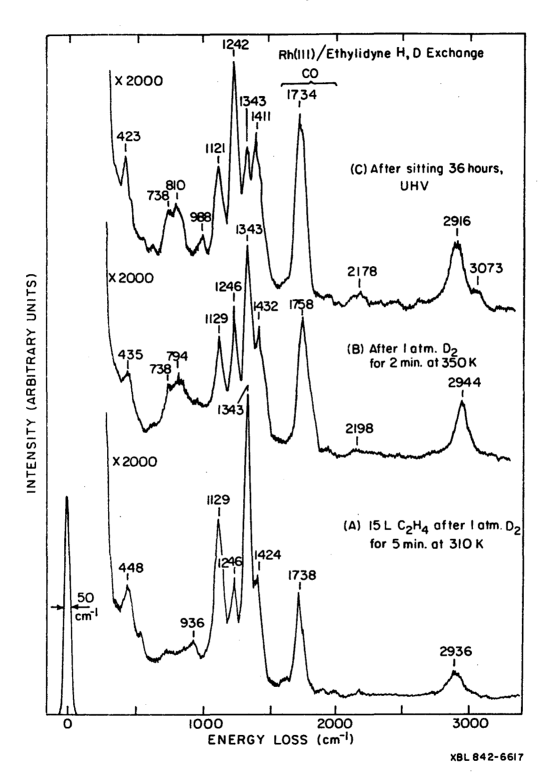


4 G

4

C

-28-

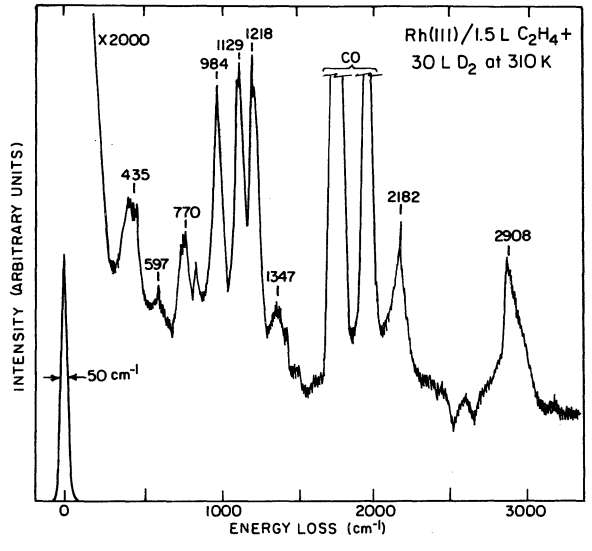


10

19- **o**

Fig. 9

-29-



XBL842-6619

`~

¢

ų,

£

Fig. 10

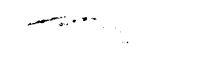
-30-

This report was done with support from the Department of Energy. Any conclusions or opinions expressed in this report represent solely those of the author(s) and not necessarily those of The Regents of the University of California, the Lawrence Berkeley Laboratory or the Department of Energy.

Ľ

ţ.

Reference to a company or product name does not imply approval or recommendation of the product by the University of California or the U.S. Department of Energy to the exclusion of others that may be suitable. TECHNICAL INFORMATION DEPARTMENT LAWRENCE BERKELEY LABORATORY UNIVERSITY OF CALIFORNIA BERKELEY, CALIFORNIA 94720



÷з. че з