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Metal [Fe(II), Cu(I), Co(II), Mn(III)]/hydroperoxideinduced activation of dioxygen (\cdot O₂ \cdot) for the ketonization of hydrocarbons: oxygenated Fenton Chemistry

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Abstract

Recent reports have established that Fenton chemistry involves the formation of hydroper-oxide (ROOH) adducts (1) of reduced transition metals [iron(II), copper(I), and cobalt(II)] via nucleophilic addition (e.g. $Fe^{II}L_x + ROOH = L_x Fe^{II}OOR$, $BH^+](I)$) (B=py or H_2O). These reactive intermediates (1) react with (a) excess catalyst ($Fe^{II}L_x$) to form $L_xFe^{III}OH(R)$; (b) excess ROOH to form O_2 , H_2O and ROH; (c) excess hydrocarbon (RH) to form ROH (Fenton chemistry); and (d) ambient dioxygen (O_2) to form adducts of 1 {e.g. $I(O_2) = [L_x Fe^{III}(O_2)OOR(BH^+)]$ (5)}. (Note: All bracketed species are speculative, but consistent with the product profiles.) The latter (5) reacts selectivity with (a) methylenic carbon centers of hydrocarbons ($-CH_2$) to form ketones [e.g. $c - C_0H_{12} = c - C_0H_{10}(O)$] (oxygenated Fenton chemistry) and (b) arylolefins via dioxygenation [e.g. cis - PhCH = CHPh = color = cis - PhCH = CHPh = color = color

species 5 exhibit catalytic turnover of O_2 [e.g. 10 mM Fe^{II}(bpy) $_2^{2+}/20$ mM t-BuOOH yields 86 mM c-C₆H₈(O) (4.3 O₂ turnovers per t-BuOOH)]. For analogous combinations with other complexes the turnover numbers are: Fe^{II}(OFPh₃) $_4^{2+}$, 3.0; Fe^{II}(PA) $_2$, 3.0; Cu^I(bpy) $_2^{+}$, 1.3; Co^{II}(bpy) $_2^{2+}$, 0.7; Mn^{III}(salen)(OAc), 8.0; Mn^{III}(bpy) $_2$ (OAc) $_3$, 7.0. With PhCH₂CH₃ as the substrate the O₂-turnover numbers are: Fe^{II}(PA) $_2$, 3.2; Fe^{II}(bpy) $_2^{+}$, 1.8; Fe^{II}(OPPh₃) $_4^{2+}$, 1.9; Cu^I(bpy) $_2^{+}$, 1.4; Co^{II}(bpy) $_2^{2+}$, 0.9; Mn^{III}(salen)(OAc) (salen = Schiff base from two salicylaldehydes and ethylenediamine), 2.0; Mn^{III}(bpy) $_2$ (OAc) $_3$, 1.6. The several species 1 react as hydroxylases (including aromatic substrates) and dehydrogenases, and species 5 react as ketonization agents and dioxygenases. With excess HOOH its ML_x-induced disproportionation to O₂ and H₂O results in production of species 5 without external sources of O₂, © 1997 Elsevier Science S.A.

1. Introduction

The reaction chemistry of coordination complexes involves substitution by a more nucleophilic ligand, e.g.

$$Fe^{II}(OH_2)_0^{2+} + 3bpy \approx Fe^{II}(bpy)_3^{2+} + 6H_2O$$
 (1)

$$Fe^{II}(bpy)_3^{2+} + H_2Y^{2-} \rightleftharpoons Fe^{II}Y^{2-} + 2bpyH^{+} + bpy$$
 (2)

$$Fe^{II}(bpy)_3^{2+} + HOOH \rightleftharpoons I(bpy)_2^{+} Fe^{II}OOH(bpyH^{+})]$$
(3)

In this set of displacement reactions bipyridine (bpy) is a stronger nucleophile than H_2O , EDTA $(H_2Y^2^-)$ stronger than bpy, and HOOH stronger than bpy, H_2O and the carboxylates of $H_2Y^2^-$. As with all ligand-substitution reactions the more nucleophilic ligand displaces an existing ligand via a single-electron-transfer (SET) mechanism (the entering ligand transfers an electron via the metal center to the displaced ligand). This is analogous to the S_N2 mechanism for nucleophilic organic reactions [1]

$$HO' + n-BuBr \xrightarrow{S_{N}2} HOBu-n + Br'$$
 (4)

$$HO^{-} + Fe^{II}Cl_{3} \rightarrow [HOFe^{II}Cl_{3}] \rightarrow HOFe^{II}Cl_{2} + Cl^{-}$$
(5)

$$2NH_3 + Ag^tCl(s) \rightarrow Ag^t(NH_3)_2^4 + Cl$$
 (6)

With saturated electrophiles (n-BuBr, Eq. (4)) the SET is to a less nucleophilic leaving group $N = +R[X](E) \rightarrow N : R + [X]$:

The nucleophilicity (Nu) of a Lewis base (nucleophile, N) is equal to its oxidation potential $[(E_{ox})_N]$ minus the bond-formation free energy $\{(-\Delta G_{BF})_{N-E'}$ for the N-E' bond that results from its reaction with an electrophile [E; electrophilicity (El) equals $(E_{red})_E$] [2]

$$N + E \rightarrow N - E'(+X) - -\Delta G_{reac}$$
 (7)

 $(-\Delta G_{BF})$ (for nucleophilic displacement reactions)

$$Nu = (E_{ox})_{N} - [(-\Delta G_{BF})_{N-E'}/23.1 \text{ kcal (eV)}^{-1} \text{ mol}^{-1}]$$
(8)

$$El = (E_{red})_{E} \tag{9}$$

The free-energy for the reaction of a nucleophile with an electrophile (Eq. (7)) can be determined with the relation

$$-\Delta G_{\text{reac}} = (El - Nu)23.1 \text{ kcal mol}^{-1} = [(E_{\text{red}})_{\text{E}} - (E_{\text{ox}})_{\text{N}}]23.1 + (-\Delta G_{\text{BF}})_{\text{N-E}},$$
(10)

Hence, the driving force for nucleophile/electrophile electron-transfer reactions is the redox potential for the nucleophile in the solution matrix (HO^-/HO^- , +0.92 V vs NHE in MeCN) [3] plus the nucleophile-substrate bond-formation free energy ($-\Delta G_{BF}$, n-Bu-OH; ~ 83 kcal mol⁻¹).

Nucleophilicity =
$$(E_{1/2})_{ox} - (-\Delta G_{BF(N-E)}/23.1)$$
 (11)
= $(+0.92 \text{ V}) - [83 \text{ kcal mol}^{-1}/23.1 \text{ kcal mol}^{-1} (\text{eV})^{-1}]$
= $(+0.92 - 3.6) \text{ V} = -2.70 \text{ V vs NHE}$
[$n - BuBr + e^- \rightarrow n - Bu + Br^-; E^{\circ\prime}, -2.0 \text{ V vs NHE}]$

Thus, in MeCN reaction Eq. (4) is exothermic by about 16 keal mol⁻¹ $[-\Delta G_{\text{reaction}} = (El - Nu) \text{ eV} = 0.7 \text{ V} \times 23.1 \text{ keal mol}^{-1} \text{ (eV)}^{-1} \approx 16 \text{ keal mol}^{-1}].$

With reaction Eq. (5) the bond-formation free energy for Cl_2Fe^{HI} OH is 62 keal mol⁻¹ and $Fe^{III}Cl_3$ is reduced at ± 0.50 V vs NHE in acetonitrile (EI). Hence, the nucleophilicity of HO in this reaction is given by the relation

Nucleophilicity =
$$(E_{1/2})_{ox} = (-\Delta G_{\Theta F(N-1)}/23.1)$$
 (12)
= $(+0.92 \text{ V}) - [62 \text{ kcal mol}^{-1}/23.1 \text{ kcal mol}^{-1} (\text{eV})^{-1}]$
= $(+0.92 - 2.68) \text{ V} = -1.76 \text{ V vs NHE}$

The reaction is exothermic by about $52 \text{ kcal mol}^{-1} \left[-\Delta G_{\text{reaction}} = (El - Nu) \text{ eV} = 2.26 \text{ V} \times 23.1 \text{ kcal mol}^{-1} \text{ (eV)}^{-1} = 52.2 \text{ kcal mol}^{-1}\right].$

Table I summarizes the oxidation potentials for nucleophiles and the reduction potentials for electrophiles in acetonitrile (MeCN) and in water [2]. The relative positions will be the same in other solvents, but nucleophiles that are stronger than HO "will be "leveled" to its nucleophilicity in aqueous media. Under all conditions hydroperoxides (e.g. HOOH, t-BuOOH) are more nucleophilic than water (and alcohols), and peroxide anions are stronger nucleophiles than other oxy anions (HO", PhO").

