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The Kinetics and Mechanism of the Oxidation of Pyruvate ion by Hypochlorous Acid

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The oxidation of pyruvic acid by hypochlorous acid was studied in the pH 1.0 – 11.0 range. The main path in the redox process is the reaction between the pyruvate ion and HOCl. It was shown that the reaction is second order in both reactants and $k_{HOCl} = 2.14 \pm 0.02 M^{-1} s^{-1} (l = 1.0 M (NaClO_4), T = 25.0 °C)$ with $\Delta H_{HOCl}^{*} = 34.6 \pm 1.2 \text{ kJmol}^{-1}$ and $\Delta S_{HOCl}^{*} = -120.3 \pm 3.6 \text{ Jmol}^{-1} \text{K}^{-1}$ for this reaction step. The relatively large negative entropy of activation is consistent with an O atom-transfer mechanism. DFT calculations support this conclusion by predicting a concerted rearrangement of the activated complex which includes the reactants and the solvent water molecule. It was also confirmed that the hydration of pyruvic acid has significant contribution to the observed kinetic features under acidic condition. The hydration reaction is a proton catalyzed process and the pseudo-first order rate constant can be interpreted by considering theprotolytic and hydration equilibria of pyruvic acid. The rate constants were determined for the non-catalytic and the proton catalyzed path: $k_{ho} = 0.24 \pm 0.01 \text{ s}^{-1}$ and $k_{h1} = 5.5 \pm 0.2 \text{ M}^{-1} \text{ s}^{-1}$.

Introduction

Disinfection processes, environmental and biochemical relevance have generated great interest in the redox chemistry of oxychlorine species.¹⁻¹⁵ In our ongoing studies we use organic substrates as initiators to generate chlorine dioxide from chlorite ion. In such systems, initial oxidation of the substrate by chlorite ion leads to the formation of intermediate species from both reactants and the final products are generated in subsequent reactions of the intermediates. Recently, we have observed very complex kinetic patterns in the chlorite ion lactic acid system. Preliminary experimental results indicate the formation of chlorine dioxide and chlorate ion as final products in variable concentration ratios depending on the pH, the concentrations and the concentration ratios of the reactants. Pyruvate ion and hypochlorous acid are formed as intermediates in the reaction and expected to play significant role in this system.¹⁶ The main objective of this work is to explore the kinetics and mechanism of the redox reaction between these two species.

In a broader context, the results also bear relevance on the redox chemistry of the reactants. The reactions of pyruvate ion have extensively been studied in recent years due to the specific role of this species in various biological redox processes.¹⁷⁻²⁰ Furthermore, the simplicity of the molecule allows modeling the oxidation of organic substrates in the presence of various oxidants.²¹⁻²⁶ Pyruvic acid is a simple substrate with a carbonyl group next to a carboxylic group. As such, its oxidation in aqueous solution may yield either acetaldehyde or acetic acid. With strong oxidants, such as the hypochlorous acid, the formation of the latter product is expected.

Hypochlorous acid is considered to be a strong disinfecting agent which is broadly used in water treatment technologies²⁷⁻³² but also plays an important role in oxidative stress in vivo systems.³³⁻⁴⁰ Due to its outstanding oxidative power, this species can significantly alter the activities and availabilities of biomolecules. Hypochlorous acid may act either as a chlorinating agent or as an O-donor oxidant leading to the formation of very different types of products. It is a key issue how these alternative reaction paths occur under environmental and biological conditions. A survey of the literature reveals that with a few exceptions^{29, 31, 41-45} only limited reliable information is available on the intimate details of the oxidation reactions of hypochlorous acid.

In this study we provide a thorough account on the kinetics of the title reaction and elaborate the mechanistic details also using DFT calculations. The results are expected to contribute to better understanding the redox chemistry of the reactants, in particular, the features of HOCl as an O-donor oxidant.

Experimental

Materials

Chloride ion free sodium hypochlorite was prepared similarly to a method described earlier.^{46, 47} Gaseous chlorine was bubbled into an aqueous slurry of yellow HgO at room temperature. The oxidant was used in adequate excess over Cl_2 and the slurry was intensively stirred for 3 hours after administering chlorine gas in order to drive the oxidation process to completion.

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$$HgO + 2 Cl_2 + H_2O = 2 HOCl + Hg_2Cl_2$$
 (1)

Unoxidized chloride was converted into $HgCl_2$ and the mixture of the HOCl/HgCl_2/HgO slurry was distilled under reduced pressure and argon atmosphere into a 0.4 M NaOH solution.

$$2 \operatorname{HOCl} = \operatorname{Cl}_2 \operatorname{O} + \operatorname{H}_2 \operatorname{O}$$
 (2)

 $Cl_2O + 2 NaOH = 2NaOCl + H_2O$ (3)

The stock solutions of NaOCl were stored at 5 °C in the dark and were standardized by iodometric titration before use.⁴⁸ The concentration of excess NaOH in the stock solution was determined by pH titration with standard HClO₄.

