Kinetic detection of osmium(VI) ester intermediates during the OsO₄-mediated aqueous dihydroxylation of chloroethylenes

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Abstract
The kinetics and mechanism of the cis dihydroxylation of cis-1,2-dichloroethylene, trans-1,2-dichloroethylene, and trichloroethylene by osmium tetroxide was studied systematically in aqueous solution. The stoichiometry of the process was determined based on the principle of continuous variation of reactant ratios with spectrophotometric detection. The results always showed 1:1 stoichiometry, which is in agreement with dihydroxylation. All three reactions were found to proceed in two distinct steps. The first step occurred on a time scale of seconds and was associated with a minor change in absorbance and was identified as the formation of a 1:1 adduct between the two reagents, which is the osmium(VI) ester that plays a decisive role in catalytic applications. This species is formed in an equilibrium that is very much shifted toward the reactants, so the osmium(VI) complex is a short-lived intermediate of the process, which is detected kinetically, but its concentration is never high enough for structural characterization. The second reaction is accompanied by major spectral changes; it involves the formation of the final products. Our results clearly show that it is possible to detect the intermediate of the process by careful kinetic studies. It is also possible that the same strategy might be successful in other OsO₄-dependent dihydroxylation processes.

KEYWORDS
aqueous chemistry, osmium (VI) ester, oxidation of alkenes, relaxation kinetics, stopped-flow

1 | INTRODUCTION

The cis dihydroxylation of carbon-carbon double bonds using osmium tetroxide is a frequently used reaction in organic chemistry, it is probably the most reliable method for this particular transformation.⁵ Osmium tetroxide has a surprisingly high number of additional practical applications such as a staining and fixative agent for treating biological samples in electron microscopy and electrochemical analysis,⁶ but it is rare and expensive. Therefore, it is often imperative to use osmium as a catalyst in the process, which can be readily achieved by adding a less expensive stoichiometric oxidizing agent that is able to transform the product osmium(VI) back...
into osmium tetroxide. Hydrogen peroxide, tert-butyl-hydroperoxide, N-methylmorpholine N-oxide, dioxygen, chlorate ion, hypochlorite ion, and periodate ion have all been reported to be suitable for this purpose under certain conditions.\cite{1,2,7,8} Despite the vastly different properties of these oxidants, it is a common aspect that they do not react with the substrate olefin directly, and their only role is to reproduce the initial form of the catalysts.\cite{1,2} Therefore, it is clear that the kinetics and properties of these oxidants, it is a common aspect that the efficiency of a catalytic method are dominantly determined by the interaction of the oxidizable substrate with osmium tetroxide, and kinetic studies on this process in the absence of an extra oxidizing agent may provide insightful information on the mechanism and the potential application of the method.

There seems to be quite general agreement on the mechanism of the cis dihydroxylation of carbon-carbon double bonds by osmium tetroxide.\cite{1,2,7,8} The formation of the catalytically active osmium(VI) ester usually assumed to contain a tetrahedral d^2 metal center by the addition of OsO_4 to the double bond is a common feature of these processes. The ester is a highly reactive intermediate that may have fleeting existence in solution but unlikely to be stable in the solid phase. In some cases, penta-coordinated dimeric monoester species Os_2O_5(O_2R)_2 and monomeric diester complexes OsO_2(O_2R)_2 were characterized structurally, and the mechanistic interpretations were also supported by quantum chemical calculations.\cite{14-22} However, these molecules might very well represent unreactive dead ends as far as the dihydroxylation process is concerned. The process can be carried out in an enantioselective way by adding coordinating chiral amines as coligands.\cite{2,8,9,11,13} These are understood to exert an influence on the process by coordinating to the osmium center directly.

