PRESSURE AS A KINETIC PARAMETER IN THE ELUCIDATION OF THE MECHANISMS OF INORGANIC, ORGANOMETALLIC AND BIOINORGANIC REACTIONS IN SOLUTION

Rudi van Eldik
Institute for Inorganic Chemistry - University of Witten/Herdecke - Stockumer Straße 10 - 5810 Witten - Germany

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The application of high pressure kinetic techniques in mechanistic studies of chemical reactions in solution, can contribute towards the elucidation of the underlying reaction mechanisms. The fundamental principles involved, the general instrumentation employed, and the construction of reaction volume profiles, are discussed. Typical examples covering various types of reactions in inorganic, organometallic and bioinorganic chemistry, that occur on a range of different time scales, are presented. The advantages of pressure as a kinetic parameter are considered and an evaluation of the gained mechanistic insight is given.

Keywords: high pressure kinetic techniques; mechanistic studies.

INTRODUCTION

The author recently had the privilege to visit a number of Universities in the State of São Paulo and to lecture on the topic of this paper. Since this topic is to some extent a new area for Brazilian chemists, the idea originated to compile a concise and generally written review to present a short overview of the principles involved, and to illustrate them with a number of examples selected from inorganic, organometallic and bioinorganic systems. This presentation requires a basic knowledge on inorganic reaction mechanisms as found in a number of general textbooks.

The development of rapid kinetic techniques for the investigation of fast chemical processes in the nanosecond to the second time range at pressures up to 300 MPa (i.e. 3 kbar), has opened the possibility to study the effect of pressure on many chemical processes that occur on transition metal complexes. The application of stopped-flow, T-jump, NMR line-broadening, flash-photolysis and pulse-radiolysis, has led to the construction of reaction volume profiles for such processes, which in turn has contributed significantly to the elucidation of the underlying reaction mechanisms. A number of examples dealing with inorganic, organometallic and bioinorganic systems will be treated here. For further information readers are referred to a series of books, conference proceedings and review papers.

In principle, it has been found that reactions that involve bond formation or bond cleavage, or major changes in electrostriction, exhibit characteristic pressure dependencies. These can along with the data obtained from chemical (concentration, pH, solvent, ionic strength) and physical (temperature) variables be used to resolve the intimate nature of the underlying reaction mechanism. The quantitative description of such pressure dependencies is done in terms of the volume of activation, $\Delta V^\ddagger$, as given in eq. (1), which represents the change in partial molar volume when the reactant species move from the ground to the transition state in terms of the Eyring theory. $\Delta V^\ddagger$ will have positive or negative values depending on whether the reaction is decelerated or accelerated by pressure, i.e. whether the volume increases or decreases from the ground to the transition state, respectively. Such data combined with partial molar volume measurements for reactant and product species, or with reaction volume data for an equilibrium process, can be employed to construct a volume profile in which the chemical process is described in terms of volume changes along the reaction coordinate as shown in Figure 1. In this example the solid line represents the expected volume profile for a bond formation process in which the transition state lies between that of the reactant and product states, whereas the dotted lines represent alternative reaction routes in which the transition state has either a higher or lower partial molar volume than both the reactant and product states.

In general, $\Delta V^\ddagger$ may be considered as the sum of at least two components: an intrinsic part ($\Delta V^\ddagger_{\text{intrinsic}}$), which represents the change in volume due to changes in bond lengths and angles, and a solvational part ($\Delta V^\ddagger_{\text{solv}}$), which represents the volume changes due to electrostriction and other effects acting on the surrounding solvent molecules during the activation process. It is principally the intrinsic contribution that is the mechanistic indicator in the case of substitution and related reactions. A schematic representation of these components for a typical bond formation or bond cleavage process, during which charge neutralization or creation may occur, is

![Diagram](image-url)

*Figure 1. Volume profile for the reaction $A + B \rightarrow [A-B]$ → $A-B$.***
given in Figure 2. The mechanistic assignment for processes in which no major solvational changes occur are on the basis of the observed $\Delta V^*$ values rather straightforward. In reactions with large changes in electrostriction, $\Delta V^*_{solv}$ may be so large that it will counteract and swamp out the $\Delta V^*_{intr}$ contribution.