Other reagents (Na-pyr*Aldrich*, KI *Reanal*, NaOH*Aldrich*) were of reagent grade quality and used without further purification. Samples were prepared using doubly deionized and ultrafiltered water obtained from a MILLI-Q RG (Millipore) water purification system.

The kinetic measurements were made at 25 ± 0.1 °C and 1.0 M ionic strength set with NaClO₄ prepared from HClO₄ (*Reanal*) and Na₂CO₃ (*Reanal*).⁴⁹ The concentration of NaClO₄ was determined by drying a known volume of the stock solution in an oven at 120 °C and weighing the mass of the recovered solid.The pH was adjusted with perchloric acid or NaOH solutions.

Instruments and methods

Iodometric and pH titrations were made with a Metrohm 721 NET Titrino system equipped with Metrohm 6.0451.100 combination platinum and Metrohm 6.0262.100 combination glass electrodes, respectively. The pH was measured within ± 0.003 unit, and it is defined as $-\log[H^+]$ in this paper.

Spectrophotometric measurements were made with a Hewlett-Packard 8543 UV/VIS diode array spectrophotometer (equipped with a built-in magnetic stirrer) using different illumination protocols to explore the possibility of unwanted photoreactions.⁵⁰ Photochemical interference was not observed in this system. Measurements were made in stoppered quartz cells of 10 mm light path. The temperature of the cell was controlled by a built-in thermoelectric Peltier device.

Conventional fast kinetic measurements were performed in aqueous solution under pseudo-first-order conditions with an Applied Photophysics DX-17MV sequential stopped-flow instrument using 10 mm optical path at 220 and 325 nm. Pseudo-first order rate constants were obtained from the average of five replicate kinetic runs. Quenched stopped-flow measurements were made with a HI-TECH RQF-63 instrument. Data fitting was made with the program packages SCIENTIST⁵¹ and ORIGIN⁵² using non-linear least-squares routines.

¹H NMR spectra were recorded by using a Bruker DRX 400 (9.4 T) NMR spectrometer equipped with a Bruker VT-1000 thermocontroller and a BB inverse *z* gradient probe (5 mm). Pyruvic acid solutions (0.5 M) were prepared in H_2O (D_2O was added to the samples in capillary). The chemical shifts are reported in ppm, using DSS (4,4-dimethyl-4-silapentane-1-sulfonic acid) as an external standard. The spectra were analyzed with the *Bruker WinNMR* software package.

DFT calculations

DFT calculations were performed with B3LYP53 and M0654 hybrid functionals. In addition, two different Pople-type basis sets have been used: the 6-31+G(d,p) double- ζ and the 6-311+G(d,p) triple- ζ basis sets with one set of diffuse and polarization functions.⁵⁵ Geometryoptimizations were carried out using the polarizable continuum model (PCM) to account for solvent effects. For PCM calculations, we used Gaussian 09's default Integral Equation Formalism variant (IEFPCM) PCM model.56, 57 The transition states were optimized by means of the synchronous transit-guided quasi-Newton method.^{58, 59} The lack of imaginary frequencies in vibrational spectral calculations were taken to verify that the calculated stationary points on the potential energy surfaces (PES) represented true minima as opposed to transition states (TS). The reactants and products are obtained from intrinsic reaction coordinate (IRC) calculations. The relative energies are Gibbs free energies obtained by frequency analysis at 298 K.The calculations were performed with the Gaussian 09 software package.60

Results and discussion

Stoichiometric experiments

The stoichiometry of the reaction was studied at different concentration ratios of pyruvate ion and hypochlorous acid. The reactants were mixed and the spectra were recorded after completion of the reaction. The composition of the spent reaction mixture was calculated by using the characteristic spectra for each major component (Fig.S1, ESI[†]). Acidic reaction mixtures were made alkaline before recording their spectra in order to enhance the sensitivity of the measurements. The stoichiometry was found invariably 1:1 for pyruvate ion and HOCl over the pH 2 -10 range (Table S1 in ESI[†]). Thus, the overall reaction can be written as follows:

$$CH_3COCOOH + HOCl = CH_3COOH + CO_2 + H_2O + Cl^{-}(4)$$

Kinetic studies in the pH 4 - 12 region

Under alkaline conditions, the reaction was studied in the excess of the oxidant using standard spectrophotometric method. Simple first-order kinetic traces were observed by monitoring the absorbance decay at the characteristic absorbance band of OCI⁻ ($\lambda_{max} = 292 \text{ nm}$).⁴⁷ The pseudo-first-order rate constant is a linear function of the hypochlorous acid concentration with zero intercept. These results confirm an overall second order rate law for the reactants.