Thus, hydroperoxides are favored relative to water in the ligand-centered

Table 1 Nucleophilicity	and electrophilic	ity of molecules	and ions (strongest or	most reactive at top of	of listing)
Nucleophile	(E _{1/2}) _{ox} (V)	vs NHE	Electrophile	$(E_{1/2})_{red}$ (V) (MeCN)	vs NHE

Nucleophile	(E _{1/2}) _{ox} (V) (MeCN)	vs NHE (H ₂ O)	Electrophile	$(E_{1/2})_{r \sim d} (V)$ (MeCN)	vs NHE (H₂O)
¢	-3.9	+2.9	H ₂ O ⁺ ·	+3.2	+2.7
(TPP~·)Co~	-1.7		$Fe^{III}(bpy)_3^{3+}$	+1.3	+1.1
(TPP-)Fe	- 1.4		но-	+0.9	+1.9
(TPP)Fe"	-0.8		Fe ^{itt} (PA) ₃	+0.4	
(TPP)Co"	-0.6		(TPP)Fe ⁱⁿ (py);	+0.4	
0,	0,7	- -0.2	(TPP)Fe ^{III} Cl	+0.2	
PhCH ₂ S	0.0		AQ (anthraquinone)	-0.6	
H00	0.0	+0.2	Coll(bpy)2.	0.6	
PhO	+0.3		O ₂	~ 0.7	-0.2
MeaN	+0.7		$Cu^{1}(bpy)_{2}^{2}$	~ 0.9	
HO.	+ 0.9	+1.9	Fell(OPPh3)24	-0.9	
MeC(O)O	+1.3		Fe ¹¹ (bpy) ₂ ²⁺	-0.9	
PhOH	+1.7		CCI ₄	-0.9	
Pyridine	+ 2.0		Fe ^{II} Cl ₂	1.0	
C1	÷ 2.2	+ 2.4	H ³ O *	-1.6	-2.1
ноон	± 2.3	+0.9 (pH 7)	n-BuI	 1.9	
H ₂ O	+3.0	+2.3 (pH 7)	n BuBr	-2.2	
*		•	n-BuCl	2.3	
			H ₂ O	- 3.9	2.9

nucleophilic substitution reactions of coordination chemistry

$$Fe^{IP}(bpv)_{1}^{N+} + H_{2}O \rightarrow (bpv)_{2}^{N+} + Fe^{III} + OH + bpvH^{+}$$
(13)

$$Fe^{III}(bpy)_3^{1*} + HOOH \rightarrow (bpy)_2^{2*}Fe^{III} - OOH + bpyH^*$$
 (14)

$$(PA)_2Fe^{111}Cl + H_2O/py \rightarrow (PA)_2Fe^{111} - OH + pyHCl$$
 (15)

$$(PA)_2 Fe^{II}CI + HOOH/py \rightarrow (PA)_2 Fe^{III} - OOH + pyHCI$$
 (16)

$$Fe^{H}(PA)_{2} + HOOH/py \rightarrow [(PA)_{2}]Fe^{H} - OOH + pyH^{4}$$
(17)

$$Fe^{II}(bpy)_{3}^{2+} + HOOH \rightarrow [(bpv)_{3}^{+}]Fe^{II} - OOH(bpyH^{+})$$
 (18)

The nucleophilic character of hydroperoxides follows from the fact that HOO is a much stronger Lewis base than HO $[(E_{ox})_{HOO} = +0.20 \text{ V}]$ vs NHE vs $(E_{ox})_{HO} = +1.89 \text{ V}$ (pH 14)] [the more negative, or less positive, the potential the more basic; with the electron (e) at the reduction potential of the solvent the ultimate Lewis base, -2.93 V vs NHE for H₂O]. Relative to the Lewis basicity of H₂O $[(E_{ox})_{H_2O,pH/5}, +2.43 \text{ V}]$, that for HOOH $[(E_{ox})_{HOOHpH/5}, +1.01 \text{ V}]$ also is much greater [1,2].

2. Fenton chemistry

The traditional formulation of the one-to-one primary step for Fenton reagents $[Fe^{II}(OH_2)_6^{2+}/HOOH$ in H_2O at pH 2] depicts the production of free hydroxyl radical $(HO\cdot)$ [4-8]

$$Fe^{II}(OH_2)_6^{2+} + HOOH \xrightarrow{k_{19}} (H_2O)_5^{2+} Fe^{III}OH + H_2O + HO$$
 (19)

$$k_{19} = 41 \text{ M}^{-1} \text{ s}^{-1}$$

With this assumption, the subsequent reactions of Fenton reagents have been based on the primary chemistry of $HO \cdot$ (generated by radiolysis of H_2O or photolysis of HOOH) [9], which reacts with iron(II) (all in aqueous media)

$$HO \cdot + Fe^{!!}(OH_2)_6^{2+} \xrightarrow{k_{20}} (H_2O)_5^{2+} Fe^{!!!}OH + H_2O$$
 (20)

$$k_{20} = 3 \times 10^8 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$$

and aliphatic (RH) and aromatic (PhR) hydrocarbons [9]

$$HO \cdot + RH \xrightarrow{k_{21}} R \cdot + H_2O \tag{21}$$

$$k_{21} = 1.1 \times 10^8 \text{ M}^{-1} \text{ s}^{-1} \text{ (CH4)}, 1.8 \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \text{ (C2H6)}, 3.7 \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \text{ (c-C5H10; cyclopentane)}$$

$$HO \cdot + PhR \xrightarrow{k_{2,2}} HOPh'R \xrightarrow{\uparrow} {}^{1}_{2}RC_{n}H_{4} - C_{n}H_{4}R$$

$$H_{2}O$$

$$(20)$$

$$k_{22} = 3.0 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$$
 (PhCH₃, 97% aryl addition), $7.5 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$ (PhCH₂CH₃, 85% aryl addition)

The resultant earbon radical (R·) can (a) dimerize to R₂, (b) react with a second HO· to form ROH, and (c) in the presence of air, couple to O₂ to form ROO· {unreactive with saturated hydrocarbons; dimerizes to [ROOOOR] \rightarrow ROOR+O₂ (when R is tertiary), $k_d = 10^3 - 10^7$ M⁻¹ s⁻¹} [6].

The kinetics for substrate reactivities with Fenton-generated "HO·" usually are determined via the relative rate of disappearance of iron(II) (Eq. (20)) to that of the substrate [4]. However, if Fenton reagents generate reactive intermediates (X) other than free HO·, the reactivity of X with iron(II) and organic substrates will be different and may not produce free carbon radicals (R·).

