

Review

A Contribution to the Experimental Microkinetic Approach of Gas/Solid Heterogeneous Catalysis: Measurement of the Individual Heats of Adsorption of Coadsorbed Species by Using the AEIR Method

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Abstract: The two first surface elementary steps of a gas/solid catalytic reaction are the adsorption/ desorption at least one of the reactants leading to its adsorption equilibrium which can be or not disturbed by the others surface elementary steps leading to the products. The variety of the sites of a conventional catalyst may lead to the formation of different coadsorbed species such as linear, bridged and threefold coordinated species for the adsorption of CO on supported metal particles. The aim of the present article is to summarize works performed in the last twenty years for the development and applications of an analytical method named Adsorption Equilibrium InfraRed spectroscopy (AEIR) for the measurement of the individual heats of adsorption of coadsorbed species and for the validation of mathematical expressions for their adsorption coefficients and adsorption models. The method uses the evolution of the IR bands characteristic of each of coadsorbed species during the increase in the adsorption temperature in isobaric conditions. The presentation shows that the versatility of AEIR leads to net advantages as compared to others conventional methods particularly in the context of the microkinetic approach of catalytic reactions.

Keywords: heats of adsorption; FTIR spectroscopy; AEIR method; Temkin model

1. Introduction

One of the aims in gas/solid heterogeneous catalysis is to correlate the global rate of the reaction (defined as the catalytic activity) including in its unit a property of the catalyst (i.e., weight, specific surface area, amount of active sites) to the elementary steps on the surface forming the detailed mechanism of the reaction. This constitutes the microkinetic approach of heterogeneous catalysis [1] which imposes the assessment of the kinetic parameters of the surface elementary steps such as the rate constants: k (pre-exponential factor and activation energy) and adsorption coefficients: K (pre-exponential factor and heat of adsorption) and the determination of the global rate of the reaction from the detailed mechanism [1]. The kinetic parameters of the surface elementary steps can be obtained either by experimental procedures on model surfaces (surface sciences approach) and conventional powdered catalysts or by theoretical methods (i.e., DFT (Density Functional Theory) calculations). From the detailed mechanism, the global rate of the reaction can be obtained by different either classical [2,3] or more recent [4–6] methods. Using, kinetic parameters from different origins may lead to ambiguities in the conclusions of a microkinetic study considering that they are dependent on the composition, morphology and structure of the catalyst (material gap) which control the type of adsorption sites (terraces, steps and corners for metal supported catalysts) and the interactions between adsorbed species. Similarly, surface sciences studies of catalytic reactions use mainly low



adsorption pressures (as compared to conventional conditions of heterogeneous catalysis) which may neglect the contribution of weakly adsorbed species (pressure gap). The use of kinetic parameters from different experimental and theoretical approaches is mainly due to the large numbers of surface elementary steps of a detailed mechanism of a catalytic reaction even for bi- and tri-atomic reactants (i.e., CO/O₂, CO/H₂, CO/NO, NH₃/NO). This is due to the fact that the number of either ruptures or creations of bonds in a surface elementary step must be limited. For instance, the number of surface elementary steps in detailed mechanisms of microkinetic studies is: 12, for NH₃ synthesis from N_2/H_2 (or its decomposition) on ruthenium catalysts [7–9]; 22, for NO/H₂ on Pt catalysts [10]; 26, for the production of C1 and C2 species from CO/H_2 on cobalt catalysts [11] and 32, for the ethylene oxidation on Ag catalysts [12]. However, among the surface elementary steps of a detailed mechanism, few of them control the global rate of the reaction: they constitute the kinetic model of the reaction [2,3]. This has led our group developing the experimental microkinetic approach (abbreviation EMA) of heterogeneous gas/solid catalytic reactions such as CO/O₂ [13,14] and CH₄ production from CO/H_2 [15,16] on Pt/Al₂O₃ catalysts. The main point of the EMA is that considering a plausible kinetic model of a catalytic reaction (based either on literature data or a formal kinetic approach), all the kinetic parameters of interest are obtained by experimental procedures on the conventional dispersed solid catalyst. This prevents using kinetic parameters coming from different sources (surface sciences, conventional catalysts, DFT) and overcomes the impacts of the material and pressure gaps. Note that it is the concurrence between the experimental catalytic activity and that from the EMA which validates the procedure; otherwise the plausible kinetic model must be reconsidered.

The first surface elementary steps of any gas/solid catalytic reaction are the adsorption with a rate R_a (rate constant k_a , activation energy E_a equal to 0 for non-activated chemisorption) followed by the desorption with a rate R_d (rate constant k_d , activation energy E_d) of at least one reactant. For $R_a - R_d = 0$, these two elementary steps lead to the adsorption equilibrium of the reactant on the sites which is characterized by the adsorption coefficient $K_a = k_a/k_d$ and a heat of adsorption $E = E_d - E_a$ (K_a determines the coverage of the adsorption sites for adsorption temperature T_a and pressure P_a). During a catalytic reaction, the adsorption equilibrium of a reactant can be disturbed or not by others surface elementary steps with a rate R_s and its coverage is controlled by the reaction equilibrium $R_a - R_d - R_s = 0$ [2,3]. However, in numerous kinetic studies of gas/solid catalytic reactions, it is often assumed that the adsorption equilibrium is not disturbed by the catalytic reaction. It is well known that the adsorption of a reactant may lead to the formation of different adsorbed species. For instance, the reactant CO of the CO/O_2 and CO/H_2 reactions on supported metal particles may lead to the formation of linear, bridged and threefold coordinated CO species which are well characterized by their IR bands in distinct wavenumber ranges [17]. Similarly, for the selective catalytic reduction of NO by NH₃ in excess O₂ (NH₃-SCR) on x% V₂O₅/y% WO₃/TiO₂ catalysts the adsorption of NH₃ leads to NH_{3ads-L} and NH₄⁺ species on the Lewis and Brønsted sites respectively which provide distinct IR bands ([18] and references therein). The role of each coadsorbed species in the reaction is a key point of microkinetic studies. For instance, their respective coverages for a composition of the reactive gas mixture are fixed by their individual heats of adsorption and adsorption coefficients implying that the determination of their values constitutes the first stage of the EMA of the reaction. Moreover, considering that the final aim of a microkinetic study is to express the global rate of the reaction as a function of the kinetic parameters of the surface elementary steps, it is of interest that experimental studies validate mathematical expressions for the adsorption coefficients and the adsorption models for each coadsorbed species formed by the reactants and others compounds of the reactive mixtures (i.e., H_2O for NH₃-SCR to consider the impact of NH₃/H₂O co-adsorption on the catalytic activity). This is the aim of an original method named Adsorption Equilibrium InfraRed spectroscopy (AEIR) which has been particularly developed in the last twenty years. This method is based on the evolutions of the IR bands characteristic of each adsorbed species during the increase in the adsorption temperature T_a in isobaric conditions. The aim of the present article is to summarize the development and the applications of this method.

