

1 **Abstract**

2 The oxidation of iodide by synthetic birnessite (δ -MnO₂) was studied in perchlorate media in the
3 pH range 4 to 8. Iodine (I₂) was detected as an oxidation product that was subsequently further
4 oxidised to iodate (IO₃⁻). The third order rate constants, second order on iodide and first order on
5 manganese oxide, determined by extraction of iodine in benzene decreased with increasing pH
6 (6.3 – 7.5) from 1790 to 3.1 M⁻² s⁻¹. Both iodine and iodate were found to adsorb significantly on
7 birnessite with an adsorption capacity of 12.7 μM/g for iodate at pH 5.7. The rate of iodine
8 oxidation by birnessite decreased with increasing ionic strength, which resulted in a lower rate of
9 iodate formation. The production of iodine in iodide-containing waters in contact with
10 manganese oxides may result in the formation of undesired iodinated organic compounds (taste
11 and odor, toxicity) in natural and technical systems. The probability of the formation of such
12 compounds is highest in the pH range 5 - 7.5. For pH < 5 iodine is quickly oxidized to iodate, a
13 non-toxic and stable sink for iodine. At pH > 7.5, iodide is not oxidized to a significant extent.

14

15 **Key words: Manganese dioxide, birnessite, iodide, iodine, iodate, kinetics, oxidation,**
16 **adsorption**

17

18

1 **Introduction**

2 *Iodine species and their occurrence in aquatic systems*

3 The complex chemistry of iodine in the environment is a result of its multiple oxidation states
4 and of its biophilic character (for reviews see Fuge and Johnson (1986) and Wong (1991)).
5 Iodine is mainly present in the environment as iodide I^- with oxidation state $-I$ and iodate IO_3^-
6 with oxidation state $+V$. A fraction of iodine is also present in the environment as iodoorganic
7 compounds that are produced by microorganisms (Wong, 1991; Gschwend et al., 1985) or by
8 chemical redox reactions (Keppler et al., 2000; Lin et al., 2008; Keppler et al., 2003; Zafiriou,
9 1975). Elemental iodine I_2 (oxidation state 0) and hypoiodous acid HOI (oxidation state $+I$) can
10 also exist as reactive intermediates i.e. metastable species at low concentrations.

11 Iodine is found in seawater at a higher concentration than in any other environment. An average
12 value of $58 \mu\text{g/L}$ is estimated for the total iodine concentration (Fuge and Johnson, 1986). In
13 seawater iodine is mostly present as iodate. Iodide maxima and thus iodate minima are often
14 found in surface waters (Wong, 1991). In contrast to seawater, iodine is present in freshwater
15 mainly as iodide and at much lower concentrations (Smith and Butler, 1979). The concentration
16 of total iodine is in the range of $0.5 - 20 \mu\text{g/L}$ in river waters (Fuge and Johnson, 1986) but
17 concentrations up to $200 \mu\text{g/L}$ were analysed in a mineral water resulting of water infiltration in
18 marine molasse (Bichsel and von Gunten, 1999a). Rainfall and seawater infiltration are also
19 common sources of iodine in freshwaters. For these reasons, total iodine concentrations are
20 higher in coastal areas.

21

22 *Manganese speciation and reactivity*

23 After iron, manganese is the second most common transition metal in the Earth's crust. The most
24 extensive deposition of Mn oxides today occurs in the oceans as Mn nodules (Murray et al.,
25 1984). In marine and freshwater environments, manganese is mainly present in its oxidation state

1 +IV (Murray et al., 1984; Zidenberg-Cherr and Keen, 1987). Manganese oxide minerals
2 participate in a variety of chemical reactions involving both organic and inorganic compounds.
3 This results in oxidation, reduction and scavenging (sorption, precipitation) processes.
4 Manganese oxides have received considerable attention in environmental studies due to their
5 ability to oxidize Co(II) to Co(III) (Manceau et al., 1997), Cr(III) to Cr(VI) (Guha et al., 2001;
6 Johnson and Xyla, 1991; Negra et al., 2005), As(III) to As(V) (Scott and Morgan, 1995), Se(IV)
7 to Se(VI) (Scott and Morgan, 1996) and S(-II) to S(0) (Yao and Millero, 1993). Based on these
8 properties, manganese oxides are also used for the removal of iron and manganese (White and
9 Asfar-Siddique, 1997) and arsenic (Scott and Morgan, 1995; Ouvreard et al., 2002) in water
10 treatment.

11

12 *Oxidation of iodide by MnO₂*

13 The oxidation of iodide by manganese oxides is a well known reaction in acidic solution. This
14 reaction is used for the analysis of oxygen according to the Winkler titration (Winkler, 1888).
15 The Mn:O stoichiometry of manganese oxide is also determined by iodometric titration (Murray
16 et al., 1984). The oxidation of iodide to iodine and/or iodate was already discussed in literature
17 based on thermodynamic data (Anschutz et al., 2000). Truesdale et al. (2001) concluded that the
18 oxidation of iodide to iodine is thermodynamically possible at pH below 6.5 – 7.0, depending on
19 the concentrations of manganous ion and the I₂/I ratios. Similar considerations for the couple
20 IO₃⁻/I indicate that the oxidation of iodide to iodate by manganese oxide is unlikely at pH > 7.75
21 (Truesdale et al., 2001). The Table 1 gives the half reactions of interest with the corresponding
22 E_H[°] and pε[°] (log K). The Figure 1 illustrates the thermodynamic calculations based on the data
23 given in Table 1. The evolution of Δlog K vs. pH for the reactions between iodine species and
24 δMnO₂ confirms that iodide could be oxidised to iodate for pH < 7 and shows that the oxidation

1 of iodide to iodine would be thermodynamically favoured compared to the oxidation of iodine to
2 iodate.

3 Thus, reactive iodine species (i.e. HOI or I₂) can be present in aquatic environmental and
4 technical systems when manganese oxides get in contact with iodide-containing waters at near
5 neutral pH values. Iodine or hypoiodous acid can further react with natural organic matter to
6 produce iodo-organic compounds (Bichsel and von Gunten, 2000). From a drinking water point
7 of view, the formation of iodo-organic compounds such as iodo-trihalomethanes is of concern
8 because these compounds lead to taste and odour problems (Bichsel and von Gunten, 2000) and
9 are more toxic than the chlorinated and brominated analogues (Plewa et al., 2004). Iodoacetic
10 acid, produced also through the reactions between NOM and reactive iodine species, has been
11 shown to be highly cytotoxic and genotoxic and to cause developmental abnormalities in mouse
12 embryos (Richardson et al., 2008).

13 The oxidation of iodide has been extensively investigated by (Bichsel and von Gunten, 1999b) in
14 water treatment conditions using oxidants such as ozone, chlorine and chloramines, however,
15 such studies have not been conducted with manganese oxides at near neutral pH.

16
17 The objective of this paper was to investigate the kinetics of the oxidation of iodide and iodine
18 by manganese dioxide under different experimental conditions (pH, ionic strength,
19 concentrations of manganese oxides, iodide and iodine). This allows assessing the potential of
20 the MnO₂-I⁻ system to form undesired iodo-organic compounds.

