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U.S. DEPARTMENT OF COMMERCE / National Bureau of Standards



Selected Specific Rates of Reactions of Transients from Water in Aqueous Solution.

III. Hydroxyl Radical and Perhydroxyl Radical and Their Radical Ions

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Selected Specific Rates of Reactions of Transients from Water in Aqueous Solution.

III. Hydroxyl Radical and Perhydroxyl Radical and Their Radical Ions

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Foreword

The National Standard Reference Data System provides access to the quantitative data of physical science, critically evaluated and compiled for convenience and readily accessible through a variety of distribution channels. The System was established in 1963 by action of the President's Office of Science and Technology and the Federal Council for Science and Technology, and responsibility to administer it was assigned to the National Bureau of Standards.

NSRDS receives advice and planning assistance from a Review Committee of the National Research Council of the National Academy of Sciences-National Academy of Engineering. A number of Advisory Panels, each concerned with a single technical area, meet regularly to examine major portions of the program, assign relative priorities, and identify specific key problems in need of further attention. For selected specific topics, the Advisory Panels sponsor subpanels which make detailed studies of users' needs, the present state of knowledge, and existing data resources as a basis for recommending one or more data compilation activities. This assembly of advisory services contributes greatly to the guidance of NSRDS activities.

The System now includes a complex of data centers and other activities in academic institutions and other laboratories. Components of the NSRDS produce compilations of critically evaluated data, reviews of the state of quantitative knowledge in specialized areas, and computations of useful functions derived from standard reference data. The centers and projects also establish criteria for evaluation and compilation of data and recommend improvements in experimental techniques. They are normally associated with research in the relevant field.

The technical scope of NSRDS is indicated by the categories of projects active or being planned: nuclear properties, atomic and molecular properties, solid state properties, thermodynamic and transport properties, chemical kinetics, and colloid and surface properties.

Reliable data on the properties of matter and materials are a major foundation of scientific and technical progress. Such important activities as basic scientific research, industrial quality control, development of new materials for building and other technologies, measuring and correcting environmental pollution depend on quality reference data. In NSRDS, the Bureau's responsibility to support American science, industry, and commerce is vitally fulfilled.

A handwritten signature in dark ink, reading "E. Ambler." The signature is fluid and cursive, with a large, sweeping initial "E" and a trailing flourish.

ERNEST AMBLER, *Acting Director*

Preface

This report is one of a series of data publications on radiation chemistry; the aim of the series is to compile, evaluate, and present the numerical results on processes occurring in systems which have been subjected to ionizing radiation. Various kinds of data are important in radiation chemistry. The quantities which were measured first were the observed radiation yields or G values (molecules formed or destroyed per 100 eV). Various indirect methods based on G values have been used to determine yields of transient species and relative rates of reactions. The spectral properties (optical, electron spin resonance) of transients have provided a direct method for their identification, and rates of the very fast reactions of transients which occur in irradiated systems have been measured directly by spectroscopic methods. Conductivity and luminescence methods have also provided a means of measuring properties of transients and their kinetics. Some reactions which occur in irradiated systems have also been studied by other methods, such as photochemistry, electric discharge, ultrasonics, chemical initiation, electron impact, etc. The emphasis in these publications is on the data of radiation chemistry, but where other pertinent data exist, they are included.

The data of radiation chemistry are voluminous; thousands of systems have been investigated. As a result there are certain collections, *e.g.* rate constants of particular types of reactions or certain properties of transients, for which tabulations of the data are considered essential, but for which critical assessment of each value is impossible. On the other hand, certain systems and properties have been studied so extensively that critical examination of these data is desirable and timely. Authors of this series of data publications have been asked to evaluate the extent to which the data can be critically assessed, to describe their criteria for evaluation, and to designate preferred values whenever possible.

Contents

	Page
Foreword.....	III
Preface.....	IV
Introduction.....	1
Table 1. Values of k used for normalizing relative rates.....	4
Table 2. Reactions of OH with transients from water (3.1–3.7).....	5
Table 3. Reactions of OH with inorganic solutes (3.8–3.122).....	7
Table 4. Reactions of OH with organic solutes (3.123–3.758).....	22
Table 5. Reactions of O^- with water, transients from water, inorganic solutes, and organic solutes (4.1–4.121).....	68
Table 6. Reactions of $HO_2(O_2^-)$ with transients from water, inorganic solutes and organic solutes (5.1–5.65).....	78
Formula index.....	87
References.....	100

Selected Specific Rates of Reactions of Transients from Water in Aqueous Solution. III. Hydroxyl Radical and Perhydroxyl Radical and their Radical Ions

Farhataziz and Alberta B. Ross

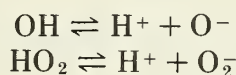
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Rates of reactions of OH and HO₂ with organic and inorganic molecules, ions and transients in aqueous solution have been tabulated, as well as the rates for the corresponding radical ions in aqueous solution (O⁻ and O₂⁻). Most of the rates have been obtained by radiation chemistry methods, both pulsed and steady-state; data from photochemistry and thermal methods are also included. Rates for over one thousand reactions are listed.

Key words: Aqueous solution; chemical kinetics; data compilation; hydroxyl radical; oxide radical ion; perhydroxyl radical; radiation chemistry; rates; superoxide ion.

1. Introduction

The short-lived products of water radiolysis for low LET radiation (cobalt-60 gamma rays, X-rays, and electrons with energies of about 30 keV and above) are e_{aq}^- , H and OH. In the presence of oxygen, hydrated electrons and hydrogen atoms are converted to other short-lived species, O₂⁻ and HO₂. The pK_a's of OH and HO₂ are 11.9 (65-0386, 66-0424) and 4.88 (70-0304), respectively; *i.e.*, O⁻ and O₂⁻ can be produced from the equilibria:



Thus, by adjusting only pH and the concentration of O₂ in water, one can produce e_{aq}^- , H, OH, HO₂, O₂⁻ and O⁻. All of these species have been characterized and their reactions with hundreds of inorganic and organic compounds have been studied. In the previous compilations, NSRDS-NBS 43 and Supplement (73-0030, 75-0002) and NSRDS-NBS 51 (75-0001), the specific rate constants for e_{aq}^- and H have been collected. The present compilation is an extension of the series and covers the specific rates for the reactions of OH, HO₂, O⁻, and O₂⁻. The literature is covered up to the latter part of 1975.

Methods

The majority of the data in this compilation are from investigations in radiation chemistry. However, data from photochemistry, Fenton's reaction, and other methods are also included. The rate constants of short-lived species are measured by *steady-state* and *pulse techniques*. The *steady-state* investigations yield ratios of rate constants which are deduced from an assumed mechanism. Values of specific rates

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from steady-state investigations may be as accurate as those measured by other methods. However, in many cases, the mechanisms are complex and the values measured with steady-state techniques must be accepted with caution. By *pulse techniques*, it is possible to study a reaction directly if one can either measure disappearance of a reactant (*decay kinetics*, d.k.) or formation of a product (*product buildup kinetics*, p.b.k.) during the course of a reaction by methods such as optical spectroscopy, electrical conductivity, etc. Unfortunately, due to experimental difficulties, such investigations are possible for only a few of the reactions compiled in these tables. The specific rates for other reactions have been measured by *competition kinetics* (c.k.) using pulse techniques.

In the present compilation an attempt was made to be as comprehensive as possible and include measurements of the rates of the same reaction by various methods, as well as measurements by the same method by different authors; values superseded by later measurements by the same authors have been omitted. Certain relative values are omitted if the standard reactions were not well characterized or if the relative values were in wide disagreement with values measured by several other methods. Examples of such data are those which have been obtained using the decolorization of organic dyes by OH or the oxidation of hydroxylaminedisulfonate ion by OH as the standard reactions.

Extended sections on radiation chemical, photochemical and chemical methods of generating OH and O⁻, and on the methods for rate determination, and the mechanisms for the reactions are found in NSRDS-NBS 46 (73-0299). The reader is also referred to that review for an analysis of complications which may occur in various systems.

Arrangement of Tables

In order to provide for internal consistency in the tables, values for rates of a number of competing reactions have been chosen to convert the rate ratios into relative rates. Those values are listed in table 1 and have been taken, whenever possible, from the review of Dorfman and Adams, NSRDS-NBS 46 (73-0299), where a critical analysis of the hydroxyl radical rate data has been made and "most reliable values" selected for a number of reactions; some of the values have been taken from an earlier paper on standardization of OH rate data by Willson, Greenstock, Adams, Wageman and Dorfman (71-0578). Many of the values chosen are the most recent directly determined rates; the references to those determinations are cited in the last column of table 1.

Reactions of OH with other transient species from water are listed in table 2. The reactions of inorganic solutes with OH are listed in table 3 in alphabetical order by main element. The reactions of organic solutes with OH are listed in table 4 in alphabetical order by name. In most cases IUPAC nomenclature has been used. The reactions of O⁻ and HO₂(O₂⁻) are listed by the same arrangement in tables 5 and 6, respectively.

The format of the tables is similar to previous parts of this series. Reactions are included in column 2 when products or mechanism have been studied. When several reactions are given, the reactions are labelled (I), (II), etc. and the rates in columns 4 and 5 are labelled (I), (II), k_I , k_{II} , etc.; if the rate in column 4 or 5 is not so labelled the value represents the sum of the rates for all contributing reactions. Reactions are second order and the rates have the units $\text{dm}^3\text{mol}^{-1}\text{s}^{-1}$ unless otherwise specified in the tables. Values for radical combination and disproportionation rates (entries 3.3, 4.5, 5.4 and 5.35) are for k (and not $2k$ as usually determined) unless it was not clear which was reported, in which case the value is the one given by the authors. Values of k which have been directly measured are given with the error limits as reported by the authors. The ratios of rate constants in the *Ratio* column are given in the form k/k_X where k_X symbolizes the rate of the competing reaction of the same short-lived species with X. In the *Comments* column ratios may be given as k/k_X or k_X/k_Y where k , k_X , and k_Y are rates of the same short-lived species with the reactant in column 2, X, and Y, respectively. For some of the entries only a ratio is given, but in most cases relative rates have been calculated from the ratios and are listed under the k column. The values of k obtained from ratios are designated as relative (rel.) and have been calculated by using the rates in table 1 (or, in a few cases, using a rate given under *Comments*). At the end of each entry for solutes which have been used in competition studies a list of entry numbers is given in which ratios involving that solute are reported.

Columns are included identifying the source of the radical and the method of measurement; other

information is given under *Comments*, such as activation energy, frequency factor, equilibrium constant, deuterium isotope effect (k_H/k_D). Temperature is 20-25°C, or assumed to be at room temperature, unless otherwise specified.

Abbreviations which have been used are tabulated at the end of this section. References are designated by number as assigned by the Radiation Chemistry Data Center; the first two digits of the number specify the year. The references are given at the end of the tables. A formula index for the solutes refers to entry numbers in the various tables and includes references to the tables of e_{aq}^- and H reactions already published.

Abbreviations and Symbols

<i>A</i>	frequency factor	ident.	identification
abs.	absorption	<i>k</i>	specific rate (in $\text{dm}^3\text{mol}^{-1}\text{s}^{-1}$ unless otherwise specified)
abstr.	abstraction	<i>K</i>	equilibrium constant
acac	acetylacetone	lum.	luminescence
ala	alanine	<i>M</i>	mol/dm^3
anal.	analysis	Me	methyl
approx.	approximate	MeOH	methanol
β -r.	beta radiolysis	μ	ionic strength
bicarb	bicarbonate ion	math.	mathematical
biol.	biological	meas.	measured
bisulf	bisulfate ion	mol.wt.	molecular weight
BzO ⁻	benzoate ion	nat	natural pH
calcd.	calculated	NB	nitrobenzene
carb	carbonate ion	obs.	observed
chem.	chemical analysis	opt.	optical spectroscopy
c.k.	competition kinetics	oxy	oxygen
concn.	concentration	PA ⁻	phenylacetate ion
condy.	electrical conductivity	p.b.k.	product buildup kinetics
cor.	corrected	perox	hydrogen peroxide
cyst	cysteamine	PhH	benzene
d.k.	decay kinetics	phot.	photolysis
detd.	determined	p <i>K</i> _a	negative logarithm of the acid dissociation constant, <i>e.g.</i> , where $\text{AH} + \text{H}_2\text{O} \rightleftharpoons \text{A}^- + \text{H}_3\text{O}^+$
<i>E</i> _a	activation energy	PNBA ⁻	<i>p</i> -nitrobenzoate ion
ε	extinction coefficient (in $\text{cm}^2\text{mol}^{-1}$ or $M^{-1}\text{cm}^{-1}$)	pol.	polarography
EDTA	ethylenediaminetetraacetate	p.r.	pulse radiolysis
en	ethylenediamine	2-PrOH	2-propanol
<i>e</i> -r.	electron radiolysis	py	pyridine
esr	esr spectroscopy	r.	radiolysis
est.	estimated	rel.	relative
Et	ethyl	RNO	<i>p</i> -nitroso- <i>N,N</i> -dimethylaniline
EtOH	ethanol	soln.	solution
Fenton	Fenton's reaction ($\text{Fe}^{2+} + \text{H}_2\text{O}_2$)	therm.	thermal
ferro	ferrocyanide ion	thym	thymine
formn.	formation	TNM	tetranitromethane
f.phot.	flash photolysis	trac.	tracer techniques
<i>G</i>	radiation yield (per 100 eV)	unpubl.	unpublished
γ -r.	gamma radiolysis	visc.	viscosimetry
gly	glycine	X-r.	X-radiolysis
hydr	hydrogen		
3HX	3-hexenedioate ion		

TABLE 1. Values of k used for normalizing relative rates

Reactant ^a	Reaction	$k(\text{dm}^3\text{mol}^{-1}\text{s}^{-1})$	Comment	Ref. ^f
<i>OH Reactions</i>				
bicarb (3.20)	$\text{OH} + \text{HCO}_3^- \rightarrow \text{OH}^- + \text{HCO}_3$ or $\text{H}_2\text{O} + \text{CO}_3^-$	3.6×10^7	b	73-1031
carb (3.21)	$\text{OH} + \text{CO}_3^{2-} \rightarrow \text{OH}^- + \text{CO}_3^-$	3.65×10^8	b,c	70-0247
CNS ⁻ (3.25)	$\text{OH} + \text{CNS}^- \rightarrow \text{CNSOH}^-$	1.1×10^{10}	b,c,d	72-0122
Fe ²⁺ (3.52)	$\text{OH} + \text{Fe}^{2+} \rightarrow \text{OH}^- + \text{Fe}^{3+}$	2.3×10^8	b	72-0354
ferro (3.54)	$\text{OH} + \text{Fe}(\text{CN})_6^{4-} \rightarrow \text{OH}^- + \text{Fe}(\text{CN})_6^{3-}$	9.3×10^9	b,c	73-1039
I ⁻ (3.66)	$\text{OH} + \text{I}^- \rightarrow \text{OH}^- + \text{I}$	1.2×10^{10}	b	72-0122
PhH (3.186)	$\text{OH} + \text{C}_6\text{H}_6 \rightarrow \text{C}_6\text{H}_5\text{OH}$	7.8×10^9	b,c	68-0304
BzO ⁻ (3.191)	$\text{OH} + \text{C}_6\text{H}_5\text{COO}^- \rightarrow (\text{OH})\text{C}_6\text{H}_5\text{COO}^-$	5.7×10^9	b,c	71-0578
EtOH (3.358)	$\text{OH} + \text{CH}_3\text{CH}_2\text{OH} \rightarrow \text{H}_2\text{O} + \text{CH}_3\text{CHOH}$	1.85×10^9	c,d	—
HCOO ⁻ (3.384)	$\text{OH} + \text{HCOO}^- \rightarrow \text{H}_2\text{O} + \text{COO}^-$	3.5×10^9	d	—
MeOH (3.511)	$\text{OH} + \text{CH}_3\text{OH} \rightarrow \text{H}_2\text{O} + \text{CH}_2\text{OH}$	9×10^8	c,d	—
NB (3.565)	$\text{OH} + \text{C}_6\text{H}_5\text{NO}_2 \rightarrow (\text{OH})\text{C}_6\text{H}_5\text{NO}_2$	3.2×10^9	b,c	68-0304
PNBA ⁻ (3.567)	$\text{OH} + p\text{-NO}_2\text{C}_6\text{H}_4\text{COO}^- \rightarrow (\text{OH})p\text{-NO}_2\text{C}_6\text{H}_4\text{COO}^-$	2.6×10^9	b,c	68-0304
RNO (3.582)	$\text{OH} + p\text{-NOC}_6\text{H}_4\text{N}(\text{CH}_3)_2 \rightarrow \text{products}$	1.25×10^{10}	b,c	69-0156
PA ⁻ (3.611)	$\text{OH} + \text{C}_6\text{H}_5\text{CH}_2\text{COO}^- \rightarrow (\text{OH})\text{C}_6\text{H}_5\text{CH}_2\text{COO}^-$	7.9×10^9	b,c	68-0304
2-PrOH (3.637)	$\text{OH} + (\text{CH}_3)_2\text{CHOH} \rightarrow \text{H}_2\text{O} + (\text{CH}_3)_2\text{COH}$	2.2×10^9	d	—
thym (3.711)	$\text{OH} + \text{C}_5\text{H}_8\text{N}_2\text{O}_2 \rightarrow \text{C}_5\text{H}_8\text{N}_2\text{O}_2\cdot\text{OH}$ (6-addn.)	5.4×10^9	d	g
<i>O⁻ Reactions</i>				
oxy (4.29)	$\text{O}^- + \text{O}_2 \rightarrow \text{O}_3^-$	3.6×10^9	b,c	69-0379
EtOH (4.65)	$\text{O}^- + \text{CH}_3\text{CH}_2\text{OH} \rightarrow \text{OH}^- + \text{C}_2\text{H}_4\text{OH}$	1.1×10^9	b,c	70-0080
3HX (4.75)	$\text{O}^- + {}^-\text{O}_2\text{CCH}_2\text{CH}=\text{CHCH}_2\text{CO}_2^- \rightarrow$ $\text{OH}^- + {}^-\text{O}_2\text{CCH}_2\text{CHCHCHCO}_2^-$	6.5×10^8	b	75-1003
MeOH (4.80)	$\text{O}^- + \text{CH}_3\text{OH} \rightarrow \text{OH}^- + \text{CH}_2\text{OH}$	5.8×10^8	b,c	70-0080
2-PrOH (4.95)	$\text{O}^- + \text{CH}_3\text{CHOHCH}_3 \rightarrow \text{OH}^- + (\text{CH}_3)_2\text{COH}$	1.5×10^9	e	—

^aNumber in parentheses indicates the number of the reaction in the following tables.

^bMost recently reported directly determined rate.

^cCited by Dorfman and Adams in NSRDS-NBS 46 (73-0299) as "most reliable values - values of which the accuracy (within the stated experimental uncertainty or lacking such a statement, within $\pm 30\%$) seems least open to question"; more than one such value is cited in NSRDS-NBS 46 for some reactions.

^dMean value of measured rates with $k(\text{OH} + \text{EtOH}) = 1.85 \times 10^9 \text{ dm}^3\text{mol}^{-1}\text{s}^{-1}$ as a secondary reference standard: Willson, Greenstock, Adams, Wageman and Dorfman (71-0578).

^eMean value of relative rates normalized for values listed in this table for competing reactions.

^fReference for the most recently reported directly measured rate.

^gRecent directly determined rates are 5.1×10^9 at natural pH (71-0578) and 5.5×10^9 at pH 9 (72-0047).

TABLE 2. Reactions of OH with transients from water

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.1	e_{aq}^- $OH + e_{aq}^- \rightarrow OH^-$	basic 3×10^{10}	—	—	—	See 1.7, 1.8, NSRDS-NBS 43.	73-0030
3.2	H $OH + H \rightarrow H_2O$	acid $\sim 2 \times 10^{10}$	—	—	—	See 2.3, NSRDS-NBS 51.	75-0001
3.3	OH $OH + OH \rightarrow H_2O_2$	7 $(4 \pm 1) \times 10^9$ (rel.)	$k/(k_{perox})^2 = 1.9 \times 10^{-6}$ $\text{mol}\cdot\text{s}/\text{dm}^3$	p.r.	chem.	assumed $k_{perox} = 4.5 \times 10^7$; obs. intensity effect on H_2O_2 and O_2 concn.	62-0052
		0.4 6×10^9 (rel.)	$k/k_H = 0.5$	p.r.	chem.	obs. $G(H_2)$; data fitting based on mechanism; assumed $2k(H + H) = k(H + OH) = 1.2 \times 10^{10}$.	63-0043
		3 6×10^9 (rel.)	—	p.r.	chem.	obs. $G(H_2)$ and $G(O_2)$ in H_2O_2 soln.; data fitting based on mechanism; assumed $k(H + OH) = 3.2 \times 10^{10}$; $k(H + H) = 1.3 \times 10^{10}$.	64-0092
		3.7 $(5.2 \pm 0.7) \times 10^9$	—	p.r.	opt.	d.k.; $\epsilon(260 \text{ nm}) = 370 \text{ cm}^2 \text{ mol}^{-1}$.	65-0010
		~ 7 5.5×10^9 (rel.)	$k/k_{ferro} = 0.59$	p.r.	opt.	c.k.; obs. $Fe(CN)_6^{3-}$ at 420 nm; data fitting based on mechanism.	66-0424
		7 $(5.2 \pm 0.5) \times 10^9$	—	p.r.	opt.	d.k.; $\epsilon(200-250 \text{ nm}) = 450-530 \text{ cm}^2 \text{ mol}^{-1}$; cor. for H and OH^- .	69-0083
3.4	O^- $OH + O^- \rightarrow HO_2^-$	>12 For other ratios see: 3.5, 3.6, 3.7, 3.12, 3.26, 3.82. $\leq 2.6 \times 10^{10}$ (rel.)	—	p.r.	opt.	c.k. with $Fe(CN)_6^{4-}$; $pK_a(OH) = 11.9 \pm 0.2$; est. based on numerous assumptions.	66-0424
3.5	HO_2 (I) $OH + HO_2 \rightarrow H_2O_3$ (II) $OH + HO_2 \rightarrow H_2O + O_2$	7 $\sim 3 \times 10^9$ (rel.)	$k \cdot k(H + H_2O_2) / k_{perox} k(H + HO_2) = 74$	p.r.	chem.	assumed $k(H + H_2O_2) = k(H + HO_2)$ and $k_{perox} = 4.7 \times 10^7$.	62-0052
		>2 6×10^9 (rel.)	$k/k_{OH} = 1$	p.r.	chem.	obs. $G(H_2O_2)$; data fitting based on mechanism; assumed $k_{OH} = 6 \times 10^9$.	63-0043
		2-3 1.4×10^{10} (rel.)	$k_I/k_{II} \cong 2.3$ $k/k_{Fe^{2+}} = 60$	e-r. p.r.	opt. chem.	obs. $G(H_2O_3)$. obs. $G(Fe^{3+})$ at $\sim 10^{22} \text{ eV g}^{-1} \text{ s}^{-1}$.	63-0075 64-0049
		3 1.5×10^{10} (rel.)	—	p.r.	chem.	obs. $G(H_2)$ and $G(H_2O_2)$; data fitting based on mechanism; assumed $k_{OH} = 6 \times 10^9$; $k(H + H) = 1.3 \times 10^{10}$.	64-0092

TABLE 2. Reactions of OH with transients from water - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.5 cont.		0.46– 6.75	7.1×10^9 (rel.)	$k/k_{\text{OH}} = 1.18$	p.r.	chem.	obs. $G(\text{H}_2\text{O}_2)$; best fit; $\text{p}K_s(\text{HO}_2) = 4.45 \pm 0.10$; assumed $k_{\text{OH}} = 6 \times 10^9$.	68-0014
3.6	H_2O_2^+ $\text{OH} + \text{H}_2\text{O}_2^+ \rightarrow$ $\text{H}_3\text{O}^+ + \text{O}_2$	0.46– 1.51	1.27×10^{10} (rel.)	$k/k_{\text{OH}} = 2.12$	p.r.	chem.	data fitting; $\text{p}k(\text{H}_2\text{O}^+ \rightleftharpoons \text{H}^+ + \text{HO}_2) = 1.2 \pm 0.3$; assumed $k_{\text{OH}} = 6 \times 10^9$.	68-0014
3.7	O_2^- $\text{OH} + \text{O}_2^- \rightarrow$ $\text{OH}^- + \text{O}_2$	2.74– 6.75	1.01×10^{10} (rel.)	$k/k_{\text{OH}} = 1.68$	p.r.	chem.	obs. $G(\text{H}_2\text{O}_2)$; data fitting; $\text{p}K(\text{HO}_2 \rightleftharpoons \text{H}^+ + \text{O}_2^-) = 4.45 \pm 0.10$; assumed $k_{\text{OH}} = 6 \times 10^9$.	68-0014
		7	$(8.0 \pm 1) \times 10^9$	—	e-r.	condy.	—	69-0547

TABLE 3. Reactions of OH with inorganic solutes

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.8	Ag^+ $\text{OH} + \text{Ag}^+ \rightarrow$ $\text{Ag}^{2+} + \text{OH}^-$	7 ~5	$(1.50 \pm 0.10) \times 10^{10}$ $(6.3 \pm 1.2) \times 10^9$ (rel.)	— $k/k_{\text{MeOH}} = 7 \pm 1$	p.r. p.r.	opt. condy.	p.b.k. c.k.; 2-fold increase in H^+ did not change rate; may be $\text{OH} + \text{Ag}^+ \rightarrow \text{AgOH}^+$	68-0436 70-0512
3.9	AsO_2^-	10.7 9	8.4×10^9 (rel.) 7.6×10^9 (rel.)	$k/k_{\text{carb}} = 23$ $k/k_{\text{RNO}} = 0.607$	p.r. γ -r.	opt. opt.	c.k. c.k.	65-0190 65-0356
3.10	$\text{Au}(\text{CN})_2^-$ $\text{OH} + \text{Au}(\text{CN})_2^- \rightarrow$ $\text{Au(II)} + \text{OH}^-$	7 2	$(4.7 \pm 0.8) \times 10^9$ 5×10^9 (rel.)	— $k/k_{\text{MeOH}} = 5.5$	p.r. p.r.	opt. opt.	p.b.k. at 330 nm. c.k.	68-0302 68-0302
3.11	BH_4^- $\text{BH}_4^- + \text{OH} \rightarrow$ $\text{BH}_4 + \text{OH}^-$	11- 12.83	1.2×10^{10}	—	p.r.	opt.	p.b.k. at 400 or 280 nm.	70-1046
3.12	Br^- (I) $\text{OH} + \text{Br}^- \rightleftharpoons$ BrOH^- $\text{BrOH}^- \rightleftharpoons \text{Br} + \text{OH}^-$ $\text{Br} + \text{Br}^- \rightleftharpoons \text{Br}_2^-$ $\text{BrOH}^- + \text{Br}^- \rightleftharpoons$ $\text{Br}_2^- + \text{OH}^-$	— 2.2 ~11 0.8	1×10^9 (rel.) — 5.8×10^8 (rel.) 1.6×10^{10} (rel.)	$k/k_{\text{EtOH}} = 0.6$ $k/k_{\text{hydr}} = 830$ $k/k_{\text{carb}} = 1.6$ $k/k_{\text{OH}} = 2.5$	γ -r. γ -r. p.r. p.r.	chem. chem. opt. calcd.	c.k. obs. $G(\text{H}_2\text{O}_2)$. c.k. obs. $G(\text{H}_2\text{O}_2)$; math. anal.; assume $k_{\text{OH}} =$ 6.4×10^9 ; method approx.	62-0053 63-0076 64-0131 64-0294
		7, 10.5	1.1×10^9 (rel.)	$k/k_{\text{BrO}^-} = 0.20$	γ -r.	trac.	c.k.; meas. $^{14}\text{CO}_2$.	65-0099
		—	5×10^8 (rel.)	$k/k_{\text{ferro}} = 0.054$	phot.	—	c.k.	65-0247
		9	1.1×10^9 (rel.)	$k/k_{\text{RNO}} = 0.089$	γ -r.	opt.	c.k.	65-0356
		7	1×10^9	—	p.r.	opt.	p.b.k. at 360 nm.	65-0382
		2	5×10^9	—	p.r.	opt.	p.b.k.; it is proposed that reaction may be $\text{OH} + (\text{Br}^- \text{---} \text{H}^+)_{\text{aq}}$ $\rightarrow \text{Br} + \text{H}_2\text{O}$.	65-0382
		7	3.9×10^9 (rel.)	$k/k_{\text{MeOH}} = 4.4 \pm$ 0.5	γ -r.	chem.	c.k. in NO-MeOH-KBr soln.	66-0118
		6	1.2×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.645$	γ -r.	—	c.k.; obs. $G(\text{H}_2\text{O}_2)$.	66-0423
		9	1.1×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.58$	γ -r.	opt.	c.k. with RNO.	66-0423
		5-9	$(1.2 \pm 0.15) \times 10^9$	—	p.r.	opt.	p.b.k. at 365 nm; k constant at this pH range but increases at low pH and decreases at higher pH.	66-0425
		~6	1×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.56 \pm$ 0.04	γ -r.	chem.	c.k.	66-0621, 67-0131
		2.7	6.5×10^9 (rel.)	$k/k_{\text{EtOH}} = 3.5 \pm$ 0.4	γ -r.	chem.	c.k.	66-0621, 67-0131
		1.3	1×10^{10} (rel.)	$k/k_{\text{EtOH}} = 5.6 \pm$ 0.4	γ -r.	chem.	c.k.	66-0621, 67-0131
		~6	1.1×10^9 (rel.)	$k/k_{\text{MeOH}} = 1.2 \pm$ 0.1	γ -r.	chem.	c.k.	66-0621, 67-0131
		2.7	8.1×10^9 (rel.)	$k/k_{\text{MeOH}} = 9.0 \pm$ 0.9	γ -r.	chem.	c.k.	66-0621, 67-0131
		1.3	1.1×10^{10} (rel.)	$k/k_{\text{MeOH}} = 12 \pm 1$	γ -r.	chem.	c.k.	66-0621, 67-0131
		2	3.1×10^{10} (rel.)	$k/k_{\text{RNO}} = 2.5$	Fenton	opt.	c.k.	67-0555
		5.5	1.2×10^9 (rel.)	$k/k_{\text{NB}} = 0.38$	r.	opt.	c.k.; obs. o -nitrophenol formn.	68-0494