We will now consider a few examples in more detail that cover various types of reactions in inorganic (coordination), organometallic and bioinorganic chemistry, viz. ligand substitution, ligation, electron-transfer, and addition and related processes.

![Figure 2. Schematic representation to illustrate the sign of the components of $\Delta V^*$](image)

**Figure 2.** Schematic representation to illustrate the sign of the components of $\Delta V^*$.

LIGAND SUBSTITUTION REACTIONS

Substitution reactions of transition metal complexes have been the topic of many mechanistic investigations because of the fundamental importance of such reactions in many chemical, biological and catalytic processes. For a general ligand substitution reaction (2), where $X$ is the leaving group and $Y$ the entering ligand, there are three simple pathways: (i) the dissociative (D) process, with an intermediate of lower coordination number; (ii) the associative (A) process, with an intermediate of higher coordination number; (iii) the interchange (I) process, in which no intermediate of lower or higher coordination number is involved, and the interchange of the two ligands can be more dissociative ($I_d$) or more associative ($I_a$) in nature, depending on whether bond breakage or bond formation is more important, respectively (Figure 3). Such ligand substitution processes should exhibit characteristic $\Delta V^*$ values depending on the degree of bond breaking or bond formation in the transition state. For solvent- and neutral ligand-exchange reactions, i.e., where no net chemical reaction occurs, changes in electrostriction will be negligible and $\Delta V^* = \Delta V^*_{intr}$. Thus $\Delta V^*$ should be a direct measure of the intrinsic volume changes that occur, such that a continuous spectrum of transition state configurations can be envisaged, ranging from a very expanded, highly dissociative one (large positive $\Delta V^*$), to a very compact, highly associative one (large negative $\Delta V^*$) as shown in Figure 4.

![Figure 3. Schematic representation of the possible ligand substitution mechanisms.](image)

![Figure 4. Volume profiles for solvent exchange mechanisms.](image)

Typical experiments on water exchange reactions on octahedral high spin divalent first row transition metal ions employing high pressure NMR techniques, indicated a definite trend along the series of ions (Figure 5). The $\Delta V^*$ data (Table I) indicate that the mechanism progressively changes from $I_a$ to $I_d$.

| Table I. Volumes of activation (cm$^3$mol$^{-1}$) for solvent S exchange on MS$^{2+}$ of the first row transition metal series$^{4b}$. |
|-----------------|-----------|-----------|-----------|-----------|-----------|-----------|
| M$^{2+}$       | $V_{3+}$  | Mn$^{2+}$ | Fe$^{2+}$ | Co$^{2+}$ | Ni$^{2+}$ | Cu$^{2+}$ |
| n/pm           | t$_{a}^2$ | t$_{a}^2$c$_{a}^2$ | t$_{a}^2$c$_{b}^2$ | t$_{a}^2$| t$_{b}^2$| t$_{b}^2$c$_{a}^2$ | t$_{b}^2$c$_{a}^2$ | t$_{b}^2$c$_{b}^2$ | t$_{b}^2$c$_{b}^2$ | t$_{b}^2$c$_{b}^2$ | |
| H$_2$O         | -4.1      | -5.4      | +3.8      | +6.1      | +7.2      | +6.1      | +7.2      | +6.1      | +7.2      | +6.1      | +7.2      |
| MeOH           | -5.0      | -5.0      | +4.3      | +7.8      | +11.4     | +8.5      | +11.4     | +8.5      | +11.4     | +8.5      | +11.4     |
| MeCN           | -7.0      | -7.0      | +3.0      | +8.1      | +8.5      | +8.5      | +8.5      | +8.5      | +8.5      | +8.5      | +8.5      |
| DMF            | +2.4      | +2.4      | +8.5      | +9.2      | +9.1      | +9.1      | +9.1      | +9.1      | +9.1      | +9.1      | +9.1      |
| NH$_3$         |           |           |           |           |           |           |           |           |           |           | +5.9      |

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Figure 5. Influence of pressure on the water exchange rate constant for octahedral high spin divalent first row transition metal ions.