21

22 **Material and methods**

23 *Chemicals*

24 Potassium iodide, iodine and potassium iodate were analytical grade (purity) and
25 purchased from Carlo Erba. Sodium perchlorate was purchased from Acros Organic (purity)

1 99%) and iodophenols were purchased from Sigma Aldrich (purity \geq 98%). All solutions were
 2 prepared with ultra pure water produced from a Milli-Q water purification system (Millipore).
 3 Mn dioxide (δ -MnO₂, similar to the naturally occurring birnessite) was synthesized according to
 4 the procedure given in Murray (1973) and concentrated to \sim 200 mM by centrifugation. The
 5 concentration of the stock δ -MnO₂ suspension was measured by the spectrophotometric
 6 determination of the manganous ion (Mn(II)) (Brewer and Spencer, 1971) after the total
 7 reduction of δ -MnO₂ by ascorbic acid. The stoichiometry of the manganese oxide particles,
 8 determined by a iodometric titration (Murray et al., 1984), was Mn:O = 1:1.89. The surface area
 9 of δ -MnO₂ particles was determined by BET analyses and was found to be 140 m²/g. The
 10 experiments were performed with the same suspension of freshly prepared δ -MnO₂ for the entire
 11 duration of the study (4 months) and no significant change in δ -MnO₂ reactivity was observed.
 12 The pK_{a2} value (eq.1) of the MnO₂ sites was determined from a potentiometric titration in a 0.01
 13 M KNO₃ solutions with 0.02 M HCl and 0.02 M NaOH. All operations were carried out under an
 14 atmosphere of argon to avoid contamination with carbon dioxide. The titrations were carried out
 15 in a 200 cm³ reactor at 25.0 (\pm 0.5°C). A pK_{a2} = 4.0 was obtained by fitting the titration data. The
 16 pH_{zpc} of birnessite was determined by Murray (1973) as 2.25.



18 The total sites concentration was determined from the acid-base titration to be 2.15 mM/g i.e. 18
 19 sites/nm². A similar surface site density of 18 sites/nm² was reported for synthetic birnessite by
 20 Yao and Millero (1996).

21 An aqueous iodine stock solution (0.5 mM) was prepared by dissolving sublimated iodine
 22 powder in ultra pure water. The exact concentration was determined by spectrophotometric
 23 measurement at a wavelength of 460 nm ($\epsilon = 746 \text{ M}^{-1} \text{ cm}^{-1}$) using a 50 mm optical cell.

24

25

1 *Experimental procedures*

2 The experiments of iodide and iodine oxidation by δ -MnO₂ were performed in 100 cm³ amber
3 bottles in a thermostated room at 25°C. The δ -MnO₂ particles (1 to 100 mM) were maintained in
4 suspension using a magnetic stir bar agitator at 300 rpm during the course of the experiment. The
5 reaction was induced by the addition of a predetermined volume of an iodide (10 mM) or iodine
6 (0.5 mM) stock solution. At given time intervals, samples were withdrawn with a 50 mL syringe
7 and filtered through a 0.2 μ m membrane filter (diameter 12 mm, regenerated cellulose, Sartorius)
8 for iodate and iodine analysis. Prior to iodide/iodine addition the pH was initially adjusted with
9 0.01 N NaOH and 0.01 N perchloric acid addition and ionic strength was fixed with NaClO₄ (0.1
10 mM to 10 mM). The solutions were not buffered to avoid interfering adsorption on manganese
11 oxides or possible reactions between reactive iodine species and organic buffers. The pH values
12 were monitored during the experiments. The oxidation of iodide and the formation of reactive
13 iodine species and iodate were studied for different concentrations of iodide (5 μ M – 100 μ M),
14 δ -MnO₂ (1 mM – 100 mM) and in the pH range 4 to 8.

15 The oxidation of iodide was also studied in the presence of benzene which was used to quickly
16 extract iodine produced during the experiment. A blank experiment with iodine confirmed that
17 extraction was quantitative. The experiments were carried out with 20 mL of an aqueous δ -MnO₂
18 suspension and 20 mL of benzene. The aqueous and the organic phases were separated at
19 different time intervals and filtered through 0.2 μ m membrane filter prior to iodide (aqueous
20 phase) and iodine (organic phase) analysis. For both aqueous and organic phases, the filtration
21 procedure did not cause any loss of iodine species.

22 Adsorption isotherms of iodate were obtained in perchlorate media at pH 5.7 by varying [IO₃⁻]
23 from 1 μ M to 100 μ M for [δ MnO₂] = 10 mM and for a contact time of 24 hours. Preliminary
24 experiments showed that the equilibrium is reached within 10 minutes.

25

1 *Analytical methods*

2 Iodide analyses were conducted with an iodide sensitive electrode (Orion 6-53 BN Thermo
3 Electro Corporation) after filtration of the δ -MnO₂ suspension. There was no interference with
4 iodate or elemental iodine.

5 Reactive iodine species (sum of I₂ and HOI) in aqueous solution were quenched with phenol and
6 analyzed as iodophenols by HPLC. Phenol (200 μ M) was added to the sample to transform
7 iodine to iodophenols. Phenol reacts quickly and quantitatively with iodine (Bichsel and von
8 Gunten, 2000). Both p-iodophenol and o-iodophenol were formed with an extent ratio of
9 approximately 15:1. Because phenol was present in high excess, the formation of di- or
10 triiodophenols was not observed. The two iodophenols were quantified relative to o- and p-
11 iodophenol standard solutions. The iodine concentration was calculated as the sum of the two
12 iodophenols. The HPLC separation was done on a Nucleosil 103-5 C18 column with an eluent
13 consisting of 50% methanol, 49.85% water, and 0.15% acetic acid. The UV detection at 231 nm
14 yielded a detection limit of 0.3 μ g I/L (Bichsel and von Gunten, 2000).

15 Iodate analysis were performed according to the method described by Salhi and von Gunten
16 (1999). A Dionex® AS19 column and AG19 guard column with 25 - 100 mM NaOH eluent
17 were used. The injection volume was reduced to 100 μ l to limit the response of perchlorate. The
18 detection limit was 20 nM with a standard deviation in the range 0.5 - 5%.

19 Iodine was analyzed in benzene by spectrophotometric measurement at a wavelength of 500 nm
20 ($\epsilon = 1000 \text{ M}^{-1} \text{ cm}^{-1}$) using a 50 mm optical cell. The detection limit was 0.5 μ M with a standard
21 deviation in the range 2 - 8%.

22

23

1 **Results and discussion**

2 *Adsorption of iodate*

3 Adsorption data plotted in Langmuir equation (Figure 2) exhibit a linear segment over the range
4 of concentrations examined at pH 5.7. This observation indicates that only one type of binding
5 site exists for iodate adsorption for our experimental conditions. The Langmuir isotherm gives a
6 concentration of sites equal to $12.7 \mu\text{mol/g}$ and an equilibrium constant of $1.73 \times 10^{-2} \text{ L}^2/\text{g}^2$ at pH
7 5.7 for an ionic strength of 1 mM. The adsorption of IO_3^- decreases with increasing pH, in
8 agreement with both surface complexation and ligand exchange mechanisms (Figure 2b). At
9 neutral pH, almost no iodate adsorption occurs on synthetic birnessite, which can be explained
10 by the low pH_{zpc} of birnessite ($\text{pH}_{\text{zpc}} = 2.25$). At pH 5.0, speciation calculation showed that
11 protonated $\equiv\text{MnOH}_2^+$ species would only be present at nanomolar concentrations for a 10 mM δ -
12 MnO_2 suspension (0.87 g/L). Figure 2b shows also that ionic strength has a very low influence
13 on iodate adsorption, which would suggest that adsorption occurs through an inner sphere
14 mechanism.

15

16 *Determination of rate constants for the oxidation of iodide to iodine*

17 The rate constants for the oxidation of iodide were determined between pH 5.35 and 7.50 in
18 presence of benzene used as extracting solvent of iodine. Both iodide (Figure 3a) and iodine
19 (Figure 3b) were followed in the aqueous and in the organic phases, respectively. The pH values
20 increased immediately after the addition of iodide and remained stable at ± 0.15 , which allowed
21 the determination of the corresponding rate constants. The results verified that one mole of
22 iodine is formed for two moles of iodide oxidized. Significant iodate formation and thus iodine
23 oxidation was only observed for pH values 5.35 and 6.32 and for reaction times ≥ 3 hours, which
24 explained the discrepancies between experimental and calculated values for iodine in Figure 3b.

1 The mass balance (i.e. $[\Gamma]_{\text{ini}} = [\Gamma]_{\text{water}} + [\text{IO}_3^-]_{\text{water}} + 2 [\text{I}_2]_{\text{benz}}$) was always verified at $98 \pm 4\%$ ($n =$
2 37).