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.12 cont.		5.5	2.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{pH} = 0.28	γ-r.	opt.	c.k. with Safranine T.	69-0279
		3.0	5.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{pH} = 0.64	γ-r.	opt.	c.k. with Safranine T.	69-0279
		2.0	1.0 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{pH} = 1.3	γ-r.	opt.	c.k. with Safranine T.	69-0279
		1-2, 6.98	4.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 1.94	γ-r.	chem.	c.k.; obs. <i>G</i> (acetone).	68-0602
		12-13	(8.9 ± 1.7) x 10 ⁸ (rel.)	—	f.phot.	opt.	c.k.; effect on decay of O ₃ ⁻ at 430 nm.	69-7340
		—	1.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 0.64	p.r.	opt.	c.k.; N ₂ O-satd.; ratio = 4 in O ₂ -satd. 1.0 <i>M</i> Br ⁻ soln.	71-0137
		9	1.75 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.14	γ-r.	opt.	c.k.; <i>E</i> _a = -6.2 ± 0.9 kcal/mol (-26 kJ/mol) (-8 to 23°C).	71-0469
		1-7	(1.06 ± 0.08) x 10 ¹⁰ (I)	—	p.r.	opt.	p.b.k. at 360 nm (Br ₂ ⁻).	72-0018
		9-11.5	—	—	p.r.	opt.	p.b.k. at 365 nm (Br ₂ ⁻); <i>K</i> ₁ = (2.86 ± 1.4) x 10 ³ dm ³ /mol.	72-0148
				For other ratios see: 3.32, 3.110, 3.394, 3.627.				
3.13	OD + Br ⁻ → OD ⁻ + Br	1.3	7.95 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 4.3 ± 0.4	γ-r.	chem.	c.k. in D ₂ O; obs. <i>G</i> (D ₂ O ₂).	68-0015
		6	6.8 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 0.37 ± 0.04	γ-r.	chem.	c.k. in D ₂ O.	68-0015
3.14	BrO ⁻ OH + BrO ⁻ → BrO + OH ⁻	11-13	4.5 x 10 ⁹ (rel.)	—	p.r.	opt.	c.k.; rel. to <i>k</i> (OH + CO ₃ ²⁻) = 4.2 x 10 ⁸ , more than two rate constants involved in analysis.	68-0153
		12-13	(1.4 ± 0.8) x 10 ⁹ (rel.)	—	f.phot.	opt.	c.k.; effect on decay of O ₃ ⁻ at 430 nm.	69-7340
3.15	BrO ₂ ⁻ OH + BrO ₂ ⁻ → BrO ₂ + OH ⁻	13	1.9 x 10 ⁹ (rel.)	—	p.r.	opt.	c.k.; relative to <i>k</i> (OH + CO ₃ ²⁻) = 4.2 x 10 ⁸ ; more than two rate constants involved in analysis; assume <i>k</i> (OH + BrO ₂ ⁻) = <i>k</i> (O ⁻ + BrO ₂ ⁻).	68-0153
		12-13	(1.4 ± 0.8) x 10 ⁹ (rel.)	—	f.phot.	opt.	c.k.; effect on decay of O ₃ ⁻ at 430 nm.	69-7340
3.16	BrO ₃ ⁻ OH + BrO ₃ ⁻ → BrO ₃ + OH ⁻	12-13	(3.9 ± 2.3) x 10 ⁶ (rel.)	—	f.phot.	opt.	c.k.; effect on decay of O ₃ ⁻ at 410 nm.	69-7340
3.17	BrO ₄ ⁻	—	< 10 ⁷	—	p.r.	opt.	d.k. (OH).	73-0106
3.18	CO OH + CO → COOH	0.4-0.7	(4.6 - 5.8) x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{Fe2+} = 2-2.5	Fenton	chem.	c.k.	57-0014
		~1	8.7 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{Fe2+} = 3.79	Fenton	chem.	c.k.	57-0014
		~1	8.3 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{Fe2+} = 3.6 ± 0.5	γ-r.	chem.	c.k.	63-0014
		7	—	<i>k</i> / <i>k</i> _{perox} = 13.0	phot.	chem.	c.k.	63-7005
		—	—	<i>k</i> / <i>k</i> _{perox} = 72	phot.	chem.	c.k.	69-7045

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.19	CO ₂	4	< 10 ⁶	—	p.r.	opt.	no abs. at 600 nm.	65-0384
3.20	OH + CO ₂ → HCO ₃	6.5	1 x 10 ⁷	—	p.r.	opt.	p.b.k. at 600 nm.	65-0384
	HCO ₃ ⁻	8.4	1.5 x 10 ⁷	—	p.r.	opt.	p.b.k. at 600 nm.	66-0139
	OH + HCO ₃ ⁻ →	nat.	(4.9 ± 0.6) x 10 ⁷	—	p.r.	opt.	p.b.k.; 3.3 x 10 ⁻³ M HCO ₃ ⁻ ;	69-0052
	H ₂ O + CO ₃ ⁻ or		(7.9 ± 1) x 10 ⁷				authors have no interpretation which value is correct.	
	HCO ₃ + OH ⁻							
		—	(4.9 ± 0.5) x 10 ⁷	—	p.r.	opt.	p.b.k.	69-0379
		—	(3.6 ± 0.3) x 10 ⁷	—	p.r.	opt.	p.b.k. at 578 nm; c.k. with 2-PrOH gave 3.8 x 10 ⁷ .	73-1031
3.21	CO ₃ ²⁻ OH + CO ₃ ²⁻ → OH ⁻ + CO ₃ ⁻		For other ratios see: 3.57, 3.64, 3.384.					
		11	3.8 x 10 ⁸ (rel.)	$k/k_1^- = 0.029 \pm 0.003$	p.r.	opt.	c.k.	65-0010
		11	3.5 x 10 ⁸	—	p.r.	opt.	p.b.k. at 580 nm.	65-0010
		10.5	~ 4.5 x 10 ⁸ (rel.)	$k/k_{\text{BrO}^-} = \sim 0.08$	γ-r.	trac.	c.k.; meas. ¹⁴ CO ₂ .	65-0099
		—	2 x 10 ⁸	—	p.r.	opt.	p.b.k.; O ₂ -satd. soln.; competing reactions may interfere.	66-0001
		<11.6	4.2 x 10 ⁸	—	p.r.	opt.	p.b.k.; k is pH dependent; calcn. is indirect.	66-0139
		10.6	(4.0 ± 0.2) x 10 ⁸	—	p.r.	opt.	p.b.k. at 600 nm.	69-0379
		11	4.7 x 10 ⁸ (rel.)	—	f.phot.	opt.	c.k.; soln. contains NO ₃ ⁻ and ethanol; rel. to $k(\text{OH} + \text{EtOH}) = 2 \times 10^9$ and $k(\text{O}^- + \text{C}_2\text{H}_5\text{OH}) = 1 \times 10^9$.	69-7218
	11	3.65 x 10 ⁸	—	p.r.	opt.	p.b.k.	70-0247	
		For other ratios see: 3.9, 3.12, 3.14, 3.15, 3.25, 3.29, 3.30, 3.63, 3.66, 3.82, 3.92, 3.93, 3.94, 3.95, 3.96, 3.97, 3.104, 3.105, 3.108, 3.109, 3.116, 3.143, 3.225, 3.351, 3.358, 3.369, 3.384, 3.403, 3.459, 3.511, 3.586, 3.615, 3.636, 3.696, 3.697, 3.698, 3.745.						
3.22	C ₂ N ₂ OH + (CN) ₂ → CNCNOH	—	≤ 10 ⁷	—	p.r.	opt.	kinetic anal. of abs. spectra of transients in N ₂ O soln. (OH and C ₂ N ₂).	71-0038
3.23	CN ⁻	9	4.5 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.36$	γ-r.	opt.	c.k.	65-0356
3.24	HCN	—	≤ 7 x 10 ⁷ (rel.)	$k/k_{\text{HCOO}^-} \leq 0.02$	γ-r.	chem.	c.k.	73-0364
3.25	CNS ⁻	—	5.8 x 10 ⁹ (rel.)	$k/k_{\text{carb}} = 16$	p.r.	opt.	c.k.	64-0131
	OH + CNS ⁻ →	7	6.6 x 10 ⁹	—	p.r.	opt.	p.b.k. at 500 nm.	65-0190
	CNSOH ⁻	2,7	1.2 x 10 ¹⁰ (rel.)	$k/k_{\text{carb}} = 33$	p.r.	opt.	c.k.	65-0190
	CNSOH ⁻ ⇌ CNS + 2-	2-2.2	9.7 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 1.80 \pm 0.25$	γ-r.	opt.	c.k.	67-0461
	OH ⁻							
	CNS ⁻ + CNS ⇌	5-5.5	1 x 10 ¹⁰ (rel.)	$k/k_{\text{thym}} = 1.95 \pm 0.30$	γ-r.	opt.	c.k.	67-0461
	(CNS) ₂							
		9	1.2 x 10 ¹⁰ (rel.)	$k/k_{\text{RNO}} = 0.95$	γ-r.	opt.	c.k.	67-0555
		2-12	(7.5 ± 0.5) x 10 ⁹	—	p.r.	opt.	p.b.k. at 500 nm.	68-0316

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.	
3.25 cont.	2-7	2.8×10^{10}	—	p.r.	opt.	p.b.k. (CNS) $_2^-$; earlier papers assumed CNS is absorbing species; for mechanism study see also 72-0126.	68-0375	
	5.5	1.2×10^{10} (rel.)	$k/k_{NB} = 3.6$	r.	opt.	c.k.; obs. <i>o</i> -nitrophenol formn.	68-0494	
	7	6.7×10^9 (rel.)	$k/k_{EtOH} = 3.6$	f.phot.	chem.	c.k.; soln. contains NO $_3^-$.	69-7218	
	—	1×10^{10} (rel.)	$k/k_{EtOH} = 5.5$ $k/k_{MeOH} = 11.6$	p.r.	opt.	c.k.; N $_2$ O-satd.; ratios 6.4 and 13.4 resp., in O $_2$ -satd. soln. contg. 0.2 <i>M</i> thiocyanate.	71-0137	
	—	$(1.08 \pm 0.10) \times 10^{10}$	—	p.r.	opt.	p.b.k. (CNS) $_2^-$ at 475 nm.	72-0122	
<i>For other ratios see: 3.33, 3.34, 3.37, 3.66, 3.71, 3.74, 3.75, 3.76, 3.77, 3.86, 3.87, 3.102, 3.103, 3.107, 3.110, 3.117, 3.121, 3.124, 3.125, 3.126, 3.129, 3.130, 3.131, 3.135, 3.138, 3.139, 3.140, 3.142, 3.143, 3.146, 3.147, 3.148, 3.151, 3.153, 3.155, 3.156, 3.157, 3.159, 3.163, 3.168, 3.169, 3.170, 3.171, 3.177, 3.178, 3.179, 3.181, 3.186, 3.191, 3.193, 3.197, 3.198, 3.202, 3.212, 3.219, 3.222, 3.223, 3.224, 3.225, 3.226, 3.227, 3.229, 3.230, 3.231, 3.232, 3.233, 3.234, 3.235, 3.237, 3.240, 3.241, 3.243, 3.245, 3.247, 3.256, 3.262, 3.263, 3.264, 3.266, 3.269, 3.270, 3.285, 3.288, 3.289, 3.290, 3.292, 3.293, 3.295, 3.296, 3.297, 3.298, 3.298, 3.300, 3.311, 3.312, 3.318, 3.323, 3.328, 3.329, 3.330, 3.331, 3.332, 3.336, 3.338, 3.339, 3.342, 3.357, 3.358, 3.361, 3.362, 3.363, 3.364, 3.366, 3.369, 3.372, 3.375, 3.381, 3.383, 3.385, 3.390, 3.396, 3.397, 3.398, 3.402, 3.403, 3.404, 3.405, 3.406, 3.407, 3.410, 3.411, 3.412, 3.413, 3.415, 3.416, 3.417, 3.418, 3.421, 3.433, 3.434, 3.442, 3.443, 3.444-6, 3.451, 3.453, 3.459, 3.473, 3.479, 3.480, 3.481, 3.483, 3.486, 3.487, 3.488, 3.489, 3.491, 3.493-3.495, 3.498, 3.503, 3.506, 3.508, 3.509a, 3.510, 3.511, 3.513, 3.520, 3.521, 3.522, 3.523, 3.524, 3.527, 3.528, 3.532, 3.535, 3.538, 3.543, 3.544, 3.545, 3.546, 3.547, 3.548, 3.554, 3.565, 3.573, 3.578, 3.580, 3.581, 3.592, 3.593, 3.594, 3.600, 3.602, 3.603, 3.604, 3.605, 3.606, 3.607, 3.613, 3.614, 3.616, 3.618, 3.621, 3.624, 3.628, 3.629, 3.630, 3.631, 3.634, 3.636, 3.637, 3.640, 3.645, 3.646, 3.649, 3.650, 3.656, 3.657, 3.659, 3.664, 3.665, 3.669, 3.670, 3.673, 3.674, 3.696, 3.703, 3.705, 3.706, 3.707, 3.709, 3.710, 3.711, 3.717, 3.718, 3.719, 3.723, 3.724, 3.726, 3.727, 3.730, 3.735, 3.737, 3.740, 3.743, 3.744, 3.746, 3.747, 3.748, 3.750, 3.751.</i>								
3.25a	Cd $^{2+}$	—	$< 5 \times 10^5$	p.r.	opt.	c.k. with Cu $^{2+}$.	75-1027	
3.25b	Cd $^+$	—	2×10^{10}	p.r.	opt., condy.	d.k. at 300 nm; Cd $^+$ from e_{aq}^- + Cd $^{2+}$.	75-1064	
3.26	OH + Cd $^+$ → OH $^-$ + Cd $^{2+}$ Ce $^{3+}$ OH + Ce $^{3+}$ → Ce $^{4+}$ + OH $^-$	0.4- 2 0.8 0 2.6- 2.95	— 3.2×10^8 (rel.) — 2.9×10^8 (rel.)	$k/k_{bisulf} = 900 \pm 300$ $k/k_{OH} = 4 \times 10^{-2}$ $k/k_{HCOOH} = 1.9 \pm 0.2$ $k/k_{EtOH} = 0.154$	p.r. p.r. γ -r. p.r.	— calcd. chem. opt.	c.k. math. anal.; assume $k_{OH} = 8.1 \times 10^8$; method approx. c.k.; 4 <i>M</i> H $_2$ SO $_4$. c.k.	60-0099 64-0294 69-0634 71-0137

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.26 cont.	0.8 —	—	$k/k_{\text{bisulf}} [\text{HSO}_4^-]$ $= 930 \pm 110 M^{-1}$ $= 30 \pm 3 M^{-1}$	γ -r.	chem.	c.k.; computer fitting based on mechanism; supercedes 57-0003.	72-0094
	4 M H_2SO_4	—	—				
	<i>For other ratios see: 3.118.</i>						
3.27	Cl^- (I) $\text{OH} + \text{Cl}^- \rightleftharpoons \text{ClOH}^-$ (II) $\text{ClOH}^- + \text{H}^+ \rightleftharpoons \text{Cl} + \text{H}_2\text{O}$ (III) $\text{Cl} + \text{Cl}^- \rightleftharpoons \text{Cl}_2^-$	1-2.5 8.9 x 10 ⁷ to 6.4 x 10 ⁸ (rel.)	$k/k_{\text{MeOH}} = 0.099$ to 0.715	p.r.	opt.	c.k.; <i>k</i> decreases with pH and is μ dependent; meas. abs. of Cl_2^- at 365 nm.	64-0149
		1-2.7 6.7 x 10 ⁷ to 1.6 x 10 ⁹ (rel.)	$k/k_{\text{ferro}} = 0.00725$ to 0.169	p.r.	opt.	c.k.; <i>k</i> decreases with pH and is μ dependent.	64-0149
		1-3 (1.16 to 2.16) x 10 ¹⁰ dm ⁶ mol ⁻² s ⁻¹	—	p.r.	opt.	p.b.k.; <i>k</i> includes [H ⁺]; not cor. for μ .	64-0149
		0-3 (0.32 to 1.84) x 10 ¹⁰ dm ⁶ mol ⁻² s ⁻¹	—	p.r.	opt.	p.b.k.; <i>k</i> includes [H ⁺]; $\mu = 0.012-1$.	64-0149
		~1-3 1.0 x 10 ¹⁰ dm ⁶ mol ⁻² s ⁻¹ (rel.)	$k/k_{\text{thym}} = 1.9 \pm 0.3$ dm ³ mol ⁻¹	γ -r.	chem.	c.k.; <i>k</i> defined for $\text{OH} + \text{H}^+ + \text{Cl}^- \rightarrow \text{Cl} + \text{H}_2\text{O}$.	65-0133
		9 < 1.25 x 10 ⁶ (rel.)	$k/k_{\text{RNO}} < 10^{-4}$	γ -r.	opt.	c.k.	65-0356
		2 5.2 x 10 ⁸ (rel.)	$k/k_{\text{RNO}} = 0.042$	Fenton	opt.	c.k.	67-0555
		~0.1 4.8 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.381$	Fenton	opt.	c.k.	67-0555
		0.8-3.4 (1.5 ± 0.3) x 10 ¹⁰ dm ⁶ mol ⁻² s ⁻¹	—	p.r.	opt.	p.b.k.; <i>k</i> refers to $\text{OH} + \text{Cl}^- + \text{H}_3\text{O}^+ \rightarrow \text{Cl} + 2\text{H}_2\text{O}$.	68-0313
		1.1 3.5 x 10 ⁷ (rel.)	$k/k_{\text{MeOH}} = 0.039$	γ -r.	chem.	c.k.; based on $k_{2-\text{PrOH}}/k_{\text{MeOH}} = 3.0$.	69-0647
		1.1 3.7 x 10 ⁷ (rel.)	$k/k_{\text{EtOH}} = 0.020$	γ -r.	chem.	c.k.; based on $k_{2-\text{PrOH}}/k_{\text{EtOH}} = 1.61$.	69-0647
		1 7 x 10 ⁸ (rel.)	$k/k_{\text{thym}} = 0.13$	Fenton	esr	c.k.; $k/k_{\text{perox}} = 9.5$; pH dependent; also with Ti(III)-H ₂ O ₂ .	69-5278
		1.3 2 x 10 ⁸ (rel.)	$k/k_{\text{MeOH}} = 0.24 \pm 0.04$	γ -r.	chem.	c.k.	71-0931
		6 < 10 ⁶	$k/k_{\text{MeOH}} < 0.001$	γ -r.	chem.	c.k.	71-0931
		~2 (4.3 ± 0.4) x 10 ⁹ (I) (2.1 ± 0.7) x 10 ¹⁰ (II) 2.1 x 10 ¹⁰ (III)	—	p.r.	opt.	d.k. at 240 nm as well as p.b.k. at 340 nm (Cl_2^-); $K_{\text{I}} = 0.70 \pm 0.13 M^{-1}$; $K_{\text{II}} = 1.6 \times 10^7$; $K_{\text{III}} = 1.9 \times 10^5 M^{-1}$.	73-1039
		~2 1.9 x 10 ¹⁰ dm ⁶ mol ⁻² s ⁻¹	—	p.r.	opt.	p.b.k. at 350 nm (Cl_2^-); <i>k</i> refers to $\text{OH} + \text{Cl}^- + \text{H}_3\text{O}^+$.	73-1089
	<i>For other ratios see: 3.291, 3.594.</i>						
3.28	$\text{OD} + \text{Cl}^- + \text{D}_3\text{O}^+ \rightarrow 2\text{D}_2\text{O} + \text{Cl}$	~2 1.6 x 10 ¹⁰ dm ⁶ mol ⁻² s ⁻¹	—	p.r.	opt.	p.b.k. (see above) $k_{\text{H}}/k_{\text{D}} = 1.17$.	73-1089
3.29	ClO^- $\text{OH} + \text{ClO}^- \rightarrow \text{ClO} + \text{OH}^-$	— 11 8.2 x 10 ⁹ (rel.)	— $k/k_{\text{carb}} = 22.5$	f.phot. p.r.	opt. opt.	estd. c.k.	71-7236 72-0301
3.30	ClO_2^- $\text{OH} + \text{ClO}_2^- \rightarrow \text{ClO}_2 + \text{OH}^-$	— (1.3 ± 0.4) x 10 ⁹	—	f.phot.	opt.	d.k. at 360 nm; assume $k(\text{OH} + \text{OH}) = 5 \times 10^9$; best fit.	71-7236
		11 5.7 x 10 ⁹ (rel.)	$k/k_{\text{carb}} = 15.7$	p.r.	opt.	c.k.	72-0301

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.31	ClO_3^-	11	$< 10^6$ (rel.)	—	p.r.	opt.	no effect on CO_3^{2-} formn. in carbon-ate soln.	72-0301
3.32	ClO_2 $\text{OH} + \text{ClO}_2 \rightarrow \text{HClO}_3$ or $\rightarrow \text{H}^+ + \text{ClO}_3^-$	—	$\leq 4 \times 10^5$	—	f.phot.	opt.	estd.	71-7236
		5.8-6.0	—	$k/k_{\text{Br}^-} \cong 1$	γ -r.	chem.	c.k. in 2-6 M ClO_4^- ; based on an assumed mechanism.	67-0019
		—	—	$k/k_{\text{perox}} > 200$ $k/k_{\text{hydr}} > 100$	γ -r.	chem.	c.k. assumed values.	67-0022
3.32a	Co^{2+}	—	$\sim 2 \times 10^6$	—	p.r.	opt.	c.k. with Cu^{2+} .	75-1027
3.33	$\text{Co}(\text{NH}_3)_6^{3+}$	—	$\leq 1.1 \times 10^8$ (rel.)	$k/k_{\text{CNS}^-} \leq 10^{-2}$	p.r.	opt.	c.k.	71-0282
3.34	$\text{Co}(\text{BzO}^-)(\text{NH}_3)_5^{2+}$	—	$(3.3 - 3.8) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = 0.3 - 0.35$	p.r.	opt.	c.k.; O_2 -satd.	71-0282
3.34a	$\text{Co}(\text{NH}_3)_5\text{py}^{3+}$	—	$(6.3-6.7) \times 10^9$	—	p.r.	opt.	p.b.k. at 345 nm.	71-0282
		5.2	6.5×10^8	—	p.r.	opt.	p.b.k. at 320 nm.	75-1088
3.35	$\text{Co}(\text{CN})_5\text{NO}^{3-}$	—	1.2×10^8 (rel.)	$k/k_{\text{RNO}} = 0.0094$	γ -r.	opt.	c.k.; depends on $k_{\text{CN}^-}/k_{\text{RNO}} = 0.42$.	71-0407
3.36	$\text{Co}(\text{acac})_3^{3+}$	1-7	4.8×10^9 (rel.)	$k/k_{\text{EtOH}} = 2.6$	r.	chem.	c.k.	70-0094
3.37	$\text{Cr}(\text{II})$ $\text{OH} + \text{Cr}(\text{II}) \rightarrow \text{OH}^- + \text{Cr}(\text{III})$	1	4.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.44$	p.r.	opt.	c.k.	72-0240
3.38	Cr^{3+} $\text{OH} + \text{Cr}^{3+} \rightarrow \text{Cr}^{4+} + \text{OH}^-$	0.4-	—	$k/k_{\text{hydr}} = 0.0082$	γ -r.	chem.	c.k.; assume $k_{\text{bisulf}}/k_{\text{hydr}} = 0.0039$.	63-0197
		1.4	—	—	—	—	—	—
		0.4-1	—	$k/k_{\text{hydr}} = 7 \pm 2$	γ -r.	chem.	c.k.; $k_{\text{bisulf}}/k_{\text{hydr}} = 0.011$.	65-0052
3.39	$\text{Cr}(\text{CN})_5\text{NO}^{3-}$	(Unexplained discrepancy in the above data.)						
		—	7.9×10^9 (rel.)	$k/k_{\text{RNO}} = 0.63$	γ -r.	opt.	c.k.; assumed $k(\text{OH} + \text{CN}^-) = 3.0 \times 10^9$.	69-0531
3.40	$\text{Cr}(\text{V})$ $\text{OH} + \text{Cr}(\text{V}) \rightarrow \text{OH}^- + \text{Cr}(\text{VI})$	—	7×10^9 (rel.)	$k/k_{\text{RNO}} = 0.56$	γ -r.	opt.	c.k.; depends on $k_{\text{CN}^-}/k_{\text{RNO}} = 0.42$.	71-0407
		—	5×10^{10}	—	γ -r.	est.	reoxidation of transient from e_{aq}^- or H reaction with chromate.	73-0186
3.41	Cu^{2+} $\text{OH} + \text{Cu}^{2+} \rightarrow \text{Cu}^{3+} + \text{OH}^-$ or $\rightarrow \text{Cu}(\text{OH})_2^{2+} + \text{Cu}(\text{OH})_2^+$	7	3.5×10^8	—	p.r.	opt.	p.b.k. at 313 nm.	65-0044,
		—	3.6×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.196$	p.r.	opt.	c.k.; meas. Cu^{3+} at 313 nm.	65-0394
		—	3.5×10^8 (rel.)	$k/k_{\text{MeOH}} = 0.385$	p.r.	opt.	c.k.; meas. Cu^{3+} at 313 nm.	65-0394
		~ 5	3×10^8 (rel.)	$k/k_{\text{t-BuOH}} = 0.67 \pm 0.07$	p.r.	condy.	c.k.; assume $k_{\text{MeOH}}/k_{\text{t-BuOH}} = 2$.	70-0512
		3-6	$(3.1 \pm 0.3) \times 10^8$	—	p.r.	opt.	p.b.k. at 300 nm.	71-0174
3.42	$\text{Cu}(\text{en})_2^{2+}$ $\text{OH} + \text{Cu}(\text{en})_2^{2+} \rightarrow \text{OH}^- + \text{Cu}(\text{en})_2^{3+}$	5.7	$(3.1 \pm 0.6) \times 10^8$	—	p.r.	opt.	p.b.k. at 300 nm.	71-0775
		6.5	$(3.0 \pm 0.6) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775
		10.2	$(5.0 \pm 1.0) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775
3.43	$\text{Cu}(\text{gly})_2^{2+}$ $\text{OH} + \text{Cu}(\text{gly})_2^{2+} \rightarrow \text{OH}^- + \text{Cu}(\text{gly})_2^{3+}$	6.1	$(1.5 \pm 0.3) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775
3.44	$\text{Cu}(\text{ala})_2^{2+}$ (see 3.43)	6.3	$(1.4 \pm 0.3) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775
3.45	$\text{Cu}(\beta\text{-ala})_2^{2+}$ (see 3.43)	5.8	$(1.2 \pm 0.2) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775
3.46	$\text{Cu}(\text{CH}_3\text{CH}_2\text{CH}(\text{NH}_3^+)\text{COO}^-)_2^{2+}$ (see 3.43)	6.1	$(2.0 \pm 0.4) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.47	$\text{Cu}(\text{CH}_3\text{CH}(\text{NH}_3^+)\text{CH}_2\text{COO}^-)_2^{2+}$ (see 3.43) 6.0	$(1.2 \pm 0.2) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775
3.48	$\text{Cu}(\text{NH}_3^+\text{CH}_2\text{CH}_2\text{COO}^-)_2^{2+}$ (see 3.43) 4.8	$(1.1 \pm 0.2) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775
3.49	$\text{Cu}((\text{CH}_3)_2\text{C}(\text{NH}_3^+)\text{COO}^-)_2^{2+}$ (see 3.43) 6.2	$(1.8 \pm 0.4) \times 10^9$	—	p.r.	opt.	p.b.k.	71-0775
3.50	$\text{Cu}(\text{EDTA})^{2-}$ ~7	4×10^9 (rel.)	$k/k_{\text{MeOH}} = 4.4$	X-r.	chem.	c.k.	73-0078
3.51	Eu^{2+} —	9×10^8	—	p.r.	opt.	d.k. (Eu^{2+}).	71-0311
	$\text{OH} + \text{Eu}^{2+} \rightarrow \text{OH}^- + \text{Eu}^{3+}$ 2	$(1.3 \pm 0.2) \times 10^9$	—	p.r.	opt.	d.k.; transient $\text{Eu}(\text{II})$ formed in $\text{Eu}(\text{III})$ soln.	73-1084
3.52	Fe^{2+} 1.2—	—	$k/k_{\text{perox}} = (2.99 \pm 0.2) \times 10^{-2}$	Fenton	chem.	c.k.; $E_a(\text{OH} + \text{H}_2\text{O}_2) - E_a(\text{OH} + \text{Fe}^{2+}) = 3.5$ kcal/mol(14.6 kJ/mol).	51-9004
	$\text{OH} + \text{Fe}^{2+} \rightarrow \text{Fe}^{3+} + \text{OH}^-$ 1.9	—	—	—	—	obs. $G(\text{H}_2)$ and $G(\text{Fe}^{3+})$; math. anal.	60-0099
	0.4 —	—	$k/k_{\text{H}} = 6.2 \times 10^{-3}$	p.r.	c.k.	p.b.k. at 305 nm; (Fe^{3+}).	64-0090
	0.3 $> 10^8$	—	—	p.r.	opt.	c.k.	64-0242
	0.4 1.7×10^9 (rel.)	—	$k/k_{\text{EtOH}} \cong 0.9$	p.r.	opt.	c.k.	66-0645,
	0.8 1.2×10^9 (rel.)	—	$k/k_{\text{PhH}} = 0.15$	γ -r., e -r.	chem.	c.k.	67-0504
	3.5 $(5 \pm 1) \times 10^8$	—	—	p.r.	opt.	p.b.k.; reported reaction is $\text{OH} + \text{Fe}_{\text{aq}}^{2+} \rightarrow \text{Fe}(\text{OH})^{2+}$.	66-0716
	2 5×10^8 (rel.)	—	$k/k_{\text{RNO}} = 0.04$	Fenton	opt.	c.k.	67-0555
	4.5— 3.4×10^8 (rel.)	—	$k/k_{\text{EtOH}} = 0.183$	p.r.	opt.	c.k.	71-0137
	6.2	—	—	—	—	—	—
	1 $(2.3 \pm 0.2) \times 10^8$	—	—	p.r.	opt.	p.b.k. at 240 nm; no temp. dependence 17-67°C.	72-0354
	For other ratios see: 3.5, 3.18, 3.56, 3.58, 3.59, 3.60, 3.77, 3.114, 3.123, 3.131, 3.149, 3.150, 3.185, 3.186, 3.190, 3.192, 3.221, 3.224, 3.239, 3.245, 3.251, 3.307, 3.320, 3.326, 3.358, 3.360, 3.365, 3.369, 3.371, 3.382, 3.404, 3.409, 3.451, 3.486, 3.491, 3.511, 3.522, 3.531, 3.546, 3.565, 3.612, 3.620, 3.636, 3.637, 3.638, 3.639, 3.642, 3.656, 3.680, 3.693, 3.694, 3.704, 3.724.						
3.53	$\text{OD} + \text{Fe}^{2+} \rightarrow \text{OD}^- + \text{Fe}^{3+}$ 1	$(9.4 \pm 0.8) \times 10^7$	—	p.r.	opt.	p.b.k. at 240 nm; in D_2O .	72-0354
3.54	$\text{Fe}(\text{CN})_6^{4-}$ 7	$(1.1 \pm 0.2) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 420 nm.	64-0213
	$\text{OH} + \text{Fe}(\text{CN})_6^{4-} \rightarrow$ 7	8.7×10^9 (rel.)	$k/k_{\text{EtOH}} = 4.7$	p.r.	opt.	c.k.	65-0007
	$\text{OH}^- + \text{Fe}(\text{CN})_6^{3-}$ 7	2.0×10^{10} (rel.)	$k/k_{\text{I}^-} = 1.67 \pm 0.018$	p.r.	opt.	c.k.; meas. abs. of I_2^- at 400 nm.	65-0010
	7, 1.2×10^{10} (rel.)	—	$k/k_{\text{BrO}^-} = 2.1 \pm 0.4$	γ -r.	trac.	c.k.; meas. $^{14}\text{CO}_2$.	65-0099
	10.7	—	—	—	—	—	—
	9 1.2×10^{10} (rel.)	—	$k/k_{\text{RNO}} = 1$	γ -r.	opt.	c.k.	65-0356
	3-7 $(1.07 \pm 0.10) \times 10^{10}$	—	—	p.r.	opt.	p.b.k. at 420 nm.	66-0424
	— 1×10^{10} (rel.)	—	$k/k_{\text{EtOH}} = 5.4$	p.r.	opt.	c.k.; N_2O -satd.; ratios 5 and 11.5, resp. in O_2 -satd. soln. contg. 0.05 M ferrocyanide.	71-0137
	— 1.1×10^{10} (rel.)	—	$k/k_{\text{MeOH}} = 12.4$	—	—	—	—
	nat. $(9.3 \pm 0.5) \times 10^9$	—	—	p.r.	opt.	p.b.k. at 410 nm.	71-0578