$$v = -\frac{d[Cl(I)]}{dt} = k_{obs}[Cl(I)][PYR]$$
(5)

where k_{obs} is a pH dependent pseudo second order rate constant and [Cl(I)] and [PYR] stand for the sum of the concentrations of the acidic and basic forms of the corresponding reactants.

The reaction became faster by decreasing the pH and eventually it could not be monitored spectrophotometrically because the spectral effect vanished. The protolytic equilibrium of the oxidant is shifted toward the acidic form under pH 7.0 and HOCl has relatively small absorbance compared to hypochlorite ion. Thus, the change was not measurable. An attempt was made to use pyruvate ion in excess and to monitor the reaction at lower wavelengths. However, due to the large absorbance of pyruvate ion, these attempts were not successful. In addition, when the reactant concentrations were increased significantly, the formation of carbon dioxide led to the evolution of small bubbles in the cell which interfered with spectrophotometric monitoring of the reaction.

In order to avoid the noted technical problems, the reaction was studied by using the quenched stopped flow method in the neutral - slightly acidic region. This kinetic technique is more laborious than the standard stopped-flow method and it has sporadically been used for studying inorganic reactions.^{61, 62} The method is based on immediate chemical quenching of the studied reaction, thus it requires the use of a quenching agent, which is capable to remove the reactant(s) instantaneously. The quenching process should be much faster than the reaction studied and should not trigger unknown side reactions. Once the reaction is stopped, the reaction mixture can easily be analyzed for the products of either the main reaction and/or the quenching reaction. In the latter case, the concentration of a species formed in equivalent amount form the reactant is determined at a given reaction time. In other words, the amounts of unreacted reactants and product can be determined at various reaction times.

The shortest cumulative dead-time of the HI-TECH RQF-63 unit used in this study is below 4 ms. This time is required to start up a reaction and subsequently freeze it by administering an appropriate quenching agent. Therefore, the instrument can conveniently be used for monitoring reactions with pseudo first order rate constants up to 100 s^{-1} . Pyruvate ion was used in large excess in the quenched stopped-flow experiments and 0.5 M NaI solution was used as quenching agent. Iodide ion instantaneously reduced hypochlorous acid to chloride ion and an equivalent amount of iodine formed. Kinetic traces were recorded by stopping the oxidation reaction at various reaction times and determining the amount of iodine formed by standard potentiometric titration with thiosulfate solution. A representative kinetic curve is plotted in Fig. 1.

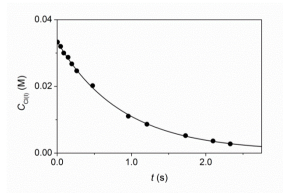


Fig. 1Typical kinetic trace obtained by the quenched stoppedflow method. $C_{PYR}^{0} = 0.500 \text{ M}$, $C_{HOCI}^{0} = 3.29 \times 10^{-2} \text{ M}$, pH = 3.88, I = 1.0 M (NaClO₄), T = 25.0 °C. Quenching agent: 0.500 M NaI.

The pseudo-second-order rate constant is shown as a function of pH in Fig. 2. The results obtained from the regular spectrophotometric and the quenched stopped-flow experiments complement each other remarkably well.

Pyruvate ion is the dominant form of the substrate above pH 4.0, thus the characteristic pH dependence of the rate constant can be interpreted by considering the fast protolytic equilibrium of hypochlorous acid (eq. 6).

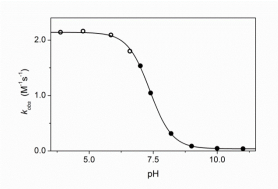


Fig. 2 The pH dependence of the pseudo-second order rate constant for the oxidation of pyruvate ion by Cl(I). Experimental rate constants were obtained by the quenched stopped-flow method (open circles) and conventional kinetic experiments (filled circles). The result of fitting to equation 7 is also shown (solid line). $I = 1.0 \text{ M} (\text{NaClO}_4)$, T = 25.0 °C.

$$OCI^{-} + H^{+} = HOCl K_{p}^{OCl} = \frac{[HOCl]}{[H^{+}][OCI^{-}]}$$
(6)