2. Context of the Development of the AEIR Method

2.1. Classical Methods for the Measurement of the Heats of Adsorption

The driving force of this development was the difficulties obtaining the data of interest for a EMA of a catalytic reaction by conventional methods such as the isosteric heats of adsorption of a gas via volumetric/gravimetric measurements, the activation energy of desorption from temperature programmed desorption experiments and the differential/integral heats of adsorption using microcalorimetry. The isosteric heat of adsorption is based on the measurement either in isothermal or isobaric conditions of the coverages of a gas [19]. This allows one determining different couples (T_a , P_a) leading to the same coverage which provide the average isosteric heat of adsorption Q_{iso} at different coverages via the Clausius-Clapeyron equation [19]:

$$\left(\frac{\partial \ln P_a}{\partial (\frac{1}{T_a})}\right)_{\theta} = \frac{-Q_{iso}}{R} \tag{1}$$

where R is the ideal gas constant. These measurements are tedious to perform and time-consuming, while the method is strongly affected by experimental uncertainties. Moreover the formation of several adsorbed species leads to average values of limited interest in line with the aims of the EMA. Moreover, Equation (1) imposes the use of a large number of experimental data often associated to successive pretreatments of the catalyst which may affect its properties (i.e., sintering of the supported metal particles). Microcalorimetry [20] provides the differential and integral heats of adsorption of a gas according to its coverage. However different experimental difficulties can be encountered such as: the presence of several adsorbed species leading to average values; the impact of gaseous impurities [21]; the non-equilibrium nature of the adsorption at low temperatures [22,23] and the contribution of parallel reactions to the adsorption at high temperatures. Moreover, isosteric methods and microcalorimetry do not provide mathematical expressions for the adsorption coefficient and the adsorption model. The difficulties of these two analytical methods explains the success of temperature programmed desorption (TPD) methods which reveal easy the presence of coadsorbed species having different activation energy of desorption via their rates de desorption [19,24,25]. TPD methods consist adsorbing a gas at a temperature low enough to obtain a very low rate of desorption and then increasing the temperature T_d in inert atmosphere to desorb progressively the different adsorbed species according to their activation energy of desorption. This leads to a succession of peaks during the increase in T_d characterized by the temperature T_m of their maximum [19,24,25]. Equations based on classical theories of the adsorption (i.e., the kinetic theory of the gases and the statistical thermodynamics) provide kinetic parameters of interest such as the activation energy of desorption from T_m [19,24,25]. However, the TPD method imposes a careful design of the experiment in line for instance with the criteria proposed by Gorte et al. [26,27] to prevent the contribution of mass and heat transfers and to neglect the readsorption. These criteria show that readsorption can be rarely prevented using conventional catalysts [28–30]. In these conditions mathematical formalisms may provide the heats of adsorption of the adsorbed species [28–31]. However, similarly to microcalorimetry difficulties in the exploitation of the TPD spectra come from the contribution of parallel surface processes such as surface reconstructions and reactions with impurities (i.e., O₂, H₂O). Moreover, for heterogeneous surfaces the TPD peak (without and with readsorption) of an adsorbed species is very broad [31] leading to strongly overlapped peaks for coadsorbed species restricting significantly the access to the kinetic/thermodynamic parameters of interest for an EMA.

2.2. Precursor Works Using IR Spectroscopy for the Measurement of the Heats of Adsorption

Different early works were dedicated to the use of IR spectroscopy in this field. These studies either assumed or established the validity of a linear relationship between (a) the amount of the adsorbed species and (b) the area of its characteristic IR band according to the Beer-Lambert law which is the basis of the quantitative exploitation of FTIR (Fourier-transform infrared spectroscopy) spectra. The first studies were made by surface sciences via Infrared reflection adsorption spectroscopy (IRAS). For instance, Kottle et al. [32] measured the isosteric heats of adsorption of a linear CO species on an evaporated golf film for T_a and P_a in the ranges of 300–383 K and 7–530 Pa using an IR band in the 2120–2115 cm⁻¹ range according to the coverage. There was a scatter in the data from Equation (1) and the authors provided the average of the heats of adsorption in two coverage ranges: 13.4 kJ/mol and 12.4 kJ/mol in the coverage ranges 0.1–0.6 and 0.3–0.6 respectively. Similarly, Richardson et al. [33] measured the isosteric heats of adsorption of the Beer-Lambert law. Moreover, after showing that the isotherms were consistent with the Langmuir adsorption model:

$$\theta(T_a, P_a) = \frac{K(T_a) P_a}{1 + K(T_a) P_a}$$
(2)

the values of the adsorption coefficient $K(T_a)$ in the temperature range of the experiments were compared to mathematical expressions from the statistical thermodynamics approach of the adsorption. This allowed the authors obtaining an estimation of the partition function of the adsorbed species. The Goodman's group [34] have used a similar procedure to study the isosteric heats of adsorption of a linear CO species adsorbed on Cu(100) characterized by an IR band in the range 2086–2064 cm^{-1} according to the coverage. They used seven isotherms for adsorption pressures in the range 10^{-3} –130 Pa showing that the isosteric heat of adsorption varied from 70 kJ/mol to 53 kJ/mol in the coverage range of 0–0.15 ML. The same group has applied the procedure to measure the isosteric heats of adsorption of a linear CO species (IR band in the range of 2096–2053 cm⁻¹ according to the coverage) adsorbed on Pd film on Ta(110) using eight isobars in the range of $\approx 10^{-7}$ –130 Pa [35]. The authors showed clearly that the isosteric heat of adsorption decreased roughly linearly with the increase in the coverage from ~96 kJ/mol to ~40 kJ/mol in the coverage range of 0–0.35 ML. Similar studies have been performed for the bridged CO species on Pd crystals (IR band in the range 1968–1947 cm⁻¹) on Pd(100) [36]: $121 \pm 8 \text{ kJ/mol}$ in the coverage range 0.45–0.55 ML and Pd(111) [37,38] with a linear decrease in the isosteric heat of adsorption from 145 kJ/mol to 103 kJ/mol in the coverage range of \approx 0–0.3 ML.