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.54 cont.		0-7	$(1.25 \pm 0.1) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 420 nm; calcd. $k(\text{OH} + \text{HFe}(\text{CN})_6^{3-}) = (9.0 \pm 0.9) \times 10^9$ and $k(\text{OH} + \text{H}_2\text{Fe}(\text{CN})_6^{2-}) = (1.7 \pm 0.5) \times 10^9$.	72-0431
		—	$(1.12 \pm 0.17) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 420 nm; c.k. with 2-PrOH gave 8.0×10^9 .	73-1031
		—	$(9.3 \pm 0.5) \times 10^9$	—	p.r.	opt.	p.b.k.; $\mu = 0.002$ to 10.	73-1039
			For other ratios see: 3.3, 3.12, 3.27, 3.62, 3.85, 3.128, 3.131, 3.134, 3.143, 3.151, 3.152, 3.168, 3.169, 3.191, 3.225, 3.231, 3.233, 3.310, 3.358, 3.367, 3.369, 3.384, 3.385, 3.394, 3.403, 3.405, 3.406, 3.473-3.473a, 3.506, 3.511, 3.527, 3.545, 3.546, 3.590, 3.614, 3.636, 3.637, 3.664, 3.686, 3.697, 3.711, 3.746.					
3.55	$\text{OD} + \text{Fe}(\text{CN})_6^{4-} \rightarrow \text{OD}^- + \text{Fe}(\text{CN})_6^{3-}$	nat	$(9.7 \pm 1.0) \times 10^9$	—	p.r.	opt.	p.b.k. at 420 nm; in D_2O .	72-0354
3.56	Fe^{3+}	acid	$(7.9 \pm 0.5) \times 10^7$ (rel.)	$k/k_{\text{Fe}^{2+}} = (3.45 \pm 0.2) \times 10^{-1}$	p.r.	—	c.k.	66-0715
			For other ratio see: 3.358.					
3.57	$\text{Fe}(\text{CN})_5\text{NO}^{2-}$	—	7.9×10^6 (rel.)	$k/k_{\text{bicarb}} = 0.22$	p.r.	opt.	c.k.; meas. abs. of CO_3^{2-} at 600 nm.	69-0052
3.58	$\text{Fe}(\text{EDTA})^-$	1	4.8×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 2.1$	X-r.	chem.	c.k.	71-0202
	$\text{OH} + \text{Fe}(\text{EDTA})^- \rightarrow \text{H}_2\text{O} + \text{prod.}$	6	1.5×10^9 (rel.)	$k/k_{\text{MeOH}} = 1.7$				
		7	9.9×10^8 (rel.)	$k/k_{\text{MeOH}} = 1.10$	X-r.	chem.	c.k.	75-0159
3.59	H_2	7	—	$k/k_{\text{perox}} = 0.94$	γ -r.	chem.	c.k.	52-0004
	$\text{OH} + \text{H}_2 \rightarrow \text{H}_2\text{O} + \text{H}$	~1	3.4×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.15$	Fenton	chem.	c.k.	57-0014
		1	2.7×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.116$	Fenton	chem.	c.k.	57-9007
		1.57	3.2×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.14$	γ -r.	chem.	c.k.	58-0004
		2.1	4.0×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.175$	γ -r.	chem.	c.k.	58-0004
		~7	—	$k/k_{\text{perox}} = 1.0$	γ -r.	chem.	c.k.	58-0012
		~1	$(3.2 \pm 0.2) \times 10^7$ (rel.)	$k/k_{\text{Fe}^{2+}} = 0.14 \pm 0.01$	Fenton	chem.	c.k.; $E_a(\text{OH} + \text{H}_2) - E_a(\text{OH} + \text{Fe}^{2+}) = 2.7 \pm 0.3$ kcal/mol (11.3 kJ/mol); $A(\text{OH} + \text{H}_2)/A(\text{OH} + \text{Fe}^{2+}) = 14 \pm 6$.	59-0028
		0.4	3.1×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.135$	γ -r.	chem.	c.k.	59-0064, 61-0100, 63-0004
		7	—	$k/k_{\text{perox}} = 0.93 \pm 0.03$	phot.	chem.	c.k.	62-7001
		2	—	$k/k_{\text{perox}} = 0.95$	γ -r.	chem.	c.k.	64-0212
		3	3.5×10^7	—	p.r.	opt.	d.k.	65-0010
		—	$(6.0 \pm 2.0) \times 10^7$	—	p.r.	opt.	d.k. at 260 nm.	66-0426
			For other ratios see: 3.12, 3.32, 3.38, 3.62, 3.82, 3.106, 3.118, 3.121, 3.291.					
3.60	D_2 $\text{OH} + \text{D}_2 \rightarrow \text{DHO} + \text{D}$	>2	$(1.2 \pm 0.1) \times 10^7$ (rel.)	$k/k_{\text{Fe}^{2+}} = 0.050 \pm 0.005$	Fenton	chem.	c.k.; $E_a(\text{OH} + \text{D}_2) - E_a(\text{OH} + \text{Fe}^{2+}) = 3.4 \pm 0.2$ kcal/mol (14.2 kJ/mol); $A(\text{OH} + \text{D}_2)/A(\text{OH} + \text{Fe}^{2+}) = 16 \pm 6$.	59-0028
3.61	D_2 $\text{OD} + \text{D}_2 \rightarrow \text{D}_2\text{O} + \text{D}$	alk.	$(1.6 \pm 0.2) \times 10^7$	—	p.r.	opt.	p.b.k. (meas. e_a^- from $\text{D} + \text{OD}^- \rightleftharpoons e_a^- + \text{D}_2\text{O}$).	68-0061
3.62	OH^- $\text{OH} + \text{OH}^- \rightarrow \text{H}_2\text{O} + \text{O}^-$	3.5-11	2.9×10^9 (rel.)	$k/k_{\text{I}^-} = 0.22$	phot.	chem.	c.k.	62-0057
		alk.	5.6×10^9 (rel.)	$k/k_{\text{ferro}} = 0.6$	p.r.	—	c.k.; estd.	65-0007
		10-14	3.0×10^8 (rel.)	$k/k_{\text{I}^-} = 0.025$	γ -r.	chem.	c.k.; pH effect on yields.	65-0219

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.62 cont.		11	$(1.2 \pm 0.3) \times 10^{10}$ (rel.)	—	p.r.	opt.	c.k. with MeOH and EtOH; soln. contains CO_3^{2-} and HCO_3^- ; assume $k(\text{O}^- + \text{H}_2\text{O}) = 1.7 \times 10^6$ and $\text{p}K_a(\text{OH}) = 11.9$.	70-0511
3.63	HO_2^- $\text{OH} + \text{HO}_2^- \rightarrow$ $\text{OH}^- + \text{HO}_2$		<i>For other ratios see: 3.107.</i>					
		13	8.3×10^9	—	p.r.	opt.	p.b.k. at 260 nm; involves various assumptions.	68-0298
		11	$1.4k + k(\text{O}^- + \text{H}_2\text{O}_2)$ $= (8 \pm 0.8) \times 10^9$ (rel.)	—	p.r.	opt.	c.k.; value relative to $k_{\text{carb}} = (4 \pm 0.2) \times 10^8$ and $\text{p}K_a(\text{OH}) = 11.9$; $\mu = 0.4$.	69-0379
		alk.	1.4×10^{10}	—	p.r.	condy.	computer anal.; more than one rate constant involved in calcn.	72-0404
3.64	H_2O_2 $\text{OH} + \text{H}_2\text{O}_2 \rightarrow$ $\text{HO}_2 + \text{H}_2\text{O}$	—	$(5 \pm 1.5) \times 10^9$ (rel.)	—	phot.	opt.	c.k.; calcd. from k/k_{RNO} at pH 7-10.52.	73-7575
		—	$(4.5 \pm 0.4) \times 10^7$	—	p.r.	opt.	Data fitting; G values.	62-0052
		3	1.2×10^7 (rel.)	—	p.r.	opt.	Data fitting; G values; rel. to $k(\text{H} + \text{H}) = 1.3 \times 10^{10}$.	64-0092
		7	$(2.6 \pm 0.8) \times 10^7$ (rel.)	$k/k_1^- = (2.2 \pm 0.7) \times 10^{-3}$	p.r.	opt.	c.k.	65-0010
		—	1.7×10^7 (rel.)	$k/k_{\text{RNO}} = 0.00136$	p.r.	opt.	c.k.	69-0156
		8.4	6.5×10^7 (rel.)	$k/k_{\text{bicarb}} = 1.8$	p.r.	opt.	c.k.	69-0379
		7	$(1.7 \pm 0.3) \times 10^7$ (rel.)	$k/k_{\text{RNO}} = 0.00136$	phot.	opt.	c.k.	73-7575
		6	4.5×10^7 (rel.)	$k/k_{\text{RNO}} = 0.0036$	phot.	opt.	c.k.	74-0052
3.65	HgCl $\text{OH} + \text{HgCl} \rightarrow \text{HgCl}^+$ $+ \text{OH}^-$ or $\text{Hg}(\text{OH})\text{Cl} + \text{H}^+$ $+ \text{Cl}^-$	<i>For other ratios see: 3.3, 3.5, 3.18, 3.32, 3.52, 3.59, 3.77, 3.115, 3.592, 3.711.</i>						
		5.0	$\sim 10^{10}$	—	p.r.	opt.	d.k. at 235 nm; reaction of e_{aq}^- or H with HgCl_2 gives HgCl .	73-0043
3.66	I^- (I) $\text{OH} + \text{I}^- \rightarrow \text{HOI}^-$ (II) $\text{HOI}^- \rightarrow \text{OH}^- + \text{I}$ (III) $\text{I} + \text{I}^- \rightleftharpoons \text{I}_2^-$	neut.	$(1.02 \pm 0.13) \times 10^{10}$	—	p.r.	opt.	p.b.k.; I_2^- is meas.; assumed that $\text{OH} + \text{I}^- \rightarrow \text{OH}^- + \text{I}$ is rate determining step.	65-0010
		10.5	1.4×10^{10} (rel.)	$k/k_{\text{BzO}^-} = 2.37 \pm 0.12$	γ -r.	trac.	c.k.; meas. $G(^{14}\text{CO}_2)$.	65-0099
		9	1.4×10^{10} (rel.)	$k/k_{\text{RNO}} = 1.14$	γ -r.	opt.	c.k.	65-0356
		—	—	$k/k_{\text{TCOO}^-} = 3.8$	γ -r.	trac.	c.k.; obs. ^3HHO .	68-0209
		9	1.2×10^{10} (rel.)	$k/k_{\text{RNO}} = 0.96 \pm 0.07$	γ -r.	opt.	c.k.; O_2 -satd.	68-0310
		—	3.4×10^{10}	—	p.r.	opt.	p.b.k.; method is indirect.	68-0375
		7	1.2×10^{10} (rel.)	$k/k_{\text{BzO}^-} = 2.1$	γ -r.	chem.	c.k.	68-0494
3.66		5.5	1.2×10^{10} (rel.)	$k/k_{\text{NB}} = 3.8$	γ -r.	opt.	c.k.; obs. σ -nitrophenol formn.	68-0494

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.66 cont.	2	1.1×10^{10} (rel.)	$k/k_{\text{PbH}} = 1.46$	γ -r.	opt.	c.k. with Safranine T.	69-0279
	3-5.5	7.4×10^9 (rel.)	$k/k_{\text{PbH}} = 0.95$	γ -r.	opt.	c.k. with Safranine T.	69-0279
	0-2	1.6×10^{10} (rel.)	$k/k_{2\text{-PrOH}} = 7.1 \pm 0.2$	γ -r.	chem.	c.k.; obs. $G(\text{acetone})$.	68-0602
	6.98	1.8×10^{10} (rel.)	$k/k_{2\text{-PrOH}} = 8.1 \pm 0.1$	γ -r.	chem.	c.k.; $\mu = 0.1 - 1.1$.	68-0602
	11	6.6×10^9 (rel.)	$k/k_{\text{carb}} = 18$	p.r.	opt.	c.k.	69-0379
	—	2.4×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 2.14$	p.r.	opt.	c.k.	70-1226
	—	4.0×10^{10} (rel.)	$k/k_{\text{NB}} = 12.6$	p.r.	opt.	c.k.	70-1226
	—	1×10^{10} (rel.)	$k/k_{\text{EtOH}} = 5.5$	p.r.	opt.	c.k.; N_2O -satd.; ratio 7 in O_2 satd. $0.1 M \text{I}^-$ soln.	71-0137
	9	1.5×10^{10} (rel.)	$k/k_{\text{RNO}} = 1.22$	r.	opt.	c.k.; $E_a = 0.7 \pm 0.3 \text{ kcal/mol}$ (2.9 kJ/mol) ($265-296 \text{ K}$).	71-0469
	~ 6	1.4×10^{10} (rel.)	$k/k_{\text{MeOH}} = 15.2 \pm 0.9$	γ -r.	chem.	c.k.	71-0931
	—	$(1.21 \pm 0.08) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 385 nm (I_2); $k_1 = k_{\text{III}}$; $k_{\text{II}} = (1.2 \pm 1.0) \times 10^8 \text{ s}^{-1}$.	72-0122
	For other ratios see: 3.21, 3.54, 3.62, 3.64, 3.91, 3.110, 3.128, 3.129, 3.131, 3.186, 3.220, 3.228, 3.343, 3.358, 3.365, 3.371, 3.384, 3.385, 3.394, 3.482, 3.511, 3.592, 3.637, 3.647, 3.711.						
3.67	IO_3^- $\text{OH} + \text{IO}_3^- \rightarrow \text{IO}_3 + \text{OH}^-$	12.4-13.6	$(9.2 \pm 0.8) \times 10^8$ (rel.)	—	f.phot.	chem. d.k. at 430 nm (O_3); value is based on $k(\text{O}^- + \text{O}_2 \rightarrow \text{O}_3) = 2.5 \times 10^9$.	70-0018
		7	$\leq 5 \times 10^7$ (rel.)	—	p.r.	opt. c.k. with EtOH and 2-PrOH; obs. decrease in abs. at 360 nm.	72-0017
		6	$(1.1 \pm 0.1) \times 10^7$	—	p.r.	opt. p.b.k. at 360 nm.	73-0027
3.68	IO_4^- $\text{OH} + \text{IO}_4^- \rightarrow \text{OH}^- + \text{IO}_4$	5.6	$(4.5 \pm 0.5) \times 10^8$	—	p.r.	opt. p.b.k. at 520 nm; computer anal.	71-0274 71-0335
3.69	Mn^{2+} $\text{OH} + \text{Mn}^{2+} \rightarrow \text{Mn}^{3+} + \text{OH}^-$	—	$\geq 1.4 \times 10^8$	—	p.r.	opt. p.b.k. at 450 nm.	65-0395
3.70	$\text{Mn}(\text{CN})_5\text{NO}^{3-}$	—	4.2×10^9 (rel.)	$k/k_{\text{RNO}} = 0.34$	γ -r.	opt. c.k.; depends on $k_{\text{CN}^-}/k_{\text{RNO}} = 0.42$.	71-0407
3.70a	$\text{Mo}(\text{CN})_8^{4-}$ $\text{OH} + \text{Mo}(\text{CN})_8^{4-} \rightarrow \text{OH}^- + \text{Mo}(\text{CN})_8^{3-}$	—	$(5.8 \pm 0.6) \times 10^9$ (rel.)	$k/k_{\text{bicarb}} = 16$	p.r.	opt. c.k.	73-1031
3.71	NH_3 $\text{OH} + \text{NH}_3 \rightarrow \text{H}_2\text{O} + \text{NH}_2$	—	1.0×10^8	—	p.r.	opt. d.k. (OH) or p.b.k. at 530 nm (NH_2).	72-0109
		11.3	1.5×10^7 (rel.) 3.6×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.00135$ $k/k_{\text{MeOH}} = 0.04$	p.r.	opt. c.k.; includes $\text{O}^- + \text{NH}_3$.	72-0460
3.72	NH_2 $\text{OH} + \text{NH}_2 \rightarrow \text{NH}_2\text{OH}$	—	9.5×10^9	—	p.r.	chem. effect of NH_3 concn. on $G(\text{NH}_2\text{OH})$.	72-0109
3.73	N_3^- $\text{OH} + \text{N}_3^- \rightarrow \text{N}_3 + \text{OH}^-$	9	1.1×10^{10} (rel.)	$k/k_{\text{RNO}} = 0.86$	γ -r.	opt. c.k.	65-0356
		9.2	1.1×10^{10} (rel.)	$k/k_{\text{BzO}^-} = 2.0$	p.r.	opt. c.k.; meas. abs of N_3 at 278 nm.	70-0649
3.74	NH_2OH	8	9.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.86$	p.r.	opt. c.k.	71-0493
3.75	NH_3OH^+	4	$\leq 5.0 \times 10^8$ (rel.)	$k/k_{\text{CNS}^-} = 0.0455$	p.r.	opt. c.k.	71-0493

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.76	NH ₂ NH ₂ 10 OH + NH ₂ NH ₂ → H ₂ O + N ₂ H ₃	1.4 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.3	p.r.	opt.	c.k.	72-0003
3.77	NH ₂ NH ₃ ⁺ ~1 OH + NH ₂ NH ₃ ⁺ → 2 H ₂ O + N ₂ H ₄ ⁺	~ 2 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{Fe²⁺} ≅ 0.09 <i>k</i> / <i>k</i> _{perox} = 1 ± 0.1	γ-r. r.	chem. chem.	c.k. calcd. assuming mechanism.	62-0136 56-7004
3.78	NOH(SO ₃) ₂ ²⁻ 6 OH + NOH(SO ₃) ₂ ²⁻ 8.4- → H ₂ O + ON(SO ₃) ₂ ²⁻ 12	1.0 x 10 ⁹ (rel.) 5.7 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.091 <i>k</i> / <i>k</i> _{pH} = 0.073	p.r. e-r.	opt. esr	c.k. obs. buildup of ON(SO ₃) ₂ ; <i>k</i> probably is concn. dependent.	72-0003 68-0471
3.79	NO(SO ₃) ₂ ²⁻ (Fremy's salt) —	4.94 x 10 ⁸ (rel.) 2.6 x 10 ¹⁰	<i>k</i> / <i>k</i> _{pH} = 0.063 —	— —	— —	calcd. calcd.	71-0596 71-0596
3.80	NO 7 OH + NO → NO ₂ ⁻ + H ⁺ 7	1.1 x 10 ¹⁰ (rel.) 8.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{MeOH} = 12.5 <i>k</i> / <i>k</i> _{EtOH} = 4.8 ± 0.6	γ-r. γ-r.	chem. chem.	c.k. (for product ident. see 70-0228). c.k.	66-0118 66-0118
	7	1.1 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 4.8 ± 0.6	γ-r.	chem.	c.k.	66-0118
	—	—	<i>k</i> / <i>k</i> _{nitrite} = 1.6 ± 0.4	f.phot.	opt.	c.k.	70-7264
	7	1 x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 220 nm (NO ₂).	73-0096
3.81	NO ₂ 9 OH + NO ₂ → HO ₂ NO	1.3 x 10 ⁹	—	p.r.	opt.	meas. buildup of abs. at 302 nm in NO ₃ ⁻ soln.; calcn. involves <i>k</i> (OH + OH) = 0.6 x 10 ¹⁰ and <i>k</i> for NO ₃ ²⁻ (+ H ₂ O) → NO ₂ + 2OH ⁻ = 5.5 x 10 ⁴ s ⁻¹ .	70-0151
3.82	NO ₂ ⁻ — OH + NO ₂ ⁻ → NO ₂ + OH ⁻ —	1.2 x 10 ¹⁰ (rel.) 1.1 x 10 ¹⁰	<i>k</i> / <i>k</i> _{carb} = 32 <i>k</i> / <i>k</i> _{OH} = 1.45	p.r. p.r.	opt. calcd.	c.k. math. anal. of data from NO ₃ ⁻ soln.; assume <i>k</i> _{OH} = 7.6 x 10 ⁹ .	64-0131 64-0294
	10.7	6.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{carb} = 18	p.r.	opt.	c.k.	65-0190
	9	8.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.65	γ-r.	opt.	c.k.	65-0356
	—	5.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{MeOH} = 6.5 ± 0.8	γ-r.	chem.	c.k. in NO-MeOH-KNO ₂ solns.	66-0118
	—	—	<i>k</i> / <i>k</i> _{hydr} = 125	γ-r.	chem.	c.k.	67-0032
	—	—	<i>k</i> / <i>k</i> _{TCCO⁻} = 3.0	γ-r.	trac.	c.k.; obs. ³ HHO.	68-0209
	9	7.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.57 ± 0.03	γ-r.	opt.	c.k.	68-0310
	11	7.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{carb} = 20	p.r.	opt.	c.k.	69-0379
	acid 1 x 10 ¹⁰ (rel.)		<i>k</i> / <i>k</i> _{MeOH} = 11.7	p.r.	condy.	c.k.	70-0254
	alk. 8.5 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{MeOH} = 9.4	p.r.	condy.	c.k.	70-0254
	>12 1 x 10 ¹⁰ (rel.)		<i>k</i> / <i>k</i> (O ⁻ + O ₂) = 4.0 ± 0.4	f.phot.	opt.	c.k.; meas. dependence of O ₃ ⁻ decay rate on OH ⁻ and NO ₂ ⁻ .	70-7264
	9	1.1 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.86	γ-r.	opt.	c.k.; <i>E</i> _a = - 1.0 ± 1.0 kcal/mol (-4.2 kJ/mol) (-8 to 23°C).	71-0469

For other ratios see: 3.80.

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.83	NO_3^- $\text{OH} + \text{NO}_3^- \rightarrow$ $\text{OH}^- + \text{NO}_3$	9	$< 5 \times 10^5$ (rel.)	—	—	—	c.k. with RNO.	66-0843
3.84	$\text{HNO}_3 (+\text{NO}_3^-)$ $\text{OH} + \text{HNO}_3 \rightarrow$ $\text{H}_2\text{O} + \text{NO}_3$	~ 0	—	—	p.r.	opt.	p.b.k. (NO_3) at 635 nm; pseudo-first order rate = $1.5 \times 10^5 \text{ s}^{-1}$ at 0.1 M HNO_3 and $4.2 \times 10^5 \text{ s}^{-1}$ at 0.4 or 1.0 M HNO_3 .	67-0002
		$\sim 0-1$	—	—	p.r.	opt.	p.b.k.; rate of formn. of NO_3 (2 to 12) $\times 10^5 \text{ s}^{-1}$; first order in H^+ and NO_3^- .	69-0417
		4 M HNO_3	—	$k[\text{H}^+][\text{NO}_3^-]/k_{\text{HCOOH}} = 0.21 \pm 0.03 \text{ M}$	γ -r.	chem.	c.k. in Ce(III)-Ce(IV)- HCOOH soln. $k_{\text{Ce(III)}}/k_{\text{HCOOH}} = 4.1$.	72-0263
			For ratio see: 3.720a					
3.84a	Ni^{2+}	—	$< 5 \times 10^5$	—	p.r.	opt.	c.k. with Cu^{2+}	75-1027
3.85	$\text{Ni}(\text{CN})_4^{2-}$ $\text{OH} + \text{Ni}(\text{CN})_4^{2-} \rightarrow$ $\text{OH}^- + \text{Ni}(\text{CN})_4^-$	—	$(9.1 \pm 0.5) \times 10^9$ (rel.)	$k/k_{\text{ferro}} = 0.98$	p.r.	opt.	c.k.; also p.b.k. at 250 nm.	74-1072
3.86	$\text{Ni}(\text{en})_n^{2+}$ $\text{OH} + \text{Ni}(\text{en})_n^{2+} \rightarrow$ $\text{OH}^- + \text{Ni}(\text{en})_n^{3+}$	8.0	$(6.0-7.2) \times 10^8$ (rel.)	$k/k_{\text{CNS}^-} = 0.055 - 0.065$	p.r.	opt.	c.k.; cor. for OH + en.	72-0461
		8.5	$(4.1-7.2) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = 0.37-0.65$				
		9.0	$(5.5-6.6) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = 0.5-0.6$				
		10.0	$(5.5-9.4) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = 0.5-0.85$				
3.87	$\text{Ni}(\text{gly})_n^{2+}$ $\text{OH} + \text{Ni}(\text{gly})_n^{2+} \rightarrow$ $\text{OH}^- + \text{Ni}(\text{gly})_n^{3+}$	10.0	$(4.9-7.7) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = 0.45 - 0.7$	p.r.	opt.	c.k.; cor. for OH + glycine.	72-0461
3.88	$\text{Ni}(\text{EDTA})^{2-}$	7	2.8×10^9 (rel.) 2.1×10^9 (rel.)	$k/k_{\text{MeOH}} = 3.1$ $k/k_{\text{HCOO}^-} = 0.61$	X-r.	chem.	c.k.	72-0173
3.88a	$\text{Os}(\text{CN})_6^{4-}$ $\text{OH} + \text{Os}(\text{CN})_6^{4-} \rightarrow$ $\text{OH}^- + \text{Os}(\text{CN})_6^{3-}$	—	$(1.03 \pm 0.12) \times 10^{10}$ 8.6×10^9 (rel.) 1.02×10^{10} (rel.)	— $k/k_{2-\text{PrOH}} = 4$ $k/k_{\text{MeOH}} = 9.3$	p.r.	opt.	p.b.k. at 330 and 410 nm; also c.k.	73-1031
3.88b	$\text{Os}(\text{NH}_3)_5\text{N}_2^{2+}$ $\text{OH} + \text{Os}(\text{NH}_3)_5\text{N}_2^{2+} \rightarrow$ $\text{H}_2\text{O} +$ $\text{Os}(\text{NH}_3)_4\text{NH}_2\text{N}_2^{2+}$	—	1×10^{10}	—	p.r.	opt.	p.b.k. at 380 nm.	75-0309, 75-1099
3.89	$(\text{NaPO}_3)_n$	—	$< 5 \times 10^6$	—	p.r.	opt.	no absorbing product formed; $n \cong 50$.	74-0036
3.90	H_3PO_4 $\text{OH} + \text{H}_3\text{PO}_4 \rightarrow$ $\text{H}_2\text{O} + \text{H}_2\text{PO}_4$	0.0	2.6×10^6 (rel.)	$k/k_{\text{MeOH}} = 0.0028$	p.r.	opt.	c.k.; obs. H_2PO_4 radical at 500 nm.	73-1049
3.91	H_2PO_4^- $\text{OH} + \text{H}_2\text{PO}_4^- \rightarrow$ $\text{H}_2\text{PO}_4 + \text{OH}^-$	~ 7	$< 1.2 \times 10^7$ (rel.)	$k/k_{\text{I}^-} < 0.001$	p.r.	opt.	c.k.; contains HPO_4^{2-} ($\text{p}K_a = 7.2$).	65-0010
		3.85-4.0	2.2×10^6 (rel.)	$k/k_{\text{MeOH}} = 0.0024$	p.r.	opt.	c.k.; obs. phosphate radical at 500 nm.	73-1049