for the early to I₄ for the later elements, which can be explained by the progressive filling of the d orbitals and the decrease in ionic radii along the series. The small positive \( \Delta V^\ddagger \) found for DMF exchange on Mn⁴⁺, a value that was measured by three independent research groups, indicates that the bulkiness of the coordinated solvent molecules also plays a crucial role in controlling the degree of bond formation/bond breakage in the transition state. In addition, \( \Delta V^\ddagger \) exhibits a remarkable sensitivity for electronic effects, for instance in the case of induced solvent exchange reactions. Typically, \( \Delta V^\ddagger \) for water exchange on Fe(H₂O)₅²⁺ has a value of -5.4 cm³mol⁻¹, compared to a value of +7.0 cm³mol⁻¹ for the 750 times faster (at 25 °C) water exchange reaction on Fe(H₂O)₅OH²⁺. Similarly, \( \Delta V^\ddagger \) for solvent exchange on Cr(H₂O)₆³⁺, viz. -9.6 cm³mol⁻¹, is significantly smaller than for the induced solvent exchange on Cr(H₂O)₅OH²⁺, viz. +2.7 cm³mol⁻¹. Thus the trans-labilization effect of coordinated hydroxide causes a changeover from I₄ to I₁ of type of mechanisms, i.e. a more positive \( \Delta V^\ddagger \). The rate enhancement observed for acetonitrile exchange in going from Ru(η⁶-C₅H₅)(CH₃CN)₃²⁺ to Ru(η⁶-C₅H₅)(CH₃CN)₂₅⁺ is also accompanied by a significant increase in \( \Delta V^\ddagger \) and a changeover in mechanism from 1 to D, respectively.

Non-symmetrical ligand substitution reactions, for which the overall reaction volume \( \Delta V \neq 0 \), exhibit pressure dependencies that correlate closely with those found for solvent exchange processes in the case of complex formation reactions. The corresponding volume profiles illustrate the looseness or compactness of the transition state during the ligand substitution process. Such volume profiles can be used to investigate a possible changeover in mechanism caused by increasing steric hindrance in substitution reactions of square planar d⁸ metal complexes, that are usually accepted to follow an associative mechanism. For instance the introduction of methyl and ethyl substituents on the dienyl-ethylenediamine ligand of Pd(II) complexes causes a decrease of up to 6 orders of magnitude in the substitution rate constant, but does not affect the nature of the mechanism since both \( \Delta S^\ddagger \) and \( \Delta V^\ddagger \) values remain strongly negative. Two representative volume profiles are given in Figure 6, from which it follows that the transition state has a significantly lower partial molar volume than either the reactant or product states, demonstrating the associative nature of the substitution process.

Ligand substitution reactions of a number of organometallic complexes and clusters have also been investigated using high pressure kinetic techniques. For instance, the substitution of coordinated CO by P(OMe)₃ on Cr(CO)₅phen, phen = 1,10 phenanthroline, is characterized by a \( \Delta V^\ddagger \) of +13.8 ± 0.5 cm³mol⁻¹, whereas the reverse reaction exhibits a \( \Delta V^\ddagger \) of +19.2 ± 0.5 cm³mol⁻¹. The volume profile for this process (Figure 7) clearly demonstrates the significantly high partial molar volume of the transition state and the operation of a dissociative (D) mechanism. The overall reaction volume is slightly negative and indicates an overall volume decrease during the substitution process. A number of studies on photo-induced substitution reactions of metal carbonyl complexes that involve the displacement of coordinated solvent molecules, or the subsequent displacement of coordinated CO when the entering ligand is a bidentate species, clearly demonstrate the sensitivity of \( \Delta V^\ddagger \), and therefore the underlying mechanism, on the

\[ \text{Reactants} \rightarrow \text{Transition State} \rightarrow \text{Products} \]

**Figure 6.** Volume profiles for the reaction Pd(L)Cl⁺ + H₂O ⇌ Pd(L)H₂O⁺ + Cl⁻.
size of the central metal and the steric hindrance on the coordinating nucleophiles. In a similar way, ligand substitution reactions that occur in the electronically excited state of organometallic complexes also exhibit very characteristic pressure dependencies\(^{10}\). In general ligand field excitation is characterized by dissociative substitution reactions and positive ΔV\(^\#\) values, whereas metal to ligand charge transfer excitation is characterized by associative substitution reactions and negative ΔV\(^\#\) values.