In parallel to the studies using IRAS, IR spectroscopy has been applied to the measurement of the heats of adsorption of CO species on conventional catalysts. These works concerned mainly weakly adsorbed species (heats of adsorption $< \approx 100 \text{ kJ/mol}$) due to the limited performances of the IR cells (working mainly in static conditions) to maintain, the catalyst at high temperatures and pressures on the IR beam. The first works were dedicated to the study of linear CO species adsorbed on the Lewis sites of metal oxides. For instance, Paukshtis et al. [39] studied the individual heats of adsorption of two coadsorbed linear CO species (IR bands in the range of 2150-2230 cm⁻¹) on different Lewis acidic sites of 16 dispersed metal oxides such as MgO, Al₂O₃, ZrO₂, TiO₂. Assuming the validity of the Beer-Lambert law for the area (named A) of the IR bands they showed that the different isotherms followed the Langmuir model and using Equation (2) they obtained the heats of adsorption of the different L CO species from the plots $[\ln(A/A_0) - 1] = f(1/T)$ for each IR bands (A₀: the area at saturation of the sites). Thus on ZrO₂ they determined that the individual heats of adsorption of two coadsorbed linear CO species characterized by IR bands at 2203 and 2183 cm⁻¹ were of 36 and 28 kJ/mol respectively. In a following study [40], the authors have studied the heats of adsorption of a pyridine species on Al_2O_3 on ZrO_2 characterized by an IR band at 1445 cm⁻¹. However, this adsorbed species did not followed the Langmuir model and the authors provided the

isosteric heats of adsorption in the range 100–160 kJ/mol by using high temperatures (range 400–700 K) and low adsorption pressures (range 1.3×10^{-2} –3200 Pa). Yates et al. [41,42] have performed similar measurements for the L CO species adsorbed the Lewis sites of Al₂O₃ and SiO₂. On Al₂O₃ they noted that two species were formed providing strongly overlapped IR bands at 2195 cm⁻¹ (main IR band) and 2213 cm⁻¹. However, using isobaric conditions (T_a in the range of 180–350 K and P_a = 659 Pa) they provided via the Langmuir model their average heat of adsorption: 20.9 kJ/mol. Garrone et al. have studied the linear CO species on the Lewis sites of different metal oxides: TiO₂ [43], ZrO₂ [44] and Na-Z5M5 [45] using isothermal conditions at $T_a \approx 300$ K for $P_a <\approx 13$ kPa. The point of interest in these studies was that the isotherms on TiO₂ and ZrO₂ were compared to the Temkin adsorption model to take into account the heterogeneity of the adsorption sites.

Considering the characterization of the kinetic/thermodynamic parameters of adsorbed species with high heats of adsorption relevant of catalytic reactions, the first applications of the IR spectroscopy were dedicated to the measurement of the activation energy of desorption. This was linked to the limited performances of the IR cells using experimental conditions representative of heterogeneous catalytic reactions [46] and reference therein. For instance, Soma-Noto and Sachtler [47] have used this procedure to measure the activation energy of the L and B CO species adsorbed on Pd/Al_2O_3 and Pd-Ag/Al₂O₃ by studying the evolution of their characteristic IR bands with the duration of the isothermal desorption in vacuum in the range 373–540 K: 113 kJ/mol and 171 kJ/mol respectively. Similarly, some authors have developed TPDIR procedures: this consisted studying the evolution of the intensities of the IR bands of the adsorbed CO species on Pt/Al_2O_3 [48] during the linear increase of the desorption temperature T_d . This provided the evolution of the coverage θ_X of a adsorbed X species with T_d giving the curves $d\theta_X/dT_d$ which were exploited according to classical TPD procedures. This means that the same difficulties than those associated to the TPD procedure may contribute to the experimental data such as the consumption of the adsorbed species by reactions with H_2O and O_2 impurities [49]. The design of microreactor IR cell using gas flow rates at atmospheric pressure and high temperatures on the IR beam has allowed the characterization of the adsorption equilibrium of adsorbed species in experimental conditions representative of heterogeneous catalysis. The difficulties in the design of these IR cells come from the association of a small optical path length (range 2–3 mm) to limit the overlap of the IR spectra of the gaseous and adsorbed species and high temperatures due to the limited thermal stability of the IR windows and their sealing materials [46] and references therein. For instance, using a microreactor IR cell, Bell et al. [50] have determined the heat of adsorption of the linear CO species on Ru/Al₂O₃ in the coverage range 1–0.85 using x% CO/H₂ gas mixtures and three isotherms at T = 498, 523 and 548 K. In this small coverage range, the experimental data were consistent with the Langmuir model (Equation (2)) indicating a heat of adsorption of 106 kJ/mol. Using a similar IR cell, Kohler et al. [51] have measured the heats of adsorption of linear CO species on unreduced (IR band at 2132 cm⁻¹) and reduced (IR band at 2090 cm⁻¹) x% Cu/SiO₂ solids with x in the range of 2–10. On the unreduced solids three isotherms at 358, 378 and 441 K with $P_a \leq 20$ kPa showed that the L CO species followed the Langmuir model leading to a heat of adsorption of 25 kJ/mol. This was confirmed by using the isosteric method showing that the heat of adsorption was independent on the coverage: ≈ 29 kJ/mol in the coverage range of 0.1–0.9. For the reduced solids, isotherms at 358, 378, 441 and 493 K showed that the coverage of the L CO species was not consistent with the Langmuir model and the isosteric method indicated that the heat of adsorption increased with the decrease in the coverage according to a profile consistent with the Freundlich model with values at low coverages varying with the copper content for $\approx 50 \text{ kJ/mol}$ to 28 kJ/mol for x = 9.5 to x = 2.1 [51]. Clarke et al. [52] have confirmed the value at low coverage (range of 0–0.18) for a reduced 7% Cu/SiO₂ by using as approximation the Langmuir model: 35 kJ/mol.

The AEIR method has been developed in line with these precursor works using the adsorption of CO on Pt containing catalysts as case study [26,53–55]. The first step was the design of a microreactor IR cell allowing a significant increase of the highest temperature (until 900 K) as compared to literature data [46] (this improvement was imposed by the high heats of adsorption at low coverages of the

L CO species on Pt particles). The aim of the experimental procedure of the AEIR method was to combine measurements at the adsorption equilibrium (i.e., this prevents the impacts of heat and mass transfers) and temperature programmed procedures (rapidity of the experiment). Considering our interests for the EMA of gas/solid catalytic reactions, the aims of the exploitation of the IR spectra were the measurement of the individual heats of adsorption of coadsorbed species via the validation of mathematical expressions for the adsorption coefficients and adsorption models provided by classical theories of the adsorption. Two applications of the AEIR method are used to support the presentation: the adsorptions of CO on Pt/Al_2O_3 and NH_3 on TiO_2 based catalysts considered as the first steps of the EMA of catalytic reactions such CO/H_2 and de- NO_x from NH_3 -SCR respectively.