In 1989 we discussed the characteristics of a Fenton reagent in an organic solvent matrix [Fe^{II}(PA)₂ (PAH = picolinic acid)/HOOH/2:1 pyridine (py)/acetic acid (HOAc) (mol/mol)] [10]. With one-to-one Fe(II)/HOOH stoichiometry (under an Ar atmosphere) the system reacted with hydrocarbons in a manner similar to that of traditional aqueous Fenton reagents [c-C₆H₁₂ \rightarrow (c-C₆H₁₁)pyl (pyl; pyridyl, C₅II₄N) (or c-C₆H₁₁OH)]. As with all Fenton systems, the dominant product is not bicyclohexyl (c-C₆H₁₁)₂, which is the major product from the reaction of HO· with cyclohexane (c-C₆H₁₂).

product profiles for substrates Table 2 the two summarizes (c-C₆H₁₂, PhCH₂CH₃) when combined with 1:20 mol-ratios of 1:1 OL ML_x(M=Fe, Co, Cu)/HOOH in the absence and the presence of O₂ (1 atm) [11-13]. The kinetic isotope effect for cyclohexane [KIE, $k_{c-C_6H_{12}}/k_{c-C_6D_{12}}$] in relation to its major products is listed for the Fenton reagents under various reaction conditions. In contrast to the 1:1 Fe^{II}(PA)₂/HOOH system [dominant product;

Table 2
Comparison of hydrocarbon (RH) reactivities for classical and oxygenated Fenton reagents with those for free hydroxyl radical (HO·)

Oxidant/solvent ^b	Primary product	(yie	ld, mM :	± 5%)*		
	c-C ₆ H ₁₂			PhCH ₂ CH ₃ $\{k_{e-C_6H_{12}}/k_{PhCH_2CH_3}\}^4$	1	
HO·/H ₂ O°	c-C ₆ H _{t1} ·		[1.0]	HOPh · CH ₂ CH ₃		{0.1}
1:1 Fe ^H (OH ₂) ₆ ⁺ /HOOH]/H ₂ O(pH 2) ^f	c-C ₆ H ₁₁ OH		[1.1]	PhCH(Me)OH		
1:1 Fe ^{II} (PA) ₂ /HOOH}/(py) ₂ HOAc ⁴	$(c-C_{b}H_{11})C_{5}H_{4}N$	4	[1.7]	PhC(O)CH ₃	2	{0.3}
1:1 Fe ^{II} (PA) ₂ /HOOH, O ₂ I/(py) ₂ HOAc	$c \cdot C_0 H_{10}(0)$	2	[2.1]	PhC(O)CH ₃	4	{0.1}
1:20 Fe ^{tt} (PA) ₂ /HOOH]/(py) ₂ HOAc ^h	$c - C_0 H_{10}(0)$	27	[2.5]	PhC(O)CH ₃	23	{0.2}
	(c-C ₀ H ₃₁)C ₅ H ₄ N	4	[1.7]	HOPhCH ₂ CH ₃	5	
1:20 Fe ^{ll} (PA) ₂ /HOOH, O ₂]/(py) ₂ HOAc	$C \cdot C_0 H_{10}(0)$	15	[2.1]	PhC(O)CH ₃	27	{0.1}
1:1 Fe"(PA)2/HOOH]/MeCN	c-C _n H ₁₁ OH	3		PhC(O)CH ₃	6	{0.1}
1:20 Fe ^H (PA) ₂ /HOOH]/McCN	$c \cdot C_0 H_{10}(O)$	2				
1:20 Fe ^{III} Cl ₄ /HOOH]/MeCN	c-C _o H ₁₁ OH,Ct	24	[2.9]	PhCH(OH,CI)CH ₃]	20	{0.2}
	$C \cdot C_0 H_{10}(O)$	8	im	PhC(O)CH ₃	15	{0,1}
1:1 Fe ^{II} (PA) ₂ /HOOH]/McCN ¹	0		•			
1:1 Fell(bpy)}* /HOOH,O ₃]/McCN	$c_{i}C_{6}H_{10}(0)$	1		PhC(O)CH ₃	2	(0.1)
1:20 Fe ⁰ (bpy)§*/HOOH]/McCN ¹	$\alpha C_0 H_{10}(O)$	5	[4.0]	PhC(O)CH ₃	14	$\{0,1\}$
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	△C ₆ H _H OH	4	[1.4]			
HHI Fell(OPPha) 14 /HOOH/HCI/McCNh	eC _a H _u Cl	4	[1.8]			
1:20 Fe ^{II} (OPPh ₃) ^{2,4} /HOOH]/MeCN ¹	e-C ₆ H _H OH	7	[1.9]	PhCH (Me)OH	21	{0,1}
, , , ,	$c = C_0 H_{10}(O)$	6	[01<]	PhC(O)CH ₃	6	(0.2)
1:20 Fe ^{it} (O ₂ bpy)}* /HOOH]/(MeCN) _a py ^t	$c=C_nH_{10}(O)$	12		PhC(O)CH ₃	14	
1:20 Co ^{II} (bpy) ₂ */HOOH]/(MeCN) ₄ py ¹	$c \cdot C_0 H_{10}(O)$	20		PhC(O)CH ₃	20	{0.2}
1:20 Cul(bpy); /HOOH]/(MeCN),pyl	$c \cdot C_0 H_{10}(O)$		[2.5]	PhC(O)CH ₃	12	(0.2)
1:20 Cu ¹ (bpy) ₂ /HOOH ₂ ,O ₂]/(MeCN) ₄ py ¹	$c \cdot C_0 H_{10}(O)$		[2.4]	PhC(O)CH ₃	12	(0.2)

^{*} Substrate and FeL, combined in 3.5 mL of solvent, followed by the slow addition of HOOH (50%, in H_2O) to give 10 or 100 mM HOOH. The product solutions were analyzed by capillary-column gas chromatography and GC-MS after a reaction time of 3 h at $24 \pm 2^{\circ}C$.

b Solvents: (py)2HOAc, 2:1 mol ratio; (MeCN)4py, 4:1 mol ratio.

² Kinetic isotope effect, $k_{e=C_0H_{42}}/k_{e=C_0H_{12}}$.

A Relative reactivity of & CoH12 versus PhCH2CH2 (per CH2-group).

e Ref. [9].

f Ref. [8].

^{* 1:1, 10} mM ML_x/10 mM HOOH.

h 1:20, 5 mM ML,/100 mM HOOR.

^{111 [}Fell(OPPh₃)2', Coll(bpy)5' and Cul(bpy)5' HOOH] also are unreactive with c-C₆H₁₂.

¹ Ref. [13].

k Ref. [14].

¹ Ref. [15].

cyclohexyl pyridyl, $(c-C_6H_{11})$ pyl], the 1:20 system yields $c-C_6H_{10}(O)$ (cyclohexanone) as the dominant product plus some $(c-C_6H_{11})$ pyl or $c-C_6H_{11}OH$. With the 1:1 systems in the presence of O_2 (1 atm), ketonization of methylenic carbon centers is dominant $[c-C_6H_{12}\rightarrow c-C_6H_{10}(O)]$.

These [11-13] and related studies of iron(II)/HCl [14] and copper(I) [15] Fenton systems have confirmed that the metal/HOOH ratio is decisive with respect to reactivity and product profile. Thus, 1:1 $ML_x/HOOH$ combinations of $Fe^{II}(bpy)_2^{2+}$, $Fe^{II}(OPPh_3)_4^{2+}$, $Co^{II}(bpy)_2^{2+}$ and $Cu^I(bpy)_2^{++}$ are not reactive with $c-C_6H_{12}$, but 1:20 combinations yield substantial quantities of $c-C_6H_{10}(O)$ as the dominant product [their 1:1 combination in the presence of O_2 also transforms $c-C_6H_{12}$ to $c-C_6H_{10}(O)$]. Clearly the latter conditions produce a different reactive intermediate than that from the 1:1 combination of classical Fenton chemistry.