The pseudo-second-order rate constant is expressed as follows:

$$k_{\rm obs} = k_{\rm HOCl} \frac{K_{\rm p}^{\rm OCl}[{\rm H}^+]}{1 + K_{\rm p}^{\rm OCl}[{\rm H}^+]} + k_{\rm OCl} \frac{1}{1 + K_{\rm p}^{\rm OCl}[{\rm H}^+]}$$
(7)

where k_{HOCI} , k_{OCI} and $K_{\text{p}}^{\text{OCI}}$ are the rate constants of the oxidation by HOCl, OCl⁻ and the protonation constant of OCl⁻, respectively. In agreement with observations in related systems, the acidic form of hypochlorous acid oxidizes the substrate more readily than the basic form. The experimental data were fitted to equation (7) and the results are listed in Table 1. The fitted value for $K_{\text{p}}^{\text{OCI}}$ from the kinetic experiments agrees well with the data reported in earlier literature⁴⁷ and obtained from pH-metric titrations in the present study.

The oxidation by HOCl is the main path of the reaction. The activation parameters for this path were determined by studying the temperature dependence of k_{obs} at pH 2.8 in the 10 - 45 °C temperature range. Under such conditions, pyruvic acid is fully deprotonated and the contribution of the OCl⁻ path is negligible, i.e. $k_{obs} \approx k_{HOCl}$. The activation parameters, ΔS^{\neq} and ΔH^{\neq} , were calculated by fitting the k_{obs} values to the Eyringequation (eq. 8) using a non-linear least squares fitting algorithm (Fig. S2, ESI[†]). The results are listed in Table 1.

$$k_{\text{HOCI}} = \frac{k'T}{h} e^{\frac{\Delta S_{\text{HOCI}}^{*}}{R}} e^{\frac{-\Delta H_{\text{HOCI}}^{*}}{RT}}$$
(8)

Kinetic studies under acidic conditions

Typical stopped-flow traces under pH 2.5 show two distinct steps, which complete within a few seconds and several minutes, respectively. When one of the reactants is used in large excess, each step of the kinetic traces can be fitted to a simple exponential expression yielding two pseudo first order rate constants. This observation is somewhat unexpected because no immediate reason can be seen why the simple kinetic pattern of the redox reaction changes at low pH. A sudden drop of the pH occurred upon mixing the reactants in these experiments and it was concluded that an additional acid induced chemical reaction may contribute to the overall process. For example, the formation of chlorine from hypochlorous acid is enhanced by increasing the acid concentration, but this reaction requires relatively large chloride ion concentration. Chloride ion free reagents were used and only small amounts of this species formed in this study when the oxidation proceeded, therefore, any interference from chlorine formation could be excluded.

Table 1	Equilibrium cor	stants and kinetic	parameters f	for the pyru	vic acid –	chlorine(I) reaction

Parameter	Value	Unit	Remark
lgK_p^{OCl}	7.40		Ref. 47
C p	7.37 ± 0.01		а
k _{OCl}	$(3.97 \pm 0.05) \ 10^{-2}$	$M^{-1}s^{-1}$	a
k _{HOCl}	2.14 ± 0.02	$M^{-1}s^{-1}$	a, slightly acidic – alkaline region
	2.20 ± 0.04	$M^{-1}s^{-1}$	a, acidic region
k _{HOCI}	$(9.1 \pm 2.0) \ 10^{-2}$	$M^{-1}s^{-1}$	a, acidic region
$\Delta H_{\rm HOCl}^{\neq}$	34.6 ± 1.2	kJmol ⁻¹	a
$\Delta S_{\text{HOCI}}^{\neq}$	-120.3 ± 3.6	$\text{Jmol}^{-1} \text{ K}^{-1}$	a
${_{\operatorname{HOCI}}} \stackrel{\neq}{\Delta H_{\operatorname{HOCI}}} \stackrel{\neq}{\Delta S_{\operatorname{HOCI}}} \stackrel{\neq}{K_{\operatorname{p}}}$	57.2 ± 3.4	M^{-1}	а

a: this work

According to literature data, $\log K_p = 1.97$ for pyruvate ion⁶³, thus, it is reasonable to assume that one of the reaction steps is somehow associated with the protonation of this species. In order to explore this possibility, kinetic traces were recorded by acidifying neutral solutions of pyruvate ion in the absence of hypochlorous acid. First order absorbance changes were observed and the pseudo-first-order rate constants agreed well with that of the faster reaction step in the oxidation experiments at the same pH and pyruvate ion concentration. However, simple proton transfer reactions are diffusion controlled and the protonation of pyruvate ion to pyruvic acid is expected to complete far within the dead-time of the stoppedflow experiments under the conditions applied. It follows that these observations cannot be interpreted by considering only the protonation step, instead, it needs to be assumed that the absorbance change is associated with a slower process triggered by the protonation of pyruvic acid.