Trichloroethylene (TCE) was in widespread industrial use for decades in the 20th century as a vapor degreasing agent for metal objects and also as an anesthetic and disinfectant in medicine. By now, its carcinogenic properties have been demonstrated, and its use was mostly phased out in developed countries.\cite{23} However, it is still present in the environment in many industrial sites, and its general lack of reactivity is still an impediment in remediation technologies.\cite{24-27} From a purely chemical point of view, it is also interesting that the C-H acidity of the molecule can be clearly detected in hydrogen isotope exchange reactions, similarly to the two isomers of 1,2-dichloroethylene.\cite{28}

This work reports our serendipitous finding that the kinetics of the formation of the key intermediate can be detected during the reaction between osmium tetroxide and three different chloroethylenes in aqueous solution. A full kinetic analysis showed that this highly reactive intermediate is indeed present in low concentrations, which precludes structural characterization, but it is definitely the species that is in the center of the catalytic activity. A formal kinetic scheme is proposed to interpret the kinetic findings with the full consideration of the various labile equilibria of osmium(VIII) species in aqueous solution.\cite{29-32}

## 2 | EXPERIMENTAL

### 2.1 | Materials

All chemicals used were commercially available and reagent grade. NaOH, TCE, trans-1,2-dichloroethylene (t-DCE), cis-1,2-dichloroethylene (c-DCE), and OsO_4 were supplied by Sigma and used without further purification. NaClO_4 (Sigma) was used to maintain constant ionic strength (denoted as I in this article) by adding an appropriate amount of 6.2 mol/dm^3 stock solution to the reaction mixture. The solubilities of chloroethylenes are somewhat limited in water (mmol/dm^3 level), and stock solutions were prepared by the addition of excess of chloroethylenes in water to ensure saturated solution conditions.

The OsO_4 stock solution was prepared by dissolving a measured amount of solid OsO_4 in NaOH solution to ensure the stability of OsO_4 in its mono/di-hydroxo form. (WARNING! Solid OsO_4 is known to be highly volatile, and its vapors are very harmful for the eyes.) Basic conditions also decrease the volatility of osmium(VIII) very substantially. The osmium concentration of this stock solution was double checked by ICP-AES measurements shortly after preparation and found to be in good agreement with the values calculated based on mass and volume measurements. Final concentrations were 0.10 mol/dm^3 for NaOH and 50 mmol/dm^3 for OsO_4. The concentration of OsO_4 in this stock solutions was also monitored regularly by spectrophotometric measurements. The solution was stable for 5 days at 4 °C. Under strongly basic conditions (2 mol/dm^3 NaOH or higher), OsO_4 can be reduced to osmium(VII) and later to osmium(VI) by water during a few days at room temperature as indicated by spectral change.\cite{29,30} No such spectral change was observed in 0.10 mol/dm^3 NaOH under our conditions.

### 2.2 | Instruments

Kinetic studies were performed on an Applied Photophysics SX-20 Stopped-flow spectrophotometer (fast complexation reactions) and Perkin-Elmer Lambda 25 spectrophotometer equipped with thermostated cell block (slow redox processes). The dead time of the stopped-flow
The instrument was experimentally determined to be 1.0 ms. Kinetic curves were evaluated, and rate constants were fitted by the respective built-in softwares of the instruments. Each point presented is the average of at least seven runs in the complexation study, while the same averaging was done for at least three runs in the redox study.

3 | RESULTS

3.1 | General observations

First, the UV-visible spectral changes of the reaction between osmium tetroxide and the three chloroethylenes were studied in basic aqueous solution. No reaction was observed with 1,1-dichloroethylene, the use of chloroethylene (better known as vinyl chloride) was not even attempted because it is a gas under normal conditions. In addition, no reaction was observed at room temperature even with the studied three chloroethylenes when the conditions were less basic than in the experiments shown in this study.

In a series of experiments, the ratio of reactants (OsO₄ vs ethylenes) was changed systematically to explore the stoichiometry of the redox reactions. Samples with different ratios were prepared and left to react for 24 hours. This period of time was sufficient for the completion of the redox process; the UV-vis spectrum of the solution was measured. The absorbance at a few selected wavelengths was plotted as a function of the reactant ratio. This plot is expected to give a sharp break point that indicates the stoichiometric ratio of reactants in the process (this technique is also known as the method of continuous variation). For all chloroethylenes, evidence for 1:1 stoichiometry was obtained. A plot based on a typical series of experiments is presented in Figure 1 for the TCE-OsO₄ system. Two more plots for the other two substrates are shown in the Supporting Information. The spectral changes indicate the reduction of osmium(VIII) to osmium(VI) clearly: The initial spectrum matches the known spectrum of aqueous osmium(VIII), while the last one is the same as the independently known spectrum of osmium(VI) in water. The appearance of the isosbestic point at about 260 nm implies that the studied process can be characterized by a single stoichiometric equation for TCE, the same was observed for c-DCE and t-DCE. The inset in Figure 1 shows absorbance as function of the TCE/Os(VIII) ratio at 320 nm. The break point occurs at 1:1 ratio, so a 1:1 TCE:Os(VIII) stoichiometry is concluded. This is in agreement with the expectation based on previous knowledge about the cis dihydroxylation reaction. However, it is also clear from the previous literature of the oxidation of chloroethylenes that the cis-dihydroxylated species hydrolyze to glyoxal and glyoxilic acid in fast reactions, which are the final detectable organic products.