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.92	HPO_4^{2-}	—	$< 5 \times 10^6$ (rel.)	$k/k_{\text{carb}} < 0.012$	p.r.	opt.	c.k.	70-0302
		9.0-12.3	$(7.9 \pm 0.4) \times 10^5$	—	p.r.	opt.	p.b.k. at 500 nm; also $k = 9 \times 10^5$ by c.k. with MeOH.	73-1049
3.93	PO_4^{3-}	—	$< 10^7$ (rel.)	$k/k_{\text{carb}} < 0.025$	p.r.	opt.	c.k.	70-0302
3.94	$\text{P}_2\text{O}_7^{4-}$	—	$< 4 \times 10^6$ (rel.)	$k/k_{\text{carb}} < 0.01$	p.r.	opt.	c.k.	70-0302
		10.3	$(9 \pm 1) \times 10^5$	—	p.r.	opt.	p.b.k. at 590 nm.	73-1049
3.95	H_2PO_2^-	10.7	1.7×10^9 (rel.)	$k/k_{\text{carb}} = 4.7$	p.r.	opt.	c.k.	65-0190
3.96	PO_3^{3-}	10.7	3.5×10^9 (rel.)	$k/k_{\text{carb}} = 9.5$	p.r.	opt.	c.k.	65-0190
3.97	$\text{OH} + \text{PdCl}_4^{2-} \rightarrow \text{Pd(III)}$	—	$(6.3 \pm 0.3) \times 10^9$ (rel.)	$k/k_{\text{t-BuOH}} = 12$	p.r.	opt.	c.k. in 0.01 M NaCl, assume $k_{\text{t-BuOH}} = 5.2 \times 10^8$; $k = 1.2 \times 10^{10}$ in 1M NaCl.	74-1087
3.98	Pr^{3+} $\text{OH} + \text{Pr}^{3+} \rightarrow \text{OH}^- + \text{Pr}^{4+}$	5.8	2×10^6 (ave.)	—	p.r.	opt.	p.b.k. at 300 nm; also detd. by c.k. with H_2O_2 or CNS $^-$.	71-0311, 72-0066
3.99	PtCl_4^{2-} $\text{OH} + \text{PtCl}_4^{2-} \rightarrow \text{Pt(III)} + \text{OH}^-$	3.5 ~11	$\sim 3.5 \times 10^6$ $(8 \pm 2) \times 10^9$	— —	p.r. p.r.	opt. opt.	p.b.k. at 290 nm. p.b.k. at 450 nm.	73-1084 69-0144
3.100	Pt(CN)_4^{2-} $\text{OH} + \text{Pt(CN)}_4^{2-} \rightarrow \text{Pt(III)} + \text{OH}^-$	~2	1.1×10^{10} (rel.)	$k/k_{\text{MeOH}} = 12$	p.r.	opt.	c.k.	69-0144
3.100a	Ru(CN)_6^{4-} $\text{OH} + \text{Ru(CN)}_6^{4-} \rightarrow \text{OH}^- + \text{Ru(CN)}_6^{3-}$	—	$(5.7 \pm 0.8) \times 10^9$	—	p.r.	opt.	p.b.k. at 330, 355 and 470 nm; c.k. with 2-PrOH gave 4.4×10^9 .	73-1031
3.101	$\text{Ru(NH}_3)_5\text{N}_2^{2+}$ $\text{OH} + \text{Ru(NH}_3)_5\text{N}_2^{2+} \rightarrow \text{OH}^- + \text{Ru(NH}_3)_5\text{N}_2^{3+}$	—	4.8×10^9	—	p.r.	opt.	p.b.k. at 440-44 nm.	71-0234
3.102	H_2S $\text{OH} + \text{H}_2\text{S} \rightarrow \text{H}_2\text{O} + \text{HS}^-$	6 2- 5.7 6 5.5	1.9×10^{10} (rel.) 2.2×10^{10} (rel.) 1.4×10^{10} (rel.) 1.5×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.7$ $k/k_{\text{CNS}^-} = 2 \pm 0.5$ $k/k_{\text{MeOH}} = 15$ $k/k_{\text{HCOO}^-} = 4.4$	p.r. p.r. p.r. p.r.	opt. opt. opt. opt.	c.k. c.k. c.k. c.k.	67-0273 67-0684 67-0684 67-0684
3.103	HS^- $\text{OH} + \text{HS}^- \rightarrow \text{OH}^- + \text{SH}$	—	9×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.82$	p.r.	opt.	c.k.; also with MeOH, formate ion.	67-0273
		10.5	8.4×10^9 (rel.)	$k/k_{\text{MeOH}} = 9.3$	p.r.	opt.	c.k.	67-0684
		10.5	9.4×10^9 (rel.)	$k/k_{\text{HCOO}^-} = 2.7$	p.r.	opt.	c.k.	67-0684
3.104	HSO_3^-	—	9.5×10^9 (rel.)	$k/k_{\text{carb}} = 26$	p.r.	opt.	c.k.	64-0131
3.105	SO_3^{2-}	—	5.5×10^9 (rel.)	$k/k_{\text{carb}} = 15$	p.r.	opt.	c.k.	64-0131
3.106	HSO_4^- $\text{OH} + \text{HSO}_4^- \rightarrow \text{HSO}_4 + \text{OH}^-$ or $\rightarrow \text{SO}_4^- + \text{H}_2\text{O}$	0.8 0.8 1 0.8 0.4 0.1-0.8 0.4-1 ~7	— — — — — — — 1.6×10^6 (rel.)	$k/k_{\text{HCOOH}} = 0.0026$ $k/k_{\text{hydr}} = 0.005$ $k/k_{\text{HCOOH}} = 0.0016$ $k/k_{\text{HCOOH}} = 0.0013$ $k/k_{\text{HCOOH}} = 0.0011$ $k/k_{\text{HCOOH}} = 0.0009$ $k/k_{\text{hydr}} = 0.0039$ $k/k_{\text{hydr}} = 0.011$ $k/k_{\text{MeOH}} = 0.0018$	γ -r. phot. γ -r. phot. phot. phot. γ -r. γ -r. p.r.	chem. chem. chem. chem. chem. chem. chem. chem. chem.	c.k.; rel to $k_{\text{Ce}^{3+}}/k_{\text{HCOOH}} = 1.7$. c.k. c.k. c.k. c.k. c.k. c.k. c.k. c.k., reaction is $\text{OH} + \text{HSO}_4^- \rightarrow \text{H}_2\text{O} + \text{SO}_4^-$ at this pH.	57-0003 62-7001 63-0048 63-0048 63-0048 63-0048 63-0197 65-0052 66-0019

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.106 cont.	~7	6.9×10^5	—	p.r.	opt.	p.b.k.; see above.	66-0019
	0-0.8	—	$k/k_{\text{hydr}} = 0.01$	γ -r.	chem.	c.k.	66-0029
	0.3-2	1.2×10^6 (rel.)	$k/k_{\text{pH}} = 1.5 \times 10^{-4}$	γ -r.	opt.	c.k. with Safranin T.	69-0279
	4 M H_2SO_4	1.5×10^6	—	p.r.	opt.	estd. from d.k. SO_4^{2-} ; $0.4k(\text{OH} + \text{SO}_4) + k(\text{H} + \text{SO}_4) = 3.3 \times 10^9$.	73-1030
<i>For other ratios see: 3.26, 3.385, 3.511, 3.637.</i>							
3.107	$\text{S}_2\text{O}_3^{2-}$ $\text{OH} + \text{S}_2\text{O}_3^{2-} \rightarrow \text{S}_2\text{O}_3^- + \text{OH}^-$	— — 1.2×10^9 (rel.) 3×10^9 (rel.) 1.6×10^9 (rel.) $(8 \pm 1.5) \times 10^9$ (rel.)	$k/k_{\text{OH}^-} = 1.2$ $k/k_{\text{MeOH}} = 1.3$ $k/k_{\text{EtOH}} = 1.9$ $k/k_{2-\text{PrOH}} = 0.8$ $k/k_{\text{CNS}^-} = 0.78$	γ -r. p.r.	chem. opt.	c.k. c.k.	71-0927 73-1027
3.108	HSO_5^- $\text{OH} + \text{HSO}_5^- \rightarrow \text{SO}_5^- + \text{H}_2\text{O}$ or $\rightarrow \text{HSO}_5 + \text{OH}^-$	— 3.5×10^8 (rel.)	$k/k_{\text{carb}} = 0.97$	p.r.	opt.	c.k.	69-0158
3.109	$\text{S}_2\text{O}_8^{2-}$	— $< 10^6$	—	p.r.	—	reaction not obs.; c.k. with CO_3^{2-} .	69-0158
3.110	H_2Se $\text{OH} + \text{H}_2\text{Se} \rightarrow \text{HSe} + \text{H}_2\text{O}$	1.0 $(1.0 \pm 0.3) \times 10^{10}$ (rel.)	—	p.r.	opt.	c.k.; rel. to $k(\text{OH} + \text{CNS}^-) = 6.6 \times 10^9$, $k(\text{OH} + \text{I}^-) = 7.0 \times 10^9$, $k(\text{OH} + \text{Br}^-) = 5.0 \times 10^9$.	69-0564
3.111	HSe^- $\text{OH} + \text{HSe}^- \rightarrow \text{HSe} + \text{OH}^-$	8.5- 11.5 $(5.5 \pm 0.1) \times 10^9$ (rel.)	—	p.r.	opt.	c.k.; meas. H_2Se_2^- at 410 nm; rel. to $k(\text{OH} + 2-\text{PrOH}) = 1.3 \times 10^9$, $k(\text{OH} + \text{HCOO}^-) = 2.5 \times 10^9$.	69-0564
3.112	SeO_3^{2-}	7 7 7 2.7×10^9 4.6×10^9 (rel.) 4.9×10^9 (rel.)	— $k/k_{\text{EtOH}} = 2.46$ $k/k_{\text{MeOH}} = 5.42$	p.r. p.r. p.r.	opt. opt. opt.	p.b.k. at 435 nm. c.k. c.k.	65-0190 65-0190 65-0190
3.113	Sm^{2+} $\text{OH} + \text{Sm}^{2+} \rightarrow \text{OH}^- + \text{Sm}^{3+}$	— 3-6 6×10^9 $(6.2 \pm 0.8) \times 10^9$	— —	p.r. p.r.	opt. opt.	d.k. (Sm^{2+}). d.k. (Sm^{2+} formed in Sm^{3+} soln.)	71-0311 73-1084
3.114	Sn^{2+} $\text{Sn}^{2+} + \text{OH} \rightarrow \text{Sn}^{3+} + \text{OH}^-$	0.8 $(1.6 \pm 0.2) \times 10^9$ (rel.)	$k/k_{\text{Fe}^{2+}} = 7 \pm 0.7$	γ -r.	chem.	c.k.	59-0007
3.115	$\text{H}_2\text{TeO}_3 + \text{HTeO}_3^-$ $\text{OH} + \text{Te(IV)} \rightarrow \text{Te(V)}$	0.4 0.4	$k/k_{\text{perox}} = 0.71$ $k/k_{\text{perox}} = 0.11$	γ -r. γ -r.	chem. chem.	c.k.; prelim. value. c.k.	67-0553 68-0356
3.116	TeO_3^{2-}	10.7 3.5×10^9 (rel.)	$k/k_{\text{carb}} = 9.5$	p.r.	opt.	c.k.	65-0190
3.117	Ti^{3+} $\text{OH} + \text{Ti}^{3+} \rightarrow \text{OH}^- + \text{Ti}^{4+}$	~1 1.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.11$	p.r.	opt.	c.k.	72-0240
3.117a	$\text{TiO}(\text{C}_2\text{O}_4)_2^{2-}$ $\text{OH} + \text{TiO}(\text{C}_2\text{O}_4)_2^{2-} \rightarrow \text{TiO}(\text{C}_2\text{O}_4) + \text{CO}_2 + \text{CO}_2^- + \text{OH}^-$	— —	$k/k_{\text{Br}^-} = 13$	γ -r.	chem.	c.k.	74-0521
3.118	Ti^+ $\text{OH} + \text{Ti}^+ (+ \text{H}^+) \rightarrow \text{Ti}^{2+} + \text{H}_2\text{O}$	0.4 0.8 0.8 — $(7.6 \pm 1) \times 10^9$	$k/k_{\text{Ce}^{3+}} = 38$ $k/k_{\text{Ce}^{3+}} = 42$ $k/k_{\text{hydr}} = 218 \pm 60$ —	γ -r. phot. γ -r. p.r.	chem. chem. chem. opt.	c.k. c.k. c.k. p.b.k. at 260 nm.	56-0004 57-7001 66-0029 66-0097

TABLE 3. Reactions of OH with inorganic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.118 cont.		1	$(1.0 \pm 0.1) \times 10^{10}$	—	p.r.	opt.	p.b.k.; cor. for H + Ti^{2+} , OH + Ti^{2+} , OH + H, etc.	74-1017
			<i>For other ratios see: 3.294.</i>					
3.119	Tm(II) OH + Tm(II) → OH ⁻ + Tm(III)	3-6	$(7 \pm 1) \times 10^9$	—	p.r.	opt.	d.k. of Tm(II) formed in Tm(III) soln.	73-1084
3.120	U(IV)	—	$\sim 2 \times 10^9$	—	γ-r.	chem.	estd. by c.k. with $\text{C}_2\text{O}_4^{2-}$; U(IV) formed in UO_2^{2+} soln.	71-0542
3.121	VO^{2+} OH + $\text{VO}^{2+} \rightarrow$ $\text{VO}_2^+ + \text{H}^+$	acid	—	$k/k_{\text{hydr}} = 11 \pm 3$	γ-r.	chem.	c.k.	66-0029
		~1	2.5×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.023$	p.r.	opt.	c.k.	72-0240
3.122	Yb^{2+}	—	3×10^9	—	p.r.	opt.	d.k. (Yb^{2+}).	71-0311
	OH + $\text{Yb}^{2+} \rightarrow$ OH ⁻ + Yb^{3+}	2	$(3.2 \pm 0.3) \times 10^9$	—	p.r.	opt.	d.k. (Yb^{2+} formed on p.r. of Yb^{3+} soln.).	73-1084
3.122a	Zn^{2+} OH + $\text{Zn}^{2+} \rightarrow$ OH ⁻ + Zn^{3+}	—	$< 5 \times 10^5$	—	p.r.	opt.	c.k. with Cu^{2+} .	75-1027

TABLE 4. Reactions of OH with organic solutes

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.123	acetaldehyde	1	5×10^8 (rel.)	$k/k_{Fe^{2+}} = 2.2$	Fenton	chem.	c.k.	49-0002
3.124	acetamide	9	1.3×10^7 (rel.)	$k/k_{EtOH} = 0.0071$	γ -r.	opt.	c.k. with RNO.	66-0423
	(I) $OH + CH_3CONH_2 \rightarrow$ $H_2O + CH_2CONH_2$	5.5	1.9×10^8 (rel.)	$k/k_{CNS^-} = 0.017$	p.r.	opt.	c.k.	70-0098
	(II) $OH + CH_3CONH_2$ $\rightarrow H_2O +$ CH_3CONH	—	1.9×10^8 (rel.)	$k/k_{CNS^-} = 0.017$	p.r.	opt.	c.k.; $k_{II} = 9.5 \times 10^7$ by anal. of transient spectra.	71-0645
3.125	2-acetamido-2-deoxy-D-galactose	—	1.6×10^9 (rel.)	$k/k_{CNS^-} = 0.147$	p.r.	opt.	c.k.	70-3081
3.126	2-acetamido-2-deoxy-D-glucose	—	3.1×10^9 (rel.)	$k/k_{CNS^-} = 0.279$	p.r.	opt.	c.k.	70-3081
3.127	acetanilide	9	5×10^9 (rel.)	$k/k_{EtOH} = 2.73$	γ -r.	opt.	c.k. with RNO.	66-0441
3.128	acetate ion	10.7	6.3×10^7 (rel.)	$k/k_{BzO^-} = 0.011$	γ -r.	trac.	c.k.; meas. $^{14}CO_2$.	65-0099
		9.0	8.8×10^7 (rel.)	$k/k_{RNO} = 0.007$	γ -r.	opt.	c.k.	65-0356
		9	7.2×10^7 (rel.)	$k/k_{EtOH} = 0.039$	γ -r.	opt.	c.k. with RNO.	66-0423
		nat.	7.0×10^7 (rel.)	$k/k_{ferro} = 0.0075$	p.r.	opt.	c.k.	71-0578
		—	8.5×10^7	—	p.r.	opt.	p.b.k. at 350 nm.	71-0578
		—	1.2×10^8 (rel.)	$k/k_{I^-} = 0.0092$	p.r.	opt.	c.k.; obs. I_2^- .	73-0020
3.129	acetic acid	1.0	1.8×10^7 (rel.)	$k/k_{I^-} = (1.4 \pm 0.2) \times 10^{-3}$	p.r.	opt.	c.k.	65-0010
		1	$(9.2 \pm 3.8) \times 10^6$	—	p.r.	opt.	d.k. at 260 nm.	65-0010
		1	2.3×10^7 (rel.)	$k/k_{CNS^-} = 0.0021$	p.r.	opt.	c.k.	65-0387
		2-2.2	2.3×10^7 (rel.)	$k/k_{thym} = 0.0043$	γ -r.	opt.	c.k.	67-0461
		1	2×10^7 (rel.)	$k/k_{thym} = 0.0037$	Fenton	esr	c.k.; $k/k_{perox} = 0.27$.	69-5278
		1	1.9×10^7 (rel.)	$k/k_{thym} = 0.0035$	Ti(III) + H_2O_2	esr	c.k.; $k/k_{perox} = 0.25$.	69-5278
		~0	—	$k/k_{acrylamide} = 0.01$	Fenton	pol.	c.k.	72-9162
		1	2.0×10^7 (rel.)	$k/k_{MeOH} = 0.022$	Fenton	chem.	c.k.; $k_{MeOH}/k_{Fe^{2+}} = 4.3$.	73-9341
3.130	acetoin	2.0	8.5×10^8 (rel.)	$k/k_{CNS^-} = 0.077$	p.r.	opt.	c.k.	65-0387
	$OH + CH_3CH(OH)COCH_3$ $\rightarrow CH_3COHCOCH_3$ + H_2O	—	1.2×10^9 (rel.)	$k/k_{CNS^-} = 0.11$	p.r.	opt.	c.k.	68-0249
3.131	acetone	7	9.0×10^7 (rel.)	$k/k_{I^-} = (7.5 \pm 0.8) \times 10^{-3}$	p.r.	opt.	c.k.	65-0010
		10.7	9.1×10^7 (rel.)	$k/k_{BzO^-} = 0.016$	γ -r.	trac.	c.k.; meas. $^{14}CO_2$.	65-0099
		6-7	9.7×10^7 (rel.)	$k/k_{CNS^-} = 0.0088$	p.r.	opt.	c.k.	65-0387
		9	7.2×10^7 (rel.)	$k/k_{EtOH} = 0.039$	γ -r.	opt.	c.k. with RNO.	66-0423
		0.8	3.8×10^7 (rel.)	$k/k_{Fe^{2+}} = 0.165$	Fenton	chem.	c.k.	66-9002
		2-2.2	7.6×10^7 (rel.)	$k/k_{thym} = 0.014 \pm 0.0015$	γ -r.	opt.	c.k.	67-0461
		9	$\sim 7 \times 10^7$ (rel.)	$k/k_{RNO} \sim 0.0056$	γ -r.	opt.	c.k.	67-0555
		nat.	1.2×10^8 (rel.)	$k/k_{ferro} = 0.0129$	p.r.	opt.	c.k.	71-0578
		1	7.2×10^7 (rel.)	$k/k_{MeOH} = 0.080$	Fenton	chem.	c.k.; $k_{MeOH}/k_{Fe^{2+}} = 4.3$.	73-9341
3.132	acetone- d_6	1	2.3×10^7 (rel.)	$k/k_{MeOH} = 0.026$	Fenton	chem.	c.k.; $k_{MeOH}/k_{Fe^{2+}} = 4.3$.	73-9341
3.133	acetonitrile	9	3.6×10^6 (rel.)	$k/k_{EtOH} = 0.0019$	γ -r.	opt.	c.k. with RNO.	66-0423
		—	7.7×10^6 (rel.)	$k/k_{HCOO^-} = 0.0022$	γ -r.	chem.	c.k.; obs. $G(CO_2)$.	73-0364
		—	2.2×10^7 (rel.)	$k/k_{PNBA^-} = 0.0085$	p.r.	opt.	c.k.	75-1003
3.134	acetophenone	9	4.8×10^9 (rel.)	$k/k_{EtOH} = 2.6$	γ -r.	opt.	c.k. with RNO.	66-0441
	$OH + C_6H_5COCH_3 \rightarrow$ $OHC_6H_5COCH_3$	7	$(6.5 \pm 0.7) \times 10^9$	—	p.r.	opt.	p.b.k. at 372 nm (hydroxyxylohexadienyl radical); cor. for (OH + OH) and (H + aromatic).	68-0304

TABLE 4. Reactions of OH with organic solutes – Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.134 cont.		nat.	5.2×10^9 (rel.)	$k/k_{\text{ferro}} = 0.56$	p.r.	opt.	c.k.	71-0578
		—	5.4×10^9	—	p.r.	opt.	p.b.k. at 372 nm.	71-0578
3.135	<i>N</i> -acetylalanine, negative ion	9.2	4.6×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.042$	p.r.	opt.	c.k.	70-0099
3.136	<i>N</i> -acetylalanyl-alanylalanine, negative ion	9.0	3.0×10^9	—	p.r.	opt.	p.b.k.	75-1004
3.137	acetylene	2.15	—	$k/k_{\text{HCOOH}} = 2.1$	γ -r.	chem.	c.k.	68-0502
3.138	<i>N</i> -acetylglucosamine	—	3.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.279$	p.r.	opt.	c.k.; unpubl. data of G.O. Phillips and N. Worthington.	68-0352
3.139	<i>N</i> -acetylglycine, negative ion	8.7	4.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.038$	p.r.	opt.	c.k.	70-0099
3.140	<i>N</i> -acetylglucylglycine, negative ion	8.6	7.8×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.071$	p.r.	opt.	c.k.	70-0099
3.141	4-(2-acetylsulfamoyl)phthalanilic acid <i>See</i> thalamyd (3.700). acriflavin	—	1.2×10^{10}	—	p.r.	opt.	d.k. at 450 nm (dye) or p.b.k. at 300-400 nm.	70-0241
3.142	acrolein $\text{OH} + \text{CH}_2=\text{CHCHO} \rightarrow$ adduct	—	7.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.64$	p.r.	opt.	c.k.	70-0165
3.143	acrylamide $\text{OH} +$ $\text{CH}_2=\text{CHCONH}_2 \sim 6$ \rightarrow adduct	10.7 7 ~ 12	3.4×10^9 (rel.) 3.3×10^9 (rel.) 4.1×10^9 (rel.) 6.2×10^9 (rel.)	$k/k_{\text{BzO}^-} = 0.59$ $k/k_{\text{CNS}^-} = 0.3 \pm 0.07$ $k/k_{\text{BzO}^-} = 0.72$ $k/k_{\text{carb}} = 17$	γ -r. p.r. r. p.r.	trac. opt. lum. opt.	c.k.; meas. $^{14}\text{CO}_2$. c.k. c.k.; salicylate detd. at 405 nm. c.k. at pH 10.9 and 12.9.	65-0099 67-0171 68-0494 70-0052
		nat.	4.7×10^9 (rel.)	$k/k_{\text{ferro}} = 0.505$	p.r.	opt.	c.k.	71-0578
		—	6.8×10^9	—	p.r.	opt.	p.b.k. at 390 nm.	71-0578
			<i>For other ratios see:</i> 3.129, 3.145, 3.247, 3.283, 3.368, 3.452, 3.563.					
3.144	acrylic acid	1	1.4×10^9 (rel.)	$k/k_{\text{MeOH}} = 1.58$	Fenton	chem.	c.k.; $k_{\text{MeOH}}/k_{\text{Fe}^{2+}} = 4.3$.	73-9341
3.145	acrylonitrile	~ 0	—	$k/k_{\text{acrylamide}} = 1.8$	Fenton	pol.	c.k.	72-9162
3.146	adenine $\text{OH} + \text{C}_5\text{H}_3\text{N}_4\text{NH}_2$ \rightarrow adduct	2-2.2 5-5.5 7.3-7.5 7	8.8×10^8 (rel.) 3.8×10^9 (rel.) 5.1×10^9 (rel.) 2.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.08$ $k/k_{\text{CNS}^-} = 0.35$ $k/k_{\text{CNS}^-} = 0.46$ $k/k_{\text{CNS}^-} = (0.25 \pm 0.05)$	p.r. p.r. p.r. p.r.	opt. opt. opt. opt.	c.k. c.k. c.k. c.k.; cor. for failure of H_2O_2 to completely scavenge e_{aq}^- .	65-0388 65-0388 65-0388 68-0316
		5.7	$(5.8 \pm 0.3) \times 10^9$	—	p.r.	opt.	p.b.k. at 450 nm.	70-3069
		6-7	5.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.464$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.147	adenosine	2-2.2	1.9×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.17$	p.r.	opt.	c.k.	65-0388
		5-5.2	3.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.35$	p.r.	opt.	c.k.	65-0388
		7.6-7.8	4.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.38$	p.r.	opt.	c.k.	65-0388
3.148	adenosine 5'-phosphate (adenylic acid)	2-2.2 5.2-5.5 9 6.9 7	1.2×10^9 (rel.) 3.0×10^9 (rel.) 4.0×10^9 (rel.) $(4.7 \pm 0.5) \times 10^9$ 4.7×10^9	$k/k_{\text{CNS}^-} = 0.11$ $k/k_{\text{CNS}^-} = 0.27$ $k/k_{\text{RNO}} = 0.32$ — —	p.r. p.r. γ -r. p.r. p.r.	opt. opt. opt. opt. opt.	c.k. c.k. c.k. p.b.k. at 350 nm. p.b.k.	65-0388 65-0388 67-0551 70-3064 73-107
3.149	adipic acid	1 2-2.2	2.9×10^8 (rel.) 1.7×10^9 (rel.)	$k/k_{\text{Fe}^{2+}} = 1.25$ $k/k_{\text{thym}} = 0.32 \pm 0.03$	Fenton γ -r.	chem. opt.	c.k. c.k.	49-000 67-046

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.150	alanine, positive ion	1	2.8 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{Fe²⁺} = 0.12	Fenton	chem.	c.k.	49-0002
		2-2.2	4.8 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.0089	γ-r.	opt.	c.k.	67-0461
		1	4.4 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.0082	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 0.59.	69-5278
3.151	alanine, zwitterion	6.0	4.6 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{ferro} = 0.005	X-r.	opt.	c.k.; not cor. for H ₂ O ₂ .	62-0023
		5.5-6	7.7 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.00697	p.r.	opt.	c.k.	65-0388
		6.8	7.9 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.0063	γ-r.	opt.	c.k.	73-0548
3.152	alanine, negative ion	9.75	6.5 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{ferro} = 0.07	X-r.	opt.	c.k.; not cor. for H ₂ O ₂ .	62-0023
3.153	alanine anhydride	5.0	1.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.164	p.r.	opt.	c.k.	71-0554
		11.0	1.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.164	p.r.	opt.	c.k.	71-0554
3.154	alanylglycine, positive ion	2-2.2	1.6 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.03	γ-r.	opt.	c.k.	65-0388
3.155	ALDH (yeast alcohol dehydrogenase)	9	1.6 x 10 ¹¹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 12.9	γ-r.	opt.	c.k.	67-0555
		—	6.7 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 6.1	p.r.	opt.	c.k.	70-1226
		—	1.5 x 10 ¹¹ (rel.)	<i>k</i> / <i>k</i> _{NB} = 48	—	—	—	—
3.155a	aldolase	5.5	1.9 x 10 ¹¹	—	p.r.	opt.	p.b.k. at 330 nm; enzyme from rabbit muscle.	75-3058
3.156	allyl alcohol OH + CH ₂ CHCH ₂ OH → CH ₂ OHCHCH ₂ OH + CH ₂ CHOHCH ₂ OH	7	2.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.18	p.r.	opt.	c.k.	65-0387
		7.0	(6.0 ± 1.5) x 10 ⁹	—	p.r.	opt.	p.b.k.	73-1070
3.157	allylammonium ion	4	8.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.785	p.r.	opt.	c.k.	70-0371
3.158	<i>p</i> -aminobenzoate ion OH + NH ₂ C ₆ H ₄ COO ⁻ → NH ₂ (OH)C ₆ H ₄ COOH	9	7.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 4.3	γ-r.	opt.	c.k. with RNO.	66-0441
3.159	<i>p</i> -aminobenzoic acid OH + NH ₂ C ₆ H ₄ COOH → NH ₂ (OH)C ₆ H ₄ COOH	6-7	1.6 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.5	p.r.	opt.	c.k.	65-0387
3.160	2-aminobutyric acid	2-2.2	3.8 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.07 ± 0.005	γ-r.	opt.	c.k.	67-0461
3.161	3-aminobutyric acid	2-2.2	7.8 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.0145 ± 0.0015	γ-r.	opt.	c.k.	67-0461
3.162	4-aminobutyric acid	2-2.2	2.2 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.040 ± 0.004	γ-r.	opt.	c.k.	67-0461
3.163	2-amino-2-deoxy-D-galactose	—	1.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.106	p.r.	opt.	c.k.	70-3081
3.164	5-aminoindole	9.0	(3.17 ± 0.31) x 10 ¹⁰ (rel.)	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> (OH + tryptophan) = 1.25 x 10 ¹⁰ .	71-0556
3.165	2-aminopyridine	9	8.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.67 ± 0.12	γ-r.	opt.	c.k.	69-0280
3.166	4-aminopyridine	9	5.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.40 ± 0.01	γ-r.	opt.	c.k.	69-0280
3.167	2-aminopyrimidine	6-7	4.0 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.032	γ-r.	opt.	c.k.; 17°C.	75-0294
3.168	amyl alcohol <i>tert</i> -amyl alcohol amylamine	<i>See</i> 1-pentanol (3.602). <i>See</i> 2-methyl-2-butanol (3.526).		—	—	—	—	—
		—	7.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{ferro} = 0.85	p.r.	opt.	c.k.; calcd. from obs. values at pH 8-13.1.	73-0016
		—	9.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.82	—	—	—	—
3.169	amylammonium ion	—	5.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{NB} = 2.8	—	—	—	—
		4	9.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.89	p.r.	opt.	c.k.	70-0371
		—	5.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{ferro} = 0.6	p.r.	opt.	c.k.; calcd. from obs. values at pH 8-13.1.	73-0016
		—	4.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.43	—	—	—	—
		—	2.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{NB} = 0.87	—	—	—	—