3 The iodide oxidation and the formation of iodine were well modelled assuming a second order
4 reaction with respect to iodide (eq. 2). A reaction second order in iodide was verified by plotting
5 $1/[\text{Iodide}]$ as a function of the reaction time with the slope $= 2.k'_{\text{obs}}$ (Figure 4). Considering that
6 the concentration of $\delta\text{-MnO}_2$ was constant and assuming a first order reaction with respect to $\delta\text{-}$
7 MnO_2 , the rate constants k_{obs} in $\text{L}^2 \text{mol}^{-2} \text{s}^{-1}$ were then calculated and are given in Table 2 (the
8 value for pH 5.35 was not considered because more than 95% of iodide was oxidized within the
9 first minutes). A second order dependence on iodide was also reported by Laurence and Ellis
10 (1972) for the oxidation of iodide by Fe(III) at acidic pH. Lin et al. (2008) determined a lower
11 order of 1.68 on iodide and a first order on PbO_2 for the reductive dissolution of PbO_2 by iodide.

12



$$14 \quad \text{with } -d[\text{I}^-]/dt = k'_{\text{obs}} [\text{I}^-]^2 = k_{\text{obs}} [\text{I}^-]^2 [\text{MnO}_2] = k [\text{H}^+]^n [\text{I}^-]^2 [\text{MnO}_2]$$

15

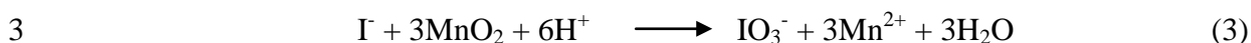
16 Values of $\log k_{\text{obs}}$ are plotted as a function of pH in Figure 5. The apparent reaction order with
17 respect to H^+ was determined as the slope of the straight line $\log k_{\text{obs}} = f(\text{pH})$ and is equal to 2.3.
18 A lower dependence of 0.66 on pH was determined by Lin et al. (2008) between pH 6.0 and 8.0
19 for the couple PbO_2/I^- . The rate constant k for reaction (2) was found to be $7.2 \times 10^{17} \text{ M}^{-4.3} \text{ s}^{-1}$.

20

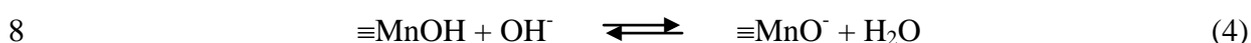
21 *Oxidation of iodide and iodine to iodate*

22 Figure 6 shows data for the oxidation of iodide for varying concentrations of $\delta\text{-MnO}_2$ (1 mM to
23 100 mM) at an initial pH of 5.0 ((a) iodide, (b) reactive iodine species, (c) iodate, (d) pH, (e)
24 mass balance on iodine species). In absence of benzene, reactive iodine species are further
25 oxidized to iodate, which is the end product of iodide oxidation. Even though 3 moles of $\delta\text{-MnO}_2$

1 are reduced per mole of iodide oxidised (eq. 3), under our experimental conditions MnO₂ was
2 always in excess of iodide.



4 For all conditions, the pH increases after the addition of iodide (Figure 6d), which is consistent
5 with the overall reaction of iodide on δ -MnO₂ where the oxidation of one mole of iodide
6 consumes 6 moles of proton. The increase of pH is lower for high concentrations of manganese
7 oxide because of the buffer capacity of manganese oxides:



9 For high δ -MnO₂ concentrations of 50 mM and 100 mM, the pH remains below 6 and iodide is
10 immediately oxidized to iodate and no iodine is detected. The resulting iodate concentrations
11 (Figure 6c) are 58 μ M and 78 μ M for 100 mM and 50 mM δ -MnO₂, respectively. The iodate
12 yield is not 100% because of the adsorption of iodate on δ -MnO₂.

13 It could be expected that iodate adsorption could decrease the rate of iodide oxidation by
14 competitive adsorption at the reactive sites. Experiments with initial iodide concentration of 50
15 μ M and 10 mM δ -MnO₂ carried out at an initial pH of 5 in absence and in presence of 50 μ M
16 iodate showed no effect of iodate on the oxidation rate of iodide (results not shown).

17
18 For δ -MnO₂ concentrations of 5 mM and 10 mM, pH values increase to 6.5 (Figure 6e). This
19 reduces the oxidation rate of iodide. Reactive iodine species are formed within the first 30
20 minutes and slowly decayed by formation of iodate. The oxidation of iodide to iodine is then
21 faster than the oxidation of iodine to iodate in agreement with the calculation of the driving
22 forces between pH 4 to 8 (Figure 1).

23 Reactive iodine species are only detected when $[\text{I}^-] > 10 \mu\text{M}$ (Figure 6a and 6b), which can be
24 related to the formation of elemental iodine according to the equilibrium:



1 According to the above equilibrium, iodine represents $\geq 85\%$ of the reactive iodine species at pH
2 6.5, and for $[I] \geq 10 \mu\text{M}$. The presence of I_2 was confirmed by the UV/visible spectra of the
3 filtered solution that exhibited a peak maximum at 460 nm. The formation of triiodide is unlikely
4 in our conditions i.e. for μmolar concentrations of iodide:

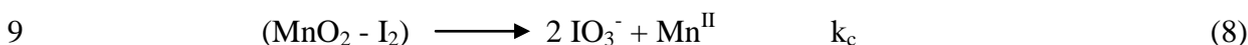
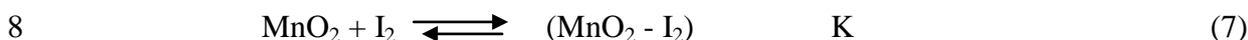


6 The mass balance of iodine (Figure 6e), calculated by considering that all reactive iodine species
7 are molecular iodine, shows that for the highest concentrations of iodine (i.e. $\text{I}_2 = 15 \mu\text{M}$ for 5
8 mM $\delta\text{-MnO}_2$ and $\text{I}_2 = 20 \mu\text{M}$ for 10 mM $\delta\text{-MnO}_2$ at 30-minutes reaction time) there is a deficit
9 of 20 to 25 μM . The deficit cannot be explained by iodate adsorption because iodate adsorption
10 is low at pH 6 and it is observed at the beginning of the reaction i.e. for low iodate concentration
11 (e.g. 6 μM for 5 mM $\delta\text{-MnO}_2$ and 30-min reaction time).

12 The oxidation of iodine to iodate was studied for an initial concentration of iodine of 48 μM and
13 10 mM $\delta\text{-MnO}_2$ at an initial pH of 5.0 (Figure 7). During the first minute of the reaction, 13.6
14 μM of iodine disappeared immediately from the solution and 9.7 μM of iodate were formed.
15 Thus, about 9 μM of iodine would be adsorbed on manganese oxides, which would explain the
16 deficit in iodine observed in the previous experiments. The loss of iodine is accompanied by an
17 increase of the pH from 5.0 to 5.9 within the first minute. After that initial pH change it remains
18 constant throughout the experiment. This observation would be consistent with a ligand
19 exchange mechanism between iodine and hydroxide ions. For the next three hours iodine is
20 further oxidised to iodate with a stoichiometric factor of 2 ($\text{I}_2:\text{IO}_3^- = 1:2$). Sites on manganese
21 oxide can then be considered as Lewis acid sites i.e. sites capable of receiving an electron pair
22 from the adsorbate. Similarly, an inner-sphere mechanism was also proposed for the oxidation of
23 sulfite on manganese dioxide (Herszage and Dos Santos Afonso, 2003). The complexation of
24 iodine by MnO_2 would result in the polarisation of the I-I bond i.e. the electron density of
25 manganese increases whereas the electron density of iodine decreases. This would enhance the

1 reactivity of iodine towards electrophilic substitution and thus the formation of iodinated organic
2 compounds. Such activation of the iodine molecule by mineral oxide catalysts is commonly used
3 in organic synthesis (Stavber et al., 2008).