The kinetics of the three processes was also studied in detail. Because the dominating absorbing species are the different forms of osmium, and chloroethylenes have negligible UV-absorption in this region; OsO₄ was selected as the substoichiometric reagent, and the chloroethylenes were used in excess. Figure 2 shows representative kinetic curves for all three substrates measured at 280 nm. They confirm that the spectral changes take at least a few hours to complete in each of the systems. All the curves...
gave excellent fits to exponential functions, which confirmed the fact that the rates of these processes show first order dependence on the concentration of the substoichiometric reagent (OsO₄). The initial absorbances in these curves matched the expectations based on the known spectrum of osmium(VIII) within experimental error. These facts, especially when considered together with the observed isosbestic point in Figure 1, gave no reason to suspect a multi-step process.

However, when the kinetic experiments were repeated on a short time scale in a stopped-flow instrument, the occurrence of a fast reaction was revealed. Figure 3 shows curves measured for two different ethylenes. These traces also gave quite acceptable fits to exponential functions, which implies pseudo-first order behavior. The amplitude of the process (ie, the difference in the initial and the final absorbance) was really tiny, smaller than 0.005 units in all cases. Of course, absolute absorbances cannot be measured reliably at this precision. However, the time dependence detected within a single curve still holds valuable information, and the pseudo-first order rate constant could be determined quite reliably. This is the consequence of the negligible random variation of the absorbance due to the fluctuation of the base line in such a short time scale. In this sense, this small signal resembles the general practice followed during relaxation measurements (temperature jump, pressure jump, and field jump).[39,40]

On the basis of these observations, it was concluded that the reaction between chloroethylenes and OsO₄ actually proceeds in two distinct steps under basic conditions. In the faster step, TCE, c-DCE, or t-DCE react with osmium(VIII) in a few seconds as evidenced by the small decrease of absorbance, which was detected in the entire spectral region above 280 nm (OsO₄ absorption). We attribute this absorbance change to the formation of the commonly assumed intermediate osmium(VI) ester complex\(^{[1,2]}\) In the slower step, the reaction proceeds via the decomposition of the complexes in each investigated system, and the appropriate form of aqueous osmium(VI) is formed from the intermediate on a time scale of a few hours.

Detailed studies were carried out on the kinetics of both the fast and slow processes in all three systems, where the reactant concentrations and the pH were changed in a systematic way to explore the rate equations of the processes. The dependence of the measured pseudo-first order rate constants on the ethylene concentrations for the fast step is shown in Figure 4. It is notable that neither the concentration nor the identity of the chloroethylene influences the value of this parameter. There is some scatter in the data, but that seems to show no particular tendency so we fully assign this scatter to random noise, which is not unreasonable given the small amplitude of the absorbance changes followed. It should also be noted that the concentration of TCE could not be increased as much as that of the two dichloroethylenes because its solubility is more limited in water.

Figure 5, on the other hand, shows that the pseudo-first order rate constant of the fast process does depend on the pH. The identity of the chloroethylene still does not make a difference here, ie, trichloroethylene and the two dichloroethylenes have very similar values at the same pH, but the pseudo-first order rate constants at lower basicities are significantly higher.

The experimental points in Figure 5 were fitted to Equation (1), which is a very general one to interpret

![FIGURE 3 Representative curves on short time scale. Conditions: [Os (VIII)] = 0.20 mmol/dm³ (trichloroethylene [TCE]) and 0.16 mmol/dm³ (cis-dichloroethylene [DCE]), [NaOH] = 0.10 mol/dm³, I = 0.10 mol/dm³, T = 25.0°C](image)