The product profiles for a Fenton reagent $[Fe^{II}(PA)_2/HOOH/2:1 py/HOAc (mol/mol)]$ with several organic substrates (RH or ArH) have been compared with those for free hydroxyl radical (HO·) in an aqueous matrix [10-13,16]. In no case is substrate dimer (R-R) (dominant product for HO·/saturated-hydrocarbon reactions in the absence of O₂) or ROOR (dominant product for HO·/saturated-hydrocarbon reactions in the presence of O₂) detected in the product solutions [9].

In the absence of O_2 , (a) the Fenton systems yield (i) R(pyl) (alkyl pyridyl) or ROH from saturated hydrocarbons (RH), (ii) PhOH from PhH, (iii) 3-hydroxypyridyl [3-HO(pyl)] from py, and (iv) PhC(O)CH₃ from PhCH₂CH₃; and (b) HO· yields (i) (R· \rightarrow R-R) from RH, (ii) (HOPhH \rightarrow Ph-Ph) from PhH, (iii) hydroxyl adduct [HOpy(o/p, 2.0) \rightarrow pyl-pyl] (bipyridyl; bpy) from py, and (iv) (HOArH \rightarrow Ar-Ar) from PhCH₂CH₃. In the presence of O_2 , (a) the Fenton system yields ketones from the methylenic centers of hydrocarbons [RH; c-C₆H₁₂ $\rightarrow c$ -C₆H₁₀(O)], and (b) HO· yields (ROO· \rightarrow ROOR +O₂) from RH.

Although HO: reacts with CH₄ ($k = 1.1 \times 10^8 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$) [9], Fenton reagents are unreactive. Hydroxyl radical reacts with Fe^{II}(bpy)₃²⁺ (bpy, 2.2'-bipyridyl) via aryl addition to give (bpy)₂²⁺ Fe^{III}(bpy – OH) (bpy-OH, hydroxyl derivative of bipyridyl) ($k = 9 \times 10^9 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$) [9], but the 1:1 combination of Fe^{II}(bpy)₂²⁺ and HOOH in MeCN is unreactive (Table 2).

In spite of the common belief that Fenton reagents (Fe^{II}L_x/HOOH) (L, ligand) produce free HO· (Eq. (19)), recent studies [11,17,18] and the results of Table 2 provide clear evidence that free HO· is not the dominant reactant, and that with highly stabilized iron(II) complexes [Fe^{II}(diethylenetriaminepentaacetate) and Fe^{II}(EDTA)] a nucleophilic adduct {(EDTA)Fe^{II}-O-OH(H₃O⁺), 1; "bound HO·"}] reacts directly with substrates [18]. Another study finds product profiles that are inconsistent with free HO· as the dominant reactive intermediate for a biological Fenton reagent [19].

The 1:1 Fe^{II}(PA)₂/HOOH combination in 2:1 py/HOAc is an effective Fenton reagent for organic substrates [10], and has reactivities and product profiles that are within the same mechanistic framework as those for traditional aqueous Fenton reagents [4]. Hence, the initial step is the reversible nucleophilic addition of HOOH to Fe^{II}(PA)₂ to give the primary reactive intermediate (1) [11-13] which reacts with (a) excess Fe^{II}(PA)₂ via path A $(k=2\times10^3 \,\mathrm{M}^{-1}\,\mathrm{s}^{-1})$ [10], (b) excess HOOH via

$$Fe^{II}(PA)_{2} + HOOH_{2} \frac{[Fe^{II}OOH(PA)_{2}(pyH)]}{[Fe^{II}OH(PA)_{2}(pyH)]} 2 (PA)_{2} Fe^{III}OH$$

$$\frac{1}{c \cdot C_{6}H_{12}} (C) \qquad (B) \qquad py$$

$$(c \cdot C_{6}H_{11})pyl + 2 H_{2}O + Fe^{II}(PA)_{2}$$

$$KIE, 1.7 \qquad O_{2} + 2 H_{2}O + Fe^{II}(PA)_{2}$$

$$(23)$$

path **B** to give O_2 and (c) excess c- C_6H_{12} via path **C** to give (c- $C_6H_{11})$ pyl (cyclohexyl pyridyl) [aqueous Fenton systems produce c- $C_6H_{11}OH$ with a kinetic-isotope-effect (KIE) of 1.1 [8], and free HO· (pulse radiolysis) produces c- C_6H_{11} · with a KIE of 1.0 [9]. Although radical traps (e.g. PhSeSePh, BrCCl₃, Me₂SO) [9] often are used to "prove" that free carbon radicals are formed by "free HO·" from Fenton reagents, these also react with nonradicals (e.g. the intermediate of path **C**, Eq. (23); Table 2). With 1M c- C_6H_{12} the process of path **C** is facilitated via formation of an intermediate (2) that produces 2-(c- $C_6H_{11})$ pyl and 4-(c- $C_6H_{11})$ pyl

$$I + c - C_0 H_{12} + py \rightarrow \{(PA)_2 Fe^{IV} (OH)[pyl(c - C_0 H_{11})]\} \rightarrow Fe^{II} (PA)_2$$

$$+ (c - C_0 H_{11})pyl + H_2 O$$
(24)

The presence of pyridine in the solvent matrix causes the primary reactant to be $[(L_x^*)Fe^BOOH(pyH^*)]$ (1), which reacts with aliphatic substrates (RH) to produce alkyl pyridyls (Rpyl) via $[(L_x^{2^*})Fe^{IV}(pyR)(OH)]$ (2). When oxidized metal complexes [e.g. $Fe^{IH}Cl_3$, $Fe^{IH}(PA)_3$, $Cu^H(bpy)_2^{2^*}$] are used, the initial event appears to be reduction by HOOH, e.g. [12,13]

$$2Fe^{H}Cl_3 + HOOH \rightarrow 2Fe^{H}Cl_2 + O_2 + 2HCl$$
 (25)

The Fe^{II}Cl₂ product in turn forms [Fe^{II}OOH(Cl₂)(H₃O⁺)] (1), which reacts with c-C₀H₁₂ and PhCH₂CH₃ via [Cl₂Fe^{IV}(OH)(R)] (2) to produce approximately 50:50 mixtures of ROH and RCl [13]. With HOOH and c-C₀H₁₂ the [KIE] value for 1 is 2.9 [13], and with t-BuOOH it is 4.3. The porphyrin catalyst {(Cl₈TPP)Fe^{II}; [tetra(2,6-dichloro-phenyl)porphyrin]iron(II)} reacts with t-BuOOH to form [(Cl₈TPP)Fe^{II}OOBu-t(H₃O⁺)] (1), which reacts with c-C₀H₁₂ via [(Cl₈TPP)Fe^{IV}(OH)(R)] (2) to produce ROH [KIE = 5.0] [12].

In summary: Fenton reagents do not produce (a) free HO_{+} , (b) free carbon radicals (R ·) or (c) aryl adducts (HO₋Ar ·). Early work [20] has demonstrated that the primary chemistry of HOOH is nucleophilic addition, even in matrices as weakly basic as water at pH 2. Hence, Fenton reagents [reduced electrophilic transition-metal complexes (Fe^HL_x, Cu^HL_x and Co^HL_x)] must have a primary step of nucleophilic addition to the metal center to give 1 (the reactive intermediate of Fenton reagents). The efficient and selective reactivity of 1 (Fenton chemistry) makes it a more reasonable cytotoxic agent than free HO₊ within the oxy-radical theory of aging and heart disease [21,22].