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.170	aniline	10.7	7.1×10^9 (rel.)	$k/k_{\text{BzO}^-} = 1.24$	γ -r.	trac.	c.k.; meas. $^{14}\text{CO}_2$.	65-0099
	$\text{OH} + \text{C}_6\text{H}_5\text{NH}_2 \rightarrow$	9	8.9×10^9 (rel.)	$k/k_{\text{EtOH}} = 4.8$	γ -r.	opt.	c.k. with RNO.	66-0441
	$\text{C}_6\text{H}_5\text{NH} + \text{H}_2\text{O}$ or	7.5-9	2.9×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 2.6$	p.r.	opt.	c.k.	69-0573
	$\rightarrow \text{OHC}_6\text{H}_5\text{NH}_2$	—	2.8×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 2.58$	p.r.	opt.	c.k.	72-0289
		8,11	$(1.4 \pm 0.3) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 355 nm, (cyclohexadienyl radical), 295 nm (anilino radical), and 500 nm.	72-0289
3.171	anilinium ion	3	5.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.49$	p.r.	opt.	c.k.	69-0573
	$\text{OH} + \text{C}_6\text{H}_5\text{NH}_3^+ \rightarrow$	~ 4	$(4.8 \pm 0.8) \times 10^9$	—	p.r.	opt.	p.b.k. at 415 nm.	72-0289
3.172	$\text{C}_6\text{H}_5\text{NH}_2 + \text{H}_2\text{O}$ or							
	$\rightarrow \text{OHC}_6\text{H}_5\text{NH}_3^+$							
3.172	anisole	9	6.0×10^9 (rel.)	$k/k_{\text{EtOH}} = 3.27$	γ -r.	opt.	c.k. with RNO.	66-0441
	$\text{OH} + \text{C}_6\text{H}_5\text{OCH}_3 \rightarrow$	7	$(12 \pm 3) \times 10^9$	—	p.r.	opt.	p.b.k. at 330 nm; cor. for (OH + OH) and (H + aromatic).	68-0304
	$(\text{OH})\text{C}_6\text{H}_5\text{OCH}_3$							
		9	5.7×10^9 (rel.)	$k/k_{\text{RNO}} = 0.45 \pm 0.04$	γ -r.	opt.	c.k.	69-0280
		—	$(5.4 \pm 0.5) \times 10^9$	—	p.r.	—	—	75-1171
3.173	anthranilic acid	—	1.1×10^{10}	—	p.r.	opt.	p.b.k.	74-1063
3.174	9,10-anthra-quinone-1-sulfonate ion	—	7.2×10^9	—	p.r.	opt.	p.b.k. ~ 460 nm; OH addn.	72-0391
3.175	9,10-anthra-quinone-2-sulfonate ion	—	5.6×10^9	—	p.r.	opt.	p.b.k. ~ 460 nm; OH addn.	72-0391
3.176	9-anthroate ion	9	8.0×10^9	—	p.r.	opt.	p.b.k.	73-0110
3.177	arginine	2-2.2	7.8×10^8 (rel.)	$k/k_{\text{thym}} = 0.145$	γ -r.	opt.	c.k.	65-0388
		2-2.2	6.7×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.061$	p.r.	opt.	c.k.	65-0388
		6.5-	3.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.32$	p.r.	opt.	c.k.	65-0388
		7.5						
		6.7	5.7×10^8 (rel.)	$k/k_{\text{RNO}} = 0.045$	γ -r.	opt.	c.k.	73-0548
3.178	ascorbate ion	7	1.3×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.17$	p.r.	opt.	c.k.	72-0266
	(H abstr.)	7	$(1.1 \pm 0.3) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 360 nm; also detd.	73-3006
							$k/k_{\text{phenylalanine}} = 1.0 \pm 0.05$.	
3.179	ascorbic acid	1	1.2×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.1$	p.r.	opt.	c.k.	65-0387
		1.5	8.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.75$	p.r.	opt.	c.k.	72-0266
3.180	asparagine	2-2.2	3.2×10^7 (rel.)	$k/k_{\text{thym}} = (6.0 \pm 0.5) \times 10^{-3}$	γ -r.	opt.	c.k.	67-0461
		6.6	4.9×10^7 (rel.)	$k/k_{\text{RNO}} = 0.0039$	γ -r.	opt.	c.k.	73-0548
3.181	aspartic acid	2-2.2	3.3×10^7 (rel.)	$k/k_{\text{thym}} = 0.0061$	γ -r.	opt.	c.k.	65-0388
		6.8-7	7.5×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.0068$	p.r.	opt.	c.k.	65-0388
		6.5	4.9×10^7 (rel.)	$k/k_{\text{RNO}} = 0.0039$	γ -r.	opt.	c.k.	73-0548
3.182	azelaic acid	2-2.2	4.6×10^9 (rel.)	$k/k_{\text{thym}} = 0.85 \pm 0.10$	γ -r.	opt.	c.k.	67-0461
3.183	Bacteriophage T ₇	—	$\sim 5 \times 10^9$ (rel.)	—	γ -r.	opt.	c.k. with ferro-cyanide; obs. G(ferri).	70-3048
3.184	benzaldehyde	9	4.4×10^9 (rel.)	$k/k_{\text{RNO}} = 0.35 \pm 0.03$	γ -r.	opt.	c.k.	69-0280
3.185	benzamide	1	1.5×10^9 (rel.)	$k/k_{\text{Fe}^{2+}} = 6.6$	Fenton	chem.	c.k.	49-0003
		9	4.4×10^9 (rel.)	$k/k_{\text{EtOH}} = 2.4$	γ -r.	opt.	c.k. with RNO.	66-0441

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.186	benzene OH + C ₆ H ₆ → C ₆ H ₆ OH	1	7.4 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{Fe2+} = 3.2	Fenton	chem.	c.k.	49-0003
		—	(4.3 ± 0.9) x 10 ⁹	—	p.r.	opt.	p.b.k. at 313 nm.	62-0020
		3	(3.3 ± 0.8) x 10 ⁹	—	p.r.	opt.	p.b.k. at 313 nm.	64-0115
		~7	3.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{I⁻} = 0.31 ± 0.03	p.r.	opt.	c.k.; obs. I ₂ ⁻ at 400 nm.	65-0010
		10.5	6.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{BzO⁻} = 1.2	γ-r.	trac.	c.k.; meas. ¹⁴ CO ₂ .	65-0099
		3	6.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{BzOH} ≅ 1.1	γ-r.	trac.	c.k.; meas. ¹⁴ CO ₂ .	65-0099
		6-7	5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.455	p.r.	opt.	c.k.	65-0387
		~1	2.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{Fe2+} = 6.7	γ-r.	chem.	c.k.	66-0645
		2-2.2	5.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.00 ± 0.08	γ-r.	opt.	c.k.	65-0388, 67-0461
		9	3.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.26	γ-r.	opt.	c.k.	67-0555
		7	(7.8 ± 1.1) x 10 ⁹	—	p.r.	opt.	p.b.k. at 313 nm; cor. for (OH + OH) and (H + aromatic).	68-0304
		~1.2	4.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 2.2	γ-r.	chem.	c.k.	68-0602
		6.98	5.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 2.3	γ-r.	chem.	c.k.	68-0602
		9	4.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.35	γ-r.	opt.	c.k.	69-0280
		7.0	7.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{PNBA⁻} = 2.9	p.r.	opt.	c.k.; formn. of PNBA ⁻ - OH adduct at 415 nm.	70-0211
3.187	benzene- <i>d</i> ₆ OH + C ₆ D ₆ → C ₆ D ₆ OH	—	8.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.745	p.r.	opt.	c.k.; <i>k</i> lowered in presence of surfactants.	71-0001, 71-0586
		1.7-1.8	4.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 2.0	Fenton	chem.	c.k.	74-9006
		—	For other ratios see: 3.12, 3.52, 3.66, 3.78, 3.106, 3.667, 3.668, 3.711.					
3.188	benzenesulfonamide	9	2.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 1.5	γ-r.	opt.	c.k. with RNO.	66-0441
		—	2.8 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO, assuming <i>k</i> (OH + sulfanilic acid) = 2.93 x 10 ⁹ .	73-0094
3.189	benzenesulfonate ion OH + C ₆ H ₅ SO ₃ ⁻ → OHC ₆ H ₅ SO ₃ ⁻	9	3.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 1.6	γ-r.	opt.	c.k. with RNO.	66-0441
		7	(4.7 ± 0.6) x 10 ⁹	—	p.r.	opt.	p.b.k. at 315 nm; cor. for (OH + OH) and (H + aromatic).	68-0304
3.190	benzenesulfonic acid	1	1.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{Fe2+} = 4.7	Fenton	chem.	c.k.	49-0003
3.191	benzoate ion OH + C ₆ H ₅ COO ⁻ → OHC ₆ H ₅ COO ⁻	6-7	5.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.5	p.r.	opt.	c.k.	65-0387
		9	4.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.3	γ-r.	opt.	c.k. with RNO.	66-0441
		6-9.4	(6.0 ± 0.7) x 10 ⁹	—	p.r.	opt.	p.b.k. at 330 nm; cor. for (OH + OH) and (H + aromatic).	68-0304
		5.5	6.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{NB} = 2.0	r.	opt.	c.k.; obs. formn. of <i>o</i> -nitrophenol.	68-0494
		9	4.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.37 ± 0.01	γ-r.	opt.	c.k.	69-0280
		—	3.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.29	p.r.	opt.	c.k.	71-0282
		nat.	5.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{ferro} = 0.581	p.r.	opt.	c.k.	71-0578
		—	5.7 x 10 ⁹	—	p.r.	opt.	p.b.k. at 330 nm.	71-0578
		—	2.5 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO assuming <i>k</i> (OH + sulfanilic acid) = 2.93 x 10 ⁹ .	73-0094
		—	For other ratios see: 3.12, 3.21, 3.54, 3.66, 3.73, 3.128, 3.131, 3.143, 3.170, 3.186, 3.192, 3.193, 3.248, 3.358, 3.384, 3.406, 3.511, 3.565, 3.607, 3.608, 3.669.					
3.192	benzoic acid OH + C ₆ H ₅ COOH → HOC ₆ H ₅ COOH	1	1.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{Fe2+} = 7.0	Fenton	chem.	c.k.	49-0003
		3	(2.1 ± 0.3) x 10 ⁹	—	p.r.	opt.	p.b.k. at 340-350 nm.	64-0115

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.192 cont.		3	5.7×10^9 (rel.)	$k/k_{\text{BzO}^-} = 1$	γ -r.	trac.	c.k.; meas. $G(^{14}\text{CO}_2)$.	65-0099
		≤ 3	$(4.3 \pm 0.8) \times 10^9$	—	p.r.	opt.	p.b.k. at 340 nm; cor. for (H + BzOH) and (OH + OH).	68-0229
3.193	benzonitrile $\text{OH} + \text{C}_6\text{H}_5\text{CN} \rightarrow$ $\text{OHC}_6\text{H}_5\text{CN}$	10.7	3.4×10^9 (rel.)	$k/k_{\text{BzO}^-} = 0.59$	γ -r.	trac.	c.k.; meas. $G(^{14}\text{CO}_2)$.	65-0099
		9	3.6×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.96$	γ -r.	opt.	c.k. with RNO.	66-0441
		7	$(4.9 \pm 0.6) \times 10^9$	—	p.r.	opt.	p.b.k. at 348 nm; cor. for (OH + OH) and (H + aromatic).	68-0304
		6.3	8.5×10^9	—	p.r.	opt.	p.b.k.	70-0657
		—	3.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.35$	p.r.	opt.	c.k.	70-0657
3.194	benzophenone $\text{OH} + \text{C}_6\text{H}_5\text{COC}_6\text{H}_5 \rightarrow$ $\text{C}_6\text{H}_5(\text{OH})\text{COC}_6\text{H}_5$	—	$(8.7 \pm 1.5) \times 10^9$	—	p.r.	opt.	p.b.k. at 380 nm.	68-0727
		—	$(9 \pm 2) \times 10^9$	—	p.r.	opt.	p.b.k. at 330 nm.	75-1125
3.195	benzoquinone	—	1.2×10^9	—	p.r.	opt.	p.b.k.(OH adduct)	67-0121
3.196	benzyl alcohol $\text{OH} + \text{C}_6\text{H}_5\text{CH}_2\text{OH} \rightarrow$ $\text{OHC}_6\text{H}_5\text{CH}_2\text{OH}$	7	$(8.4 \pm 1.2) \times 10^9$	—	p.r.	opt.	p.b.k. at 320 nm; cor. for (OH + OH) and (H + aromatic).	68-0304
		9	4.6×10^9 (rel.)	$k/k_{\text{RNO}} = 0.37 \pm 0.02$	γ -r.	opt.	c.k.	69-0280
3.197	benzylammonium ion	4	1.2×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.14$	p.r.	opt.	c.k.	70-0371
3.198	β -benzylglucoside	~ 7	4.2×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 3.79$	p.r.	opt.	c.k.	71-0480
3.198a	benzyl methyl ether	1.7-1.8	1.2×10^{10} (rel.)	$k/k_{2-\text{PrOH}} = 5.5$	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.199	benzylpenicillin	—	7.1×10^9 (rel.)	$k/k_{\text{PA}^-} = 0.9$	γ -r.	opt.	c.k. with RNO.	73-0134
3.200	benzylpenicilloic acid	—	7.1×10^9 (rel.)	$k/k_{\text{PA}^-} = 0.9$	γ -r.	opt.	c.k. with RNO.	73-0134
3.201	benzyltrimethylammonium ion	5.0	6.8×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 3.1$	r.	chem.	c.k.	68-0205
3.202	biacetyl $\text{OH} + \text{CH}_3\text{COCOCH}_3 \rightarrow$ $\text{H}_2\text{O} + \text{CH}_2\text{COCOCH}_3$	—	1.6×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.015$	p.r.	opt.	c.k.	68-0249
3.202a	biphenyl	—	$(9.0 \pm 1.0) \times 10^9$	—	p.r.	opt.	p.b.k.	75-1096
3.203	4-biphenylcarboxylate ion	9	6.8×10^9	—	p.r.	opt.	p.b.k.	73-0110
3.204	2,2'-biphenyldicarboxylate ion (diphenate ion)	9	7.0×10^9	—	p.r.	opt.	p.b.k.	73-0110
3.205	4,4'-biphenyldicarboxylate ion	9	8.3×10^9	—	p.r.	opt.	p.b.k.	73-0110
3.206	2,2'-bipyridine	9.3	6.2×10^9	—	p.r.	opt.	p.b.k.	71-0582
3.207	4,4'-bipyridine	9.3	5.3×10^9	—	p.r.	opt.	p.b.k.	71-0582
3.208	bromoacetate ion	9	4.4×10^7 (rel.)	$k/k_{\text{EtOH}} = 0.024$	γ -r.	opt.	c.k. with RNO.	66-0423
3.209	<i>p</i> -bromobenzoate ion	9	3.1×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.68$	γ -r.	opt.	c.k. with RNO.	66-0441
3.210	2-bromoethanol	—	7.8×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.42$	γ -r.	opt.	c.k. with RNO.	67-0050
3.211	5-bromoindole	9.0	$(1.57 \pm 0.18) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.212	5-bromoorotate ion	7	3×10^9	—	p.r.	opt.	p.b.k. at 340 nm.	73-0002
		7	6.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.6$	p.r.	opt.	c.k.	73-0002
3.212a	1-(<i>p</i> -bromophenyl)ethanol	1.7-1.8	7.0×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 3.2$	Fenton	chem.	c.k.	74-9006
3.213	<i>m</i> -bromophenyl- β -D-glucopyranoside	—	3.2×10^9 (rel.)	—	γ -r.	opt.	c.k. with RNO; $k(\text{OH} + \text{X}) = 4.4 \times 10^9$ (X = phenyl- β -D-glucopyranoside) as standard.	71-0056

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.214	2-bromopropionate ion 8.5	2.2×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.12$	γ -r.	opt.	c.k. with RNO.	67-0050
3.215	3-bromopropionate ion 8.5	2.2×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.12$	γ -r.	opt.	c.k. with RNO.	67-0050
3.216	2-bromopyridine 9	2.4×10^9 (rel.)	$k/k_{\text{RNO}} = 0.19 \pm 0.01$	γ -r.	opt.	c.k.	69-0280
3.217	3-bromopyridine 9	1.1×10^9 (rel.)	$k/k_{\text{RNO}} = 0.09 \pm 0.01$	γ -r.	opt.	c.k.	69-0280
3.218	α -bromotetrate ion 7 $\text{OH} + \text{C}_4\text{H}_2\text{BrO}_3^- \rightarrow \text{HBr} + \text{C}_4\text{H}_2\text{O}_4^-$	7.7×10^9	—	p.r.	opt.	d.k. at 258 nm as well as p.b.k. at 360 nm.	74-1053
3.219	5-bromouracil (BU) 9	4.2×10^9 (rel.)	$k/k_{\text{RNO}} = 0.34$	r.	opt.	c.k.	67-0555
	$\text{OH} + \text{BU} \rightarrow \text{BUOH}$ 7.0	3.6×10^9	—	p.r.	opt.	p.b.k. at 335 nm; complex kinetics.	69-0558
	7	4.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.364$	p.r.	opt.	c.k.	72-0049
	7	5.6×10^9	—	p.r.	opt.	p.b.k. at 340 nm.	72-0049
3.220	1,3-butadiene —	11	5.8×10^9	p.r.	opt.	p.b.k. at 340 nm.	72-0049
		7.7 $\times 10^9$ (rel.)	$k/k_1 = 0.64$	p.r.	opt.	c.k.; obs. formn. of I_2^- at 400 nm.	67-0041
		—	—	—	—	—	—
3.221	1,2-butanediol 1	4.4×10^6 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.019$	therm.	chem.	c.k.; persulfate oxidation.	49-0002
3.222	1,3-butanediol 7	2.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.197$	p.r.	opt.	c.k.	65-0387
		2.4×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.3$	γ -r.	opt.	c.k. with RNO.	66-0423
3.223	1,4-butanediol 7	3.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.29$	p.r.	opt.	c.k.	65-0387
		3.0×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.6$	γ -r.	opt.	c.k. with RNO.	66-0423
3.224	2,3-butanediol 1	2.3×10^6 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.010$	therm.	chem.	c.k.; persulfate oxidation.	49-0002
	(I) $\text{OH} + (\text{MeCHOH})_2 \rightarrow \text{MeCHOHCHOHMe} + \text{H}_2\text{O}$ 7	1.3×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.12$	p.r.	opt.	c.k.	65-0387
	(II) $\text{OH} + (\text{MeCHOH})_2 \rightarrow \text{CH}_2\text{CHOHCHOHMe} + \text{H}_2\text{O}$ —	—	$k_{\text{II}}/k_1 = 0.41$	p.r.	opt.	detd. % α -alcohol radical by reaction with TNM; $\leq 0.1\%$ alkoxy radical detd. by reaction with I^-	73-0126
3.225	1-butanol 2-2.2	4.6×10^9 (rel.)	$k/k_{\text{thym}} = 0.85 \pm 0.08$	γ -r.	opt.	c.k.	67-0461
	(I) $\text{OH} + \text{C}_3\text{H}_7\text{CH}_2\text{OH} \rightarrow \text{H}_2\text{O} + \text{C}_3\text{H}_7\text{CHOH}$ 5-5.5	4.6×10^9 (rel.)	$k/k_{\text{thym}} = 0.85 \pm 0.10$	γ -r.	opt.	c.k.	67-0461
	(34%, 69-0522) 7	4.0×10^9 (rel.)	$k/k_{\text{carb}} = 11$	p.r.	opt.	c.k.	65-0190
	(II) $\text{OH} + \text{C}_3\text{H}_7\text{CH}_2\text{OH} \rightarrow \text{H}_2\text{O} + \text{CH}_3\text{CH}_2\text{CHCH}_2\text{OH}$ nat. 7	3.7×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.34$	p.r.	opt.	c.k.	65-0387
	$\rightarrow \text{H}_2\text{O} + \text{CH}_3\text{CH}_2\text{CHCH}_2\text{OH}$ 9	3.7×10^9 (rel.)	$k/k_{\text{EtOH}} = 2$	γ -r.	opt.	c.k. with RNO.	66-0423
	$\text{CH}_3\text{CH}_2\text{CHCH}_2\text{OH}$ nat. —	4.0×10^9 (rel.)	$k/k_{\text{ferro}} = 0.43$	p.r.	opt.	c.k.	71-0578
	$+ \text{CH}_3\text{CHCH}_2\text{CH}_2\text{OH}$ —	—	$k_{\text{III}}/k_1 \leq 0.1$	p.r.	opt.	detd. % α -alcohol and alkoxy radicals by reaction with TNM and I^- , resp.	73-0126
	$+ \text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{OH}$ —	—	$k_{\text{II}}/k_1 = 1.4$	—	—	—	—
	(III) $\text{OH} + \text{C}_3\text{H}_7\text{CH}_2\text{OH} \rightarrow \text{H}_2\text{O} + \text{C}_3\text{H}_7\text{CH}_2\text{O}$ —	6×10^8 (rel.)	$k/k_{\text{MeOH}} = 0.62$	Ti(III) + H_2O_2	esr	c.k.	73-5253
	For other ratios see:		3.310.	—	—	—	—
3.226	2-butanol 2-2.2	2.7×10^9 (rel.)	$k/k_{\text{thym}} = 0.50 \pm 0.05$	γ -r.	opt.	c.k.	67-0461
	$\text{OH} + \text{C}_2\text{H}_5\text{CHOHCH}_3 \rightarrow \text{H}_2\text{O} + \text{C}_2\text{H}_5\text{COHCH}_3$ 7	3.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.28$	p.r.	opt.	c.k.	65-0387
	$\text{C}_2\text{H}_5\text{COHCH}_3$ 9	2.4×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.3$	γ -r.	opt.	c.k. with RNO.	66-0423
	(53%, 69-0522) + $\text{CH}_3\text{CHCHOHCH}_3$, etc. —	—	—	—	—	—	—
3.227	<i>tert</i> -butanol —	See 2-methyl-2-propanol (3.546).		—	—	—	—
	2-butanone 6-7	9.0×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.082$	p.r.	opt.	c.k.	65-0387

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.228	1-butene —	7.7×10^9 (rel.)	$k/k_1^- = 0.64$	p.r.	opt.	c.k.; obs. formn. of I_2^- at 400 nm.	67-0041
3.229	1-butene-3-one —	8.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.77$	p.r.	opt.	c.k.	70-0165
3.230	<i>N-tert</i> -butylacetamide 5-6	1.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.103 \pm 0.01$	p.r.	opt.	c.k.	71-0414
3.231	<i>tert</i> -butyl alcohol butylamine —	See 2-methyl-2-propanol (3.546).		p.r.	opt.	c.k.; calcd. from obs. values at pH 8-13.1.	73-0016
		7.3×10^9 (rel.)	$k/k_{\text{ferro}} = 0.79$				
		8.3×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.75$				
		5.4×10^9 (rel.)	$k/k_{\text{NB}} = 1.7$				
3.232	<i>tert</i> -butylamine 12 $\text{OH} + (\text{CH}_3)_3\text{CNH}_2 \rightarrow \text{H}_2\text{O} + \bullet\text{CH}_2(\text{CH}_3)_2\text{CNH}_2 + (\text{CH}_3)_3\text{CNH}$	6.0×10^9 (rel.)	—	p.r.	opt.	c.k., extrapolated value based on $k/k_{\text{CNS}^-} = 3.64 \times 10^{-1}$ (obs.) at pH 10.9.	71-0585
3.233	butylammonium ion 4	5.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.5$	p.r.	opt.	c.k.	70-0371
	—	2.8×10^9 (rel.)	$k/k_{\text{ferro}} = 0.3$	p.r.	opt.	c.k.; calcd. from obs. values at pH 8-13.1.	73-0016
		3.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.28$				
		2.05×10^9 (rel.)	$k/k_{\text{NB}} = 0.64$				
3.234	<i>tert</i> -butylammonium ion 4	2.4×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.022$	p.r.	opt.	c.k.	70-0371
	3.2	7.0×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0636$	p.r.	opt.	c.k.	71-0585
	$\text{OH} + (\text{CH}_3)_3\text{CNH}_3^+ \rightarrow \text{H}_2\text{O} + \bullet\text{CH}_2(\text{CH}_3)_2\text{CNH}_3^+$						
3.235	butyleneoxide-1,2 <i>tert</i> -butyl mercaptan 7	See 1,2-epoxybutane (3.352).		p.r.	opt.	c.k.	69-0553
		1.9×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.7$				
	$\text{OH} + (\text{CH}_3)_3\text{CSH} \rightarrow \text{H}_2\text{O} + (\text{CH}_3)_3\text{CS}$						
3.236	<i>p-tert</i> -butylphenol 9	1.9×10^{10} (rel.)	$k/k_{\text{RNO}} = 1.49 \pm 0.26$	γ -r.	opt.	c.k.	72-0837
3.237	butyraldehyde 2.0	3.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.35$	p.r.	opt.	c.k.	65-0387
3.238	butyrate ion 9	1.85×10^9 (rel.)	$k/k_{\text{EtOH}} = 1$	γ -r.	opt.	c.k. with RNO.	66-0423
3.239	butyric acid 1	1.6×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.72$	Fenton	chem.	c.k.	49-0002
	2-2.2	1.9×10^9 (rel.)	$k/k_{\text{thym}} = 0.35 \pm 0.03$	γ -r.	opt.	c.k.	67-0461
3.240	carbon disulfide 7.6	8.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.73$	p.r.	opt.	c.k.; meas. abs. increase at 280 nm (CS_2OH) or at 500 nm.	67-0687, 73-1015
	$\text{OH} + \text{CS}_2 \rightarrow \text{CS}_2\text{OH} \rightleftharpoons \text{CSO}^- + \text{H}^+$						
3.241	carboxymethyl-cellulose (polyanion) —	2.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.24$	p.r.	opt.	c.k.; concn. in monomer units.	68-0352
3.242	carboxypeptidase A 7.8	$(6.9 \pm 1.0) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 320 nm; contains ~ 15% H reaction product.	73-1060
3.243	catalase —	1.4×10^{11} (rel.)	$k/k_{\text{CNS}^-} = 12.58$	p.r.	opt.	c.k.; mol. wt. $\cong 2.5 \times 10^5$.	66-0499
3.244	cellobiose 6.5	3.6×10^9 (rel.)	$k/k_{\text{RNO}} = 0.29 \pm 0.01$	γ -r.	opt.	c.k.	69-0580
3.245	chloral hydrate 1	1.1×10^9 (rel.)	$k/k_{\text{Fe}^{2+}} = 4.7$	Fenton	chem.	c.k.	49-0002
	—	3.15×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.285$	p.r.	opt.	c.k.	73-0062
	$\text{OH} + \text{CCl}_3\text{CH}(\text{OH})_2 \rightarrow \text{H}_2\text{O} + \text{CCl}_3\text{C}(\text{OH})_2$						
3.246	chloroacetate ion 9	5.5×10^7 (rel.)	$k/k_{\text{EtOH}} = 0.0296$	γ -r.	opt.	c.k. with RNO.	66-0423
	(I) $\text{OH} + \text{ClCH}_2\text{COO}^-$ 6.5	6×10^9 (I) (rel.)	$k_1/k_{\text{II}} \cong 4.0$	γ -r.	chem.	c.k.	69-0422
	$\rightarrow \text{H}_2\text{O} + \text{ClCHCOO}^-$	1.5×10^8 (II) (rel.)	$k_{\text{II}}/k_{\text{MeOH}} \cong 0.08$				
	(II) $\text{OH} + \text{ClCH}_2\text{COO}^- \rightarrow \text{Cl}^- + \text{products}$						