![FIGURE 4 Dependence of \(k_{\text{obs}}\) on the chloroethylene concentrations in the fast step of their reactions with osmium (VIII). Conditions: [Os(VIII)] = 0.20 mmol/dm³, [NaOH] = 0.10 mol/dm³, I = 0.10 mol/dm³, T = 25.0°C](image)
pH-dependences\cite{41}, although the equilibrium process characterized by the pK in the formula is not identified at this point, this will only be done in Section 4. Rate constants \( k_{\text{acid}} \) and \( k_{\text{base}} \) correspond to the reactions of the acidic and basic forms of the same reactant, respectively.

\[
k_{\text{obs}} = \frac{[H^+]k_{\text{acid}} + 10^{-pK}k_{\text{base}}}{[H^+] + 10^{-pK}}.
\]

The fit was carried out in a global way, so common parameter values were forced for the three different substrates. \([H^+]\) in this formula was calculated from the hydroxide ion concentrations with the known value of the ionic product of water (if defined by concentrations rather than activities\cite{42}, \( pK_w = 13.58 \) at \( I = 0.10 \) mol/dm\(^3\) and 298.15 K\cite{43}). It was observed that the estimated value of \( k_{\text{base}} \) was very low and substantially smaller than its standard error. This is an indication that the basic pathway does not contribute significantly to the observed rate. Therefore, \( k_{\text{base}} \) was fixed to zero. The remaining two parameters were determined as \( k_{\text{acid}} = 4.3 \pm 0.1 \) s\(^{-1}\) and \( pK = 12.0 \pm 0.1 \). The best fit is shown as a line in Figure 5. Although the fit of the data points is far from perfect, this is still seen as a reasonable one given the small signal followed and the fact that three different reactions were considered in a common fit.

As was already mentioned, the slow process was also very well described by an exponential curve. Figure 6 shows the dependence of the pseudo-first order rate constants on the substrate concentrations in all systems. Unlike in the fast step, the different substrates give noticeably different rate constant and reaction rates. \( t\)-DCE is the most reactive, whereas TCE takes the longest time to complete the reaction under otherwise identical conditions. The plots in Figure 6 also show that the pseudo-first order rate constants are directly proportional to the substrate concentration (which was used in excess). These observations are normally indicative of a simple first order process with respect to the substrate, and therefore an overall second order reaction. The slopes of the straight lines give the second order rate constants in each system, they were determined as \( k_{\text{2nd}} = 0.036 \pm 0.001 \) dm\(^3\) mol\(^{-1}\) s\(^{-1}\) (TCE), \( k_{\text{2nd}} = 0.111 \pm 0.008 \) dm\(^3\) mol\(^{-1}\) s\(^{-1}\) (c-DCE), and \( k_{\text{2nd}} = 0.53 \pm 0.01 \) dm\(^3\) mol\(^{-1}\) s\(^{-1}\) (t-DCE).

The pH-dependence of the slow pseudo-first order rate constant was studied in all three systems as well. The results are shown in Figure 7. It was mentioned earlier that the different substrates show characteristically different reactivities on the slow time scale. A change in the pH influences the measured pseudo-first order rate constants in all three cases. However, unlike to the fast process, the
values tend to increase as the pH increases. The experimental points measured in all three cases were fitted to the function given in Equation (1). Again, this formula gave an acceptable interpretation of the observed dependence. The parameters were determined separately in each case: \( k_{\text{acid}} = (2.3 \pm 1.0) \times 10^{-5} \text{ s}^{-1} \), \( k_{\text{base}} = (6.4 \pm 0.9) \times 10^{-5} \text{ s}^{-1} \), \( pK = 12.4 \pm 0.4 \) (TCE), \( k_{\text{acid}} = (1.5 \pm 0.3) \times 10^{-4} \text{ s}^{-1} \), \( k_{\text{base}} = (3.4 \pm 0.1) \times 10^{-4} \text{ s}^{-1} \) and \( pK = 11.9 \pm 0.2 \) (c-DCE), and \( k_{\text{acid}} = (2.5 \pm 2.3) \times 10^{-4} \text{ s}^{-1} \), \( k_{\text{base}} = (1.44 \pm 0.08) \times 10^{-3} \text{ s}^{-1} \) and \( pK = 11.9 \pm 0.3 \) (t-DCE). It is seen that \( k_{\text{acid}} \) for t-DCE is barely larger than its standard error, which indicates that the corresponding pathway has a very limited contribution to the overall reaction progress under the experimental conditions.