3. The Adsorption Equilibrium InfraRed Spectroscopy Method

3.1. IR Cell Microreactor for the Application of the AEIR Method

The AEIR method has been developed using a homemade stainless steel microreactor IR cell in transmission mode working at atmospheric pressure [46]. It has been designed (see Figure 1 in [46]) taking into account previous models and literature data. Briefly, a short path length (\approx 2.2 mm) limits the contribution of the gas phase to the IR spectra of the adsorbed species allowing using adsorption pressure of CO until \approx 20 kPa. The originality of the IR cell is that the two CaF₂ IR windows delimiting the path length in the heating part of the cell, are positioned on polished flat flanges without sealing materials (the two windows was maintained by using vacuum on one of their faces). This permits using temperatures in the range of 300–900 K with an heating rate of \approx 0.1–20 K/min [46]. The powdered catalyst (weight in the 40–200 mg range) was compressed into a disk (diameter = 18 mm) positioned on the IR beam between the two CaF₂ windows. In a recent work, it has been shown that a DRIFT cell can be used for the AEIR method (using the pseudo absorbance mode) taking into account that according to its design, heat transfers may create some difficulties to know the exact temperature of the fraction of the solid submitted to the IR beam for T > \approx 623 K [56].

3.2. Experimental Procedure of the AEIR Method

After pretreatment of the catalyst at high temperatures (i.e., H₂ reduction at 713 K for 2.9% Pt/Al_2O_3 and O_2 oxidation for TiO₂ based solids) it is cooled to 300 K. Then the switches H₂ (or O₂) \rightarrow He \rightarrow *x*% G/He (i.e., G either CO or NH₃) lead to the adsorption of G at the adsorption pressure $P_a = x \, 10^3$ Pa. After the stabilization of the IR bands of the adsorbed species, indicating the attainment of the adsorption equilibrium, the adsorption temperature T_a is increased ($\alpha \approx 10$ K/min) in the presence of x% G/He following the changes in the IR spectra of the adsorbed species. This provides the change in the intensities (in absorbance mode) of the IR bands characteristic of each adsorbed species X_{ads} at the adsorption equilibrium as a function of T_a in isobaric condition. It has been shown that the gas/solid system evolves by a succession of adsorption equilibriums taking into account that the high adsorption pressure and the moderate heating rate lead to a fast change from an adsorption equilibrium at (T_a, P_a) to that at $(T_a + dT_a, P_a)$ [30]. Similarly to the classical methods dedicated to the measurement of the heats of adsorption, surface processes parallel to the adsorption (i.e., surface reconstruction, CO dissociation) may contribute to the change of the intensity of the IR bands of the adsorbed species. However, the AEIR method permits to take into account these contributions by comparing the intensities of the IR bands at different adsorption temperatures during the first heating (i.e., 713 K) and cooling (i.e., 300 K) cycle in x% G/He. Often differences are observed due to reconstruction [53–55] and CO dissociation [57]. However, these processes are ended after the first heating/ cooling cycle in x% G/He as attested by the repeatability of the intensities of the IR bands of the adsorbed species during a second heating/cooling cycle in x% G/He: this means only the IR spectra of the second cycle (stabilized surface) are exploited via the AEIR method. The intensities of the IR bands of the adsorbed species can be modified by another process as observed for the adsorption of CO on supported Ag° particles [58]. After a first heating/cooling cycle, it has been observed that the IR band of the B CO species (at 1994 cm⁻¹) increases during the heating stages in parallel to the decrease in the IR band of the L CO species (at 2044 cm⁻¹). The reverse situation is observed during the cooling stages [58]. A similar process has been described by Müslehiddinoglu and Vannice [59] during the isothermal desorption at 300 K of the adsorbed CO species on Ag° particles. According to literature data [58] and references therein, this has been ascribed to an intensity transfer (in the 1/1 ratio) from the IR band of the B CO species to that of the L CO species. This transfer does not contribute significantly to the observations for different situations either if the amount of B CO species is low as compared to that of the L CO species (i.e., Pt/Al₂O₃) or if the two adsorbed species have different heats of adsorption allowing the significant decrease in the coverage of one of them without affecting that of the second species. For others situations the AEIR method does not apply.

As example of experimental data of the AEIR method, the inset of Figure 1 shows that the adsorption of 1% CO/He at 300 K on 2.9% Pt/Al₂O₃ leads to an IR spectrum with three IR bands at 2073, 1878 and 1835 cm⁻¹ ascribed [53–55] to linear, bridged and three fold coordinated CO species (named L, B and 3FC CO species respectively). Figure 1 gives the evolution of the IR band of the L CO species during the second increase in T_a for 1% CO/He. Similar spectra are obtained for the B and 3FC CO species.



Figure 1. Evolution of the IR band of the linear CO species adsorbed on a reduced 2.9% Pt/Al₂O₃ catalyst with the adsorption temperature T_a at $P_a = 1$ kPa. (a–f) $T_a = 378, 453, 533, 633, 693$ and 783 K. **Insert:** IR bands of the different adsorbed CO species on 2.9% Pt/Al₂O₃ for $T_a = 300$ K and $P_a = 1$ kPa.

3.3. Exploitation of the IR Spectra According to the AEIR Method

Considering the Beer-Lambert law, the amount of each adsorbed species X is proportional to the intensity of its IR band. This allows one obtaining the experimental evolutions of the coverage of each adsorbed species: θX_{ex} , with T_a , in isobaric conditions from the change in its IR band as for the L CO species in Figure 1:

$$\theta X_{ex} = \frac{A_X(T_a)}{A_{XM}} \tag{3}$$

where $A_X(T_a)$ and A_{XM} are the area of the IR band (in absorbance mode with values lower than ≈ 1.1) characteristic of the X species at T_a and at saturation of the sites respectively. The value of A_{XM} is obtained by ascertaining that the area of the IR band is not modified by either the increase in T_a in isobaric conditions or the increase in P_a in isothermal conditions. This is often the situation for the adsorbed CO species on metal particles at $T_a = 300$ K for P_a in the range 1–10 kPa due to their high heats of adsorption at full coverage. However, for weakly adsorbed species such as the linear CO