3. Oxygenated Fenton chemistry

When excess HOOH (or t-BuOOH) is combined with transition-metal complexes, it becomes the dominant substrate for the initially formed Fenton intermediate (1, Eq. (23)). This reaction facilitates the disproportionation of hydroperoxides via species 3 (rapid in the case of HOOH; and much slower in the case of t-BuOOH) [12,13,15].

$$[Fe^{II}OOR(PA)_{2}^{-1}(BH^{*})] + HOOR \longrightarrow [(PA)_{2}Fe^{IV}(OH)(OOR)]$$

$$1 \qquad ROH, B$$

$$R = H \text{ or } t\text{-Bu}$$

$$O_{2} + Fe^{II}(PA)_{2}$$

$$(26)$$

Hence, a system with excess HOOH initially produces its own O_2 -atmosphere. For example the 5 mM Fe^{II}(PA)₂/100 mM HOOH/I M c-C₆H₁₂ system yields 27 mM c-C₆H₁₀(O) and 4 mM (c-C₆H₁₁)pyl (the respective [KIE]-values are 2.5 and 1.7. Table 2). More than half of the HOOH is decomposed to O_2 (Eq. (26)). When t-BuOOH is used in place of HOOH the system yields 7 mM c-C₆H₁₁OOBu-t ([KIE] = 8.4), 11 mM c-C₆H₁₀(O) ([KIE]=7.6) and 19 mM (c-C₆H₁₁)pyl ([KIE]=4.6). Again, almost half of the t-BuOOH is decomposed to O_2 via Eq. (26). Combinations of Fe^{II}(PA)₂ and HOOH in the presence of O_2 transform c-C₆H₁₂ to c-C₆H₁₀(O) ([KIE], 2.1) as the only detectable product (Table 2).

Although 1:1 combinations of $[Fe^{II}(bpy)_2^{2+}, Fe^{II}(O_2bpy)_2^{2+}, Fe^{II}(OPPh_3)_4^{2+}, Co^{II}(bpy)_2^{2+}]$ or $Cu^I(bpy)_2^{2+}]/HOOH$ in the absence of O_2 are unreactive with c- C_6H_{12} , they readily react in its presence to produce c- $C_6H_{10}(O)$. When their ratio is 1:20 ML $_2/HOOH$, most of the HOOH is transformed to O_2 via Eq. (26), which results in analogous Fenton-induced activation of O_2 for reaction with c- C_6H_{12} and $PhCH_2CH_3$ (Table 2).

The 9 mM Fe^{II}(PA)₂/9 mM *t*-BuOOH system in the absence of O_2 and substrate reacts via a Fenton process (path A, Eq. (23)) to give $(PA)_2Fe^{III}OH$. In the presence of O_2 (with or without 1M c-C₆H₁₂) there is no evidence for free Fe^{II}(PA)₂ in the reaction matrix (no electrochemical oxidation, but a two-electron per iron irreversible reduction) [12]. These observations are compelling evidence that species 1 (formed from *t*-BuOOH) produces an O_2 -adduct (5)

$$I + O_2 \rightarrow [Fe^{III}(OOBu - t)(PA)_2^{-}(O_2)(pyH^+)]5[I(O_2)]$$
 (27)

which reacts with excess c-C₆H₁₂ to produce c-C₆H₁₀(O). The dioxygen adduct (5) appears to be the steady state primary reactive intermediate rather than species 1 on the basis of the enhanced [KIE] value for ketone formation {8.5 [for initial formation of 6] vs 4.6 [for formation of (c-C₆H₁₁)pyl]}. Table 3 summarizes the product profiles for several ML_x/t-BuOOH/O₂ (1 atm) combinations with c-C₆H₁₂ and c-C₆H₁₀. In all cases the oxygen atoms that are incorporated in the product species come from O₂ [12].

The production of 16 mM PhC(O)Me by the 5 mM Fe^{II}(PA)₂/5 mM t-BuOOH/O₂/1M PhCH₂CH₃ system [12] indicates that (a) most

Motes [ML]; Me = Fe, Cu, Co, Mr] - Brook-indused arrivation of O. (i after for the oxygenation of cyclohexane (e-C₆H₁₃) and cyclohexene (e-C₆H₁₆); Oxygenated Person chansity

(0)°(26)′(26)′(20)′(20)′(20)′(20)′(20)′(20)′(20)′(20		r-Bucoh Salvert	Solvent	Products (EMES)	f±5₹; «C,H ₁ 0H	E facy ^b	c-C _e H _{1d} (1M)	,-С ₆ Н ₉ ОН	c-C ₆ H ₁₀ (0) (epoxide)	Effncy ^b (%)
20 21 FY HOA: 12 0 60 45 20 21 FY HOA: 1 66 85 0 20 MeCN 4-15 60 60 20 MeCN 5 4 6 50 60 20 MeCN 6 6 22 20 At MeCN 57 1 1 20 10 20 MeCN 6 6 199 (26) 20 MeCN 7 6 1 20 20 At MeCN 7 6 1 1 20 20 MeCN 6 0 159 (26) 20 Zi FY HOA: 6 0 159 (26) 20 Zi FY HOA: 6 0 159 (29) 20 Zi FY HOA: 6 0 22	Sam Ferry	6	1 '		- -	89	15	0	0	300
29 2.1 p. HOAc : 156 85 0 20 MeCN 4-pp 6 60 20 MeCN 5 4 6 86 (0) ^p 20 MeCN 5 4 45 71 6 MeCN 6 6 30 22 20 4:1 MeCN pp 6 0 30 25 10 MeCN 6 0 139 (26) ^p 20 MeCN 6 0 139 (26) ^p 20 MeCN 6 0 139 (26) ^p 20 Zi py HOAc 6 6 0 22	Omul Periods	#		+· ((°m)	9	\$ 5	C1	0	235
20 MeCN 4-1-7 5 (0)7 45 86 (0)7 25 MeCN 5 4 45 71 26 MeCN 5 4 45 71 6 MeCN 5 4 45 71 6 MeCN 6 0 36 22 80 41 MeCN 5 6 0 35 10 1 20 10 20 MeCN 6 0 139 (26) 20 21 py HOAc 6 0 159 (29) 20 21 py HOAc 6 0 22	Dail Felth	ន			10 10 10 10 10 10 10 10 10 10 10 10 10 1	85	0	54	0	25
35 4:1 MeCN 79: 4 6 50 60 36 MeCN 6 45 71 6 MeCN 6 36 22 20 4:1 MeCN 79: 6 0 30 25 10 4:1 MeCN 79: 1 1 20 10 20 MeCN 0 139 (26) 20 MeCN 0 159 (29) 20 2:1 py HOAc 6 0 22	O mid Ference	8	MeCh	9. 4.	(2 (2) (2) (4)	45	86 (0) ⁵	60 (2.5)	0	730
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6 MeCN 6 8 36 22 29 4:1 MeCN py 6 0 30 25 10 4:1 MeCN py 6 10 20 MeCN 6 0 139 (26)' 20 MeCN 6 0 159 (29)' 20 2:1 py HOAc 6 6 0 22	DEM EGG	H	Mech	٧,	••	5 5	71	69	0	700
29 4:1 MeCN p5 6 0 30 25 10 4:1 MeCN p5 1 20 10 20 MeCN 0 0 139 (26) 20 MeCN 0 0 159 (29) 20 2:1 py HOAc 6 0 22	I mM Falterii	9	MeCN	6	9	36	22	3		
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29 MeCN 0 0 139 (26) ^f 26 MeCN 0 0 159 (29) ^f 20 2:1 py HOAc 0 6 0 22	FORM COTCESS!		#I MEOUS	41-4		20	01	C 1	0	120
26 MeCN 8 9 0 159 (29) ⁷ 20 2:1 py HOAc 8 6 0 22	S MM Mary by 1.0 Act.	S	MeCN	CD2	Φ	0	139 (26)	41 (14)	\$ (1) ₍	925 (200)
26 24 pv HOAc \$\dirth{\phi}\$ \$\dirth{\phi}\$ \$0 22	SmM Mar welen Oace	71	NEC.	4571	©	0	159 (29)	48 (12) ^f	6(1)	1070 (210) ^c
	5 mM Mm (2, 22) (0.82)	97	2:1 py HOA:	(C)	ij	0	22	61	18	295
20 mM Ma ²⁷ (sites) (6Act) 20 2:1 py HOAc 6 6 0 24 25	26 mM Ma ^C (s.ien.) (OAc)	8	2.1 py HOAc	energy Co.	•	0	24	25	30	395

The product solitons were analyzed by capillary-column gas cinomategraphy after a reaction time of 3 h at 24 ± 2 °C.