4 | DISCUSSION

First, it must be pointed out that this study has a highly unusual aspect, which is finding the fast process. In routine high quality kinetic studies, this process would probably not have been recognized as the easily detectable slow process is first order with respect to both of its reagents, and even a comparison of expected and detected initial absorbances\(^{[44]}\) does not necessarily imply the need for including (or indeed, searching for) a fast process.\(^{[45]}\)

So the discovery of the fast part of the reactions seems serendipitous.

To interpret the kinetic findings, the highly labile equilibria in the solutions of OsO\(_4\) have to be considered first. Osmium(VIII) is present as molecular OsO\(_4\) in neutral aqueous solutions and shows very peculiar vibrational fine structure in its UV-vis spectrum.\(^{[30]}\) In basic medium, osmium(VIII) forms two perosmate ions, OsO\(_4\)(OH)\(^{-}\) and OsO\(_4\)(OH)\(^{2-}\) in a stepwise manner:\(^{[29–31]}\)

\[
\text{OsO}_4 + \text{H}_2\text{O} = \text{OsO}_4\text{OH}^- + \text{H}^+ \quad K_{a1}, \tag{2}
\]

\[
\text{OsO}_4\text{OH}^- + \text{H}_2\text{O} = \text{OsO}_4\text{(OH)}_2^{2-} + \text{H}^+ \quad K_{a2}. \tag{3}
\]

The corresponding \( K_{a1} \) and \( K_{a2} \) values were reported in the literature as \( 8.69 \times 10^{-13} \text{ M} (pK_{a1} = 12.2) \) and \( 7.58 \times 10^{-15} \text{ M} (pK_{a2} = 14.4)\).\(^{[29]}\) In agreement with these values, the single negative aqueous form of osmium(VIII) dominates under our conditions, which explains why vibrational fine structure\(^{[30]}\) is not detected in the spectra shown in Figure 1. At the lower end of the pH range of these studies, the neutral OsO\(_4\) species is in a slight excess over the mononegative ion. These equilibria only involve simple proton transfer processes, which are understood to be very fast (equilibrium reached within 1 μs under all practical conditions), so only the equilibrium constants need to be considered in our kinetic studies.

We found that all our kinetic findings are interpreted by the following kinetic scheme where the notation E stands for the oxidizable chloroethylene derivative:

\[
\text{OsO}_4 + \text{E} \xrightleftharpoons{k_{\text{a}}}{\text{OsO}_4\text{(O}_2\text{E)}} \quad (4)
\]

\[
\text{OsO}_4\text{OH}^- + \text{E} \xrightleftharpoons{k_{\text{b}}}{\text{OsO}_4\text{(O}_2\text{E})\text{OH}^-} \quad (5)
\]

\[
\text{OsO}_4\text{(O}_2\text{E}) \xrightarrow{k_{\text{a}E}} \text{OsO}_4 + \text{E} \quad (6)
\]

\[
\text{OsO}_4\text{(O}_2\text{E})\text{OH}^- \xrightarrow{k_{\text{b}E}} \text{OsO}_4\text{OH}^- + \text{E} \quad (7)
\]

\[
\text{OsO}_4\text{(O}_2\text{E}) + \text{H}_2\text{O} = \text{OsO}_4\text{(O}_2\text{E})\text{OH}^- + \text{H}^+ \quad K_{aE} \quad (8)
\]

\[
\text{OsO}_4\text{(O}_2\text{E}) + 4\text{H}_2\text{O} \xrightarrow{k_{2a}} \text{OsO}_4\text{(OH)}_4^{2-} + \text{E(OH)}_2^+ + 2\text{H}^+ \quad (9)
\]

\[
\text{OsO}_4\text{(O}_2\text{E})\text{OH}^- + 3\text{H}_2\text{O} \xrightarrow{k_{2a}} \text{OsO}_4\text{(OH)}_4^{2-} + \text{E(OH)}_2^+ + \text{H}^+ \quad (10)
\]

This scheme may seem complicated at first sight, but in fact it is quite simple. The essence is that the osmium (VI) ester intermediate forms in a reversible way form osmium(VIII) and the chloroethylene (Equations 4 and 6) then undergoes decomposition giving the final cis diol product (Equation 9). All of these reactions have their counterpart where the osmium center is coordinated to a hydroxide ion (Equations 5, 7, and 10). Finally, not only the pre-equilibrium shown in Equation (2) is
established between the two osmium(VIII) aqua forms but also the two different osmium(VI) esters (the one with a hydroxide ion coordinated and the one without) interconvert in a very rapid process (Equation 8). In effect, this means that the scheme has one reversible reaction (adduct formation between the reactants) and one irreversible (formation of the final products), but these are also possible in the same way when a hydroxide ion is attached to the osmium center.