Ligand substitution reactions of cobalamin (Vitamin B\(_{12}\)) has also attracted attention of high pressure kineticists. In these systems the usually kinetic inert Co(III) center is considerably labilized by the corrin ring, and there has been some disagreement in the literature concerning the mechanism of these substitution reactions\(^{16}\). The observed ΔV\(^\#\) for the forward and reverse reactions in (3) did not allow an unequivocal assignment of the mechanism in terms of I\(_d\) or D. For the reaction with pyridine (py) the observed rate constant

\[
B_{12}-H_2O^+ + L^- \rightleftharpoons B_{12}-L(1+\delta)^- + H_2O
\]  

(3)

reached a limiting value at high pyridine concentrations, which has now been interpreted in terms of a precursor formation step followed by a rate-determining ligand exchange process, i.e. an I\(_d\) mechanism, for which the corresponding volume profile is presented in Figure 8\(^{16}\). The significantly higher partial molar volume of the transition state compared to the reactant and product states, demonstrates the dissociative character of the reaction.

![Figure 8. Volume profile for the reaction B\(_{12}\)-H\(_2\)O\(^+\) + py \rightleftharpoons B\(_{12}\)-py\(^+\) + H\(_2\)O.](image)

The development of high-pressure pulse-radiolysis techniques\(^{16}\) has enabled the possibility to study the formation of metal-carbon bonds and the associated volume changes. Such processes are closely related to ligand substitution reactions since the aliphatic free radical (produced via pulse-radiolysis) must substitute a solvent molecule (usually water) on the metal center to produce the metal-carbon σ bond. A volume profile for the reaction of Co\(^{II}\)(nta)(H\(_2\)O)\(_2\) (nta = nitrotriacetic) with +CH\(_3\) to produce Co\(^{III}\)(nta)(H\(_2\)O)CH\(_3\)\(^{2-}\) clearly indicated that the process is controlled by solvent exchange on the Co\(^{II}\)(nta)(H\(_2\)O)\(_2\) species that follows an I\(_d\) mechanism. Subsequently, the effect of pressure on the formation of a series of organo-chromium(III) species were studied in an effort to obtain mechanistic information on solvent exchange and complex formation reactions of Cr(H\(_2\)O)\(_6\)\(^2+\). This reaction was studied for 10 different aliphatic radicals +R, which all exhibited very similar complex formation rate constants and small positive ΔV\(^\#\) values with an average of +4.3 ± 1.0 cm\(^3\)mol\(^{-1}\), independent of the nature of R\(^{11}\). These values suggested that Cr-C bond formation is controlled by solvent exchange on Cr(H\(_2\)O)\(_6\)\(^2+\), which follows an I\(_d\) mechanism, and a typical volume profile is reported in Figure 9. Thus breakage of the Cr\(^{II}\)-H\(_2\)O bond accounts for the volume increase in going to the transition state, which is followed by a significant volume collapse due to Cr-C bond formation and the contraction due to Cr\(^{II}\)-R → Cr\(^{II}\)-R\(^*\).

![Figure 9. Volume profile for the reaction Cr(H\(_2\)O)\(_6\)\(^2+\) + (CH\(_2\))\(_n\)OH \rightleftharpoons Cr\(\{\text{H}_2\text{O}\} \text{(CH}_2\text{)}\text{OH}\) + H\(_2\)O.](image)

**LIGATION REACTIONS**

The mechanistic understanding of the binding of small molecules such as O\(_2\), CO and NO to ferrous hemes and hemoproteins has received significant attention from kineticists. Such processes control the overall transport of such molecules in biological systems. The flash photolysis techniques outlined in (4) have been used to study ligand binding to two model heme systems, viz. protoheme dimethyl ester (PHDME) and monochelated protoheme [MCPH]\(^{12,13}\), and the effect of pressure on such processes is reported in Table II.