species on the Lewis sites of metal oxides: ZrO_2 [60] and TiO_2 [61] the saturation of the adsorption sites is not obtained at 300 K for the highest adsorption pressure available with the IR cell. For this situation, an estimation of A_{XM} is obtained according to the procedure of Kohler et al. [51]: assuming that the adsorption follows the Langmuir's model (Equation (2)) for non dissociative chemisorption, then the plot of $(1/A_X(300 \text{ K}))$ vs. $(1/P_a)$ provides A_{XM} . For a gas/solid system leading mainly to one adsorbed species, the validity of Equation (3) has been ascertained by different authors [33,35,36,39,50,51] and we have confirmed this point for the L CO species on the reduced metal particles of Cu/Al_2O_3 [62,63] and Ir/Al_2O_3 [64]. If the adsorption leads to different adsorbed species such as L, B and 3FC CO species on Pt/Al₂O₃ (inset Figure 1), the ascertainment of the Beer-Lambert law presents difficulties, due to the fact that at 300 K, volumetric/gravimetric methods provide the total amount of adsorbed species: $Q_T = \Sigma Q_X$ whereas A_X depends of a specific X species. For the adsorption of CO on Pt particles, FTIR spectroscopy shows that for 1% CO/He and $T_a > 550$ K, mainly the L CO species is present on the surface due to the difference in the heats of adsorption of the L, B and 3FC CO species. In these conditions, the validity of Equation (3) has been ascertained performing carbon mass balance with a mass spectrometer at the introduction of 1% CO/He taking into account that CO is involved in different processes: adsorption; dissociation and reaction with OH groups of the support forming CO_2 and H_2 [65]. These results and literature data have led us applying the Beer-Lambert law for all adsorbed species X_{ads} providing from Equation (3) the experimental curve $\theta X_{ex} = f(T_a)$ at different adsorption pressures P_a . For instance, symbols \blacksquare in Figure 2 provide the evolution of the coverage of the linear CO species θL_{ex} on the reduced 2.9% Pt/Al₂O₃ catalyst for $P_a = 1$ kPa considering the data in Figure 1. Similarly, symbols ▲ and ● give the experimental coverage of the B and 3FC CO species (inset Figure 1) for $P_a = 1$ kPa from experiments similar to Figure 1 with a higher amount of catalyst to improve the accuracy of the measurements and after the decomposition of their overlapped IR bands [54]. Note that for the B CO species, A_M in Equation (3) is obtained at 300 K for $P_a \ge 10$ kPa [34]. The experimental data in Figure 2 are compared to theoretical curves providing the individual heats of adsorption and the mathematical expressions of interest for the adsorption equilibriums of L, B and 3FC CO species.



Figure 2. Comparison between experimental and theoretical curves $\theta_X = f(T_a)$ at $P_a = 1$ kPa for the different X CO species on 2% Pt/Al₂O₃: \blacksquare , \bullet , and \blacktriangle experimental data for the L, 3FC and B CO species respectively; (a), (b) and (c) theoretical curve according to Equations (4) and (5) for the L, 3FC and B CO species (see the text for the E₀ and E₁ values of each adsorbed species).

3.4. Exploitation of the Experimental Curves $\theta X_{ex} = f(T_a, P_a)$ According to the AEIR Method

Considering our interest for the development of the EMA of catalytic reactions, the aims of the exploitation of the experimental data were the measurement of the individual heats of adsorption of the coadsorbed species via the validation of mathematical expressions for the adsorption coefficients and adsorption models provided by the classical theories of adsorption. It is a fact that the Langmuir model (Equation (2)) is rarely representative of experimental data for strongly adsorbed species on conventional catalysts due to the heterogeneity of the adsorption sites/adsorbed species. However, its extension to heterogeneous surfaces, via the integral approach and the distribution functions [15,16] and referenced therein, provides different equations for others adsorption models. For instance assuming a linear decrease in the heat of adsorption of an adsorbed species with the increase in its coverage, the integral approach leads to the generalized Temkin equation model [66] for non dissociative adsorption:

$$\theta_{th} = \frac{\mathrm{RT}_{\mathrm{a}}}{\Delta \mathrm{E}} \times \ln(\frac{1 + K(\mathrm{E}_{0}) - \mathrm{P}_{\mathrm{a}}}{1 + K(\mathrm{E}_{1}) - \mathrm{P}_{\mathrm{a}}}) \tag{4}$$

where E_0 (K(E_0)) and E_1 (K(E_1)) are the heats of adsorption (adsorption coefficient) at low and high coverages and $\Delta E = E_0 - E_1$. The statistical thermodynamics and the absolute rate theory provide the adsorption coefficient as a function of the partition functions of the gaseous molecule, the adsorbing site and adsorbed molecule [19,67–69]. In the temperature range of gas/solid catalytic reactions \approx 300–900 K, the partition function of a gaseous molecule is dominated by those of translation, rotation and vibration whereas that of the localized adsorbed molecule is dominated by those of rotation and vibration. In many cases, the ratio of the partition functions of rotation and vibration of the gaseous and adsorbed species can be reasonably approximated to \approx 1 [30,70] leading to the mathematical expression of the adsorption coefficient:

$$K(T_{a}) = \frac{h^{3}}{(2 \pi m k)^{3/2}} \frac{1}{T_{a}^{5/2}} \exp(\frac{E_{d} - E_{a}}{R T_{a}})$$
(5)

where h is Planck's constant, k is Bolztmann's constant, m is the mass of the molecule, E_d and E_a are the activation energies of desorption and adsorption respectively, while $E = E_d - E_a$ is the heat of adsorption. Note that as commented by Tompkin [19], the attainment of the adsorption equilibrium implicates a surface diffusion of the adsorbed species: this is compatible with a localized adsorbed species considering that localized means that the lifetime of the adsorbed species at the site is longer than its time in flight on the surface.

To obtain the individual heats of adsorption of coadsorbed species, the experimental evolutions of the coverage of each species (Figure 2) are compared to theoretical curves obtained by using Equations (4) and (5) and selecting a couple of E_0 and E_1 values leading to an overlap between experimental and theoretical curves (note that for $E_0 \approx E_1$ the theoretical curve obtained from Equation (4) is overlapped with that from the Langmuir model (Equation (2)). For instance in Figure 2, the curves a, b, c which overlap the experimental data for the L, B and 3FC CO species are obtained considering the following couples of heats of adsorption (E_0 , E_1) in kJ/mol at low and high coverages: (206, 115), (94, 45) and (135, 104). This shows that the AEIR method allows one determining the heats of adsorption of an adsorbed species from a single isobar. The practice shows that the choice of the E_1 and E_0 values is limited to short ranges ($\approx \pm 5$ kJ/mol) otherwise the experimental and theoretical curves are clearly distinct. This accuracy is due to the fact that E_1 and E_0 determine the temperature leading to the decrease in the coverage from 1 and the slope of the linear section of the isobar respectively [53].