The first, reversible step occurs on the fast time scale. Note that a tiny absorbance decrease is detected for this process. This is in agreement with the generally agreed formulation of the active intermediate as an osmium (VI) ester. If it contains a d^2 metal center, its spectral characteristics must resemble those of the product aqueous osmium(VI) and not the initial osmium(VIII). In agreement with previous literature data, Figure 1 shows that osmium(VI) has generally much smaller light absorption above 260 nm than osmium(VIII), the decrease in absorbance during the fast process is quite understandable. Furthermore, the tiny absorbance change of the process is caused by the fact that it reaches equilibrium at very low conversion. The small absorbance decrease of the curves in Figure 3 relative to the total measured absorbance implies that the conversion is not more than 2% to 3% in equilibrium.

The measured kinetics of the process, and especially the independence of the measured pseudo-first order rate constants of the substrate concentration (Figure 4) deserves some additional attention here. From the theory of relaxation experiments,[39,44] it is well known that the lifetime in an equilibrium that is heavily shifted toward the initial reagents is set by the rate constant of the reverse reaction even if the process is advancing in the forward direction. This is a pivotal point in the present study, so we also report a derivation by a method that is independent of the kinetic relaxation theory[39] because the product complex is absent at the initial time, which is very atypical in relaxation.

The rate equation of a reversible reaction in which E and O come together to form the adduct EO takes the following form:

\[
\frac{d[EO]}{dt} = k_+([E]_0 - [EO])([O]_0 - [EO]) - k_- [EO]. \tag{11}
\]

It is shown in the Supporting Information that this equation can be solved analytically (the solution of a more general version of this rate equation is also found in the literature[45]). The precise solution under conditions where EO is not present in the initial mixture is given as:

\[
[EO] = \frac{A}{2k_+} - \frac{B(A + B + Ae^{-Rt} - Be^{-Rt})}{2k_+ (A + B - Ae^{-Rt} + Be^{-Rt})}.
\]

\[
A = k_+ [E]_0 + k_+ [O]_0 + k_- B = \sqrt{A^2 - 4k_+^2 [E]_0 [O]_0}. \tag{12}
\]

As given above, A and B are combination parameters in this formula that have been introduced to shorten the mathematical expressions. When the equilibrium is shifted toward the initial reagents, the value of the reverse rate constant is large and dominates in the definition of A because \(k_- > k_+ [E]_0 + k_+ [O]_0\). As derived in the Supporting Information, Equation (12) can be simplified into the following, exponential form under these conditions:

\[
[EO] \approx \left( \frac{[E]_0}{2} + \frac{[O]_0}{2} - \frac{k_-}{2k_+} - \left( \frac{[E]_0}{2} + \frac{[O]_0}{2} + \frac{k_-}{2k_+} \right)^2 - [E]_0 [O]_0 \right) \left( 1 - e^{-k_- t} \right). \tag{13}
\]

So indeed, the pseudo-first order rate constant of the process will simply be \(k_-\), ie, the first order rate constant of the reverse process.

This effect explains why Figure 4 shows that the pseudo-first order rate constant of the fast process is independent of the substrate concentrations. It is more interesting to consider why it is independent even of the identity of the substrate. Here, it is probably best to consider that the reaction is the dissociation of the osmium (VI) ester where the reactive center (the metal atom) is not in the direct vicinity of the substituents that were on the double bond in the original substrate. Therefore, it is arguable that the rate of the dissociation of this ester is not very much influenced by the exchange of hydrogens to chlorine on the carbon atoms. It is also clear how the pH dependence shown in Figure 5 arises: the OsO₂(O₂E) and OsO₂(O₂E)OH⁻ have different dissociation rate constants as the coordination of the hydroxide ion occurs directly on the metal center. In terms of elementary rate constants shown in the scheme of Equations (4) to (10), the pseudo-first order rate constant of the fast process can be given as:

\[
k_{obs} = \frac{[H^+] k_{1a} + K_{ae} k_{1b}}{[H^+] + K_{ae}}. \tag{14}
\]

At this point, it should be remarked that the three different chloroethylenes studied here might as well have three different pK_{ae} values. However, as the pseudo-first order rate constants measured for the fast process do
not depend on the identity of the substrate significantly, neither does $pK_{aE}$. It can also be observed that this $pK_{aE}$ is in fact not significantly different from $pK_{a1}$, which implies that the affinity of the metal center toward hydroxide ion does not even depend on whether the bonded oxygens are involved in further ester bonds or not.