![Flash photolysis reaction](image)

(4)

The data show a clear correlation between ΔV\(^#\) and k\(_{on}\), viz. for the slower reactions recombinination of the separated pair is rate-limiting (negative ΔV\(^#\) value), whereas for the faster reactions the process becomes diffusion controlled in toluene and slows down with increasing pressure due to the increase.
Table II. $\Delta V^*$ data for the bimolecular addition of various neutral ligands to five-coordinate ferrous model heme complexes in toluene as solvent$^{12a}$

<table>
<thead>
<tr>
<th>Heme complex</th>
<th>L</th>
<th>$k_{oo}(25{^\circ}C)$ M$^{-1}$s$^{-1}$</th>
<th>$\Delta V^*$ cm$^{-3}$mol$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCPH</td>
<td>CO</td>
<td>1.1 x 10$^7$</td>
<td>-19.3 ± 0.4</td>
</tr>
<tr>
<td>MCPH</td>
<td>O$_2$</td>
<td>1.1 x 10$^7$</td>
<td>-11.3 ± 1.0</td>
</tr>
<tr>
<td>(MeNC) PHDME</td>
<td>MeNC</td>
<td>3.9 x 10$^8$</td>
<td>+11.6 ± 0.8</td>
</tr>
<tr>
<td>(t-BuNC) PHDME</td>
<td>t-BuNC</td>
<td>2.5 x 10$^8$</td>
<td>+9.9 ± 1.0</td>
</tr>
<tr>
<td>(1-Melm) PHDME</td>
<td>1-Melm</td>
<td>1.5 x 10$^8$</td>
<td>+10.9 ± 3.1</td>
</tr>
</tbody>
</table>

in viscosity of toluene. When the reaction of CO with MCPH is studied as a function of pressure in a highly viscous medium, viz. 90/10 (v/v) mineral oil/toluene, the observed reaction rate constant shows a changeover in rate-determining step with increasing pressure$^{12b}$. In the low pressure range bond formation is rate-determining and $\Delta V^*$ is negative, whereas in the high pressure range diffusion becomes rate-determining and $\Delta V^*$ is positive.

Similar techniques were used to study the pressure dependence of the binuclear association rate constant for the reaction of sperm whale myoglobin with O$_2$ and CO. Surprisingly, the slower binding of CO (5 x 10$^5$ M$^{-1}$s$^{-1}$ at 25 °C) exhibits a $\Delta V^*$ of -10 cm$^3$mol$^{-1}$, but the faster binding of O$_2$ (1 x 10$^7$ M$^{-1}$s$^{-1}$ at 25 °C) exhibits a $\Delta V^*$ of +8 cm$^3$mol$^{-1}$$^{12a}$. This apparent discrepancy motivated us to perform a detailed volume profile analysis for both reactions$^{12a,d}$, T-jump techniques were used to study the binding of O$_2$ and CO to deoxyxmyoglobin (Mb), whereas the reverse reactions, i.e. the release of O$_2$ and CO, were studied using a stopped-flow technique in which O$_2$ and CO were rapidly removed from the equilibrium mixture in a chemical way. In addition, the pressure dependence of the overall equilibrium constant was studied using UV-VIS techniques. From the $\Delta V^*$ data for the "on" and "off" reactions the overall reaction volume could be calculated, viz. $\Delta V = \Delta V_{on}\Delta V_{off}$, which turned out to be in good agreement with the directly determined values. These data could now be employed to construct a volume profile for the reaction of myoglobin with O$_2$ and CO, which have been combined in Figure 10. The volume profile for the binding of O$_2$ is characterized by the unexpected volume increase in going from the reactant to the transition state, as observed in the flash photolysis experiments, followed by a significant volume collapse in going to the product state. The observed volume increase has been ascribed to rate-determining movement of O$_2$ through the protein to the heme pocket, which may involve desolvation, some opening up of the protein as well as significant hydrogen bond formation with the distal histidine$^{12c}$. This is followed by rapid bond formation with the Fe(II) center, during which the metal center changes from high spin to low spin, the Fe(II) moves into the porphyrin plane, and some conformational changes on the protein may occur. These effects all add up to a large volume collapse of -23 cm$^3$mol$^{-1}$ observed from the transition state to the product state. The overall reaction volume of -18 cm$^3$mol$^{-1}$ demonstrates the large volume collapse caused by the bonding of O$_2$. The volume profile for the binding of CO shows the expected volume decrease for rate-determining bond formation going from the reactant to the transition state. Surprisingly, the reverse bond cleavage process is accompanied by a volume decrease, which may be related to the significantly different bonding mode of CO compared to O$_2$. It is known that the porphyrin iron binds O$_2$ under an angle of 115 ° and exhibits hydrogen bonding to histidine E7, which results in a sterically favoured orientation of O$_2$ in the porphyrin pocket. In contrast, CO does not show any hydrogen bonding, and its favoured linear bonding geometry is not possible due to histidine E7. This causes the heme pocket to widen significantly as compared to deoxyxmyoglobin due to steric tension. It follows that Fe-CO bond cleavage will be accompanied by a decrease in steric tension and an associated volume collapse due to reorganization of the protein pocket as CO leaves the iron coordination site. This significantly different bonding mode of CO must also account for the much higher partial molar volume of MbCO and the much smaller absolute reaction volume observed for the binding of CO, viz. -6 cm$^3$mol$^{-1}$, than for the binding of O$_2$.