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH _c	k	Ratio	Source	Method	Comment	Ref.
3.247	chloroacetic acid	2-2.2	8.1 x 10 ⁷ (rel.)	$k/k_{\text{thym}} = 0.015 \pm 0.0015$	γ -r.	opt.	c.k.	67-0461
		1	4.3 x 10 ⁷ (rel.)	$k/k_{\text{CNS}^-} = 0.00394$	p.r.	opt.	c.k.	65-0387
		~0	—	$k/k_{\text{acrylamide}} = 0.012$	Fenton	pol.	c.k.	72-9162
3.248	chlorobenzene	10.7	6.3 x 10 ⁹ (rel.)	$k/k_{\text{BzO}^-} = 1.10$	γ -r.	trac.	c.k.; meas. ¹⁴ CO ₂ .	65-0099
		9	4.5 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.36 \pm 0.05$	γ -r.	opt.	c.k.	69-0280
3.248a	<i>o</i> -chlorobenzoate ion	7	6.8 x 10 ⁹ (rel.)	$k/k_{2-\text{PrOH}} = 3.1$	γ -r.	chem.	c.k.	74-0167
3.248b	<i>m</i> -chlorobenzoate ion	7	6.3 x 10 ⁹ (rel.)	$k/k_{2-\text{PrOH}} = 2.8$	γ -r.	chem.	c.k.	74-0167
3.249	<i>p</i> -chlorobenzoate ion	9	3.2 x 10 ⁹ (rel.)	$k/k_{\text{EtOH}} = 1.75$	γ -r.	opt.	c.k. with RNO.	66-0441
	OH + ClC ₆ H ₄ COO ⁻ → Cl(OH)C ₆ H ₄ COO ⁻	6-9.4	(5.0 ± 0.8) x 10 ⁹	—	p.r.	opt.	p.b.k. at 345 nm; cor. for (OH + OH) and (H + aromatic).	68-0304
		7	7.3 x 10 ⁹ (rel.)	$k/k_{2-\text{PrOH}} = 3.3$	γ -r.	chem.	c.k.	74-0167
3.250	2-chloroethanol	—	9.25 x 10 ⁸ (rel.)	$k/k_{\text{EtOH}} = 0.5$	γ -r.	opt.	c.k. with RNO.	67-0050
3.251	chloroform	0.4	1.0 x 10 ⁷ (rel.)	$k/k_{\text{Fe}^{2+}} = 0.043$	β -r.	chem.	c.k.	60-0016
	OH + CHCl ₃ → H ₂ O + CCl ₃	0.4	5.3 x 10 ⁶ (rel.)	$k/k_{\text{Fe}^{2+}} = 0.023$	Fenton	chem.	c.k.	60-0016
		9	1.4 x 10 ⁷ (rel.)	$k/k_{\text{EtOH}} = 0.0077$	γ -r.	opt.	c.k. with RNO.	66-0423
		—	7.4 x 10 ⁶ (rel.)	—	r.	chem.	c.k. with Fe ²⁺ .	66-9002
		5.5-5.8	~ 5 x 10 ⁶ (rel.)	—	γ -r.	chem.	est. from effect of Fe ²⁺ on G(Cl ⁻).	70-0013
3.252	5-chloroindole	9.0	(1.91 ± 0.04) x 10 ¹⁰ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.253	<i>m</i> -chlorophenol	9	7.2 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.58 \pm 0.05$	γ -r.	opt.	c.k.	72-0837
3.254	<i>o</i> -chlorophenol	9	8.2 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.66 \pm 0.12$	γ -r.	opt.	c.k.	72-0837
3.255	<i>m</i> -chlorophenyl- β -D-glucopyranoside	—	3.2 x 10 ⁹ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to phenyl- β -D-glucopyranoside (X); $k(\text{OH} + \text{X}) = 4.4 \times 10^9$.	71-0056
3.256	<i>p</i> -chlorophenyl- β -D-glucopyranoside	—	3.4 x 10 ⁹ (rel.)	—	γ -r.	opt.	c.k. with RNO; relative to phenyl- β -D-glucopyranoside (X); $k(\text{OH} + \text{X}) = 4.4 \times 10^9$.	71-0056
		—	5.7 x 10 ⁹ (rel.)	$k/k_{\text{CNS}^-} = 0.515$	p.r.	opt.	c.k.	71-0056
3.257	2-chloropropionate ion	8.5	2.4 x 10 ⁸ (rel.)	$k/k_{\text{EtOH}} = 0.13$	γ -r.	opt.	c.k. with RNO.	67-0050
3.258	3-chloropropionate ion	8.5	3.1 x 10 ⁸ (rel.)	$k/k_{\text{EtOH}} = 0.17$	γ -r.	opt.	c.k. with RNO.	67-0050
3.259	2-chloropyridine	9	1.75 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.14 \pm 0.01$	γ -r.	opt.	c.k.	69-0280
3.260	4-chloropyridine	9	3.1 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.25 \pm 0.01$	γ -r.	opt.	c.k.	69-0280
3.261	chlorotrifluoromethane	—	~ 5 x 10 ⁸ (rel.)	—	—	—	c.k. with BzO ⁻ ; cited from unpubl. data.	70-0407
3.262	5-chlorouracil	7	5.2 x 10 ⁹ (rel.)	$k/k_{\text{CNS}^-} = 0.472$	p.r.	opt.	c.k.	72-0049
		7	5.5 x 10 ⁹	—	p.r.	opt.	p.b.k. at 340 nm.	72-0049
		11	5.8 x 10 ⁹	—	p.r.	opt.	p.b.k. at 340 nm.	72-0049
3.263	chondroitin 4-sulfate 1	—	8.0 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.073$	p.r.	opt.	c.k.; concn. of polyanion in hexose units.	70-3081
3.264	chondroitin 6-sulfate 1	—	6.8 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.062$	p.r.	opt.	c.k.; concn. in hexose units.	70-3081

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.265	α -chymotrypsin	6.6	3.5×10^{10} (rel.)	$k/k_{\text{RNO}} = 2.8$	γ -r.	opt.	c.k.	73-0548
3.266	citric acid	2-2.2	5.4×10^8 (rel.)	$k/k_{\text{thym}} = 0.10 \pm 0.051$	γ -r.	opt.	c.k.	67-0461
3.267	collagen	1 —	5.0×10^7 (rel.) 4.0×10^{11} (rel.)	$k/k_{\text{CNS}^-} = 0.00455$ —	p.r. p.r.	opt. opt.	c.k. c.k. with CNS^- ; reference rate not given; mol. wt. 360,000.	65-0387 68-3007
3.268	<i>o</i> -cresol	9	1.1×10^{10} (rel.)	$k/k_{\text{RNO}} = 0.90 \pm 0.15$	γ -r.	opt.	c.k.	72-0837
3.269	<i>p</i> -cresol	9	1.3×10^{10} (rel.)	$k/k_{\text{RNO}} = 1.04 \pm 0.09$	γ -r.	opt.	c.k.	72-0837
		5.5	$(1.2 \pm 0.2) \times 10^{10}$ (rel.)	$k/k_{\text{CNS}^-} = 1.1$	p.r.	opt.	c.k.	73-0003
3.270	crotonaldehyde	—	5.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.53$	p.r.	opt.	c.k.	70-0165
3.271	crotonic acid	1	2.7×10^9 (rel.)	$k/k_{\text{MeOH}} = 2.96$	Fenton	chem.	c.k.; $k_{\text{MeOH}}/k_{\text{Fe}^{2+}} = 4.3$.	73-9341
3.272	cynoacetate ion	9	1.6×10^7 (rel.)	$k/k_{\text{EtOH}} = 0.0084$	γ -r.	opt.	c.k. with RNO.	66-0423
3.272a	cyanocobalamin	—	6.5×10^9	—	p.r.	opt.	p.b.k. at 310-330 nm.	74-1105
3.273	5-cyanoindole	9.0	$(1.06 \pm 0.24) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.274	1-(<i>p</i> -cyano-phenyl)-ethanol	1.7-1.8	3.3×10^9 (rel.)	$k/k_{\text{RNO}} = 1.3$	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.275	<i>p</i> -cyanophenyl- β -D-glucopyranoside	—	3.5×10^9 (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{X}) = 4.4 \times 10^9$, X = phenyl- β -D-glucopyranoside.	71-0056
3.276	cyclobutane-carboxylate ion	9	3.0×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.6$	γ -r.	opt.	c.k. with RNO.	66-0423
3.277	cycloheptanol	1.7-1.8	2.0×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 0.91$	Fenton	chem.	c.k.	74-9006
3.278	cycloheptanol-1- <i>d</i>	1.7-1.8	1.5×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 0.70$	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.279	cycloheptatriene	—	$(7 \pm 2) \times 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS^- ; reference rate not given.	71-0710
3.280	1,3-cyclohexadiene $\text{OH} + \text{C}_6\text{H}_8 \rightarrow \text{H}_2\text{O} + \text{C}_6\text{H}_7$ and $\text{C}_6\text{H}_8\text{OH}$	7.0	1×10^{10} (rel.)	$k/k_{\text{PNBA}^-} = 3.8$	p.r.	opt.	c.k.; formn. of PNBA^- -OH adduct at 415 nm; 30% H abstraction.	70-0211
3.281	1,4-cyclohexadiene $\text{OH} + \text{C}_6\text{H}_8 \rightarrow \text{H}_2\text{O} + \text{C}_6\text{H}_7$ and $\text{C}_6\text{H}_8\text{OH}$	7.0	7.7×10^9 (rel.)	$k/k_{\text{PNBA}^-} = 2.96$	p.r.	opt.	c.k.; formn. of PNBA^- -OH adduct at 415 nm; 45% H abstraction.	70-0211
3.282	cyclohexanecarboxylate ion	9	5.4×10^9 (rel.)	$k/k_{\text{EtOH}} = 2.9$	γ -r.	opt.	c.k. with RNO.	66-0423
3.283	<i>trans</i> -1,2-cyclohexanediamine-tetraacetic acid	~ 0	—	$k/k_{\text{acrylamide}} = 2.0$	Fenton	pol.	c.k.	72-9162
3.284	cyclohexene $\text{OH} + \text{C}_6\text{H}_{10} \rightarrow \text{H}_2\text{O} + \text{C}_6\text{H}_9$	7.0	8.8×10^9 (rel.)	$k/k_{\text{PNBA}^-} = 3.4$	p.r.	opt.	c.k.; formn. of PNBA^- -OH adduct at 415 nm.	70-0211

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.285	cyclohexylammonium ion	4	1.0 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.96	p.r.	opt.	c.k.	70-0371
3.286	cyclopentane OH + C ₅ H ₁₀ → H ₂ O + C ₅ H ₉	—	3.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{PNBA⁻} = 1.15	p.r.	opt.	c.k.	74-1052
3.287	cyclopentanecarboxylate ion	9	4.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.2	γ-r.	opt.	c.k. with RNO.	66-0423
3.288	cyclopentene	—	7.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.64	p.r.	opt.	c.k.	74-1052
3.289	cysteamine (cyst)	1.4	1.6 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.44	p.r.	opt.	c.k.	67-0554
	OH +	6.5, 9	1.4 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.29	p.r.	opt.	c.k.	67-0554
	NH ₂ CH ₂ CH ₂ SH → NH ₂ CH ₂ CH ₂ S + H ₂ O	1	1.9 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{thym} = 3.5	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 251.	69-5278
		—	—	<i>k</i> / <i>k</i> _{uracil} = 3.45	p.r.	esr	c.k.	72-3003
			<i>For other ratios see: 3.627.</i>					
3.290	cysteine	1	1.3 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.2	p.r.	opt.	c.k.	65-0387
		2-2.2	5.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.10 ± 0.10	γ-r.	opt.	c.k.	67-0461
		—	<i>ca.</i> 3.4 x 10 ⁹	—	p.r.	opt.	p.b.k.	69-0638
		1	8.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.53	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 112.	69-5278
		7	4.0 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{thym} = 7.42	p.r.	esr	c.k.	72-3003
		7	—	<i>k</i> / <i>k</i> _{uracil} = 6.21	p.r.	esr	c.k.	72-3003
		0.4	1.7 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.5	p.r.	opt.	c.k.; ± 15%;	73-0090
		5.8	1.9 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.7			not cor. for	
		9.8	1.8 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.6			ionization; p <i>K</i> _a =	
		10.8	1.8 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.6			1.8, 8.3, 10.8.	
3.291	cystine	2	—	<i>k</i> / <i>k</i> _{Cl⁻} = 24	γ-r.	chem.	c.k.	63-0127
		2-2.2	6.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.03	γ-r.	opt.	c.k.	65-0388
		—	<i>ca.</i> 3-4 x 10 ⁹	—	p.r.	opt.	p.b.k.	69-0638
		1	9.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.76	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 130.	69-5278
		6.5	2.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.168	γ-r.	opt.	c.k.	73-0548
3.292	cytidine	2-2.2	3.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.303	p.r.	opt.	c.k.	65-0388
		5.2-5.4	5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.45	p.r.	opt.	c.k.	65-0388
		7.2-7.4	4.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.42	p.r.	opt.	c.k.	65-0388
		5.6	(6.4 ± 0.2) x 10 ⁴	—	p.r.	opt.	p.b.k. at 350 nm.	70-3069
		7	5.8 x 10 ⁹	—	p.r.	opt.	p.b.k. (OH addn.)	73-1071
3.293	cytidine-5'-phosphate (5'-cytidylic acid)	2-2.2	2.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.23	p.r.	opt.	c.k.	65-0388
		7.4-7.6	4.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.4	p.r.	opt.	c.k.	65-0388
		7.3	(4.9 ± 0.2) x 10 ⁹	—	p.r.	opt.	p.b.k. at 425 nm.	70-3069
		7	4.7 x 10 ⁹	—	p.r.	opt.	p.b.k. at 425 nm (OH adduct).	73-1071
3.294	cytochrome C (ferri)	5-10	—	<i>k</i> / <i>k</i> _{hydr} = 500	X-r.	chem.	c.k.	62-3002
		—	5.5 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{Tl⁺} = 5.5	γ-r.	opt.	c.k.; absorbance change at 550 nm; assume <i>k</i> _{Tl⁺} = 10 ¹⁰ ; <i>k</i> (OH + Fe ²⁺ cytC) = 4.6 x 10 ¹⁰	67-3020
		—	1.4 x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 550 nm; cor. for H + H, H + OH, H + cyt C by computer anal.	72-1002
		5.4, 7	2.7 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{thym} = 5 ± 1	γ-r.	opt.	c.k.	72-3071
		6.3	6.1 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 4.1	γ-r.	opt.	c.k.	73-0548
3.295	cytosine	2-2.2	3.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.28	p.r.	opt.	c.k.	65-0388
		5-6	4.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.41	p.r.	opt.	c.k.	65-0388
		7.4-7.6	4.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.447	p.r.	opt.	c.k.	65-0388
		11.4	≥ 7 x 10 ⁹	—	p.r.	opt.	p.b.k. at 335 nm.	68-0597

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.295 cont.		5.8	$(6.2 \pm 0.4) \times 10^9$	—	p.r.	opt.	p.b.k. at 450 nm.	70-3069
		7	$(6.8 \pm 0.5) \times 10^9$	—	p.r.	opt.	p.b.k.	73-1071
		6-7	4.0×10^9 (rel.)	$k/k_{\text{RNO}} = 0.32$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.296	deoxyadenylic acid	2-2.2	1.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.124$	p.r.	opt.	c.k.	65-0388
		6.4-6.6	3.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.32$	p.r.	opt.	c.k.	65-0388
3.297	deoxycytidylic acid	2-2.2	3.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.27$	p.r.	opt.	c.k.	65-0388
		4.3-4.5	3.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.35$	p.r.	opt.	c.k.	65-0388
		6.7-7	5.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.455$	p.r.	opt.	c.k.	65-0388
		7	4.9×10^9	—	p.r.	opt.	p.b.k.	73-1071
3.298	deoxyguanylic acid	2-2.2	4.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.42$	p.r.	opt.	c.k.	65-0388
		6.5-7	6.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.62$	p.r.	opt.	c.k.	65-0388
3.299	deoxyribose	—	1.9×10^9	—	—	—	—	66-0845
3.300	2-deoxy-2-sulfoamino-D-glucose	—	2.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.192$	p.r.	opt.	c.k.	70-3081
3.301	dextran	7	$> 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS^- , BzO^- , RNO; k varies with chain length; k per monomer unit.	70-0394
3.301a	diamide <i>See</i> N,N,N',N'-tetramethyl-1,2-diazenedicarboxamide (3.696). di- <i>tert</i> -butyl disulfide $\text{OH} + \text{C}_4\text{H}_9\text{SSC}_4\text{H}_9 \rightarrow \text{OH}^- + \text{RSSR}^+$	—	$(6.5 \pm 1.5) \times 10^9$	—	p.r.	opt.	p.b.k.	75-1089
3.302	1,1-dichloroethene $\text{OH} + \text{CH}_2=\text{CCl}_2 \rightarrow \text{CH}_2\text{OHCCl}_2$	—	$(4.1 \pm 0.4) \times 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS^- ; reference rate not given.	71-0709
3.303	1,2-dichloroethene $\text{OH} + \text{CHCl}=\text{CHCl} \rightarrow \text{CHCl}(\text{OH})\text{CHCl}$	—	$(5.0 \pm 0.4) \times 10^9$	—	p.r.	condy.	p.b.k. (Cl^-); $(\text{CHClOHCHCl} \rightarrow \text{H}^+ + \text{Cl}^- + \text{CHOCHCl})$.	71-0709
		—	$(4.4 \pm 0.4) \times 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS^- ; reference rate not given.	71-0709
3.304	1,4-dicyanobenzene $\text{OH} + \text{C}_6\text{H}_4(\text{CN})_2 \rightarrow \text{C}_6\text{H}_4\text{OH}(\text{CN})_2$	—	$(7.2 \pm 0.7) \times 10^8$ (rel.)	$k/k_{\text{MeOH}} = 0.8$	p.r.	opt.	c.k.; obs. buildup of OH adduct at 370 nm.	73-0121
3.305	1,2-diethoxyethane	9	2.2×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.2$	γ -r.	opt.	c.k. with RNO.	66-0423
3.306	diethoxymethane	9	1.6×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.84$	γ -r.	opt.	c.k. with RNO.	66-0423
3.307	diethylammonium ion	1	9.2×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.4$	Fenton	chem.	c.k.	49-0002
3.307a	diethyl disulfide $\text{OH} + \text{C}_2\text{H}_5\text{SSC}_2\text{H}_5 \rightarrow \text{OH}^- + \text{RSSR}^+$	—	$(1.4 \pm 0.5) \times 10^{10}$	—	p.r.	opt.	p.b.k.	75-1089
3.308	diethyleneglycol	9	2.0×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.1$	γ -r.	opt.	c.k. with RNO.	66-0423
3.309	diethyleneglycol diethyl ether	9	3.1×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.7$	γ -r.	opt.	c.k. with RNO.	66-0423
3.310	diethylenetriaminepentaacetic acid $\text{OH} + \text{DTPA} \rightarrow \text{H}_2\text{O} + \text{CO}_2 + \text{prod.}$	6.0	1.6×10^9 (rel.)	$k/k_{\text{ferro}} = 0.17$ $k/k_{\text{BuOH}} = 1.36$ $k/k_{\text{t-BuOH}} = 8.6$	γ -r.	chem.	c.k.; obs. $G(-\text{DTPA})$.	72-0169
3.311	diethyl malonate	6-7	6.5×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0591$	p.r.	opt.	c.k.	65-0387
3.312	diethyl succinate	6-7	7.8×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.071$	p.r.	opt.	c.k.	65-0387
3.313	1,2-difluorobenzene	—	4.5×10^9 (rel.)	—	p.r.	opt.	c.k. with CNS^- ; reference rate not given.	73-0054

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.314	1,4-difluorobenzene —	6×10^9 (rel.)	—	p.r.	opt.	c.k. with CNS ⁻ ; reference rate not given.	73-0054
3.315	dihydro-6-methyluracil —	2.3×10^9	—	p.r.	opt.	p.b.k. at 500 nm; true rate should be lower.	74-1085
	—	1×10^9	—	p.r.	opt.	c.k. with <i>tert</i> -BuOH, CNS ⁻ and EtOH.	74-1085
3.316	dihydroorotate ion 7	3.0×10^9	—	p.r.	opt.	p.b.k. (OH adduct).	70-0567
3.317	5,6-dihydrothymine ~7 ~12.4	2.2×10^9 0.4×10^9	— —	p.r.	opt.	p.b.k.; obs. transients at 400 (pH = 7) and 320 (pH = 12.4) nm.	68-0312
	7	1.6×10^9	—	—	—	cited from 69-0012.	70-0567
	—	$<2.2 \times 10^9$	—	p.r.	opt.	p.b.k. at 500 nm.	74-1085
3.318	dihydrouracil 6-8	1.5×10^9 (rel.)	$k/k_{\text{RNO}} = 0.12$	γ -r.	opt.	c.k.; 17°C.	75-0294
	7	$<(2.1 \pm 0.5) \times 10^9$	—	p.r.	opt.	p.b.k. at 420 nm.	69-0571
	7	$(1.2 \pm 0.2) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = (0.11 \pm 0.02)$	p.r.	opt.	c.k.; cor. for incomplete scavenging of e_{aq}^- by H ₂ O ₂ .	69-0571
	7	1.3×10^9	—	—	—	cited from 69-0012.	70-0567
	—	$<2.0 \times 10^9$	—	p.r.	opt.	p.b.k. at 500 nm.	74-1085
	<i>m</i> -dihydroxybenzene	<i>See m</i> -hydroxyphenol (3.455).					
	<i>o</i> -dihydroxybenzene	<i>See o</i> -hydroxyphenol (3.456).					
	<i>p</i> -dihydroxybenzene	<i>See</i> hydroquinone (3.446).					
3.319	2,5-dihydroxy-2,5-dimethyl-3-hexyne 1	3.1×10^9 (rel.)	$k/k_{\text{MeOH}} = 3.38$	Fenton	chem.	c.k.; $k_{\text{MeOH}}/k_{\text{Fe}^{2+}} = 4.3$.	73-9350
3.320	4,5-dihydroxy-2,7-naphthalenedisulfonic acid 0.1	8.5×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.37$	γ -r.	chem.	c.k.	67-0025
3.320a	1,2-dimethoxybenzene —	$(5.2 \pm 0.5) \times 10^9$	—	p.r.	—	—	75-1171
3.320b	1,3-dimethoxybenzene —	$(7.2 \pm 0.7) \times 10^9$	—	p.r.	—	—	75-1171
3.320c	1,4-dimethoxybenzene —	$(7.0 \pm 0.7) \times 10^9$	—	p.r.	—	—	75-1171
3.321	1,2-dimethoxyethane 9	1.6×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.85$	γ -r.	opt.	c.k. with RNO.	66-0423
3.322	dimethoxymethane 9	5.7×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.31$	γ -r.	opt.	c.k. with RNO.	66-0423
3.323	<i>N,N</i> -dimethylacetamide 5.5	3.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.32$	p.r.	opt.	c.k.	70-0098, 71-0645
	OH + CH ₃ CON(CH ₃) ₂ → H ₂ O + CH ₃ CON(CH ₃)CH ₂						
3.324	dimethylammonium ion 1	$\sim 10^6$ (I)	—	e -r.	esr	estd. from drop in aminium radical signal on addn. of <i>tert</i> -BuOH.	72-5118
	(I) OH + (CH ₃) ₂ NH ₂ ⁺ → H ₂ O + (CH ₃) ₂ NH ⁺						
	(II) OH + (CH ₃) ₂ NH ₂ ⁺ → H ₂ O + CH ₂ (CH ₃)NH ₂ ⁺						
3.325	<i>N,N</i> -dimethylaniline 9	8.9×10^9 (rel.)	$k/k_{\text{EtOH}} = 4.8$	γ -r.	opt.	c.k. with RNO.	66-0441
	—	1.3×10^{10}	—	p.r.	opt.	p.b.k. at 455 and 330 nm.	72-0289
3.326	<i>N,N</i> -dimethylanilinium ion 1	1.5×10^9 (rel.)	$k/k_{\text{Fe}^{2+}} = 6.6$	Fenton	chem.	c.k.	49-0003

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.327	3,3-dimethylbutyrate ion	9	1.7×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.9$	γ -r.	opt.	c.k. with RNO.	66-0423
3.327a	dimethyl disulfide $\text{OH} + \text{CH}_3\text{SSCH}_3 \rightarrow \text{OH}^- + \text{RSSR}^+$	—	$(1.7 \pm 0.3) \times 10^{10}$	—	p.r.	opt.	p.b.k.	75-1089
3.328	<i>N,N</i> -dimethylformamide	5.5	1.7×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.155$	p.r.	opt.	c.k.	70-0098
3.329	1,1-dimethylhydrazine	9.2	1.6×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.45$	p.r.	opt.	c.k.	72-0003
3.330	1,2-dimethylhydrazine	10.1	1.4×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.3$	p.r.	opt.	c.k.	72-0003
3.331	1,1-dimethylhydrazinium ion	3.5	8.1×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.074$	p.r.	opt.	c.k.	72-0003
3.332	1,2-dimethylhydrazinium ion	3.5	7.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.065$	p.r.	opt.	c.k.	72-0003
3.333	1,2-dimethylindole	9.0	$(1.25 \pm 0.02) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.334	1,3-dimethylindole	9.0	$(1.01 \pm 0.08) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.335	2,3-dimethylindole	9.0	$(1.26 \pm 0.01) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.336	2,4-dimethylphenyl- β -D-glucopyranoside	—	4.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.41$	p.r.	opt.	c.k.	71-0056
3.337	3,4-dimethylphenyl- β -D-glucopyranoside	—	3.6×10^9 (rel.)	—	γ -r.	opt.	c.k. with RNO, rel. to $k(\text{OH} + \text{X}) = 4.4 \times 10^9$, X = phenyl- β -D-glucopyranoside.	71-0056
3.337a	2,2-dimethyl-1-phenyl-1-propanol	1.7-1.8	1.1×10^{10} (rel.)	$k/k_{2\text{-PrOH}} = 5.2$	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.338	dimethyl phosphate ion $\text{OH} + (\text{CH}_3\text{O})_2\text{PO}_2^- \rightarrow \text{H}_2\text{O} + \text{CH}_2\text{O}(\text{CH}_3\text{O})\text{PO}_2^-$	—	1.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.011$	p.r.	opt.	c.k.	72-3008
3.339	<i>N,N</i> -dimethylpivalamide	5-6	3.9×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.357$	p.r.	opt.	c.k.	71-0414
3.340	dimethyl sulfide <i>See</i> methyl sulfide (3.552).	9	3.1×10^9 (rel.)	$k/k_{\text{RNO}} = 0.25 \pm 0.01$	γ -r.	opt.	c.k.	69-0280
3.341	2,6-dimethylpyridine	9	3×10^9 (rel.)	$k/k_{\text{RNO}} = 0.24 \pm 0.01$	γ -r.	opt.	c.k.	69-0280
3.341a	dimethyl sulfone	1.5	$< 6 \times 10^6$	—	Ti(III) + H_2O_2	esr	estd. rel. to $k(\text{OH} + \text{Ti(III)}) = 3 \times 10^9$.	75-5237
3.342	dimethyl sulfoxide	—	7.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.64$	p.r.	opt.	c.k.	67-0186
		—	5.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.53$	p.r.	opt.	c.k.	73-1077
	<i>For other ratios see: 3.348.</i>							

For other ratios see: 3.348.