4
5 The reduction of metal oxides by iodide involves several reaction steps including adsorption of
6 iodine on the metal oxide surfaces, electron transfer between iodine and the metal oxide and
7 release of the oxidized species and the reduced metal ions:



10 Such reaction mechanisms can be described by a Langmuir-Hinshelwood or Michaelis–Menten
11 type rate law where the apparent reaction order vary between 0 and 1 depending on the rate
12 limiting step. When adsorption is the rate limiting step, the reaction is first order with respect to
13 the concentration of I^- or I_2 and first order with respect to the metal oxide concentration (MnO_2).
14 When electron transfer is the rate limiting step, the rate is proportional to the concentration of
15 surface complexes ($\text{MnO}_2\text{-I}_2$). When the concentration of the reductive species reaches a
16 concentration where all the reactive sites are saturated, the concentration of the surface complex
17 is constant and the reaction rate is zero order. Before saturation, the rate is also first order with
18 respect to the reductive species and the metal oxide surface. The second order rate constant k_{app}
19 of iodine oxidation is the product of the equilibrium constant K and the rate constant of the
20 decomposition of the surface complex k_c (eq. 9).

$$21 \quad -d[\text{I}_2]/dt = k_c [\text{MnO}_2 - \text{I}_2] = k_c K [\text{MnO}_2] [\text{I}_2] = k_{\text{app}} [\text{MnO}_2] [\text{I}_2] \quad (9)$$

22
23 Because $[\text{MnO}_2]$ was in excess compared to $[\text{I}_2]$, the plot of $\ln[\text{I}_2]/[\text{I}_2]_0$ versus reaction time after
24 the initial adsorption phase gave a straight line with an apparent first order rate constant $k'_{\text{app}} =$
25 $2.57 \times 10^{-4} \text{ s}^{-1}$ ($r^2 = 0.993$, $n = 14$). By taking into account both iodine concentration in solution

1 and the I_2 concentration adsorbed on MnO_2 (i.e. $[MnO_2 - I_2]$), a rate constant $k_c = 1.0 \times 10^{-3} \text{ s}^{-1}$
2 was then calculated for the decomposition of the surface complex (eq. 10).

$$3 \quad -d[I_2]/dt = k'_{app} [I_2] = k_c [MnO_2 - I_2] , \quad k_c = \frac{k'_{app} \cdot [I_2]}{[MnO_2 - I_2]} \quad (10)$$

4

5 *Effect of pH*

6 Figure 8 shows the effect of pH on the kinetics of the oxidation of iodide and the formation of
7 iodate and active iodine species. As expected from the driving force (ΔpE) (see Figure 1) and
8 from previous experiments conducted in presence of benzene, the rates of iodide oxidation
9 (Figure 8a) and iodate formation (Figure 8b) strongly increase with decreasing pH. Reactive
10 iodine species are formed within the first 30 minutes except for pH 7 where a slower formation
11 was observed (Figure 8c). For an initial $pH \leq 6$, the formation of iodine on $\delta\text{-MnO}_2$ is
12 accompanied by an increase in pH for the first 30 minutes of the reaction (Figure 8d). At an
13 initial pH of 8, about 6 μM of iodide disappear immediately after $\delta\text{-MnO}_2$ addition probably by
14 simple sorption onto the manganese dioxide surface. Nevertheless, slow oxidation of iodide also
15 occurs at pH 7.3-7.4 (initial pH 8). An iodate formation rate of 0.7 $\mu\text{M/h}$ was determined. For
16 this condition, iodine could not be detected in solution. Iodine was quantified for all the other pH
17 conditions (Figure 8c). Results showed that the iodine exposure i.e. the integral of the iodine
18 concentration versus time curve is higher in the pH range 6 – 7 than at readily acidic pH because
19 at $pH < 6$ iodide is rapidly converted to iodate.

20

21 *Effect of ionic strength*

22 The effect of ionic strength on iodide oxidation and iodine and iodate formation was studied for
23 ionic strength from 0.1 to 10 mM NaClO_4 . Results in Figures 9 show that for an increase in ionic
24 strength, the stability of reactive iodine species in solution increased (Figure 9a) and thus the rate

1 of iodate formation decreased (Figure 9b). The apparent first order rate constants of oxidation of
2 iodine determined from data plotted in Figure 9a linearly decreased with $I^{1/2}$ (Figure 10). The
3 mass balance for iodine at 30 minutes was 72% for 1 mM NaClO₄, 64% for 5 mM and 57% for
4 10 mM. Because iodate adsorption is negligible at pH 6.2 – 6.5, this suggests that the adsorption
5 of reactive iodine species increases when the ionic strength increases.

6 One possible explanation of the negative effect of ionic strength on the oxidation of reactive
7 iodine species could be the change of speciation between I₂ and HOI. When ionic strength
8 increases, the hydrolysis of iodine (reaction 7) is shifted to HOI, which is expected to be more
9 stable than I₂ in presence of manganese oxide. However, the ionic strength corrections using the
10 Debye-Hückel equation shows that the HOI/I₂ ratio only increases from 0.174 to 0.193 for 0.1
11 mM to 10 mM ionic strength (calculations were done for pH = 6.5 and [I] = 10 μM). Thus, the
12 speciation of HOI/I₂ cannot only account for the effect of ionic strength observed in our
13 conditions.

14 Ionic strength can also directly affect the reactivity of manganese dioxide. In aqueous solution,
15 freshly prepared δ-MnO₂ particles are aggregates of size ranging from about 0.2 and 0.4 μm.
16 The size grows to 1.0 μm over period of months, which was explained by coagulation (Stone and
17 Morgan, 1984). Thus, the coalescence of particles was proposed to be partly responsible of the
18 decrease of reactivity (e.g. rates of reduction by phenolic compounds) observed during aging of
19 manganese oxides (Stone and Morgan, 1984). In our experiments, increasing ionic strength also
20 increases the size of MnO₂ aggregates by coagulation, which might explain the decrease of the
21 rate of iodate formation. Increasing ionic strength decreases also the water solubility and favors
22 hydrophobic interactions. Thus, and because of the strong hydrophobicity of iodine, the increase
23 of iodine adsorption with increasing ionic strength might be due to hydrophobic interactions
24 between iodine and manganese oxide particles. Hydrophobic interactions have already been
25 proposed to explain the adsorption of certain organic compounds on MnO₂ (Bernard et al.,

1 1997). The hydrophobic interactions are long range interactions, which might also cause a
2 reduction of the rate of electron transfer between iodine and the MnO₂ surface when ionic
3 strength increases.

4

5 **Implications for water treatment**

6 Our results show that iodine can be produced at near neutral pH when iodide containing waters
7 are in contact with manganese oxide. In water treatment, manganese oxides are commonly used
8 for Mn removal. Even though the residence times in MnO₂ filters are only 10 - 15 minutes and
9 iodide concentrations are usually in the ppb range, the formation of low amount of iodine might
10 result in the formation of iodinated organic compounds. Our results give also evidence that
11 iodine readily adsorbs on synthetic birnessite. In absence of organic compounds, the
12 complexation of iodine is followed by electron transfer with formation of iodate. Iodate is the
13 desired sink for water utilities, because it is inert and non-toxic (Burgi et al., 2001). The results
14 suggest that the formation of iodinated organic compounds can be reduced by 1) maintaining
15 high pH values > 7.5 to limit iodine formation or 2) by lowering the pH around 5.0 to rapidly
16 convert iodide to stable iodate. Mineral content of natural waters would also influence the
17 formation of iodinated organic compounds because reactive iodine species have been found
18 more stable when ionic strength increases from 0.1 mM to 10 mM. The presence of specific
19 mineral species such as carbonate and sulfate might also affect the formation of iodinated
20 organic compounds by competing with iodine for adsorption/activation on manganese dioxide.

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1 **Conclusion**

2 The oxidation of iodide to iodate by synthetic birnessite was studied between pH 4 and 8 for
3 different conditions.

4 ○ Iodate slightly adsorbs on birnessite at pH 5.7 with adsorption capacity of 12.7 $\mu\text{mol/g}$.