**ELECTRON-TRANSFER REACTIONS**

Electron-transfer reactions of transition metal complexes are accompanied by a change in the oxidation state of the metal atom and the overall charge on the complex ion. This causes both intrinsic and solvational volume changes, such that it is reasonable to expect that electron-transfer reactions should exhibit characteristic $\Delta V^*$ values. In the case of symmetrical intermolecular reactions, the $\Delta V^*$ data can also be accounted for theoretically on the basis of the Marcus-Hush-Stranks treatment$^{13}$. A typical example is the self-exchange reactions of Fe(H$_2$O)$_6^{2+}$ and Fe(H$_2$O)$_6^{3+}$/Fe(H$_2$O)$_6^{2+}$ for which $\Delta V^*$ has the values -11 and +1 cm$^3$mol$^{-1}$, respectively$^{13}$. These values can be accounted for essentially quantitatively on the basis of an adiabatic intermolecular mechanism for the former and a hydroxo-bridged intramolecular mechanism that involves the release of a solvent molecule for the latter process.

In the case of non-symmetrical intermolecular reactions between complexes of the type Co$^{III}$((NH$_3$)$_3$X)$_{2+}$ and Fe(CN)$_4^{3-}$, $\Delta V^*$ usually has large positive values, which can be accounted for in terms of a large decrease in electrostriction due to charge neutralization, Co$^{III} + CO$ and Fe(CN)$_4^{3-}$ $\rightarrow$ Fe(CN)$_3^{2-}$X$^{3-}$. Similarly, the oxidation of L-ascorbic acid and the ascorbate ion by Fe(CN)$_3^{3+}$ exhibits $\Delta V^*$ values of ca. -16 cm$^3$mol$^{-1}$, which can again be accounted for in terms of the decrease in electrostriction when Fe(CN)$_3^{3+}$ is reduced to Fe(CN)$_3^{4+}$. Similar studies for the oxidation of L-ascorbic acid by Fe(H$_2$O)$_6^{3+}$ and Fe(H$_2$O)$_6$OH$^{2+}$ indicate $\Delta V^*$ values of +14 and +5 cm$^3$mol$^{-1}$, respectively$^{14a}$. For the less labile Fe(H$_2$O)$_6^{3+}$ species it was suggested that the redox process follows an intermolecular route, such that the significant volume increase will be related to the reduction of Fe(H$_2$O)$_6^{3+}$ to Fe(H$_2$O)$_6^{2+}$. In the case of the more labile Fe(H$_2$O)$_6$OH$^{2+}$ species, it was suggested that complex formation is rate-deter-

**Figure 10. Comparison of the volume profiles for the reactions**

$\text{Mb} + O_2 \rightarrow \text{MbO}_2$ and $\text{Mb} + CO \rightarrow \text{MbCO}$. 

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mining, followed by a rapid intramolecular electron-transfer reaction, such that ∆Vf correlates with that found for solvent exchange on Fe(H₂O)₃OH²⁺. Recent work has shown that long-distance electron-transfer reactions between Ru(II) complexes and ferri cytochrome c exhibit ∆Vf values between -16 and -18 cm³ mol⁻¹ for both intermolecular and intramolecular processes⁵c, 14c. These values are once again assigned to the significant increase in electrostriction when Ru(III) is oxidized to Ru(II) 14c.