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.343	1,4-dioxane	7	2.8×10^9 (rel.)	$k/k_1^- = 0.23 \pm 0.02$	p.r.	opt.	c.k.; obs. I_2^- formn. at 400 nm.	65-0010
	$\text{OH} + \text{C}_4\text{H}_8\text{O}_2 \rightarrow$ $\text{H}_2\text{O} + \text{C}_4\text{H}_7\text{O}_2$	2-2.2	2.0×10^9 (rel.)	$k/k_{\text{thym}} = 0.37 \pm 0.04$	γ -r.	opt.	c.k.	65-0388, 67-0461
		9	1.8×10^9 (rel.)	$k/k_{\text{EtOH}} = 1$	γ -r.	opt.	c.k. with RNO.	66-0423
		7	—	$k/k_{t\text{-BuOH}} = 3.5$	Ti(III) + H_2O_2	esr	c.k.	74-5144
	<i>For other ratios see: 3.627.</i>							
3.344	diphenylacetate ion	9.1	4×10^9	—	p.r.	opt.	p.b.k. at 340 nm.	72-0047
3.345	diphenylamine	9	1.3×10^{10} (rel.)	$k/k_{\text{RNO}} = 1.04 \pm 0.04$	γ -r.	opt.	c.k.	69-0280
3.345a	di-2-propyl disulfide	—	$(2.0 \pm 1.0) \times 10^{10}$	—	p.r.	opt.	p.b.k.	75-1089
	$\text{OH} + \text{C}_3\text{H}_7\text{SSC}_3\text{H}_7 \rightarrow$ $\text{OH}^- + \text{RSSR}^+$							
3.346	2,2'-dithiobis-(ethylamine)	1	1.4×10^{10} (rel.)	$k/k_{\text{thym}} = 2.6$	Fenton	esr	c.k.; $k/k_{\text{perox}} = 186$.	69-5278
3.347	dithiothreitol	7	$(1.5 \pm 0.5) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 300 nm; ratio with phenylalanine = 2.	73-1020
	$\text{OH} +$ $\text{SHCH}_2\text{CHOHCHOHCH}_2\text{S}^-$ $\rightarrow \text{H}_2\text{O} +$ $\cdot\text{SCH}_2\text{CHOHCHOHCH}_2\text{S}^-$							
3.348	DNA	—	$\sim 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS^- .	65-0388
	$\text{OH} + \text{DNA} \rightarrow$ transient	9	$\sim 1.2 \times 10^{13}$ (rel.)	$k/k_{\text{RNO}} \cong 10^3$	γ -r.	opt.	c.k.; mol. wt. 5×10^6 .	67-0555
		~ 7	0.6×10^9	—	p.r.	opt.	p.b.k.; obs. transients at 400 (pH = 7) and 320 (pH = 12.4) nm; assume nucleotides (mol. wt. 350) react independently.	68-0312
		~ 12.4	0.6×10^9	—				
		—	$< 2.6 \times 10^8$ (rel.)	—	p.r.	opt.	c.k. with CNS^- ; based on nucleotide concn.;	68-0845
		7.5	1.3×10^{13}	—	p.r.	opt.	p.b.k. at 310 and 420 nm; $k = 8 \times 10^8$ per nucleotide base group.	69-0018
		7	$(4 \pm 1) \times 10^8$	—	p.r.	opt.	p.b.k. at 340 nm; k per base unit.	73-1071
		—	3×10^8 (rel.)	—	γ -r.	trac.	c.k.; effect of <i>tert</i> -BuOH, EtOH, 2-PrOH, <i>iso</i> -BuOH, isoamyl alcohol and dimethyl sulfoxide on binding of ^{14}C -nitrofurazone to DNA.	73-1077
		—	5.2×10^8	—	p.r.	opt.	d.k. as well as p.b.k. at 400 nm; rate in terms of nucleotide concn. (mol. wt. 360).	73-3016
		7.0	1.8×10^8 (rel.)	—	γ -r.	trac.	c.k. assuming $k(\text{OH} + \text{dimethyl sulfoxide}) = 6 \times 10^9$; effect on binding of	73-3080

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.348 cont.		—	—	—	p.r.	opt.	¹⁴ C-nitrofurazone in N ₂ O satd. 0.3% DNA; k per base group of mol. wt. \cong 330. k for DNA-bound proflavine $\sim 2 \times 10^9$.	75-3094
3.349	dodecyl sodium sulfate	—	7.6×10^9 (rel.)	—	p.r.	opt.	c.k. with CNS ⁻ ; k_{CNS^-} not given; $k = 5.0 \times 10^8$ for concn. $> 8.1 \times 10^{-3} M$.	71-0586
3.350	egg white	—	—	—	p.r.	opt.	p.b.k. at 420 nm (cystine anion radical); OH half-life $< 5 \times 10^{-9}$ s.	73-1059
3.351	eosin, dianion	—	1.8×10^{10} (I)(rel.)	$k_1/k_{\text{carb}} = 50$	p.r.	opt.	c.k.	66-0501
	(I) OH + S \rightarrow charge transfer	10.5	1.2×10^{10} (I + II) (rel.)	$k_1 + k_{\text{II}}/k_{\text{carb}} = 34$	—	—	c.k.; cor for presence of HCO ₃ ⁻ .	67-0038
	(II) OH + S \rightarrow addn.	9.0	$k_{\text{I}} = 1.7 \times 10^9$	—	p.r.	opt.	X abs. at 450 nm.	68-0309
		9.0	$k_{\text{II}} = 6 \times 10^8$	—	p.r.	opt.	adduct abs. at 600 nm.	68-0309
3.352	1,2-epoxybutane	9	7.6×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.41$	γ -r.	opt.	c.k. with RNO.	66-0423
3.353	1,2-epoxypropane	9	2.4×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.13$	γ -r.	opt.	c.k. with RNO.	66-0423
3.354	2,3-epoxypropanol	9	4.6×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.246$	γ -r.	opt.	c.k. with RNO.	66-0423
3.355	erythritol	9	2.0×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.1$	γ -r.	opt.	c.k. with RNO.	66-0423
3.356	ethane	1.2	—	$k/k_{\text{HCOOH}} = 10 \pm 1$	γ -r.	chem.	c.k.	66-0265
3.357	ethanesulfonate ion	—	1.0×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0091$	p.r.	opt.	c.k.	68-0352
3.358	ethanol (EtOH)	1	8.7×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 3.8$	Fenton	chem.	c.k.	49-0002
	(I) OH + EtOH \rightarrow H ₂ O + CH ₃ CHOH	6.6, 10.5	1.1×10^9 (rel.)	$k/k_{\text{ferro}} = 0.12$	X-r.	opt.	c.k.; not cor. for H ₂ O ₂ .	62-0023
	(II) OH + EtOH \rightarrow H ₂ O + CH ₂ CH ₂ OH	— 7	1.3×10^9 (rel.) 1.9×10^9 (rel.)	$k/k_{\text{carb}} = 3.6$ $k/k_{\text{ferro}} = 0.21$	p.r. p.r.	opt. opt.	c.k. c.k.	64-0131 65-0007
	(III) OH + EtOH \rightarrow H ₂ O + CH ₃ CH ₂ O	7	9.1×10^8 (rel.)	$k/k_{\text{I}^-} = 0.076 \pm 0.007$	p.r.	opt.	c.k.; I ₂ ⁻ formn. meas. at 400 nm.	65-0010 67-0041
		3, 10.5	1.6×10^9 (rel.)	$k/k_{\text{BaO}^-} = 0.29$	γ -r.	trac.	c.k.; meas. ¹⁴ CO ₂ .	65-0099
		—	1.5×10^9 (rel.)	$k/k_{\text{ferro}} = 0.16$	phot.	—	c.k.	65-0247
		9.0	1.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.146$	γ -r.	opt.	c.k.; meas. at 400 nm.	65-0356
		2-2.2	2.0×10^9 (rel.)	$k/k_{\text{tbyrn}} = 0.37 \pm 0.035$	γ -r.	opt.	c.k.	65-0388, 67-0461
		5-5.5	1.8×10^9 (rel.)	$k/k_{\text{tbyrn}} = 0.33 \pm 0.035$	γ -r.	opt.	c.k.	65-0388, 67-0461
		7, 10.7	$\sim 1.8 \times 10^9$ (rel.)	$k/k_{\text{carb}} \cong 4.8$	p.r.	opt.	c.k.	65-0190
		7	1.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.164$	p.r.	opt.	c.k.	65-0190
		2	1.65×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.15$	p.r.	opt.	c.k.	65-0387
		7	1.9×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.17$	p.r.	opt.	c.k.	65-0387
		—	1.9×10^9 (rel.)	$k/k_{\text{MeOH}} = 2.1$	p.r.	opt.	c.k. with HSO ₄ ⁻ , meas. abs. at 450 nm. (SO ₄ ⁻).	66-0019
		—	—	$k/k_{\text{TCOO}^-} = 0.53$	γ -r.	trac.	c.k.	68-0209
		—	1.8×10^9 (rel.)	$k/k_{\text{BaO}^-} = 0.31$	p.r.	opt.	c.k.; obs.	68-0304
		—	1.8×10^9 (rel.)	$k/k_{\text{PA}^-} = 0.23$			hydroxycyclo-	
		—	1.8×10^9 (rel.)	$k/k_{\text{PNBA}^-} = 0.70$			hexadienyl radical buildup.	
		5.5	1.7×10^9 (rel.)	$k/k_{\text{NB}} = 0.54$	r.	opt.	c.k.	68-0494
		7	1.7×10^9 (rel.)	$k/k_{\text{ReO}^-} = 0.29$	r.	lum.	c.k.	68-0494

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH		<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.358 cont.	~1.2	1.4 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{2-PrOH} = 0.63	γ-r.	chem.	c.k.	68-0602
	6.98	1.3 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{2-PrOH} = 0.61	γ-r.	chem.	c.k.	68-0602
	—	1.6 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{RNO} = 0.128	p.r.	opt.	c.k.	69-0156
	1	2.1 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{thym} = 0.395	Fenton	esr	c.k.;	69-5278
	1	1.9 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{thym} = 0.35	Ti(III) - H ₂ O ₂	esr	<i>k</i> / <i>k</i> _{perox} = 28.9. c.k.;	69-5278
	—			<i>k</i> _I / <i>k</i> _{II} ≅ 8.6	γ-r.	chem.	c.k. with H ₂ O ₂ .	70-0338
	11	2.0 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{carb} = 5.5	p.r.	opt.	c.k.; assume p <i>K</i> _a (OH) = 11.9.	70-0511
	—			<i>k</i> _I / <i>k</i> _{II} = 6.0 ± 0.2	γ-r.	chem.	c.k. with H ₂ O ₂ ; <i>k</i> _H / <i>k</i> _D (I) = 1.24 ± 0.04; <i>k</i> _H / <i>k</i> _D (II) = 1.55 ± 0.06 or 2.28 ± 0.20.	71-0081
	9	6.2 x 10 ⁸ (rel.)		<i>k</i> / <i>k</i> _{RNO} = 0.05	γ-r.	opt.	c.k.; <i>E</i> _a = -4.0 ± 1.1 kcal/mol (-16.7 ± 4.5 kJ/mol) at -8 to 23°C.	71-0469
	nat.	1.9 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{ferro} = 0.2	p.r.	opt.	c.k.	71-0578
	0.82	1.7 x 10 ⁹ (I) (rel.) 1.6 x 10 ⁸ (II) (rel.)		<i>k</i> _I / <i>k</i> _{Fe²⁺} = 7.32 <i>k</i> _{II} / <i>k</i> _{Fe²⁺} = 0.69	Fenton	chem.	c.k.; also re- ported <i>k</i> / <i>k</i> _{Fe³⁺} = 7.0.	71-9132
	—	—		<i>k</i> _{II} / <i>k</i> _I = 0.16	p.r.	opt.	detd. % of	73-0126
	—	—		<i>k</i> _{III} / <i>k</i> _I = 0.03			α-alcohol and ethoxy radicals by reaction with TNM and I ⁻ , resp.	
	—	1.65 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{ferro} = 0.18	p.r.	opt.	c.k.	73-1046
7,10.6	1.5 x 10 ⁹ (rel.)		<i>k</i> / <i>k</i> _{RNO} = 0.12	phot.	opt.	c.k.; H ₂ O ₂ soln; assume <i>k</i> _{H₂O₂} / <i>k</i> _{RNO} << 1.	73-7575	
For other ratios see: 3.12, 3.13, 3.25, 3.26, 3.27, 3.36, 3.41, 3.52, 3.54, 3.66, 3.80, 3.107, 3.112, 3.124, 3.127, 3.128, 3.131, 3.133, 3.134, 3.158, 3.170, 3.172, 3.185, 3.188, 3.189, 3.191, 3.193, 3.203, 3.209, 3.210, 3.214, 3.215, 3.222-3, 3.225, 3.226, 3.238, 3.246, 3.249, 3.250, 3.251, 3.257, 3.258, 3.272, 3.276, 3.282, 3.287, 3.305, 3.306, 3.308, 3.309, 3.321, 3.322, 3.325, 3.327, 3.352, 3.353, 3.354, 3.355, 3.359, 3.360, 3.369, 3.370, 3.371, 3.378, 3.380, 3.403, 3.405, 3.406, 3.408, 3.439, 3.440, 3.449, 3.502, 3.509, 3.511, 3.512, 3.514, 3.515, 3.516, 3.522, 3.523, 3.524, 3.526, 3.529, 3.530, 3.534, 3.545, 3.546, 3.549, 3.565, 3.566, 3.567, 3.573, 3.592, 3.593, 3.598, 3.601, 3.602, 3.607, 3.609, 3.610, 3.611, 3.620, 3.622, 3.634, 3.635, 3.636, 3.637, 3.638, 3.641, 3.669, 3.673, 3.690, 3.693, 3.695, 3.698, 3.711, 3.712, 3.720a, 3.723, 3.724, 3.731, 3.732, 3.752.								
3.359	ethanol- <i>d</i> ₅ OH + C ₂ D ₅ OH → HDO + CD ₃ CDOH	6	1.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 0.62	γ-r.	chem.	c.k. with Br ⁻ .	66-0423
3.360	2-ethoxyethanol	9	1.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 0.9	γ-r.	opt.	c.k. with RNO.	66-0423
3.361	ethyl acetate	1	2.5 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{Fe²⁺} = 1.1	Fenton	chem.	c.k.	49-0002
		6-7	4.0 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.0364	p.r.	opt.	c.k.	65-0387
		2.0- 2.2	2.8 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.052 ± 0.004	γ-r.	opt.	c.k.	67-0461
3.362	ethylamine	12	1.3 x 10 ¹⁰ (rel.)	—	p.r.	opt.	c.k.; calcd. from <i>k</i> / <i>k</i> _{CNS⁻} = 1.0 at pH 11.2.	71-0585
		—	3.2 x 10 ⁹ (rel.) 6.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{NB} = 1.3 <i>k</i> / <i>k</i> _{CNS⁻} = 0.58	p.r.	opt.	c.k.; calcd. from values obs. at pH 8-13.1.	73-0016

TABLE 4. Reactions of OH with organic solutes – Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.363	ethylammonium ion	2	7.8×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.0071$	p.r.	opt.	c.k.	70-0371
		4	5.1×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0464$	p.r.	opt.	c.k.	70-0371
		3.1	3.0×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0273$	p.r.	opt.	c.k.	71-0585
		—	3.8×10^8 (rel.)	$k/k_{\text{NB}} = 0.085$	p.r.	opt.	c.k.; calcd. from	73-0016
			6×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.54$			values obs. at	
						pH 8-13.1.		
3.364	ethyl butyrate	6-7	1.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.146$	p.r.	opt.	c.k.	65-0387
3.365	ethylene	—	4.9×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 2.13$	γ -r.	opt.	c.k.	67-0037
	(I) $\text{OH} + \text{CH}_2=\text{CH}_2 \rightarrow \bullet\text{CH}_2\text{CH}_2\text{OH}$	—	4.8×10^9 (I + II) (rel.)	$k_{\text{I+II}}/k_{\text{I}^-} = 0.402$	p.r.	opt.	c.k.; meas. I_2^- at 400 nm.	67-0041
	(II) $\text{OH} + \text{CH}_2=\text{CH}_2 \rightarrow \text{CH}_2=\text{CH}\bullet + \text{H}_2\text{O}$	—	$(1.0 \pm 0.2) \times 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS^- and HCO_3^- ; details not given.	67-0269
	7		$k_{\text{II}}/k_{\text{I}} = 0.3$	γ -r.	chem.	meas.	67-0522	
							$G(\text{alcohols})$.	
3.366	ethylenediamine	4	$\sim 3.5 \times 10^7$ (rel.)	—	—	—	c.k. with EtOH.	66-0401
		8.0	$(5.3 \pm 1.0) \times 10^8$ (rel.)	$k/k_{\text{CNS}^-} = 0.0265$	p.r.	opt.	c.k.; at pH 8.5, 9.0, 10.0 ratio is 0.225, 0.3 and 0.5, resp.	72-0461
3.367	ethylenediamine tetraacetate ion	9	2.6×10^9 (rel.)	$k/k_{\text{ferro}} = 0.28$	γ -r.	opt.	c.k. with RNO assuming $k_{\text{RNO}} = k_{\text{ferro}}$.	67-0555
		7	1.2×10^9 (rel.)	$k/k_{\text{RNO}} = 0.096$	phot.	opt.	c.k.; H_2O_2 soln; assume $k_{\text{H}_2\text{O}_2}/k_{\text{RNO}} \ll 1$.	73-7575
		10	4.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.38$				
		10.3	5.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.46$				
		10.6	6.3×10^9 (rel.)	$k/k_{\text{RNO}} = 0.5$				
3.368	ethylenediamine-tetraacetic acid	—	5.3×10^9 (rel.)	$k/k_{\text{MeOH}} = 5.9$	X-r.	chem.	c.k.	72-0056
		~ 0	—	$k/k_{\text{acrylamide}} = 2.2$	Fenton	pol.	c.k.	72-9162
3.369	ethyleneglycol $\text{OH} + \text{CH}_2\text{OHCH}_2\text{OH} \rightarrow \text{H}_2\text{O} + \text{CH}_2\text{OHCHOH}$	1	8.3×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.36$	therm.	chem.	c.k.; persulfate oxidation.	49-0002
		7	1.6×10^9 (rel.)	$k/k_{\text{carb}} = 4.3$	p.r.	opt.	c.k.	65-0190
		7	1.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.13$	p.r.	opt.	c.k.	65-0387
		9	1.5×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.82$	γ -r.	opt.	c.k. with RNO.	66-0423
		6	1.1×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.62$	γ -r.	chem.	c.k. with Br^- .	66-0423
		2-2.2	1.7×10^9 (rel.)	$k/k_{\text{thym}} = 0.32 \pm 0.03$	γ -r.	opt.	c.k.	67-0461
		nat.	1.5×10^9 (rel.)	$k/k_{\text{ferro}} = 0.158$	p.r.	opt.	c.k.	71-0578
		—	—	—	p.r.	opt.	> 0.1% alkoxy radical detd. by reaction with I^- .	73-0126
3.370	ethylene oxide	—	2.1×10^9 (rel.)	$k/k_{\text{ferro}} = 0.225$	p.r.	opt.	c.k.	73-1046
		9	6.7×10^7 (rel.)	$k/k_{\text{EtOH}} = 0.036$	γ -r.	opt.	c.k. with RNO.	66-0423
3.371	ethyl ether	1	1.5×10^9 (rel.)	$k/k_{\text{Fe}^{2+}} = 6.4$	Fenton	chem.	c.k.	49-0002
		~ 7	4.9×10^9 (rel.)	$k/k_{\text{I}^-} = 0.38 \pm 0.04$	p.r.	opt.	c.k.; meas. I_2^- formn. at 400 nm.	65-0010
		9	1.4×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.74$	γ -r.	opt.	c.k. with RNO.	66-0423
		2	2.8×10^9 (rel.)	$k/k_{\text{thym}} = 0.52 \pm 0.04$	γ -r.	opt.	c.k.	67-0461
		~ 1.2	2.5×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 1.15$	γ -r.	chem.	c.k.	68-0602
3.372	ethyl formate	6-7	3.8×10^8 (rel.)	$k/k_{2-\text{PrOH}} = 1.20$	γ -r.	chem.	c.k.	68-0602
		6-7	3.8×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.035$	p.r.	opt.	c.k.	65-0387
3.373	N-ethylmalamic acid	6.0	7.0×10^9	—	p.r.	opt.	p.b.k.	72-0144

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.374	<i>N</i> -ethylmaleimide	—	4.3 × 10 ⁹ (rel.)	$k/k_{\text{thym}} = (0.8 \pm 0.1)$	X-r.	opt.	c.k.	69-0562
		6.0	9.0 × 10 ⁹	—	p.r.	opt.	p.b.k.	72-0144
3.375	4-ethyl-5-hydroxy-2-methylpyridine	6.5	1.4 × 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.11 \pm 0.01$	γ-r.	opt.	c.k.	69-0580
3.375a	1-(<i>p</i> -ethyl-phenyl)ethanol	1.7-1.8	1.5 × 10 ¹⁰ (rel.)	$k/k_{2\text{-PrOH}} = 6.7$	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.376	ethyl propionate	6-7	8.7 × 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.079$	p.r.	opt.	c.k.	65-0387
	Flagyl	See 1-(2-hydroxyethyl)-2-methyl-5-nitroimidazole (3.452a).						
3.377	fluorescein	—	(1.4 ± 0.2) × 10 ⁹	—	p.r.	opt.	computer anal.	68-0172
	OH + dye → adduct	—	(1.6 ± 0.3) × 10 ⁹	—	p.r.	opt.	computer anal.	68-0172
	OH + dye → X• + OH ⁻						(X• = semi-oxidized fluorescein.	
		10	1.2 × 10 ¹⁰	—	p.r.	opt.	d.k. as well as p.b.k.	73-6068, 74-1063
3.378	fluoroacetate ion	9	3.0 × 10 ⁷ (rel.)	$k/k_{\text{EtOH}} = 0.016$	γ-r.	opt.	c.k. with RNO.	66-0423
3.379	fluorobenzene	—	8 × 10 ⁹ (rel.)	—	p.r.	opt.	c.k. with CNS ⁻ ; reference rate not given.	73-0054
3.380	<i>p</i> -fluorobenzoate ion	9	3.5 × 10 ⁹ (rel.)	$k/k_{\text{EtOH}} = 1.88$	γ-r.	opt.	c.k. with RNO.	66-0441
3.381	5-fluorouracil	7	5.2 × 10 ⁹ (rel.)	$k/k_{\text{CNS}^-} = 0.472$	p.r.	opt.	c.k.	72-0049
		7	5.5 × 10 ⁹	—	p.r.	opt.	p.b.k. at 340 nm.	72-0049
		11	6.0 × 10 ⁹	—	p.r.	opt.	p.b.k. at 340 nm.	72-0049
3.382	formaldehyde	1	6.9 × 10 ⁸ (rel.)	$k/k_{\text{Fe}^{2+}} = 3.0$	Fenton	chem.	c.k.	49-0002
	OH + HCHO → H ₂ O + CHO	1.3		$k/k_{\text{oxalic acid}} = 40$	r.	chem.	c.k.	68-0503
3.383	formamide	5.5	< 5.0 × 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} < 0.045$	p.r.	opt.	c.k.	70-0098
3.384	formate ion	5.8-	~ 1.9 × 10 ⁹ (rel.)	$k/k_{\text{ferro}} \cong 0.2$	X-r.	opt.	c.k.; not cor.	62-0023
	OH + HCOO ⁻ → H ₂ O + COO ⁻	10.1					for H ₂ O ₂ .	
		7	2.9 × 10 ⁹ (rel.)	$k/k_{\text{I}^-} = 0.24 \pm 0.002$	p.r.	opt.	c.k.; I ₂ formn. at 400 nm.	65-0010
		10.7	3.5 × 10 ⁹ (rel.)	$k/k_{\text{BzO}^-} = 0.62$	γ-r.	trac.	c.k.; meas. ¹⁴ CO ₂ .	65-0099
		—	2.4 × 10 ⁹ (rel.)	$k/k_{\text{ferro}} = 0.26$	phot.	—	c.k.	65-0247
		9.0	4.0 × 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.32 \pm 0.02$	γ-r.	opt.	c.k.	65-0356
		2-5	3.4 × 10 ⁹ (rel.)	$k/k_{\text{thym}} = 0.63 \pm 0.06$	γ-r.	opt.	c.k.; calcd. <i>k</i> on basis of rates obtained with formic/formate systems as function of pH.	67-0461
		7	3.8 × 10 ⁹ (rel.)	$k/k_{\text{BzO}^-} = 0.66$	r.	chem.	c.k.	68-0494
		6.98	2.9 × 10 ⁹ (rel.)	$k/k_{2\text{-PrOH}} = 1.34$	γ-r.	chem.	c.k.	68-0602
		—	2.2 × 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.176$	p.r.	opt.	c.k.	69-0156
		11	4.0 × 10 ⁹ (rel.)	$k/k_{\text{carb}} = 10.6$	p.r.	opt.	c.k. with CO ₃ ²⁻ (μ = 0.4); assumed p <i>K</i> _a (OH) = 11.9	69-0379
		8.4	2.9 × 10 ⁹ (rel.)	$k/k_{\text{bicarb}} = 80$	p.r.	opt.	c.k. meas. CO ₃ ⁻ .	69-0379
		nat.	2.8 × 10 ⁹ (rel.)	$k/k_{\text{ferro}} = 0.301$	p.r.	opt.	c.k.	71-0578
		For other ratios see: 3.24, 3.88, 3.102, 3.103, 3.111, 3.133, 3.137, 3.460, 3.592, 3.643, 3.682, 3.742.						
		For ratios with ³ HCO ₂ ⁻ see: 3.66, 3.82, 3.357, 3.511, 3.637.						
3.385	formic acid	2.5	6.5 × 10 ⁸ (rel.)	$k/k_{\text{ferro}} \cong 0.07$	X-r.	opt.	c.k.; not cor.	62-0023
	OH + HCOOH → H ₂ O + COOH	1.0	1.6 × 10 ⁸ (rel.)	$k/k_{\text{I}^-} = (1.3 \pm 0.2) \times 10^{-2}$	p.r.	opt.	c.k.; obs. formn. of I ₂ at 400 nm.	65-0010
		1	1.3 × 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.0114$	p.r.	opt.	c.k.	65-0387

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.	
3.385 cont.		2-5	1.6 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.03 ± 0.03	γ-r.	opt.	c.k.; calcd. <i>k</i> on basis of rates obtained with formic/formate system as a function of pH.	67-0461	
		~1.2	1.3 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 0.06	γ-r.	chem.	c.k.	68-0602	
		1	1.5 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.028	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 1.95.	69-5278	
		1	1.5 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.028	Ti(III) - H ₂ O ₂	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 1.98.		
		0.8	—	<i>k</i> / <i>k</i> _{bisulf} [HSO ₄ ⁻] = 690 ± 80	γ-r.	chem.	c.k.; computer anal.	72-0094	
		0	—	= 14.6 ± 0.6					
			For other ratios see: 3.26, 3.84, 3.106, 3.137, 3.356.						
3.386	fumaric acid	1	1.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{MeOH} = 1.24	Fenton	chem.	c.k.; <i>k</i> _{MeOH} / <i>k</i> _{Fe²⁺} = 4.3.	73-9341	
3.387	Furadantin	7	9.3 x 10 ⁹	—	p.r.	opt.	p.b.k. as well as d.k.	73-1018	
3.388	2-furaldehyde	9	7.75 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.62 ± 0.04	γ-r.	opt.	c.k.	73-0301	
3.389	Furamazone	7	1.03 x 10 ¹⁰	—	p.r.	opt.	p.b.k. as well as d.k.	73-1018	
3.390	furan	—	3.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.35	p.r.	opt.	c.k.	71-0360	
	OH + C ₄ H ₄ O → (OH)C ₄ H ₄ O	9	1.45 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 1.16 ± 0.05	γ-r.	opt.	c.k.	73-0301	
3.391	furfuryl alcohol	9	1.5 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 1.19 ± 0.10	γ-r.	opt.	c.k.	73-0301	
3.392	2-furoate ion	9	1.15 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.92 ± 0.16	γ-r.	opt.	c.k.	73-0301	
3.393	gelatin	—	9.1 x 10 ¹⁰ (rel.)	—	p.r.	opt.	c.k. with CNS ⁻ ; reference rate not given; mol. wt. 100,000.	68-3007	
3.394	glucose	—	7.4 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{ferro} ≅ 0.08	phot.	—	c.k.	65-0247	
	(I) OH + C ₆ H ₁₂ O ₆ → deoxyglucose	—	1.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{I⁻} = 0.1	p.r.	opt.	c.k.	65-0391	
	(II) OH + C ₆ H ₁₂ O ₆ → malondialdehyde	2-2.2	2.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.40 ± 0.03	γ-r.	opt.	c.k.	67-0461	
		6.5	3.8 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.03 ± 0.01	γ-r.	opt.	c.k.	69-0580	
		8.8	—	<i>k</i> _I / <i>k</i> _{Br⁻} = 0.865	p.r.	chem.	c.k.	70-0251	
		—	—	<i>k</i> _{II} / <i>k</i> _{Br⁻} = 0.642	p.r.	chem.	c.k.	70-0251	
3.395	glucosephosphate	6.5	1.6 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.013 ± 0.003	γ-r.	opt.	c.k.	69-0580	
3.396	D-glucuronate ion	—	3.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.27	p.r.	opt.	c.k.	70-0509, 70-3081	
3.397	glucuronic acid	acid	1.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.12	p.r.	opt.	c.k.	70-0509	
3.398	D-glucuronolactone	—	1.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.15	p.r.	opt.	c.k.	70-0509	
3.399	glutamic acid	2-2.2	1.4 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.0255	γ-r.	opt.	c.k.	65-0388	
		6.5	2.3 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.018	γ-r.	opt.	c.k.	73-0548	
3.400	glutamine	2-2.2	1.6 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.029 ± 0.003	γ-r.	opt.	c.k.	67-0461	
		6.0	5.4 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.043	γ-r.	opt.	c.k.	67-0461	
3.401	glutaric acid	2-2.2	7.0 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.13 ± 0.01	γ-r.	opt.	c.k.	67-0461	
3.402	glutathione (reduced)	1	1.4 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.3	p.r.	opt.	c.k.	65-0387	
		1	1.4 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{thym} = 2.6	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 186.	69-5278	
	(oxidized)	1	1.1 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.98	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 143.	69-5278	