5 The adsorption decreases with increasing pH. At circumneutral pH the adsorption is not
6 significant.

7 ○ The rate of iodide oxidation decreases with increasing pH with an H^+ dependence of 2.3.

8 Third order rate constants, second order in iodide and first order in $\delta\text{-MnO}_2$, were
9 determined and varied from 1790 $\text{M}^{-2} \text{s}^{-1}$ at pH 6.30 to 3.1 $\text{M}^{-2} \text{s}^{-1}$ at pH 7.50.

10 ○ Iodine was detected as a reactive intermediate species in the oxidation of iodide to iodate,
11 especially for the lowest MnO_2/I^- ratios and for a pH range between 6.0 and 7.0. For a
12 high MnO_2/I^- ratio and $\text{pH} < 6$, iodide is rapidly converted to iodate and therefore iodine
13 was not observed.

14 ○ Increasing the ionic strength from 10^{-4} to 10^{-2} M reduces the rate of iodate formation by
15 decreasing the oxidation rate of iodine. Therefore the iodine exposure is higher for high
16 ionic strengths.

17 ○ Iodine significantly adsorbs on synthetic birnessite that reacts as a Lewis acid by
18 receiving an electron pair. Complexation of iodine on manganese oxides might contribute
19 to the activation of iodine through to the polarization of the I – I bond. Complexed or free
20 iodine can then react with natural organic matter in the environment and in water
21 treatment processes to produce iodinated organic compounds in the pH range 6.0 – 7.0.

22 ○ Further research is needed to evaluate the potential formation of iodinated organic
23 compounds like iodo-THMs (iodoform) when iodide-containing natural waters are
24 treated by manganese oxides.

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Table 1. Half reactions, E_H° and $p\varepsilon^\circ$ ($\log K$) for Mn and iodine species. The E_H° and $p\varepsilon^\circ$ ($\log K$) is for the standard state condition (1M in all species including H^+).

Table 2. Third order rate constants k_{obs} in $L^2 \text{ mol}^{-2} \text{ s}^{-1}$ for the oxidation of iodide by $\delta\text{-MnO}_2$ (25°C, 1 mM NaClO_4)

Table 1.

Half reactions	E_H° in V	$p\varepsilon^\circ$ (log K)
$\frac{1}{2} \text{MnO}_2 + 2 \text{H}^+ + \text{e}^- \rightleftharpoons \frac{1}{2} \text{Mn}^{2+} + \text{H}_2\text{O}$	1.29	21.86
$\frac{1}{2} \text{I}_{2(\text{aq})} + \text{e}^- \rightleftharpoons \text{I}^-$	0.615	10.42
$\frac{1}{5} \text{IO}_3^- + \frac{6}{5} \text{H}^+ + \text{e}^- \rightleftharpoons \frac{3}{5} \text{H}_2\text{O} + \frac{1}{10} \text{I}_{2(\text{aq})}$	1.178	19.96
$\frac{1}{6} \text{IO}_3^- + \text{H}^+ + \text{e}^- \rightleftharpoons \frac{1}{2} \text{H}_2\text{O} + \frac{1}{6} \text{I}^-$	1.085	18.38

Table 2.

pH	k_{obs}
6.32	1790 ± 280
6.73	250 ± 14
6.80	72 ± 3
7.39	5.5 ± 0.8
7.50	3.1 ± 0.2

Figure 1. $\Delta \log K$ vs. pH plots for redox reactions between iodine species and $\delta\text{-MnO}_2$. Calculations were made using $p\epsilon^\circ$ in Table 1 and for $\{\text{I}_2\} = \{\text{IO}_3^-\} = \{\text{I}^-\} = 1$. Low activity of Mn^{2+} of 10^{-7} was used for calculation because Mn^{2+} was considered to be catalytically reoxidized by oxygen in presence of MnO_2 . Negative log K values indicate unfavourable reactions.

Figure 2. Adsorption of iodate on $\delta\text{-MnO}_2$ (a) adsorption isotherm ($[\delta\text{-MnO}_2] = 10 \text{ mM}$, NaClO_4 1 mM, pH 5.7) (b) effect of pH and ionic strength ($[\delta\text{-MnO}_2] = 10 \text{ mM}$; $[\text{IO}_3^-]_0 = 100 \text{ }\mu\text{M}$)

Figure 3. Kinetics of (a) iodide oxidation and (b) iodine formation (experimental data (symbols) and third order kinetic model (line); $[\delta\text{-MnO}_2] = 10 \text{ mM}$, $[\text{I}^-]_0 = 100 \text{ }\mu\text{M}$, $\text{I} = 1 \text{ mM NaClO}_4$, the pH values are constant pH values ± 0.15 for reaction times $> 30 \text{ min.}$).

Figure 4. Plot of $1/[\text{I}^-]$ versus time for iodide oxidation by $\delta\text{-MnO}_2$ ($[\delta\text{-MnO}_2] = 10 \text{ mM}$, $[\text{I}^-]_0 = 100 \text{ }\mu\text{M}$, $\text{I} = 1 \text{ mM NaClO}_4$)

Figure 5. Dependence of the third order rate constant k_{obs} on pH ($\text{I} = 1 \text{ mM NaClO}_4$)

Figure 6. Evolution of iodine species and pH for varying concentrations of $\delta\text{-MnO}_2$ (NaClO_4 1 mM, initial pH 5, $[\text{I}^-] = 100 \text{ }\mu\text{M}$). (a) iodide, (b) reactive iodine species (i.e. $\text{I}_2 + \text{HOI}$), (c) iodate, (d) pH, (e) mass balance for iodine species.

Figure 7. Reaction of $\delta\text{-MnO}_2$ with iodine (I_2) ($[\delta\text{-MnO}_2] = 10 \text{ mM}$, $[\text{I}_2]_0 = 48 \text{ }\mu\text{M}$, NaClO_4 1 mM)

Figure 8. Effect of pH on the kinetics of the oxidation of iodide (a), the formation of iodate (b) and reactive iodine species (c), (d) shows the evolution of the pH ($[\delta\text{-MnO}_2] = 10 \text{ mM}$, $[\text{I}]_0 = 100 \mu\text{M}$, $\text{NaClO}_4 1\text{mM}$).

Figure 9. Effect of ionic strength on iodate and iodine formation ($[\delta\text{-MnO}_2] = 10 \text{ mM}$, $[\text{I}]_0 = 100 \mu\text{M}$, initial pH 5.0, final pH 6.2-6.5)

Figure 10. Effect of ionic strength on the apparent first order rate constant of iodine oxidation ($[\delta\text{-MnO}_2] = 10 \text{ mM}$, $[\text{I}]_0 = 100 \mu\text{M}$, pH 6.2 – 6.5)

Figure 1.