^{*} Efficiency for product formation, mM of products per now of 1-Bu00H (169% represents one product species per 1-Bu00H).

Under un argon atmosphise.

^{*} Product was feelight, 1991.

R.R. dimer.

Under air 16.2 aim O.1

of the oxygen in the product comes from O₂ and (b) the reaction is initiated by species 5, but (c) the catalytic cycle is carried by species 6 [(PA)₂Fe^{IV}(OH)(OOR)] (three times as much product as initial *t*-BuOOH).

Cyclohexene (c-C₆H₁₀) has similar reactivity in the 10 mM Fe^{II}(bpy)₂²⁺/20 mM t-BuOOH/O₂ system with at least 4 O₂ turnovers per t-BuOOH (Table 3).

Scheme 1 outlines a set of reaction paths that are consistent with the product profiles for Fenton chemistry and oxygenated Fenton chemistry [1:1 $ML_x/HOOH$ (or $t\text{-BuOOH})/*O_2$] with cyclohexane ($c\text{-}C_6H_{12}$) as the substrate. The species 3 that is formed from t-BuOOH (rather than HOOH) is about 100 times longer lived [decomposes to O_2 and t-BuOH (Eq. (26))], and thereby can react with $c\text{-}C_6H_{12}$ to form $c\text{-}C_6H_{11}OOBu\text{-}t$. Because the *KIE* value for its formation is 8.4, the reactive intermediate must involve a pathway other than that for Fenton chemistry (path C [KIE, 4.6] and Table 3). The product is only observed with t-BuOOH (and not HOOH), which is consistent with the longer lifetime of 3 when formed from excess t-BuOOH.

With 20:1 HOOH(Bu-t)/Fe^{II}(PA)₂ ratios, substantial fractions of the HOOH(Bu-t) are decomposed to O₂ via species 3 (Eq. (26)) (rapidly for HOOH and slowly for t-BuOOH). This internally generated O₂ in turn combines with 1 to form 5, which accounts for the production of ketone (rather than ROOBu-t) in O₂-free systems of t-BuOOH. Electrochemical results [12] confirm that excess t-BuOOH with Fe^{II}(PA)₂ undergoes a sustained disproportionation to O₂ and formation of 5 [same reduction peak as for 1:1 Fe^{II}(PA)₂/t-BuOOH in the presence of O₂]. A similar set of observations and rationalizations has been presented for incorporation of O₂ derived from t-BuOOH in a Fe(III)/t-BuOOH/c-C_BH₁₀/(10:1 py/HOAc) system [23].

Other iron(H) complexes $[Fe^{II}(bpy)_2^{2+}, Fe^{II}(OPPh_3)_4^{2+}, Fe^{II}Cl_2]$ undergo initial nucleophilic addition by HOOH to form an analogue of species 1. For the $Fe^{II}L_x^{2+}$ complexes in pure McCN this is a cationic reactive intermediate $[(L_x^+)Fe^{II}OOH(H_3O^+)]$ that reacts with excess HOOH to form 3, which decomposes to O_2 and reacts with substrates. In the presence of O_2 1 reacts with $c-C_6H_{12}$ to form 6, which reacts with a second $c-C_6H_{12}$ to produce $c-C_6H_{10}(O)$ and $c-C_6H_{11}OH$ (Scheme 1).

With excess HOOH or t-BuOOH, the primary reactive intermediates (1) disproportionate HOOH (rapidly) and t-BuOOH (slowly) via path B and species 3 (Scheme 1). For the conditions of excess t-BuOOH and substrate (RH), species 3, reacts with RH to produce ROOBu-t (the [KIE] values for c-C₆H₁₂ range from 5.4 to 8.4 [13]). The reactivity parameters for [Co^{II}(bpy)₂²⁺] [13] and [Cu^I(bpy)₂⁺] [15] are similar and in accord with the proposition that all of these complexes activate HOOH initially via a species 1, which reacts with hydrocarbon substrates (RH) via path C to form species 2. In general, the reactivity parameter [KIE] has smaller values for the path-C step than for the path-D step of Scheme 1.

With excess O_2 most of the species 1 react with substrates (RH) via path **D** to form species 6, which in the case of c- C_6H_{12} , reacts initially either with excess HOOH

$$Fe^{II}(PA)_{2} + HOOH \xrightarrow{k} [L_{2}Fe^{II}OOH(pyH^{+})] \xrightarrow{Fe^{II}L_{2}} L_{2}Fe^{III}OFe^{III}L_{2}$$

$$k = 2 \times 10^{3} \, M^{T}s^{-1} \qquad (D) \qquad (C) \qquad (B) \qquad py, H_{2}O \qquad HOOH$$

$$O_{2} \quad c \cdot C_{6}H_{12} \qquad py, H_{2}O \qquad py$$

$$c\text{-}C_6H_{10}(\text{O}) + c\text{-}C_6H_{11}(\text{O}) + \text{Fe}^{11}L_2$$

$$\text{Fe}^{11}(\text{bpy})_2^{2+} + \text{HOOH} + c\text{-}C_6H_{12} \underbrace{\begin{array}{c} O_2 \\ \text{MeCN} \end{array}}_{\text{H}_2\text{O}} \underbrace{\begin{array}{c} (L_2^{2+}\text{Fe}^{1V}\text{OH}(\text{OOC}_6H_{11})) \\ \frac{|c\text{-}C_6H_{10}(\text{O}) + c\text{-}C_6H_{10}(\text{O}) + c\text{-}C_6H_{10}(\text{O}) + c\text{-}C_6H_{10}(\text{O}) + c\text{-}C_6H_{10}(\text{O}) + c\text{-}C_6H_{10}(\text{O}) \\ \text{KIE}, 4.0; \quad 1.4 \quad + \text{Fe}^{11}L_2^{2+} \\ \text{With $t\text{-}Bu(\text{O})$} + \frac{|c\text{-}C_6H_{10}(\text{O}) + c\text{-}C_6H_{10}(\text{O}) + c\text{-}C_6H_{10}(\text{$$

With t-BuOOLE

$$\begin{split} \{L_{2} Fe^{II}OOBu(py\Pi^{*})\} + c\cdot C_{n}\Pi_{12} & \longrightarrow py(C_{n}\Pi_{11}) + t\cdot BuO\Pi + H_{2}O + Fe^{II}L_{2} \\ 1 & KIE, 4.6 \\ \{L_{2} Fe^{IV}(OH)(OOBu)\} + c\cdot C_{n}\Pi_{12} & \longrightarrow c\cdot C_{n}\Pi_{11}OOBu\cdot t + H_{2}O + Fe^{II}L_{2} \\ 3 & KIE, 8.4 \\ 3 + t\cdot BuOOH + c\cdot C_{n}\Pi_{12} & \longrightarrow 6 + 2t\cdot BuOH \\ & \frac{|c\cdot C_{n}H_{12}|}{2t\cdot BuOOH} - 6 + c\cdot C_{n}H_{10}(O) + 2t\cdot BuOH \\ & 1 + O_{2} + c\cdot C_{n}H_{12} & \longrightarrow 6 + t\cdot BuOH + py \\ & \frac{|c\cdot C_{n}H_{12}|}{|c\cdot C_{n}H_{12}|} - c\cdot C_{n}H_{10}(O) + c\cdot C_{n}H_{11}OH + Fe^{II}L_{2} \end{split}$$