3.5. Heats of Adsorption from the AEIR Method and Isosteric Heats of Adsorption

Using three isobars similar to Figure 1 for P_{CO} = 1000, 100 and 10 Pa [53], the validity of the AEIR procedure has been ascertained for the L CO species by showing that the E₀ values obtained

from Equations (4) and (5) are consistent with the isosteric heats of adsorption (Equation (2)) which is independent on the adsorption model. Similar conclusions have been obtained for different adsorbed CO species on metal particles [54,62,64]. This indicates that Equations (4) and (5) provide a very well representation of the properties of adsorbed CO species whereas as compared to the isosteric heats of adsorption a single isobar is needed using the AEIR method to obtain the individual heats of adsorption of coadsorbed species.

It is a fact that the same adsorption model (localized adsorbed species and Temkin's model) allows fitting numerous experimental data dedicated to the heats of adsorption of L and B CO species formed by the adsorption of CO on supported metal particles on metal oxides as indicated in Table 1.

Metal Particles on Alumina	Heat of Adsorption of Adsorbed CO Species in kJ/mol				
	Linear CO Species		Bridged CO Species		Ref
	E ₁	E ₀	E ₁	E ₀	- 1001.
Pt°	115	206	45	94	[53-55]
Pd°	54	92	92	168	[71]
$ m Rh^{\circ}$	103	195	75	125	[72]
Ir°	115	225			[64]
$ m Ru^{\circ}$	115	175			[73]
Cu°	57	82	78	125	[62]
Au°	47	74			[61]
Ag°	58	76	84	88	[58]
Ni°	100	153	106	147	[74]
Fe°	79	105			[75]
Co°-C *	93	165			[57]

Table 1. Heats of adsorption at low (E_0) and high (E_1) coverages of the Linear and Bridged CO species adsorbed on different metal supported particles on metal oxides by using the AEIR method.

* Co° sites modified by C deposition from the CO dissociation.

The versatility of the Temkin model is probably due to the fact that it is the best representation of the heterogeneity of the surface for strongly adsorbed species on conventional catalysts. Classically the heterogeneity for adsorbed species is ascribed to either a difference in the adsorption properties of the sites (biographical or intrinsic heterogeneity) or an interaction between adsorbed species (induced heterogeneity) [66]. The Temkin model is one of the proposals [76,77] to represent by a equation the evolution of the coverage of a gas on a heterogeneous surface as a function of the adsorption temperature and pressure. Different studies have considered the contribution of each type of heterogeneity on the modeling of the coverage in particular the induced heterogeneity due to lateral interaction (Ref. [78] and references therein). However, Temkin [66] noted that the two types of heterogeneities can be simultaneously operant and that a single equation must be representative of this situation to prevent an excessive mathematical complexity. This has been justified by different authors [79–81]. Moreover, considering that the heats of adsorption of an adsorbed species at low (E_0) and high (E_1) coverages have limited values such as 206 kJ/mol and 115 kJ/mol for the L CO species on Pt particles (Table 1), the comparison of a linear (Temkin model) and an exponential (Freundlich model) decrease in the heats of adsorption with the increase in the coverage according to $E_T(\theta) = [E_0 - (E_0 - E_1)\theta]$ and $E_F(\theta) = E_0 \exp [-\theta \ln(E_0/E_1)]$ respectively, shows that the highest difference (6.6 kJ/mol at coverage 0.5) is in the range of the accuracy of the measurements.

3.6. Development of the AEIR Method

The AEIR method has been applied to different gas/catalyst systems such as: NO on 2.7% Pt/Al₂O₃ [82]; aromatic hydrocarbons on SiO₂ [70] and NH₃ [18,83,84] and H₂O [85] on different TiO₂ based solids. For this last application it has been observed that some IR bands provide $\theta_{ex} = f(T_a)$

selected for the measurement is due to the contributions of two adsorbed species having different heats of adsorption. In this situation $\theta_{ex} = f(T_a)$ gives the evolution of the average coverage of the two adsorbed species. A development of the AEIR method allows one obtaining the individual heats of adsorption of the two species as shown for the adsorption of NH₃ on the Lewis sites (named NH_{3ads-L} species) of TiO₂ P25 from Degussa [18] which is of particular interest because different IR bands can be used for the measurement of the individual heats of adsorption of the coadsorbed NH_{3ads-L} species supporting the development of the method. For instance, Figure 3 gives the evolution of the IR bands in the range 2000–1100 cm⁻¹ of the NH₃ species adsorbed on TiO₂ with the increase in T_a for 0.1% NH₃/He.



Figure 3. Impact of the adsorption temperature T_a on the IR bands of the adsorbed NH₃ species on TiO₂-P25 using 0.1% NH₃/He: (a–e) T_a = 300, 373, 473, 573 and 673 K. **Inset:** Decomposition of the δ_s IR bands of the NH_{3ads-L2} and NH_{3ads-L1} species at 300 K.

At 300 K, the overlapped IR bands at 1142 and 1215 cm⁻¹ in Figure 3 are ascribed to the δ_s deformations of two adsorbed NH₃ species on different Lewis sites L1 and L2 named NH_{3ads-L1} and NH_{3ads-L2} respectively [18] whereas their δ_{as} deformations contribute to the IR band at 1596 cm⁻¹. Moreover, in Figure 3, the broad IR band at 1477 cm⁻¹ and the shoulder at 1680 cm⁻¹ are ascribed to the antisymmetric and symmetric deformation of NH₄⁺ species formed by the adsorption of NH₃ on Brønsted sites [18] and references therein. The increase in T_a to 713 K for 0.1% NH₃/He leads to the decrease in the different IR bands: those of the NH₄⁺ species disappear at ≈423 K indicating weakly adsorbed species whereas those of the NH_{3ads-L2} species are present at 713 K. The individuals heats of adsorption of the NH_{3ads-L1} and NH_{3ads-L2} species have been obtained after decomposition of the two δ_s IR bands as shown in the inset of Figure 3 for T = 300 K. Considering similar IR absorption coefficients for the two NH_{3ads-L} species and taking into account that they are at full coverage at 300 K for 0.1% NH₃/He, the decomposition indicates that the L1 and L2 sites represent 70% and 30% of the Lewis sites of TiO₂ P25 respectively.