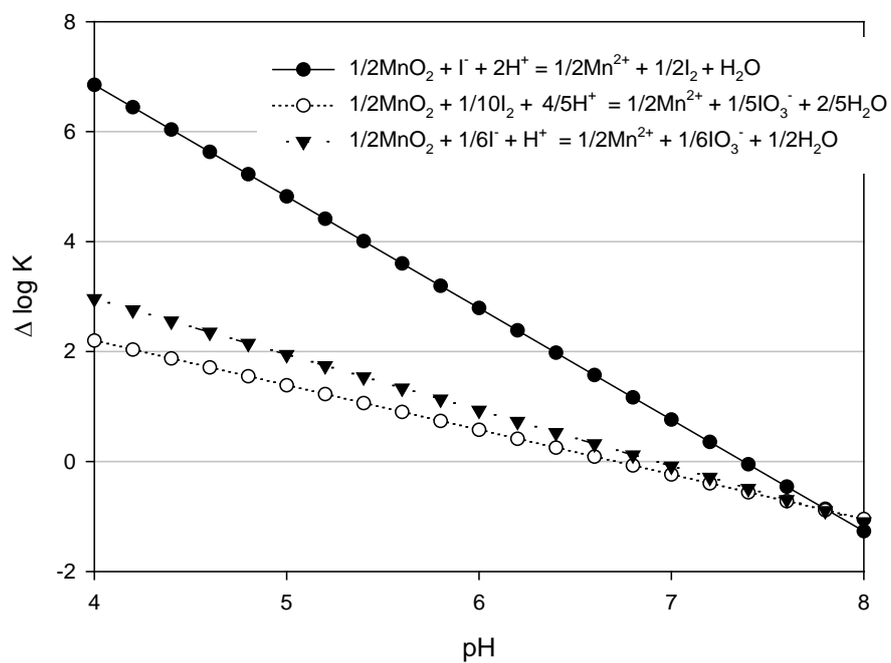


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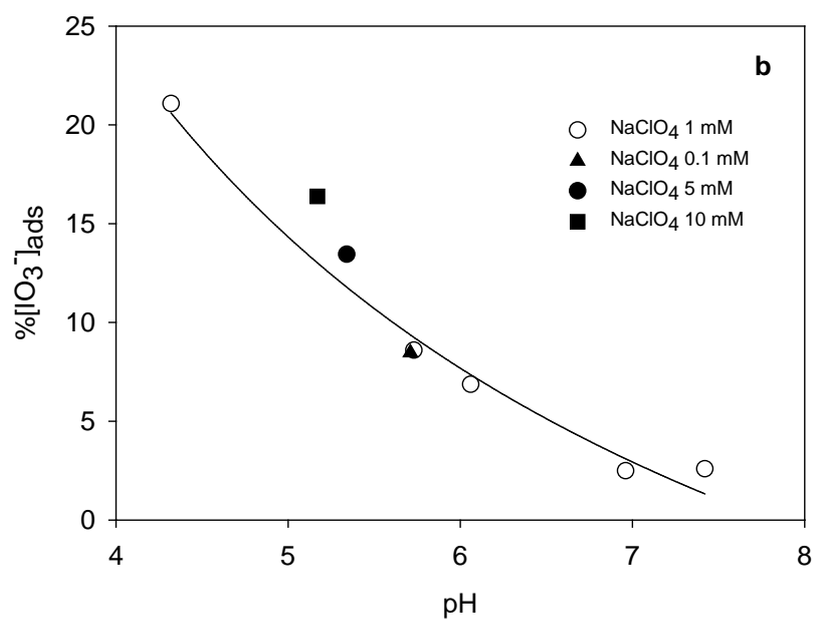
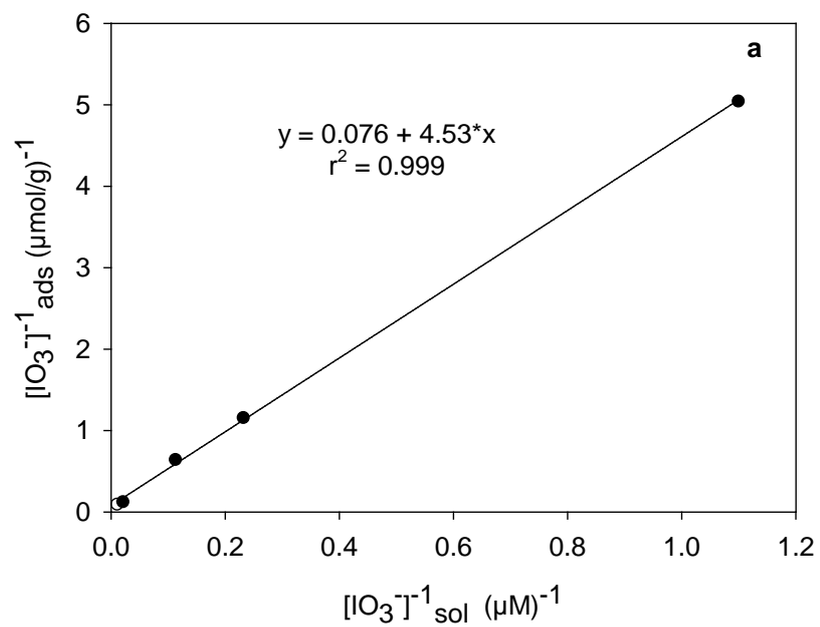


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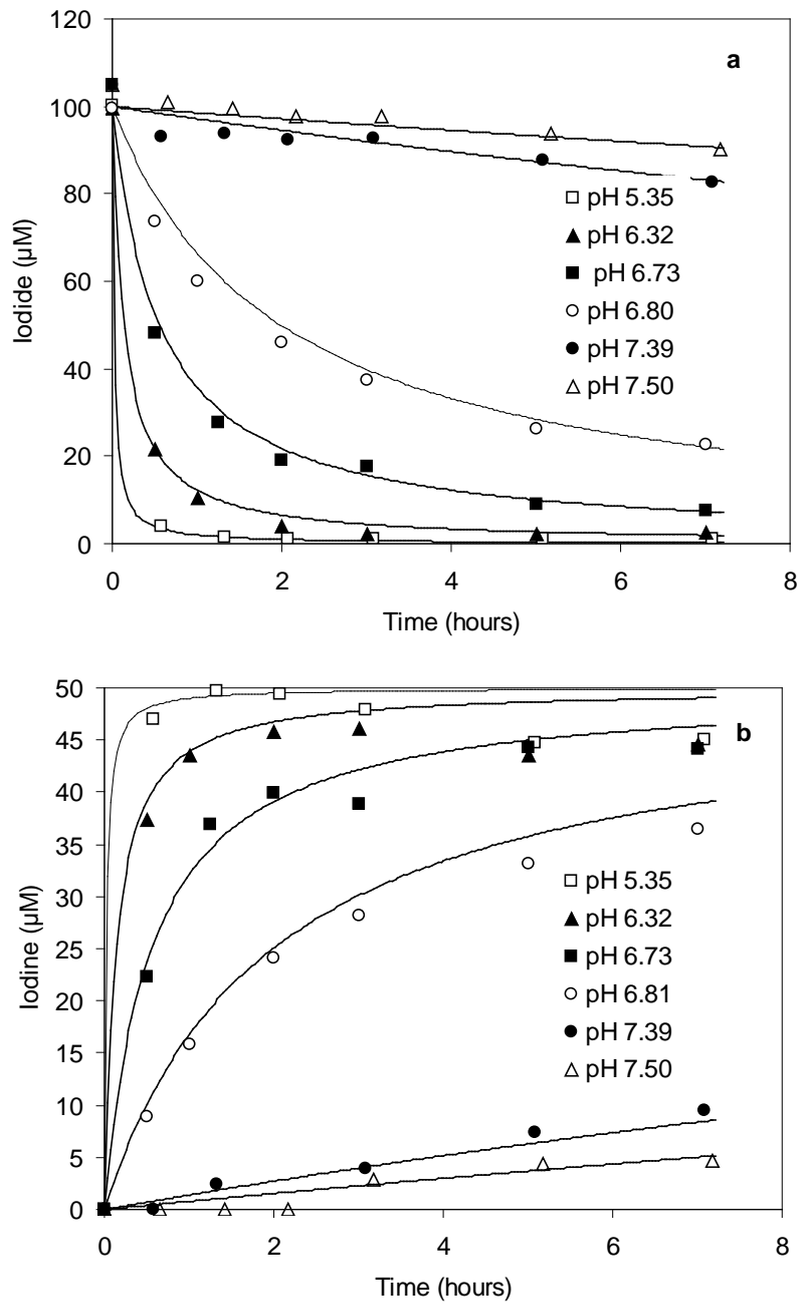