The kinetics of the slow step can be interpreted in a more commonly used sequence of thought\textsuperscript{[41,44]} based on the proposed scheme. On this time scale, the two reversible reactions in Equations (4) to (7) already reach equilibrium, as this equilibrium is shifted toward the initial reagents, the concentrations of the osmium(VI) esters can be given using the concentrations of the reactants at every time instant:

$$[\text{OsO}_2(\text{O}_2\text{E})] = \frac{k_{1a}}{k_{1a-}}[\text{OsO}_4][E],$$

$$[\text{OsO}_2(\text{O}_2\text{E})\text{OH}^-] = \frac{K_{aE}}{[\text{H}^+]}[\text{OsO}_2(\text{O}_2\text{E})] = \frac{k_{1a}K_{aE}}{k_{1a-}[\text{H}^+]}[\text{OsO}_4][E].$$

(15)

(16)

It must also be considered that the total amount of aqueous osmium(VIII) is distributed between two different forms, the neutral molecule and the mononegative ion:

$$[\text{Os(VIII)}] = [\text{OsO}_4] + [\text{OsO}_4(\text{OH})^-] = [\text{OsO}_4] + \frac{K_{a1}}{[\text{H}^+]}[\text{OsO}_4].$$

(17)

The formation rate of the cis-diol is then:

$$\frac{d[E(\text{OH})_2]}{dt} = k_{2a}[\text{OsO}_2(\text{O}_2\text{E})] + k_{2b}[\text{OsO}_2(\text{O}_2\text{E})\text{OH}^-] = (k_{2a} + \frac{k_{2b}K_{aE}}{[\text{H}^+]}k_{1a-})[\text{OsO}_4][E] = \frac{k_{2a}[\text{H}^+]}{[\text{H}^+]} + \frac{k_{2b}K_{aE}k_{1a}}{[\text{H}^+]} + \frac{k_{1a}}{k_{1a-}}[\text{Os(VIII)}][E].$$

(18)

Given that the substrate E is used in large excess over osmium(VIII) in the experiments, the pseudo-first order rate constant of the slow processes is expected to be:

$$k_{obs} = \frac{k_{2a}[\text{H}^+]}{[\text{H}^+] + K_{a1}} + \frac{k_{2b}K_{aE}k_{1a}}{k_{1a-}}[E].$$

(19)

This is in excellent agreement with the experimental observations shown in Figures 6 and 7. It should be noticed that $K_{a1} \cong K_{aE}$ was already established at the analysis of the fast step, so the analogy with Equation (1) is complete with $k_{\text{acid}} = [E]k_{2a}k_{1a}/k_{1a-}$ and $k_{\text{base}} = [E]k_{2b}k_{1b}/k_{1b-}$. It is also remarkable that, unlike for the fast process, the pseudo-first order rate constant of the slow phase depends on the identity of the substrate very significantly. The $k_{1a-}$ and $k_{1b-}$ values already led to the argument that for the properties of the osmium (VI) ester, the substituents farther away from the metal center do not seem to exert much kinetic influence. This same line of thought leads to the estimate that $k_{2a}$ and $k_{2b}$ are probably close to independent of the identity of the substrate. Therefore, the reason for the kinetic difference in the slow process between the three studied substrates must lie primarily in the $k_{1a}$ values, which characterize the rate of association of osmium tetroxide with the oxidizable chloroethylene.

An essential point in our interpretation is that the adduct OsO$_2$(O$_2$E) is a minor intermediate because its formation equilibrium is shifted towards the reactants. In other words, the thermodynamic stability of the osmate ester is low. At first sight, this might seem to contradict well established quantum chemical calculations,\textsuperscript{[46,47]} which showed that the osmate ester formation is highly exergonic. The calculated energy changes for ethylene as an oxidizable alkene were $-21.2$ kcal mol$^{-1}$ ($=-88.7$ kJ mol$^{-1}$)$^{[46]}$ and $-18.1$ kcal mol$^{-1}$ ($=-75.7$ kJ mol$^{-1}$).\textsuperscript{[47]} Without the intention to question the accuracy of these results, the next paragraphs will show that the results of these quantum chemical calculations are irrelevant for the present study.