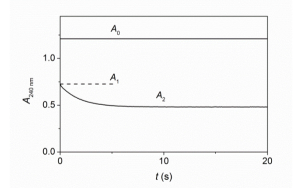
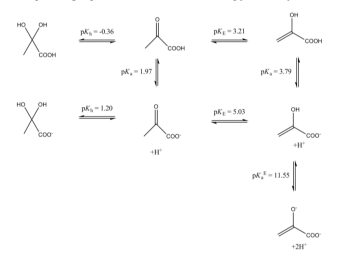


Fig.3 Typical absorbance change upon mixing the solution of pyruvate ion with perchloric acid (A_2). For comparison, the absorbance obtained by dilution of the same pyruvate ion solution with neutral background electrolyte solution is also shown (A_0). Upon acidifying the pyruvate ion solution, the absorbance drops to A_1 within the dead-time of the instrument. $C_{PYR}^{0} = 5.00 \times 10^{-3}$ M, pH = 1.50, *I* = 1.0 M (NaClO₄), *T* = 25.0 °C.

A thorough inspection of the kinetic traces confirms this assumption. As shown in Fig. 3, there is a sudden absorbance drop from A_0 to A_1 within the dead-time of the instrument followed by a slower process associated with an additional absorbance change saturating at A_2 .

It is well documented that α -ketocarboxylic acids are involved in keto-enol and hydration equilibria.⁶³⁻⁶⁶Scheme 1 shows all the possible equilibrium steps together with the corresponding equilibrium constants for the pyruvate system.⁶³



Scheme 1 Acid – base, keto – enol and hydration equilibria in aqueous solution of pyruvate ion. Reproduced from Ref. [63].

According to the data in Scheme 1, the enolic form of pyruvic acid and pyruvate ion is present in less than 0.1 and 0.01 %, respectively and the hydrated form of pyruvate ion is also negligible. In contrast, the hydration of pyruvic acid is substantial, and the keto and hydrated forms are present in comparable concentrations under acidic conditions. ¹H NMR studies support this overall model and the pH dependence of the CH₃ peak of pyruvic acid is consistent with the simultaneous presence of the keto and hydrated forms (Fig. S3, ESI†). These features offer a possibility for coherent interpretation of the experimental observations by the following reaction steps.

$$pyr^{-} + H^{+} = Hpyr K_{p}^{PYR} = \frac{[Hpyr]}{[H^{+}][pyr^{-}]}$$
(9)

Hpyr + H₂O = Hpyr(H₂O)
$$K_{\rm h} = \frac{[\text{Hpyr}(\text{H}_2\text{O})]}{[\text{Hpyr}]} k_{\rm h}, k_{\rm -h}$$
 (10)

where K_p^{PYR} , K_h , k_h , and k_{-h} are the protonation constant, hydration equilibrium constant, forward and backward hydration rate constants, respectively. Thus, the sudden drop of the absorbance is due to the fast protonation of pyruvate ion. This is a pre-equilibrium followed by the hydration of the ketoform of pyruvic acid. The hydration stabilizes the acidic form of pyruvate and an apparent protonation constant, K_p ', can also be defined as follows:

$$K'_{p} = \frac{[\text{Hpyr}] + [\text{Hpyr}(\text{H}_{2}\text{O})]}{[\text{H}^{+}][\text{pyr}^{-}]} = K_{p}^{\text{PYR}}(1 + K_{h})$$
(11)

The plots of A_1 and A_2 as a function of pH are consistent with such protolytic steps and allow the calculation of K_p and K_h by fitting A_1 and A_2 simultaneously to the following expressions:

$$A_{1} = \frac{\varepsilon_{\text{pyr}} + \varepsilon_{\text{Hpyr}} K_{\text{p}}^{\text{PYR}}[\text{H}^{+}]}{1 + K_{\text{p}}^{\text{PYR}}[\text{H}^{+}]}$$
(12)

$$A_{2} = \frac{\varepsilon_{\text{pyr}} + \varepsilon_{\text{Hpyr}} K_{\text{p}}^{\text{PYR}}[\text{H}^{+}] + \varepsilon_{\text{Hpyrh}} K_{\text{p}}^{\text{PYR}} K_{\text{h}}[\text{H}^{+}]}{1 + K_{\text{p}}^{\text{PYR}}[\text{H}^{+}](1 + K_{\text{h}})}$$
(13)

where ε_{pyr} , ε_{Hpyr} and ε_{Hpyrh} are the molar absorbance of pyr, Hpyr and Hpyr(H₂O), respectively. The experimental data were fitted simultaneously to equations 12 and 13 (Fig. 4) and the estimated parameters are listed in Table 2.