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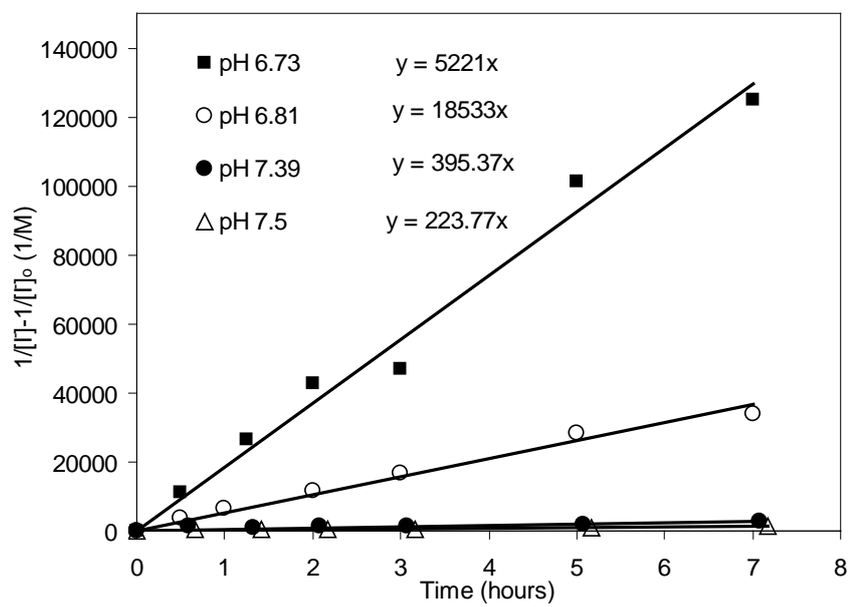


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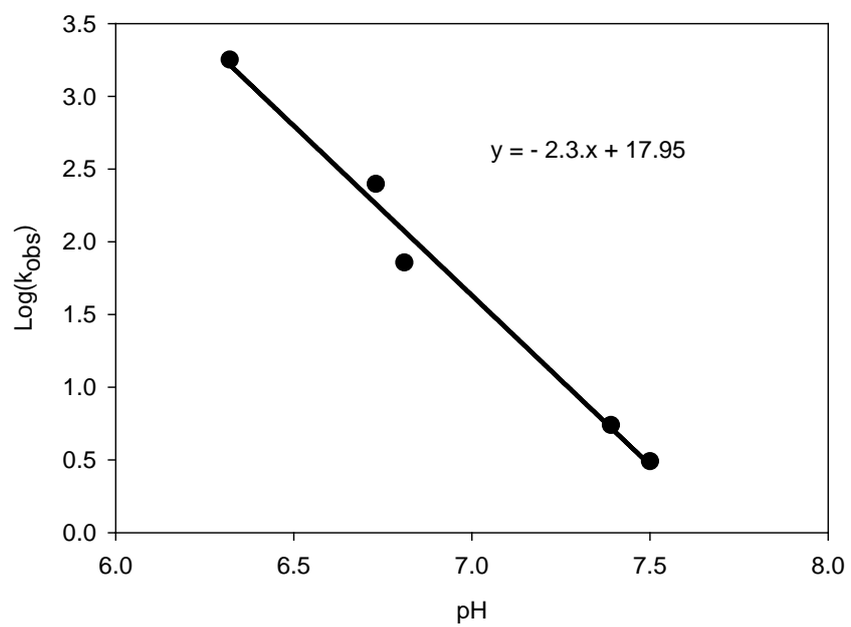


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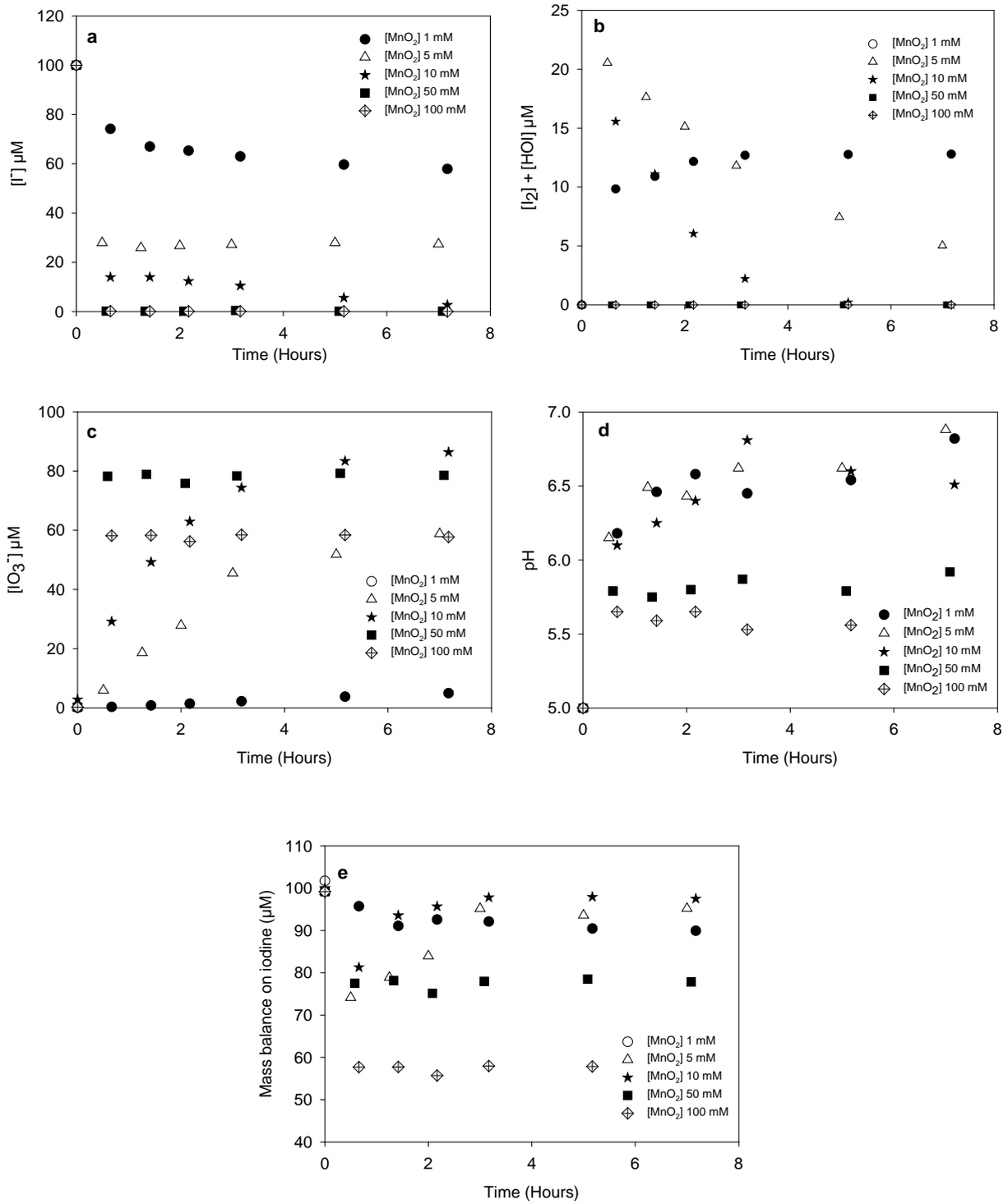