After decomposition at each adsorption temperature, the square and triangle symbols in Figure 4 give from Equation (3), the $\theta_{ex} = f(T_a)$ curves in isobaric conditions of NH_{3ads-L1} and NH_{3ads-L2} respectively.

Curves a and b which overlap the experimental data are obtained using Equations (4) and (5) with the following couples of (E₀ and E₁) values in kJ/mol (112, 56) and (160, 104) for the NH_{3ads-L1} and NH_{3ads-L2} species respectively. The circle symbols in Figure 4 give $\theta_{ex} = f(T_a)$ using the δ_{as} IR band at 1596 cm⁻¹ at 300 K which is common to the two NH_{3ads-L} species. Equations (4) and (5) do not allow one obtaining a theoretical curve overlapped with the experimental data in the full coverage whatever the set of E₀ and E₁ values. This is consistent with the fact that the two NH_{3ads-L} species have significantly different heats of adsorption. However, the individual heats of adsorption of the NH_{3ads-L1} and NH_{3ads-L2} species can be determined by comparison of the experimental data with the theoretical average coverage provided by [18]:

$$\theta_{\rm th}(T_a, P_a) = x_1 \ \theta_{\rm L1}(T_a, P_a) + x_2 \ \theta_{\rm L2}(T_a, P_a) \tag{6}$$

where $\theta_{L1}(T_a, P_a)$ and $\theta_{L2}(T_a, P_a)$ are the theoretical coverages of NH_{3ads-L1} and NH_{3ads-L2} respectively provided by Equations (4) and (5) and x_1 and x_2 represent the contribution (in fraction) of each NH_{3ads-L} species to the IR band at saturation of the L1 and L2 sites. For instance, curve c in Figure 4, which overlaps the experimental data is obtained using in Equation (6): $x_1 = 0.73$ and $x_2 = 0.27$, $E_{L1}(1) = 56$ kJ/mol, $E_{L1}(0) =$ 105 kJ/mol, $E_{L2}(1) = 105$ kJ/mol, $E_{L2}(0) = 160$ kJ/mol. The heats of adsorption are consistent with those obtained using the δ_s IR bands (curves a and b in Figure 4) whereas x_1 and x_2 are consistent with the values provided by decomposition of the δ_s IR band at 300 K.



Figure 4. Heats of adsorption of the adsorbed NH₃ species on TiO₂-P25 using the AEIR method. ▲ and □ coverages of the NH_{3ads-L2} and NH_{3ads-L1} adsorbed species respectively using their δ_s IR bands; full lines: (a) and (b) theoretical coverages of the NH_{3ads-L1} and NH_{3ads-L2} species respectively using Equations (4) and (5) with E_{L1}(1) = 56 kJ/mol, E_{L1}(0) = 102 kJ/mol, E_{L2}(1) = 102 kJ/mol and E_{L2}(0) = 160 kJ/mol; • average coverage of the NH_{3ads-L1} and NH_{3ads-L2} species using their common δ_{as} IR band at 1596 cm⁻¹; (c) theoretical evolution of the average coverage of the NH_{3ads-L1} and NH_{3ads-L2} species respectively using Equation (6) considering $x_1 = 0.73$ and $x_2 = 0.27$ for the NH_{3ads-L1} and NH_{3ads-L2} species respectively and with heats of adsorption of E_{L1}(1) = 56 kJ/mol, E_{L1}(0) = 105 kJ/mol, E_{L2}(1) = 105 kJ/mol and E_{L2}(0) = 160 kJ/mol (see the text for more details).

This procedure of the AEIR method using Equation (6) is of particular interest for the measurement of the individual heats of adsorption of NH_{3ads-L} species on sulfated TiO₂ containing catalysts because the strong ν (S=O) IR band of the sulfate groups prevents using the δ_s IR band of the NH_{3ads-L} species and only their common δ_{as} IR band can be exploited by the AEIR method [83]. Similarly the deposition of V_2O_5 and/or WO₃ on TiO₂-P25 [84] decreases the IR transmission of the solid below $\approx 1200 \text{ cm}^{-1}$

preventing using the decomposition of the δ_s IR band of the NH_{3ads-L} species as shown in Figure 5 for the adsorption of 0.1% NH₃/He on 6% WO₃/TiO₂. Moreover, this deposition leads to the presence of strong Brønsted and Lewis sites as shown by the evolutions of the IR bands of the NH_{3ads-L} (i.e., IR band at 1602 cm⁻¹) and NH₄⁺ (1445 cm⁻¹) species in Figure 5 during the increase in T_a indicating that the two species are present at $T_a > 673$ K. The IR spectra in Figure 5 and the AEIR method give from Equation (6) the individual heats of adsorption of (a) two NH_{3ads-L} species using the IR band at 1602 cm⁻¹ and (b) two NH₄⁺ species using the IR band at 1445 cm⁻¹ [84].



Figure 5. IR bands of adsorbed NH₃ species after adsorption of 0.1% NH₃/He on 6% WO₃/TiO₂-P25 pretreated at 713 K in helium as a function of adsorption temperature T_a : (a–e) T_a = 300, 373, 498, 573 and 673 K.

3.7. Application of the AEIR Method to Different Topics Relevant of Heterogeneous Catalysis

The AEIR method can be applied using either IR transmission or reflectance mode. In IR transmission, the microreactor IR cell must associate a low path length and high temperatures [46]. In DRIFT mode, temperature gradients in the solid sample and thermal/chemical stability of the sealing material/IR windows (i.e., presence of H_2O) must be taken into account [56,75]. Indeed, the level of the performances of the IR cells is dependent on the heats of adsorption of the adsorbed species of interest: strongly adsorbed species such as L CO species on Pt particles impose using high temperatures to observe the decrease in their coverage in isobaric conditions. Note that the AEIR method does not impose a variation of the decrease in the range 1–0: experiments in the temperature range corresponding to the beginning of the decrease in the coverage with spectra representative of the linear section of the isobar (Figures 2 and 4) provide the heats of adsorption of the adsorbed species [53].