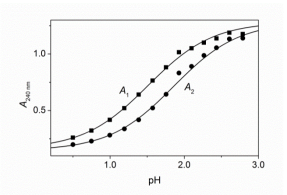


Fig. 4 The characteristic absorbance values upon mixing the solution of pyruvate ion with perchloric acid, cf. Figure 3. The experimental data (dots and squares) were fitted simultaneously to equations 12 and 13 (solid lines) $C_{PYR}^{0} = 5.00 \times 10^{-3}$ M, I = 1.0 M (NaClO₄), T = 25.0 °C.

Table 2 Equilibrium constants and kinetic parameters for the protolytic equilibria and hydration of pyruvic acid	Table 2Equilibrium	i constants and kinetic	parameters for the	protolytic ec	uilibria and h	vdration of	pyruvic acid
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Parameter	Value	Unit	Remark
K _p ^{PYR}	93.3	M^{-1}	Ref. [63]
P	32.5 ± 2.0	M^{-1}	a
K _h	1.2 ± 0.1		a
K _p	72.2	M^{-1}	a, calculated value, eq. 11
r	57.2 ± 3.4	M^{-1}	a, fitted value, eq. 16
k _{h0}	0.24 ± 0.01	s^{-1}	a
k _{h1}	5.5 ± 0.2	$M^{-1}s^{-1}$	a
$\varepsilon_{ m pyr}$	256.6 ± 3.4	$M^{-1}cm^{-1}$	a
E _{Hpyr}	31.8 ± 2.1	$M^{-1}cm^{-1}$	a
E _{Hpyrh}	27.3 ± 2.8	$M^{-1}cm^{-1}$	a

a: this work

The calculated protonation constant of pyruvate ion is about three times smaller than the value reported earlier.⁶³This is partly due to the difference in the experimental conditions applied, however, other factors should also be considered. The protonation constant was estimated by introducing various approximations in the evaluation of the experimental data in the earlier study. This could easily lead to biased results. In our case, the equilibrium constants were calculated directly from stopped-flow photometric experimental data which are expected to provide more reliable results.

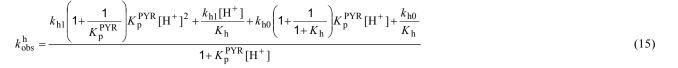
The pseudo-first-order rate constant of hydration increases with increasing $[H^+]$ indicating that the reaction of pyruvic acid

is acid catalyzed (Fig. 5). The corresponding rate constants can be expressed as follows:

$$k_{\rm h} = k_{\rm h0} + k_{\rm h1}[{\rm H}^+] \qquad k_{\rm -h} = k_{\rm -h0} + k_{\rm -h1}[{\rm H}^+] \qquad (14)$$

where k_{h0} , k_{-h0} and $k_{h1}k_{-h1}$ are the forward and reverse rate constants for the non-catalytic and acid catalysed paths of hydration, respectively.

The protonation equilibrium of pyruvate ion is expected to be diffusion controlled and the pH dependence of the pseudo first order rate constant of hydration, k_{obs}^{h} , is given by the following expression:



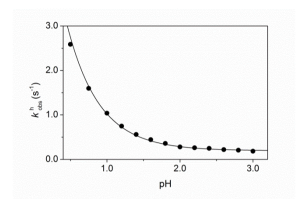
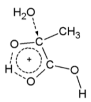


Fig. 5 The pH dependence of the pseudo-first-order rate constant for the hydration of pyruvic acid: experimental data (circles) and the fitted results on the basis of equation 15 (solid line). $\lambda = 240$ nm, $C_{PYR}^{0} = 5.00 \times 10^{-3}$ M, I = 1.0 M (NaClO₄), T = 25.0 °C.

The forward and reverse rate constants of hydration were calculated by fitting k_{obs}^{h} to equation (15) using fixed values for K_{p} and K_{h} . The results are listed in Table 2. In an earlier study, the rate of dehydration of hydrated pyruvic acid was found to be dependent on the pyruvate ion concentration.⁶⁴ This is in sharp contrast with our finding because the rate constant is independent of pyruvate ion concentration in the 5.0×10^{-4} – 1.0×10^{-2} M concentration range. Furthermore, according to our results the proton catalyzed step is considerably slower and the non-catalytic reaction is much faster than it was reported earlier. It is difficult to identify the main reasons for such differences but several factors should be considered. The technical setup of the kinetic experiments was less sophisticated in the cited study and it is not clear how the relatively fast reactions could be monitored with any acceptable reliability. In addition, the kinetic runs were made with pre-acidified pyruvic acid solutions which may undergo slow decomposition. If the reagent solutions were prepared long before the kinetic experiments, the products of the acidic decomposition could easily interfere with the kinetic studies. Such problems were circumvented in this study by using neutral pyruvate ion solutions as a reactant.