**ADDITION AND RELATED PROCESSES**

Up to now we have seen that bond formation processes are characterized by negative ∆Vf values, and it is therefore not surprising that oxidative addition reactions are characterized by significantly negative ∆Vf values, partly due to bond formation and partly due to a decrease in the size of the metal center during the oxidation process⁶b. Typical ∆Vf values can be as large as -30 to -40 cm³ mol⁻¹, and exhibit a strong solvent dependence due to the significant electrostriction contribution.

It was recently shown that [2+2] cyclo-addition reactions on the coordinated ligand of pentacarbonyl carbene complexes of Cr and W, as well as insertion reactions into the metal-carbene bond of pentacarbonyl(carbene)chromium and -tungsten complexes, all exhibit ∆Vf values between -15 and -25 cm³ mol⁻¹ 15a,b. It follows that such processes are in general accelerated by pressure, which could have some synthetic application ⁴c. Addition reactions of α,β-unsaturated carbene complexes shown in (5) exhibit ∆Vf values around -16 cm³ mol⁻¹ in acetonitrile, which become more negative in going to less polar solvents 15c. These data exhibit a good correlation with the solvent parameter qₚ (i.e. the pressure derivative of q) as shown in Figure 11, from which it follows that the intercept, ∆Vfₚₚ₀, has a value of ca. -14 cm³ mol⁻¹. A similar study for the addition of a series of p-substituted anilines to a Fischer carbene complex as shown in (6), resulted in the data summarized in Table III 15d.

The observed second order rate constant shows a good correlation with the increase in basicity of the amine, which is accompanied by a significant decrease in ∆Hf and an increase in ∆Vf to more positive values. These trends demonstrate that the slow reactions (high ∆Hf) occur via a late transition state (strongly negative ∆Vf), whereas the fast reactions (low ∆Hf) occur via an early transition state (more positive ∆Vf). This demonstrates the usefulness of combined ∆Hf and ∆Vf data to obtain information on the location of the transition state in terms of “early” or “late”, as illustrated schematically in Figure 12.

![Figure 11. Plot of ∆Vf versus qₚ for reaction (5) with M = W, R¹ = Ph and R² = Et. Solvents: acetonitrile (1), 1,2-dichlorobenzene (2), chlorobenzene (3), benzene (4), n-heptane (5).](image)

![Figure 12. Schematic presentation of the concept of an “early” and “late” transition state in terms of enthalpy and volume effects along the reaction coordinate.](image)

**CONCLUSIONS**

The analysis of a chemical process in terms of volume changes along the reaction coordinate can help us to visualize the nature and structure of the transition state in terms of intrinsic and solvational changes in partial molar volume. With the data presently available, intrinsic volume changes in
organic, organometallic and bioinorganic systems are fairly well understood and can be interpreted with confidence. This is, however, not the case for solvational volume changes and much more work is required to resolve the origin of all the observed effects. There are many instances in which it is experimentally impossible to construct a volume profile, for instance due to the occurrence of subsequent reactions or the irreversibility of the process as found in many electron-transfer and photo-induced reactions. Nevertheless, the volumes of activation for such processes can still be employed very successfully to gain information on the nature of the transition state.

It is important to realize that the presented interpretations of $\Delta V^\circ$ data are all based on a simplified version of the transition state theory, which has its limitations and restrictions, and various modifications are being discussed. Notwithstanding this complication that also concerns the interpretation of the temperature dependence of a chemical reaction, it is clear from the given examples that the additional physical parameter pressure has added a decisive dimension to mechanistic studies of inorganic, organometallic and bioinorganic reactions in solution. The fact that the rate-determining step of a particular process exhibits a characteristic pressure dependence, creates the possibility to tune the reactivity of particular systems via the application of moderate pressures. This can for instance lead to the selective synthesis of particular reaction products in cases where product distribution proves to be pressure dependent, or to the optimization of the design of industrial chemical reactions.

It is the authors personal opinion that when dealing with mechanistic studies in general, one should investigate as many chemical and physical variables as possible in order to obtain as much indirect information as possible on the nature of the underlying reaction mechanism. Only then can the suggested mechanism come close to the “real” mechanism, which is a goal set by many kineticists but not always accomplished. If we can contribute in this way to a better understanding of the mechanism of chemical reactions in solution, then we have fulfilled our educational commitments and have added a drop to the bucket of existing knowledge.

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REFERENCES AND NOTES

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