Scheme 1. Fenton Chemistry and the activation of dioxygen (Oxygenated Fenton Chemistry)

via path E or excess c-C₀H₁₂ via path F, and finally with excess Fe^HL₃ via path G (Scheme 1). Thus, the various species 1 (Fenton intermediates) catalyze the incorporation of O_2 into the ketone and alcohol products [e.g. $Fe^{it}(bpy)_2^{2+}/t - BuOOH$, O₂/e-C₀H₁₂, Table 3].

$$2c - C_0H_{12} + O_2^* + t - BuOOH \xrightarrow{Fe^{H}L_{\chi}} c - C_0H_{10}(O^*) + c - C_0H_{11}O^*H + t$$

$$- BuOH + H_2O$$
(29)

Results [23] for a Fe^{III}(NO₃)₃/t-BuOOH/^{IS}O₂/c-C₈H₁₆ system in acetonitrile establish that the O-atoms in the c-C₈H₁₄(O) and c-C₈H₁₅OH products are from O₂. This supports the stoichiometry of Eq. (29) [10:20 Fe^{II}L_x/t-BuOOH systems are 20–80% efficient (ketone per t-BuOOH, Table 3)].

For substrates with weak C-H bonds in their CH₂ groups {PhCH₂CH₃ and the allylic carbons of cyclohexene (c-C₆H₁₀)}, species 6 becomes a catalyst for the activation of O₂. When the reaction efficiency for such substrates is > 100% (ketone per t-BuOOH, Eq. (29), Table 3), species 6 must activate O₂ for reaction with the substrate [turnovers per Fe^{II}L_x \geq (t-BuOOH/Fe^{II}L_x)]

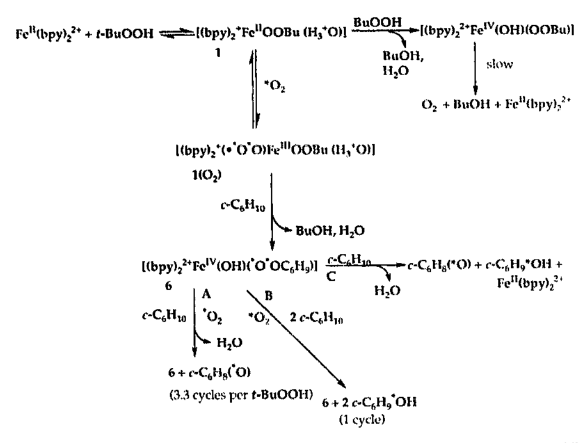
$$[L_x Fe^{IV}(OH) (OOC_6H_0)] \xrightarrow{c-C_6H_{10}} 6b + c + C_6H_8(O) + H_2O$$

$$= c + c + C_6H_0 + c + C_6H_0 + C_6H_0 + C_6H_0$$

$$= c + c + C_6H_0 + c + C_6H_0 + C_6H_0$$

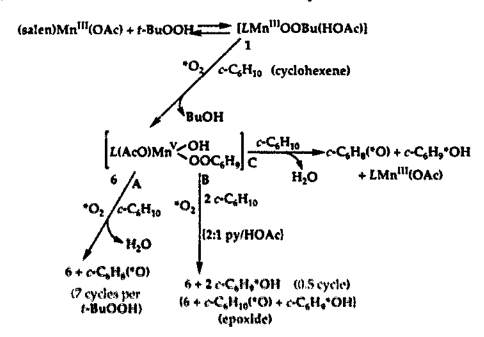
Hence, the 10 mM Fe^{II}(bpy) $_2^{2+}/20$ mM t-BuOOH/c-C $_6$ H $_{10}$ system has at least 3 O $_2$ turnovers via path E, which is similar to the 1.4 O $_2$ turovers per copper for the 5 mM Cu^I(bpy) $_2^{+}/5$ mM t-BuOOH/PhCH $_2$ CH $_3$ system [15]. Likewise, the 10 mM Fe^{II}(OPPh $_3$) $_4^{2+}/20$ mM t-BuOOH/IM c-C $_6$ H $_{10}$ system has almost 2 O $_2$ turnovers via Eq. (30) [12]. Scheme 2 outlines a set of reaction paths that are consistent with the product profiles with c-C $_6$ H $_{10}$ as the substrate.

Perhaps the most dramatic example of oxygenated Fenton chemistry is the



Scheme 2. Proposed pathways and intermediates for the metal/hydroperoxide-induced activation of O_2 for the ketonization of cyclohexane (e- C_6H_{10}).

Mn^{III}L_x/t-BuOOH-induced activation of O₂ for reaction with c-C₆H₁₀ (Table 3) [24]. With (salen)Mn^{III}OAc (salen: Schiff base of two salicylaldehydes and ediplemediamine) in MeCN more that 10 product molecules are produced per t-BuOOH. When the solvent is changed to 2:1 py/HOAc, the overall efficiency is reduced by a factor of three. However, whereas the ketone [c-C₆H₈(O), 2-cyclohexene-one] is the dominant product in MeCN, in py/HOAc approximately equal amounts of epoxide, alcohol and ketone are produced (about 4 product molecules per t-BuOOH). A set of reaction paths that are consistent with the product profiles and reaction dynamics are outlined in Scheme 3. Species 1 is unreactive with c-C₆H₁₂, and the system is ineffective when t-BuOOH is replaced by HOOH. The Mn(II) analogues of the complexes in Table 3 are not effective Fenton catalysts.



Scheme 3. Mn^{III}L_x/t-BuOOH-induced activation of O₂ for the oxygenation of olelins.

Thus, transition-metal complexes $[ML_x; Fe^{II}L_x, Mn^{III}L_x, Cu^I(bpy)_2^+]$ and $Co^{II}(bpy)_2^+]$ undergo nucleophilic addition by hydroperoxides (HOOH or t-BuOOH) to form $[L_xMOOH(BH^+)]$ (1), which in the presence of O_2 oxygenates hydrocarbons and related organic substrates via species 6 (oxygenated Fenton chemistry; Scheme 1, Scheme 2 and Scheme 3). Analysis of the results in Table 2 and Table 3 confirms that the catalytic efficiency for the various species 6 depends on the metal, its ligand, the hydroperoxide (HOOH or t-BuOOH), the solution matrix, and the substrate. Hence, the $Fe^{II}(PA)_2/HOOH/(2 py/HOAc)$ system provides the most potent reactive intermediates (1 and 5; smallest [KIE], 1.7 and 2.1, respectively for c- C_6H_{12}), but has low reaction efficiencies because HOOH acts as a competitive substrate. In contrast, the $Mn^{III}(salen)(OAc)/t$ -BuOOH/MeCN system is not an effective Fenton reagent, nor an oxygenated Fenton reagent with c- C_6H_{10} as the substrate it is by far the most efficient oxygenated Fenton reagent (8.2 ketones [c- $C_6H_8(O)]$ and 2.4 alcohols [c- $C_6H_9OH]$ produced per t-BuOOH). Table 4 summarizes the relative reaction efficiencies (ketone per hydroper-