Figure 7

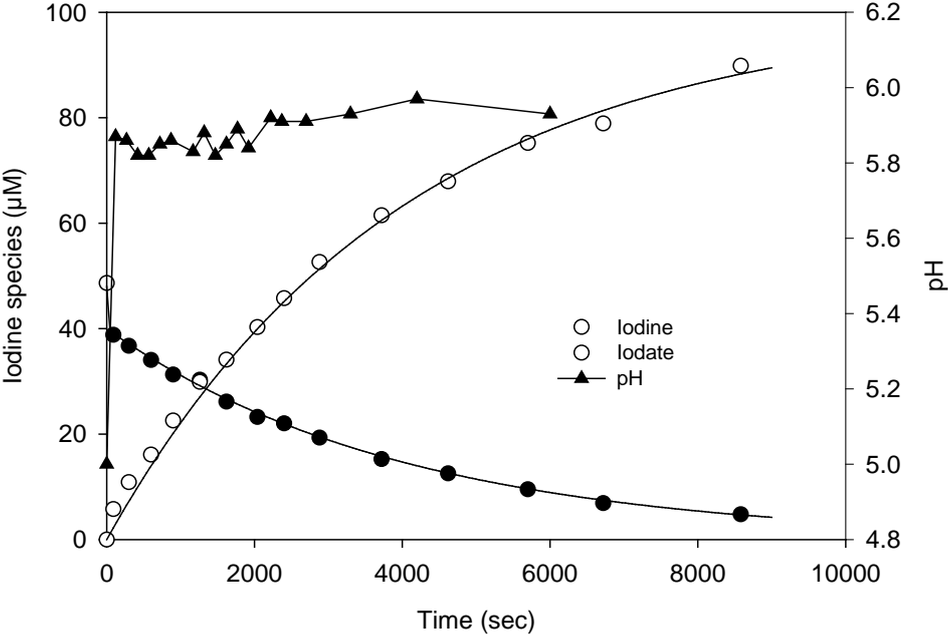


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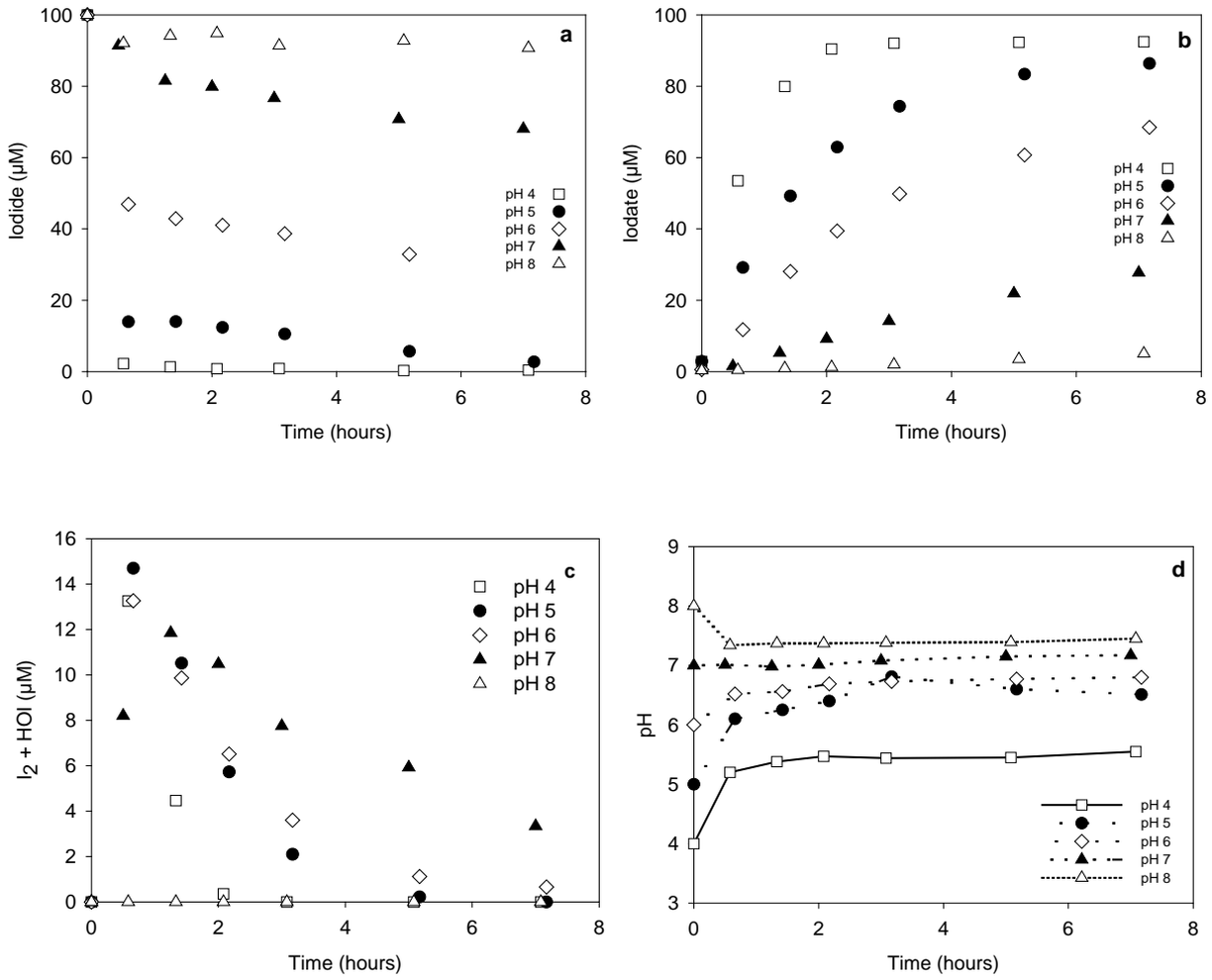


Figure 9

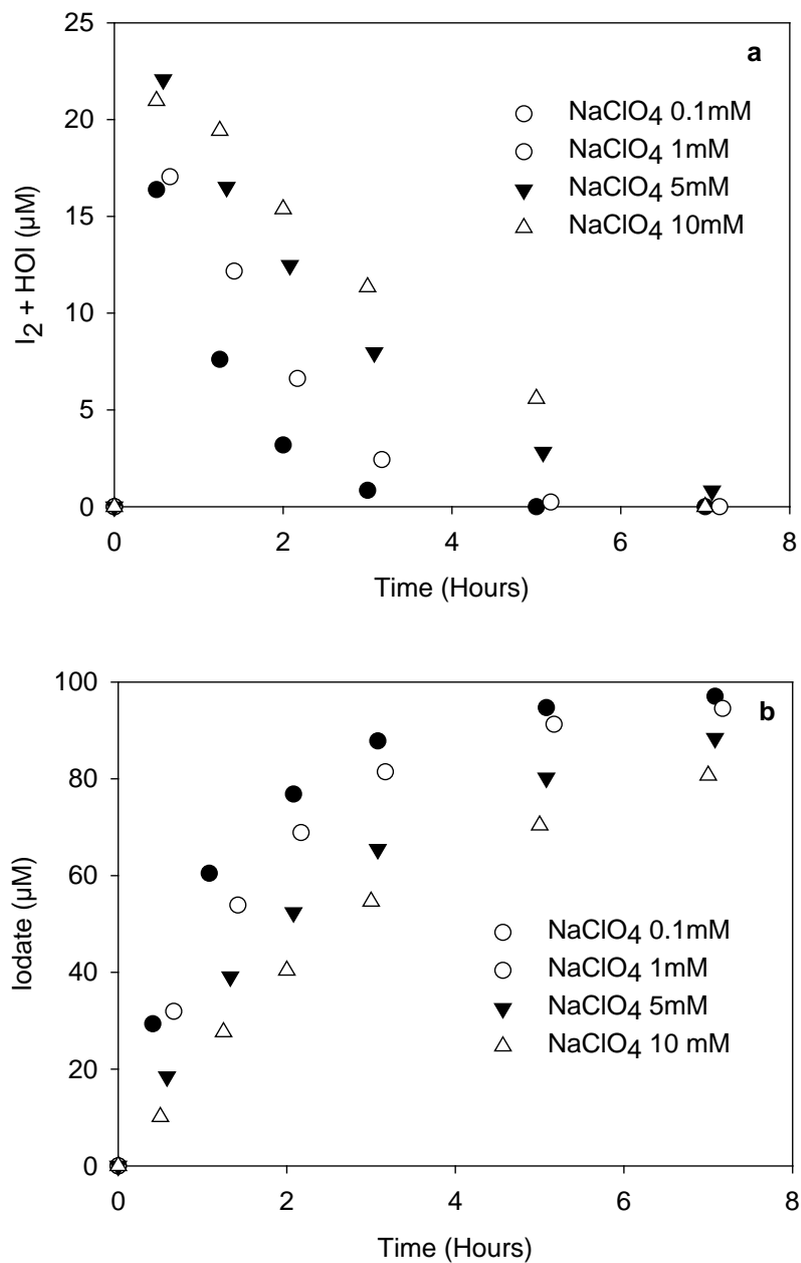


Figure 10.