First, it is pointed out that the reported energy values are not Gibbs free energies and are therefore unsuitable for predicting the actual driving force of the adduct formation directly, separate considerations about the entropy changes would be needed. As this is an adduct formation reaction, the entropy change of the process must certainly be highly negative, which would result in a Gibbs free energy that is considerably less negative than $-88.7$ kJ mol$^{-1}$ or $-75.7$ kJ mol$^{-1}$. However, there seems to be insufficient data to make this consideration truly quantitative.

A much more important point is that the process was experimentally studied in aqueous solution in this work. In this phase, thermodynamic data are available in the literature to develop some truly quantitative and meaningful thermodynamic analysis. The starting point is the following aqueous standard two-electron reduction potential, which is known experimentally: $^{[30,48,49]}$

\[ \text{OsO}_4 + 2 \text{H}_2\text{O} + 2 e^- \rightarrow \text{OsO}_2(\text{OH})_4^{2-} \quad E^\circ = 0.46 \text{ V}. \]

At first sight, this reduction potential may seem very low and it might also be somewhat counterintuitive that OsO$_4$ is only a mild oxidant in water. As a qualitative confirmation of this weak oxidizing power, we carried out experiments with iodide ion ($E^\circ = 0.54$ V). No
reaction was observed at all, which is in agreement with the literature value of the standard reduction potential for the aqueous osmium(VIII)/osmium(VI) couple.

Further thermodynamic data can be found in standard tables. The standard Gibbs free energy of formation for gaseous ethylene is $\Delta_G^\circ = 68.4$ kJ mol$^{-1}$, for liquid ethylene glycol $\Delta_G^\circ = -327.2$ kJ mol$^{-1}$, for liquid water $\Delta_G^\circ = -237.1$ kJ mol$^{-1}$, whereas the Henry constant for the dissolution of ethylene in water is 21.7 kPa m$^3$ mol$^{-1}$. All the thermodynamic data are given for 298.15 K.

A detailed derivation in the Supporting Information is presented to show that these standard thermodynamic data lead to an estimate of the standard free energy change of the dihydroxylation of ethylene with OsO$_4$ in aqueous solution:

$$\text{OsO}_4(aq) + \text{C}_2\text{H}_4\text{H}_2\text{O}(l) \rightarrow \text{OsO}_2(\text{OH})_4^{2-}(aq) + \text{C}_2\text{H}_6\text{O}_2(l) + 2\text{H}^+(aq)$$

$$\Delta G^\circ = -17.8 \text{ kJ mol}^{-1}.$$ 

It should be emphasized that this estimate is based on firm experimental data only. As the driving force of the entire aqueous reaction is quite small, it is certainly impossible to have a highly exergonic intermediate in it. This is in agreement with our experimental findings (the osmate ester is a minor intermediate) and the model proposed in this work. Again, we wish to emphasize that the analysis presented here does not necessarily imply that the accuracy of the gas phase quantum chemical calculations are questionable, it is rather a warning that drawing very general conclusions from such data for solution phase reactions has serious limitations.

## 5 | CONCLUSION

In conclusion, this work reports that the generally accepted, but seldom in fact detected osmium(VI) ester intermediate of the cis dihydroxylation of olefins with osmium tetroxide can be detected by kinetic methods if di- and trichloroethylenes are used as substrates in aqueous medium. The kinetic studies have clearly confirmed that this intermediate plays a key role in the chemical transformation. In the catalytic form of this method, less expensive oxidants are employed to convert back the product osmium(VI) into to the original osmium tetroxide, but these stoichiometric oxidants do not react with the olefin, so the results reported here are fully transferable to catalytic systems as well. It is also important that the possibility of the kinetic detection of the intermediate was not obvious from the studies of the overall process: neither the simple second order rate equation nor the spectral analysis of spectrophotometric studies provided any clue for the existence of the fast process. Therefore, it is quite possible that the same strategy will lead to the successful kinetic detection of the intermediate in many other systems as well.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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