Table 4
Relative reaction efficiencies (ketone/hydroperoxide) for the ketonization of cyclohexane (c- C_6H_{12}), cyclohexene (c- C_6H_{10}) and ethyl benzene (PhCH₂CH₃) via metal/hydroperoxide-induced activation of O_2^a

c-C ₆ H ₁₂ :HO0	OH:Fe ^{II} (PA) ₂ :	>Co ^{ll} (bpy) ₂ +	>Cu ¹ (bpy) ₂ ⁺ >	Fe ^{II} (bpy) ₂ ² ∼ F	e ¹¹ (OPPh ₃) ₄ ²⁺		
(K/HP)	0.4 [2.1] ^b	0.2 [2.9] ^b	0.1 [2.4] ^b	0.05 [4.0] ^b	0.05 [>10] ^b		
(Alc/HP)	0.0	0.0	0.07 [1.1] ^b	0.04 [1.4] ^b	0.07 [1.9] ^b		
t-BuOOH:	$Fe^{11}(PA)_2 > C$	$u^1(bpy)_2^+ > Fe$	$^{0}(bpy)_{2}^{2+} \sim Fe^{0}$	$(O(.2h_3)_4^{2+} > C$	o ^{II} (bpy)2+		
(K/HP)	0.8 [7.6] ^b	0.3 [8.8] ⁶	$0.2 [> 10]^{b}$	0.2 [10] ^b	0.1 [9.6] ^b		
(Alc/HP)	0.0	$0.1 [>8]^{b}$	0.25 [4.8] ^b	0.31 [5.4] ^b	0.03 [6.3] ^b		
$c - C_6 H_{10}$:HO0	OH:Co ^{II} (bpy) ₂	$^+ > \text{Fe}^{\text{II}}(\text{PA})_2$	$\sim Cu^{l}(bpy)_{2}^{+} >$	$Fe^{tt}(bpy)_2^{2+} > F$	e ^{ll} (OPPh ₃) ₄ ²⁺		
(K/HP)	0.2	0.1	0.1	0.05	0.03		
t – BuOOH	:Mn ^{tit} (salen)(0	OAc)>Mn ^{III} (l	$(OAc)_3 > 1$	An ^{III} (OPPh ₃) ₄ (OAc)3 > Fell (bp)	/) ² " >	
$Fe^{it}(PA)_2 >$	$Cu^{\dagger}(bpy)_{2}^{+} > 0$	Co ⁿ (bpy) ₂ +Fe	⁽¹⁾ (OPPh ₃) ₄ ²⁺				
(K/HP)	8.0	7.0	5.6	4.3	3.0	1.3	0.7
(Alc/HP)	2.4	2.1	0.6	3.0	1.8	0.0	
PhCH ₂ CH ₃ :H	OOH:Fe ^{il} (PA)	$_2 > Co^{it}(bpy)_2^2$	$^{\prime} > \text{Fe}^{\text{II}}(\text{bpy})_2^2$	>Cu ¹ (bpy) ₂ ⁺ >	Fe ¹¹ (OPPh ₃) ₄ *		
(K/HP)	0.8	0.2	0.15	0.1	0.05		
t-BuOOH	$: Fe^{11}(PA)_2 > M$	fn ^{an} (salen)(O/	Ac) > Fe ^{II} (OPPh	$_{3})_{4}^{2+} > \text{Fe}^{11}(\text{bpy})$	$\frac{3}{2}$ > $Mn^{III}(bpy)$	2(OAc)	,>
Cu ^t (bpy) ₂ ^t :	> Co ^u (Երչ) ^ի ՝	Mn ^{III} (OPPh ₃),	(OAc)3				
(K/HP)	3.2	2.0	1.9	1.8	1.6	1.4	0.9
(Alc/HP)	0.0	0.0	0.8	0.6	0.0	0.0	0.2

^{*} Data from Refs. [11-13].

oxide) for the several systems with cyclohexanc, cyclohexane and ethyl benzene. Although the various species 5 and 6 formed via HOOH are more reactive than those formed via t-BuOOH (the former have much smaller KIE values), the latter have much larger efficiencies (HOOH is a much more reactive competitive substrate than t-BuOOH).

4. Summary

Hydroperoxides (ROOH) in solution react as nucleophiles towards electrophilic substrates [SO₂; RX; HOCl; transition-metal complexes (Fe^{II}L_x, Cu^IL_x, Mn^{III}L_x, Co^{II}L_x)] [25]. In contrast, dialkyl peroxides (ROOR) [with a weaker O-O bond ($\Delta H_{\rm DBE}$ about 40 kcal mol⁻¹) than that for ROOH (about 50 kcal mol⁻¹)] are much less reactive and must be activated via homolytic dissociation of their O-O bond. Because all coordination complexes of metals (ML_x) have electrophilic metal centers with nucleophilic ligands, the primary chemistry of hydroperoxides towards them is nucleophilic addition and substitution. When the complex includes a reduced transition metal [e.g. Fe^{II}(PA)₂], nucleophilic addition yields the reactive intermediate for Fenton chemistry (1) and the precursor to the reactive intermediate for oxygenated Fenton chemistry {5 [1(O₂)]}. Table 5 lists proposed formulations for the reactive

^b Kinetic-isotope-effect; $[KIE] = k_{c-C_0B_{12}}/k_{c-C_0D_{12}}$.

Table 5 Proposed formulations for the reactive intermediates for Fenton chemistry (1) and oxygenated Fenton chemistry $\{5 \ [1(0_2)] \ and \ 6\}$

2py/HOAc {[(PA);](· OO)Fe ^{tt} OOH(Bu)[pyH *]}	MeCN
((DA) = 1(OO) EalifOOU (Day) = (U+1)	
{[(PA) ₂ ['OO)re"OOn{Bu}[pyn]}	$\{[(Cl)_2^+](\cdot OO)Fe^{HOOH(Bu)}[H_3O^+]\}$
$\{(bpy)_2^+(\cdot OO)Fe^{itt}OOH(Bu)[H_3O^+]\}$	$\{(bpy)_2(O_2)Cu^{III}OOH(Bu)[H_3O^+]\}$
{(bpy); (·OO)Co ^{lll} OOH(Bu)[pyH*]}	{(bpy) ₂ (AcO) ₂ (·OO)Mn ^{IV} OOBu[HOAc]}
	{(salen) (·OO)Mn ^{IV} OOBu[HOAc]}
$[(PA_2Fe^{iV}(OH)(OOR)][Cl_2Fe^{iV}(OH)(OOR)]$	[(bpy)3*Fe ^{IV} (OH)(OOR)]
	$[(bpy)_2^2 + Co^{(V)}(OH)(OOR)]$
[(bpy) ₂ (AcO) ₃ Mn ^V (OH)(OOR)]	[(salen)(AcO)Mn ^v (OH)(OOR)]
	CH ₃ (ethyl benzene)
base (ethylenediamine plus 2 salicylaldehydes)	
	{(bpy) ₂ *(·OO)Fe ^{III} OOH(Bu)[H ₃ O*]} {(bpy) ₂ *(·OO)Co ^{III} OOH(Bu)[pyH*]} [(PA ₂ Fe ^{IV} (OH)(OOR)] [Cl ₂ Fe ^{IV} (OH) (OOR)] [(bpy) ₂ *Cu ^{III} (OH)(OOR)] [(bpy) ₂ (AcO) ₃ Mn ^V (OH)(OOR)] ₂ (cyclohexane), c-C ₆ H ₁₀ (cyclohexene), PhCH ₂ I nic acid (2-carboxyl pyridine)

intermediates for Fenton chemistry (1) and oxygenated Fenton chemistry {5 [1(O₂)] and 6} within the present study. Each of these reactive intermediates (1, 5, 6) includes a stabilized hydroxyl group (HO) that (a) is less reactive than free HO and (b) gives products from organic substrates via the internal formation of an intermediate with an iron-carbon bond [or Fe(pyl)-carbon bond, or an FeOO-carbon bond] that are different from those for free HO. Thus, the metal-induced activation of hydroperoxides via nucleophilic addition (Fenton chemistry) is a highly disciplined "distant cousin" to the radical chemistry of free HO. Hence, the specific reactivities of Fenton reactive intermediates [1, 5, 6] are affected by the metal (Fe, Cu, Mn or Co), the ligand and the solution matrix. The more limited reactivity of Fenton systems is more than compensated for by their selectivity and unique ability to produce pure products.

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