The design of the experiments associated to the AEIR method is simple as compared to others analytical procedures dedicated to the measurement of the heats of adsorption. This permits the application of the method to study the impacts of different parameters associated to the catalyst preparation on the individual heats of adsorption of coadsorbed species. For instance, for the adsorption of CO on metal particles supported on metal oxides, the AEIR method permits to study the impacts of: the precursors of the metal particles [86,87]; the nature of the support [88,89]; the metal dispersion [55] and the deposition

of additives (i.e., K on Pt/Al₂O₃ [90]). Similarly the AEIR method allows studying the geometric and electronic effects due to the formation of bimetallic particles via the changes in the nature of the adsorbed CO species and in their heats of adsorption respectively. For instance, the AEIR method reveals that the insertion of Sn in Pd particles [91] leads to the total disappearance of the bridged CO species due to a geometric effect. Moreover, the heat of adsorption of the L CO species on Pd° sites, which varies linearly with its coverage from $E_0 = 92 \text{ kJ/mol}$ to $E_1 = 54 \text{ kJ/mol}$ on monometallic particles, is slightly modified on the Pd-Sn bimetallic particles: $E_0 = 90 \text{ kJ/mol}$ and $E_1 = 50 \text{ k/mol}$, indicating a very small electronic effect. Similarly, Meunier et al. have used the AEIR method in DRIFT mode to study the impacts of the insertion of Zn on Pd° sites of a Pd/CeO₂ catalyst [92]. The insertion on Zn suppresses the B CO species for Pd-Zn particles reduced at 773 K varies from $E_0 = 105 \pm 5 \text{ kJ/mol}$ to $E_1 = 68 \pm 5 \text{ kJ/mol}$ revealing a modest electronic effect of Zn. The same group has used the AEIR method to study the electronic effect of the insertion on Sn in Pt° particles supported on Al₂O₃ revealing a strong electronic effect with a heat of adsorption at low coverage half of that on the monometallic particles [93].

The AEIR method is well adapted to the study of the individual heats of adsorption of adsorbed CO species on metal particles which is of interest either for the characterization of the solid using CO as a probe or for the EMA of catalytic reactions involving CO as reactant. This explains the interest of different groups for its application. For instance, Collins et al. [94] have applied the method to measure the individual heats of adsorption of one L and two B CO species on the Pd sites of a 2% Pd/SiO₂ catalyst: the values of the heats of adsorption of the main B CO species $E_0 = 168 \text{ kJ/mol}$ and $E_1 = 62 \text{ kJ/mol}$ are consistent with those measured on a Pd/Al₂O₃ catalyst [71] (see Table 1). Similarly, Chen et al. [95] have used the AEIR procedure in DRIFT mode to measure the heats of adsorption of two coadsorbed L_1 and L_2 CO species on Cu particles supported on SiO₂: IR bands at 2134 and 2119 cm⁻¹ respectively, in relationship with the water-gas shift reaction. The E_0 and E_1 values are of 51 kJ/mol and 39 kJ/mol for the L1 and 70 kJ/mol and 46 kJ/mol for L_2 CO species. These values are consistent with those measured on a Cu/Al₂O₃ catalyst [62] (see Table 1). Rioux et al. [96] have used the AEIR method in diffuse reflectance mode for the measurement of the heat of adsorption of the L CO species on 2.69% Pt/SiO₂ (mesoporous silica: SBA-15, Pt particles 2.9 nm) using 10% CO/He showing a linear decrease with the increase in the coverage from $E_0 = 167 \text{ kJ/mol}$ to $E_1 = 125 \text{ kJ/mol}$. The E_1 value is consistent with that measured on 2.9% Pt/Al₂O₃ (Table 1) while that a low coverage is slightly lower probably due to the nature of the support and the type of Pt particles due to the preparation method.

Diemant et al. [97] have used Equations (4) and (5) for the exploitation of the coverage of a L CO species on planar Au/TiO₂ model catalysts with different particle sizes obtained from Polarization Modulation-IRAS (PM-IRAS) spectra. They show that the evolutions of the experimental coverages in isobaric condition ($P_{CO} \approx 10$ mbar) are consistent with the theoretical curves. They reveal the significant impact of the particles size Φ on the heats of adsorption of the L CO species: i.e., E₀ decreases from 74 kJ/mol to 62 kJ/mol for Φ in the range of 2–4 nm [97]. The values at low and high coverages for $\Phi = 2$ nm: 74 kJ/mol and 40 kJ/mol are consistent with those determined on a conventional Au/TiO₂ catalyst [61] (see Table 1). This shows that the AEIR method may allow us studying the impacts of the material gap between conventional catalysts and model surfaces on a key parameter of catalytic reactions.

Thus the AEIR method is often used to reinforce studies using the adsorption of a gas as a probe of the surface properties of a catalyst by providing via the individual heats of adsorption of the adsorbed species a quantification of the strength of the sites. For instance, this was one of the aims studying the modifications of the Lewis and Brønsted acidic sites by the deposition of WO₃ and V₂O₅ groups on sulfated and sulfate free TiO₂ supports species [18,83,84]. However, the AEIR method has been clearly developed as contribution to the EMA of catalytic reactions. In this field one of its interest is that it allows one studying the impact of the presence of a second gas (reactant or not) such as O₂, H₂ and H₂O on the heats of adsorption of adsorbed CO species on Pt/Al₂O₃ [98]. Particularly, the method provides experimental data on the change in the coverages of the different adsorbed species due to

the coadsorption. These data can be modeled using theoretical coverages obtained from the Temkin formalism for competitive chemisorption without [15] and with [16] transformation of the reactants which is a key step of the EMA of a catalytic reaction such as for CO/H_2 on Pt/Al_2O_3 [15,16].

4. Conclusions

The AEIR method developed and applied during the last twenty years for the characterization of individual heats of adsorption of coadsorbed species formed by the adsorption of a gas on a solid catalyst, constitutes a tool for the development of the experimental microkinetic approach of gas/solid heterogeneous catalysis using conventional powdered catalysis. For each adsorbed species, the method allows, from the evolution of their characteristic IR bands in isobaric conditions to measure their individual heats of adsorption at different coverages via the validation of mathematical expressions of the adsorption coefficients and adsorption models. These data allow an accurate modeling of the two first surface elementary steps of any gas/solid catalytic reaction taking into account the diversity of the adsorption sites on a conventional catalyst. The design of the experiment for the AEIR method is easy as compared to others classical methods: a single isobar is needed using either a pellet of catalyst in IR transmission mode or sized catalyst particles for IR reflectance mode. This facilitates the use of the method for the study of the impacts of the modifications of the catalyst on the heats of adsorption such as the natures of the precursors, supports and additives and the formation of bimetallic particles. Moreover, the fact that the procedure can be applied on model surfaces [97] permits studying the impact of the material gap on an important thermodynamic parameters controlling the coverage of the surface during a catalytic reaction.

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