The pH dependence of the rate of the hydration of pyruvic acid can be interpreted by considering the structural features of α -keto carboxylic acids. The acid catalysis confirms that pyruvic acid is involved in the formation of a protonated species along the hydration process. The protonation most likely occurs at the carbonyl group of pyruvic acid, and the formation of an internal hydrogen bond to the carboxylic group may stabilize this intermediate. Such a structure may enhance the possibility of a nucleophile attack by a water molecule at the partially positive carbon atom of the carbonyl group (Scheme 2).

Under acidic conditions, the fast step in the pyruvate ion – HOCl reaction is consistent with the results obtained for the hydration of pyruvic acid. It follows that the slower step in Figure 6 corresponds to the oxidation of pyruvate ion. The hydration and protonation reactions should be considered as fast pre-equilibria with respect to the redox process. The pseudo first order rate constant is linearly dependent on the concentration of both pyruvate ion and HOCl concentration when they are used in large excess over the other reactant (Fig. S4, ESI[†]). Thus, the redox reaction remains first order for each reactant under acidic condition. The pH dependence of k_{obs} clearly demonstrates that the reaction between the deprotonated form of the substrate and HOCl is the main reaction path in the redox process and the reaction with Hpyr has minor contribution (Fig. 7).



Scheme 2. The proposed structure of the possible intermediate in the proton catalyzed hydration of pyruvic acid.

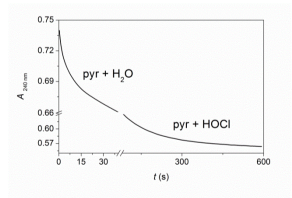


Fig. 6 Typical kinetic trace for the oxidation of pyruvic acid by hypochlorous acid under acidic conditions. $C_{PYR}^{0} = 1.00 \times 10^{-2}$ M, $C_{HOCI}^{0} = 1.00 \times 10^{-3}$ M, pH = 1.50, *I* = 1.0 M (NaClO₄), *T* = 25.0 °C.

Since the hydration of pyruvic acid is much faster than the redox reaction, the pH profile shown in Figure 7 can be interpreted by considering the formal protonation constant of pyruvic acid, K_p '. The pseudo-first order rate constant, k_{obs} , was fitted to the following expression:

$$k_{\text{obs}} = \frac{1}{1 + K'_{p}[\text{H}^{+}]} k_{\text{HOCl}} + \frac{K'_{p}[\text{H}^{+}]}{1 + K'_{p}[\text{H}^{+}]} k'_{\text{HOCl}}$$
(16)

where k_{HOCI} is the rate constant for the reaction of Hpyr with HOCI. The fitted values for k_{HOCI} , k_{HOCI} and K_{p} are listed in Table 1. These data are in reasonable agreement with the results obtained from other experiments. (Tables 1 and 2)

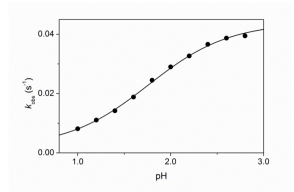


Fig. 7 The pH dependence of the pseudo-first-order rate constant for the oxidation of pyruvic acid by hypochlorous acid under acidic conditions: experimental data (circles) and the fitted results on the basis of equation 18 (solid line).). $\lambda = 325$ nm, $C_{\rm PYR}^{0} = 2.00 \times 10^{-2}$ M, $C_{\rm HOCI}^{0} = 1.00 \times 10^{-3}$ M, I = 1.0 M (NaClO₄), T = 25.0 °C.

Mechanistic considerations and DFT calculations

The simplest way to envision the oxidation of pyruvate ion by hypochlorous acid is to assume an O-atom transfer step Because of the simplicity of the between the reactants. reaction, there are limited options to confirm experimentally the validity of this assumption. With obvious limitations and cautiousness, the activation parameters can be used to test the mechanism. The comparison of activation entropies for related reactions is particularly informative, but the significance of the conclusions obtained in this way should never be overrated. In a recent study, activation parameters were reported for the oxidation of amaranth by hypochlorous acid.⁶⁷ These literature data could be an obvious choice for comparative evaluation of our results. However, an inspection of that paper reveals that fast preequilibrium between hypochlorous acid and its basic form was ignored during the evaluation of the kinetic data. This led to false interpretation of the kinetic results and the corresponding activation parameters need to be termed as erroneous.

Recently, we have demonstrated that O-transfer steps are associated with relatively large negative entropies of activation in simple oxidation reactions of oxone.⁶⁸ This is most likely the consequence of significant geometrical rearrangements associated with the reaction. An O-transfer step would also require massive geometrical changes in the pyruvate ion hypochlorous acid reaction and, therefore, be consistent with a large negative entropy of activation which, in fact, was observed in this system (Table 1). This indicates that a simple O-transfer is probably also operative in the pyruvate ion – hypochlorous acid reaction. The formation of acetic acid requires that the incoming oxygen attacks the carbonyl group and the actual reaction path may include some sort of an intramolecular interaction between the reactants, which is followed by concerted rearrangement of the chemical bonds.

The DFT calculations lend support to this assumption. Energy profiles were calculated for the reaction between pyruvate ion and HOCl. Alternatively, one water molecule was also added to the model checking the possible role of the solvent molecule. The optimized transition states are shown in Figure 8. Substantial differences between the structural arrangements of the reactants in the transition state were not found in the absence and presence of a water molecule. The calculations predict that the reaction is thermodynamically highly favored (Table S2, ESI†). The oxidation is a concerted process where the bond-lengths of the reactants are changing simultaneously. Thus, the HO–Cl and the $C_{carbonyl}$ – $C_{carbonyl}$ bonds are elongated somewhat while the $C_{carbonyl}$ –OH bond is shortened along the reaction coordinate and the water molecule assists this process via hydrogen bonds.

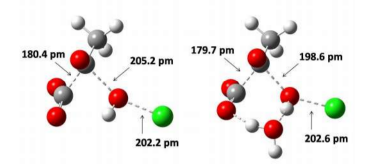


Fig. 8 The transition states in the pyruvate ion – HOCl reaction in the absence and presence of one water molecule calculated at the M06/6-311+G(d,p) level.

As expected, the oxidation involves the attack of the Oatom of HOCl on the carbon atom of the carbonyl group. The results can easily be conceptualized by considering that the carbon atom is partially positive and as such it is prone to a nucleophile attack by the partially negative moiety of the oxidant. The structure of the transition state suggests a very plausible interpretation of the mechanism of the reaction. The rearrangement of the activated complex can easily lead to simultaneous formation of a C–O bond at the carbonvl site, the breakage of the C-C bond, the release of the chloride ion and a proton transfer from HOCl to the water molecule. This concerted rearrangement of the primary and secondary chemical bonds explains the relatively large entropy of activation observed experimentally. Calculations at various theoretical levels do not predict significant differences in the activation Gibbs free energies (ΔG^{\neq}) (Table 3). The added water molecule decreases the ΔG^{\neq} values and makes more exergonic the process. This effect is more pronounced when we use M06 functional and larger basis sets. The structures of the transition states and the related imaginary frequencies are given in the ESI[†] (Table S3).

Conclusions

The results presented here confirm that the oxidation of pyruvic acid by hypochlorous acid is a strongly pH dependent process. The main oxidation path involves HOCl and the deprotonated form of pyruvic acid. The kinetic observations can be interpreted by a simple rate equation which is first order in both reactants. Under acidic conditions, the kinetic traces feature two distinct first order steps. The faster process is associated with the hydration of the substrate and can be treated as a fast preequilibrium compared to the oxidation process. The entropy of activation for the oxidation is consistent with an O-transfer process, and the results of DTF calculations confirm this conclusion.

Functional	Basis sets	pyruvate ion – HOCl	pyruvate ion – HOCl – H2O
B3LYP	6-31+G(d,p)	133.7	129.2
	6-311+G(d,p)	124.6	121.9
M06	6-31+G(d,p)	127.3	113.7
	6-311+G(d,p)	119.8	101.9

Table 3The calculated activation Gibbs free energies of the pyruvate ion – HOCl and the pyruvate ion – HOCl – H_2O systems (in kJmol⁻¹)

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Notes

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[†] Electronic Supplementary Information (ESI) available: Figures showing the spectra of the reactants, the temperature dependence of k_{obs} at pH 2.8, the ¹H NMR spectra of pyruvic acid as a function of pH, k_{obs} as the function of HOCl concentration at pH 1.0; tables on the stoichiometry of the reaction, and DFT results on the Gibbs free energies and coordinates of the transition state.. See DOI: 10.1039/b000000x/

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