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.403	glycerol	—	1.9×10^9 (rel.)	$k/k_{\text{carb}} = 5.3$	p.r.	opt.	c.k.	64-0131
		7	1.8×10^9 (rel.)	$k/k_{\text{carb}} = 4.8$	p.r.	opt.	c.k.	65-0190
		10.7	1.9×10^9 (rel.)	$k/k_{\text{carb}} = 5.1$	p.r.	opt.	c.k.	65-0190, 65-0387
		9.0	2.0×10^9 (rel.)	$k/k_{\text{RNO}} = 0.164 \pm 0.008$	γ -r.	opt.	c.k.	65-0356
		7	1.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.14$	p.r.	opt.	c.k.	65-0387
		9	1.85×10^9 (rel.)	$k/k_{\text{EtOH}} = 1$	γ -r.	opt.	c.k. with RNO.	66-0423
		2-2.2	1.9×10^9 (rel.)	$k/k_{\text{tbym}} = 0.35 \pm 0.03$	γ -r.	opt.	c.k.	67-046
		nat.	1.9×10^9 (rel.)	$k/k_{\text{ferro}} = 0.204$	p.r.	opt.	c.k.	71-0578
		—	2.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.18$	p.r.	opt.	c.k.	73-1077
		—	1.6×10^9 (rel.)	$k/k_{\text{ferro}} = 0.17$				
3.404	glycine, positive ion	1	8×10^6 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.035$	Fenton	chem.	c.k.	49-0002
		1	1.6×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.0015$	p.r.	opt.	c.k.	65-0387
		2.8	8.1×10^6 (rel.)	$k/k_{\text{tbym}} = 0.0015$	γ -r.	opt.	c.k.	65-0388
		2.8-3	1.0×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.00091$	p.r.	opt.	c.k.	65-0388
3.405	glycine, zwitterion	7.0	5.6×10^6 (rel.)	$k/k_{\text{ferro}} = 0.0006$	X-r.	opt.	c.k.; not cor. for H_2O_2 .	62-0023
		—	2.6×10^8 (rel.)	$k/k_{\text{ferro}} = 0.028$	phot.	—	c.k.	65-0247
		5.8-6	1.6×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.0015$	p.r.	opt.	c.k.	65-0388
		5	4.6×10^6 (rel.)	$k/k_{\text{EtOH}} = 0.0025$	γ -r.	opt.	c.k. with RNO.	66-0423
		6.7	1.7×10^7 (rel.)	$k/k_{\text{RNO}^-} = 0.00135$	γ -r.	opt.	c.k.	73-0548
3.406	glycine, negative ion	9.45	8.4×10^8 (rel.)	$k/k_{\text{ferro}} = 0.09$	X-r.	opt.	c.k.; not cor. for H_2O_2 .	62-0023
		10.5	2.8×10^9 (rel.)	$k/k_{\text{ferro}} = 0.3$	X-r.	opt.	c.k.; not cor. for H_2O_2 .	62-0023
		10.5	2.7×10^9 (rel.)	$k/k_{\text{BzO}^-} = 0.47$	γ -r.	trac.	c.k.; meas. $^{14}\text{CO}_2$.	65-0099
		9.5-9.7	1.9×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.17$	p.r.	opt.	c.k.	65-0388
		12	2.0×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.1$	γ -r.	opt.	c.k. with RNO.	66-0423
		10.0	5.3×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.485$	p.r.	opt.	c.k.	72-0461
		5.0	1.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.11$	p.r.	opt.	c.k.	71-0554
3.407	glycine anhydride	11.0	1.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.11$	p.r.	opt.	c.k.	71-0554
		9	7.1×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.386$	γ -r.	opt.	c.k. with RNO.	66-0423
3.408	glycolate ion	5.5, 7.0	$(8.6 \pm 0.7) \times 10^8$ (rel.)	$k/k_{\text{CNS}^-} = 0.078$	p.r.	opt.	c.k.	75-1053
3.409	glycolic acid	1	4.6×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 2.0$	Fenton	chem.	c.k.	49-0002
		2-2.2	4.6×10^8 (rel.)	$k/k_{\text{tbym}} = 0.085 \pm 0.005$	γ -r.	opt.	c.k.	67-0461
3.410	glycylalanine	2-2.2	1.8×10^8 (rel.)	$k/k_{\text{tbym}} = 0.0339$	γ -r.	opt.	c.k.	65-0388
		5.5-6	3.5×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.032$	p.r.	opt.	c.k.	65-0388
3.411	glycylglycine, positive ion	2-2.2	1.4×10^8 (rel.)	$k/k_{\text{thym}} = 0.0252$	γ -r.	opt.	c.k.	65-0388
		2.2-2.4	1.6×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0144$	p.r.	opt.	c.k.	65-0388
3.412	glycylglycine, zwitterion	6-7	2.6×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.024$	p.r.	opt.	c.k.	65-0387
		5.5-6	2.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0197$	p.r.	opt.	c.k.	65-0388
		4.2	4.4×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.04$	p.r.	opt.	c.k.	70-0099
3.413	glycylglycine, negative ion	10.5	5.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.47$	p.r.	opt.	c.k.	70-0099
3.414	glycylglycine amide	3.3	2.7×10^8	—	p.r.	opt.	p.b.k.	75-1004
3.415	glycylglycylglycine, positive ion	2-2.2	1.6×10^8 (rel.)	$k/k_{\text{thym}} = 0.029$	γ -r.	opt.	c.k.	65-0388
		2.8-3	2.4×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.022$	p.r.	opt.	c.k.	65-0388
3.416	glycylglycylglycine, zwitterion	5.5-6	1.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.16$	p.r.	opt.	c.k.	65-0388
		8.5-8.7	1.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.16$	p.r.	opt.	c.k.	65-0388
		5.4	7.3×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.066$	p.r.	opt.	c.k.	70-0099

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.417	glycylglycylglycine, negative ion 10.6	5.0 x 10 ⁹ (rel.)	$k/k_{\text{CNS}^-} = 0.455$	p.r.	opt.	c.k.	70-0099
3.418	glycylglycylglycylglycine, positive ion 2-2.2	2.4 x 10 ⁸ (rel.)	$k/k_{\text{thym}} = 0.045$	γ-r.	opt.	c.k.	65-0388
	2.4-2.6	3.5 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.032$	p.r.	opt.	c.k.	65-0388
	5.5-6	4.5 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.041$	p.r.	opt.	c.k.	65-0388
	7.7-7.9	1.2 x 10 ⁹ (rel.)	$k/k_{\text{CNS}^-} = 0.11$	p.r.	opt.	c.k.	65-0388
	negative ion 9.5-9.7	3.0 x 10 ⁹ (rel.)	$k/k_{\text{CNS}^-} = 0.27$	p.r.	opt.	c.k.	65-0388
3.419	glycylisoleucine positive ion 2-2.2	2.4 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 0.452$	γ-r.	opt.	c.k.	65-0388
3.420	glycylleucine positive ion 2-2.2	2.6 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 0.484$	γ-r.	opt.	c.k.	65-0388
3.421	glycylmethionine positive ion 2-2.2	4.4 x 10 ⁸ (rel.)	$k/k_{\text{thym}} = 0.081$	γ-r.	opt.	c.k.	65-0388
	2-2.2	1.1 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.00985$	p.r.	opt.	c.k.	65-0388
	zwitterion 5-5.2	2.2 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.0197$	p.r.	opt.	c.k.	65-0388
3.422	glycylphenylalanine, positive ion 2-2.2	8.9 x 10 ⁸ (rel.)	$k/k_{\text{thym}} = 0.165$	γ-r.	opt.	c.k.	65-0388
3.423	glycylproline, positive ion 2-2.2	1.5 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 0.27$	γ-r.	opt.	c.k.	65-0388
3.424	glycylserine, positive ion 2-2.2	5.9 x 10 ⁸ (rel.)	$k/k_{\text{thym}} = 0.11$	γ-r.	opt.	c.k.	65-0388
3.425	glycyltyrosine, positive ion 2-2.2	9.7 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 1.8$	γ-r.	opt.	c.k.	65-0388
3.426	glycylvaline, positive ion 2-2.2	1.2 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 0.226$	γ-r.	opt.	c.k.	65-0388
3.427	glyoxal OH + CHOCHO → H ₂ O + COCHO 1.3	—	$k/k_{\text{oxalic acid}} = \frac{46}{46}$	r.	chem.	c.k.	68-0503
3.428	guanine —	1.05 x 10 ¹⁰	—	—	—	—	66-0845
	10.0	9.2 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.74$	γ-r.	opt.	c.k.; 17°C.	75-0294
3.429	guanosine 9	7.6 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.61$	γ-r.	opt.	c.k.	67-0555
3.430	guanylic acid 6.7	(4.7 ± 0.2) x 10 ⁹	—	p.r.	opt.	p.b.k. at 325 nm.	70-3069
3.431	hemin —	~ 1.0 x 10 ¹⁰	—	—	—	—	66-0844
3.432	hemoglobin —	3.6 x 10 ¹⁰	—	—	—	—	66-0844
3.433	heparin —	3.6 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.033$	p.r.	opt.	c.k.; concn. of polyanion in hexose units.	68-0352, 70-3081
3.434	heparin, desulfated —	8.0 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.073$	p.r.	opt.	c.k.; concn. of polyanion in hexose units.	70-3081
3.435	1-heptanol 2-2.2	6.2 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 1.15 \pm 0.10$	γ-r.	opt.	c.k.	67-0461
3.436	hexadecyltrimethylammonium bromide —	1.1 x 10 ¹⁰ (rel.)	$k/k_{\text{MeOH}} = 11.8$	p.r.	opt.	c.k.; meas. Br ₂ ⁻ at 360 nm; concn. < 9 x 10 ⁻⁴ M; at higher concn. ratio = 2.4.	71-0001 71-0586
3.437	2,4-hexadien-1-ol 7.0	(9.8 ± 1.0) x 10 ⁹	—	p.r.	opt.	p.b.k.	73-1070
3.438	hexafluorobenzene —	2 x 10 ⁹ (rel.)	—	p.r.	opt.	c.k. with CNS ⁻ ; 280 nm. abs. grows in at same rate as condy. (F ⁻).	73-0054
3.438a	OH + C ₆ F ₆ → addn. → F ⁻ + H ⁺ + •C ₆ F ₅ =O	~7	7.2 x 10 ⁹	—	—	—	—
3.438a	hexamethylbenzene ~7	7.2 x 10 ⁹	—	p.r.	opt.	p.b.k.; OH addn. as well as H abstr. (< 50%).	75-1009
3.439	1,6-hexanediol 9	4.6 x 10 ⁹ (rel.)	$k/k_{\text{EtOH}} = 2.46$	γ-r.	opt.	c.k. with RNO.	66-0423

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.440	hexanoate ion 9	3.9×10^9 (rel.)	$k/k_{\text{EtOH}} = 2.1$	γ -r.	opt.	c.k. with RNO.	66-0423
3.441	1-hexanol 2-2.2	5.9×10^9 (rel.)	$k/k_{\text{thym}} = 1.10 \pm 0.10$	γ -r.	opt.	c.k.	67-0461
3.442	histidine 2-2.2	1.9×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.174$	p.r.	opt.	c.k.	65-0388
	6-7	5.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.455$	p.r.	opt.	c.k.	65-0388
	6.7	4.3×10^9 (rel.)	$k/k_{\text{RNO}} = 0.34$	γ -r.	opt.	c.k.	73-0548
3.443	histidylhistidine 5.5-6.5	9.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.82$	p.r.	opt.	c.k.	65-0388
3.444	hyaluronic acid —	1.3×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.116$	p.r.	opt.	c.k.; based on disaccharide unit.	67-0730
	—	6.7×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.061$	p.r.	opt.	c.k.; concn. of polyanion in hexose units.	68-0352, 70-3081
3.445	hyaluronic acid, sulfated —	6.0×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.055$	p.r.	opt.	c.k.; concn. of polyanion in hexose units.	70-3081
3.446	hydroquinone 6-7	2.1×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.88$	p.r.	opt.	c.k.	65-0387
	$\text{OH} + \text{HOC}_6\text{H}_4\text{OH} \rightarrow \text{C}_6\text{H}_4(\text{OH})_3$ —	1×10^{10} (rel.)	$k/k_{\text{MeOH}} = 10.8$	p.r.	opt.	c.k. with HSO_4^- ; obs. decreased abs. at 450 nm.	66-0019
	9	5.5×10^9 (rel.)	$k/k_{\text{RNO}} = 0.44 \pm 0.12$	γ -r.	opt.	c.k.	72-0837
3.447	hydroxocobalamin —	$\sim 10^{10}$	—	—	—	c.k. with RNO.	72-3046
3.447a	hydroxyacetamide 8.5	$(1.1 \pm 0.1) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = 0.1$	p.r.	opt.	c.k.	75-1053
	<i>o</i> -hydroxybenzaldehyde See salicylaldehyde (3.668a).						
3.448	<i>p</i> -hydroxybenzaldehyde 9	1.0×10^9 (rel.)	$k/k_{\text{RNO}} = 0.82 \pm 0.2$	γ -r.	opt.	c.k.	72-0837
	<i>o</i> -hydroxybenzoate ion See salicylate ion (3.669).						
3.449	<i>p</i> -hydroxybenzoate ion 9	5.5×10^9 (rel.)	$k/k_{\text{EtOH}} = 2.95$	γ -r.	opt.	c.k. with RNO.	66-0441
	7	$(9 \pm 2) \times 10^9$	—	p.r.	opt.	p.b.k. at 375; cor. for (OH + OH) and (H + aromatic).	68-0304
	$\text{OH} + \text{OHC}_6\text{H}_4\text{COO}^- \rightarrow (\text{OH})_2\text{C}_6\text{H}_4\text{COO}^-$						
3.450	2-hydroxybutyric acid 9	8.7×10^9 (rel.)	$k/k_{\text{RNO}} = 0.70 \pm$	γ -r.	opt.	c.k.	72-0837
	1	6.9×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 3.0$	Fenton	chem.	c.k.	49-0002
3.451	2-hydroxyethyl acetate —	8.5×10^8 (rel.)	$k/k_{\text{CNS}^-} = 8.25 \times 10^{-2}$	p.r.	opt.	c.k.	75-1126
3.452	2-hydroxyethyl-ethylenediamine-triacetic acid ~ 0	—	$k/k_{\text{acrylamide}} = 1.9$	Fenton	pol.	c.k.	72-9162
3.453	1-(2-hydroxyethyl)-2-methyl-5-nitroimidazole —	4.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.37$	p.r.	opt.	c.k.; d.k. at 320 nm gave $k \cong 10^9$.	74-1135
	—	$(5.5 \pm 0.2) \times 10^9$	—	p.r.	opt.	p.b.k. at 420 nm.	75-1067
3.454	2-hydroxyethyl-sulfide ion 11	4.0×10^9 ($\pm 15\%$)	—	p.r.	opt.	p.b.k. at 410-420 nm (RSSR $^-$).	69-0553
	$\text{OH} + \text{OH}(\text{CH}_2)_2\text{S}^- \rightarrow \text{OH}^- + \text{OH}(\text{CH}_2)_2\text{S}^\bullet$						
3.454a	1-(2-hydroxy-3-methoxypropyl)-2-nitroimidazole —	$(7.1 \pm 0.3) \times 10^9$	—	p.r.	opt.	p.b.k. at 475 nm.	75-1067
3.455	5-hydroxyindole 9.0	$(1.67 \pm 0.10) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.456	<i>m</i> -hydroxyphenol 9	1.2×10^{10} (rel.)	$k/k_{\text{RNO}} = 0.97 \pm 0.12$	γ -r.	opt.	c.k.	72-0837
3.457	<i>o</i> -hydroxyphenol 9	1.1×10^{10} (rel.)	$k/k_{\text{RNO}} = 0.89 \pm 0.14$	γ -r.	opt.	c.k.	72-0837
	<i>p</i> -hydroxyphenol See hydroquinone (3.446).						

TABLE 4. Reactions of OH with organic solutes – Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.458	<i>p</i> -hydroxyphenyl- β-D-glucopyrano- side	—	2.7 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> = 4.4 x 10 ⁹ for phenyl β-D-glucopyrano- side.	71-0056
3.459	<i>p</i> -hydroxyphenyl- propionate ion	10.6	2.1 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{carb} = 58.5	p.r.	opt.	c.k.	68-0062
		6.3	(1.2 ± 0.2) x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.1	p.r.	opt.	c.k.	73-0003
		11.0	(1.6 ± 0.2) x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.45	p.r.	opt.	c.k.	73-0003
3.460	<i>p</i> -hydroxyphenyl- propionic acid	4.6	1.3 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{HCOO⁻} = 3.7	p.r.	opt.	c.k.; p <i>K</i> _a = 4.6, 10.1.	68-0062
3.461	hydroxyproline	2-2.2	3.6 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.066	γ-r.	opt.	c.k.	65-0388
		6.8	3.2 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.0255	γ-r.	opt.	c.k.	73-0548
3.461a	2-hydroxypro- pionamide	4.5, 7.0	(1.3 ± 0.3) x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.12	p.r.	opt.	c.k.	75-1053
3.461b	2-hydroxypurine	6-7	5.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.4	γ-r.	opt.	c.k.; 17°C.	75-0294
3.462	2-hydroxypyridine, anion	9	4.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.38 ± 0.01	γ-r.	opt.	c.k.	69-0280
3.463	3-hydroxypyridine	6.5	6.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.54 ± 0.03	γ-r.	opt.	c.k.	69-0280
3.464	3-hydroxypyridine, anion	9	5.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.43 ± 0.02	γ-r.	opt.	c.k.	69-0280
3.465	4-hydroxypyridine anion	9	2.75 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.23 ± 0.01	γ-r.	opt.	c.k.	69-0280
3.466	α-hydroxytetronate ion	7	4.7 x 10 ⁹ (rel.)	—	p.r.	opt.	p.b.k. at 360 nm.	74-1053
3.466a	hypoxanthine	6-7	2.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.22	γ-r.	opt.	c.k.; 17°C.	75-0294
3.467	imidazole	3.4	5.5 x 10 ⁹	—	p.r.	opt.	p.b.k.; OH addn.;	75-1066
		6.8	8.7 x 10 ⁹				p <i>K</i> _a = 7.1, 14.5.	
		10.9	1.2 x 10 ¹⁰					
3.468	indole	9.0	(1.37 ± 0.05) x 10 ¹⁰ (rel.)	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> (OH + tryptophan) = 1.25 x 10 ¹⁰ .	71-0556
		9.0	(3.18 ± 0.25) x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 320 nm.	71-0556
		—	1.3 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 5.9	γ-r.	chem.	c.k.	72-0541
3.469	indole-3-acetic acid	—	1.1 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 5.0	γ-r.	chem.	c.k.	72-0541
3.470	indole-5-acetic acid	9.0	(0.79 ± 0.07) x 10 ¹⁰ (rel.)	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> (OH + tryptophan) = 1.25 x 10 ¹⁰ .	71-0556
3.471	indole-3-propionic acid	—	1.4 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 6.5	γ-r.	chem.	c.k.	72-0541
3.472	indoline	9.0	(2.02 ± 0.14) x 10 ¹⁰ (rel.)	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> (OH + tryptophan) = 1.25 x 10 ¹⁰ .	71-0556
3.473	inositol	9.0	(3.83 ± 0.48) x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 320 nm.	71-0556
		6.5	1.0 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.8 ± 0.1	γ-r.	opt.	c.k.	69-0580
		—	1.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.13	p.r.	opt.	c.k.	73-1077
3.473a	iodoacetic acid	1	1.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{ferro} = 0.19	p.r.	opt.	c.k.	74-5286
3.474	iodobenzene	9	5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.40 ± 0.02	γ-r.	opt.	c.k.	69-0280
3.475	2-iodobenzoate ion	9	4.5 x 10 ⁹ (rel.)	—	—	—	c.k. with RNO.	66-0843
3.476	3-iodobenzoate ion	9	2.9 x 10 ⁹ (rel.)	—	—	—	c.k. with RNO.	66-0843

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.477	4-iodobenzoate ion	9	2.5×10^9 (rel.)	—	—	—	c.k. with RNO.	66-0843
3.478	iodomethane $\text{OH} + \text{CH}_3\text{I} \rightarrow$ $\text{CH}_3\text{OH} + \text{I}$	—	1.4×10^9 (rel.)	$k/k_{\text{MeOH}} = 1.54$	γ -r.	chem.	c.k.; meas. I_2 yields.	69-0019
3.479	3-iodopropionic acid	—	$(1.2 - 4.0) \times 10^8$ (rel.)	$k/k_{\text{CNS}^-} = (1.1 - 3.6) \times 10^{-2}$	p.r.	opt.	c.k.	70-1226
			1.6×10^8 (rel.)	$k/k_{\text{NB}} = 0.051$	p.r.	opt.	c.k.	70-1226
3.480	isoamyl alcohol isoamylammonium ion	See 3-methyl-1-butanol (3.527). 4	7.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.714$	p.r.	opt.	c.k.	70-0371
3.481	isobutanol isobutylammonium ion	See 2-methyl-1-propanol (3.545). 4	3.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.278$	p.r.	opt.	c.k.	70-0371
3.482	isobutylene	—	5.9×10^9 (rel.)	$k/k_{\text{I}^-} = 0.49$	p.r.	opt.	c.k.; meas. I_2^- at 400 nm.	67-0041
3.483	isobutyramide	5-6	1.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.143$	p.r.	opt.	c.k.	71-0414
3.483a	isoguanine	11.0	1.23×10^{10} (rel.)	$k/k_{\text{RNO}} = 0.98$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.484	isoleucine	2-2.2	1.8×10^9 (rel.)	$k/k_{\text{thym}} = 0.34$	γ -r.	opt.	c.k.	65-0388
		6.6	1.7×10^9 (rel.)	$k/k_{\text{RNO}} = 0.14$	γ -r.	opt.	c.k.	73-0548
3.485	isoorotate ion	7	4.0×10^9	—	p.r.	opt.	p.b.k. (OH adduct).	70-0567
		6-7	2.5×10^9 (rel.)	$k/k_{\text{RNO}} = 0.2$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.486	isopropanol isopropyl acetate	See 2-propanol (3.637). 1	2.9×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 1.25$	Fenton	chem.	c.k.	49-0002
		6-7	4.5×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.041$	p.r.	opt.	c.k.	65-0387
		2.0	4.3×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0395$	p.r.	opt.	c.k.	65-0387
3.487	isopropylamine $\text{OH} + (\text{CH}_3)_2\text{CHNH}_2 \rightarrow$ $\text{H}_2\text{O} + (\text{CH}_3)_2\text{CNH}_2$	12.0	1.3×10^{10} (rel.)	—	p.r.	opt.	c.k.; value extrapolated from obs. $k/k_{\text{CNS}^-} = 8.2 \times 10^{-1}$ at pH 10.8.	71-0585
3.488	isopropylammonium ion $\text{OH} + (\text{CH}_3)_2\text{CHNH}_3^+ \rightarrow$ $\text{H}_2\text{O} + \text{CH}_2(\text{CH}_3)\text{CHNH}_3^+$	3.0 4	5.0×10^8 (rel.) 4.7×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0455$ $k/k_{\text{CNS}^-} = 0.0429$	p.r. p.r.	opt. opt.	c.k. c.k.	71-0585 70-0371
3.489	keratan sulfate	—	7.9×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.072$	p.r.	opt.	c.k.	71-0067
3.490	lactate ion	9	4.8×10^9 (rel.)	—	—	—	c.k. with RNO.	66-0843
		—	7×10^8	—	p.r.	—	prelim. value.	74-1007
3.491	lactic acid	1	3.9×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 1.7$	Fenton	chem.	c.k.	49-0002
		2-2.2	6.5×10^8 (rel.)	$k/k_{\text{thym}} = 0.12 \pm 0.01$	γ -r.	opt.	c.k.	67-0461
		1	4.3×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0393$	p.r.	opt.	c.k.	65-0387
3.492	lactose	6.5	2.4×10^9 (rel.)	$k/k_{\text{RNO}} = 0.19 \pm 0.02$	γ -r.	opt.	c.k.	69-0580
3.493	leucine, positive ion	2-2.2	1.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.14$	p.r.	opt.	c.k.	65-0388
		2-2.2	2.0×10^9 (rel.)	$k/k_{\text{thym}} = 0.37$	γ -r.	opt.	c.k.	65-0388
3.494	leucine, zwitterion	5.5-6	1.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.15$	p.r.	opt.	c.k.	65-0388
		6.9	1.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.145$	γ -r.	opt.	c.k.	73-0548
3.495	leucine, negative ion	9.7-9.9	3.7×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.34$	p.r.	opt.	c.k.	65-0388
3.496	luminol	9.5	8.7×10^9	—	p.r.	opt.	p.b.k.	73-1068
3.497	lysine	2-2.2	6.5×10^8 (rel.)	$k/k_{\text{thym}} = 0.12$	γ -r.	opt.	c.k.	65-0388
		6.6	3.5×10^8 (rel.)	$k/k_{\text{RNO}} = 0.028$	γ -r.	opt.	c.k.	73-0548
3.497a	lysine vasopressin	~6	$(1.4 \pm 0.2) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 325 nm.	74-1102
3.498	lysozyme	9	1.9×10^{10} (rel.)	$k/k_{\text{RNO}} = 1.49$	γ -r.	opt.	c.k.	67-0555
		5.6	5.2×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 4.7$	p.r.	opt.	c.k.; mol. wt. 15,000; k is upper limit.	68-0683
		7.4	4.9×10^{10}	—	p.r.	opt.	p.b.k. at 350 nm.	69-3039

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.498 cont.	7.4	—	$k/k_{t-BuOH} = 56$	—	—	c.k.	69-3039
	6.4	4.2×10^{10} (rel.)	$k/k_{RNO} = 3.4$	γ -r.	opt.	c.k.	73-0548
3.499	malate ion	—	—	p.r.	—	prelim. value.	74-1007
3.500	maleic acid	1	4.6×10^8 (rel.)	$k/k_{MeOH} = 0.515$	Fenton	chem. c.k.; $k_{MeOH}/k_{Fe^{2+}} = 4.3$.	73-9341
3.501	malic acid	2-2.2	5.4×10^8 (rel.)	$k/k_{thym} = 0.10 \pm 0.01$	γ -r.	opt. c.k.	67-0461
3.502	malonate ion	9	5.5×10^7 (rel.)	$k/k_{EtOH} = 0.0296$	γ -r.	opt. c.k. with RNO.	66-0423
3.503	malonic acid	6-7	3.0×10^8 (rel.)	$k/k_{CNS^-} = 0.0273$	p.r.	opt. c.k.	65-0387
		2-2.2	2.0×10^7 (rel.)	$k/k_{thym} = 0.0037$	γ -r.	opt. c.k.	67-0461
		1	2.6×10^7 (rel.)	$k/k_{MeOH} = 0.017$	Fenton	chem. c.k.; $k_{MeOH}/k_{Fe^{2+}} = 4.3$.	73-9341
3.504	melibiose	6.5	3.8×10^9 (rel.)	$k/k_{RNO} = 0.3 \pm 0.1$	γ -r.	opt. c.k.	69-0580
3.505	menaquinone (Vitamin K ₃)	—	5.5×10^9	—	—	—	73-0026
3.506	2-mercaptoacetate ion	<i>See</i> thioglycolate ion (3.705).					
	2-mercaptoethanol	6-7	8.5×10^9 (rel.)	$k/k_{CNS^-} = 0.773$	p.r.	opt. c.k.	65-0387
	OH + OH(CH ₂) ₂ SH →	7	2.7×10^{10} (rel.)	$k/k_{CNS^-} = 2.43$	p.r.	opt. c.k.	69-0553
	H ₂ O + OH(CH ₂) ₂ S•	6.5	6.0×10^9 (rel.)	$k/k_{ferro} = 0.65$	p.r.	opt. c.k.	71-0175
		6	3.3×10^9 (rel.)	$k/k_{NB} = 1.04$	p.r.	opt. c.k.; cor. for H.	71-0175
		6	1.8×10^{10} (rel.)	$k/k_{CNS^-} = 1.68$	p.r.	opt. c.k.	71-0175
	<i>See also</i> 2-hydroxyethylsulfide ion (3.453).						
	2-mercaptoethylamine	<i>See</i> cysteamine (3.286).					
	2-mercaptopropionate ion	<i>See</i> thiolactate ion (3.706).					
3.508	3-mercaptopropionate ion	6.0	3.0×10^{10} (rel.)	$k/k_{CNS^-} = 2.7 \pm 0.4$	p.r.	opt. c.k.; $pK_a = 4.3$, 10.3 for the acid.	73-0090
		10.7	2.1×10^{10} (rel.)	$k/k_{CNS^-} = 1.9 \pm 0.3$			
3.509	2-mercaptovaline	<i>See</i> penicillamine (3.596).					
	methane	9	2.4×10^8 (rel.)	$k/k_{EtOH} = 0.13$	γ -r.	opt. c.k. with RNO.	66-0423
	OH + CH ₄ →	5.5	$(1.21 \pm 0.4) \times 10^8$	—	p.r.	opt. d.k. (OH) at 250 nm.	72-0445
3.509a	methanesulfonic acid	—	1.3×10^9 (rel.)	$k/k_{CNS^-} = 0.114$	p.r.	opt. c.k.	75-1072
3.510	methanethiol	7	3.3×10^{10} (rel.)	$k/k_{CNS^-} = 3.04$	p.r.	opt. c.k.	69-0553
	OH + CH ₃ SH →						
	H ₂ O + CH ₃ S						
	<i>See also</i> methylsulfide ion (3.553).						
3.511	methanol (MeOH)	1	5.3×10^8 (rel.)	$k/k_{Fe^{2+}} = 2.3$	Fenton	chem. c.k.	49-0002
	(I) OH + CH ₃ OH →	7	6.0×10^8 (rel.)	$k/k_I = 0.046 \pm 0.004$	p.r.	opt. c.k.; meas. I ₂ at 400 nm.	65-0010
	H ₂ O + CH ₂ OH						
	(II) OH + CH ₃ OH →	10.5	9.7×10^8 (rel.)	$k/k_{BzO^-} = 0.17$	γ -r.	trac. c.k.; formn. of ¹⁴ CO ₂ .	65-0099
	H ₂ O + CH ₃ O						
		7	8.8×10^8 (rel.)	$k/k_{carb} = 2.4$	p.r.	opt. c.k.	65-0190
		10.7	8.4×10^8 (rel.)	$k/k_{carb} = 2.3$	p.r.	opt. c.k.	65-0190
		7.0	7.7×10^8 (rel.)	$k/k_{CNS^-} = 0.07$	p.r.	opt. c.k.	65-0190
		—	5.7×10^8 (rel.)	$k/k_{ferro} = 0.061$	phot.	— c.k.	65-0247
		9.0	1.1×10^9 (rel.)	$k/k_{RNO} = 8.6 \pm 0.4) \times 10^{-2}$	γ -r.	opt. c.k.	65-0356
		2	7.4×10^8 (rel.)	$k/k_{CNS^-} = 0.067$	p.r.	opt. c.k.	65-0387
		7	8.0×10^8 (rel.)	$k/k_{CNS^-} = 0.073$	p.r.	opt. c.k.	65-0387
		4.5	3×10^8	—	γ -r.	chem. est. from yields in carboxylation of methanol.	65-0375
		2-2.2	8.6×10^8 (rel.)	$k/k_{thym} = 0.16 \pm 0.015$	γ -r.	opt. c.k.	65-0388, 67-0461
		5-5.5	9.4×10^8 (rel.)	$k/k_{thym} = 0.175 \pm 0.015$	γ -r.	opt. c.k.	65-0388, 67-0461

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.511 cont.		—	—	$k/k_{\text{bisulf}} = 550$	p.r.	opt.	c.k.; obs. formn. of SO_4^- at 450 nm.	66-0019
		6	8.6×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.464$	γ -r.	chem.	c.k. with Br^- .	66-0423
		9	1.0×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.555$	γ -r.	opt.	c.k. with RNO.	66-0423
		—	5.5×10^8 (rel.)	$k/k_{\text{I}^-} = 0.046$	p.r.	opt.	c.k.; meas. I_2^- at 400 nm.	67-0041
		2	2.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.22$	Fenton	opt.	c.k.	67-0555
		—	—	$k/k_{\text{TCOO}^-} = 0.3$	γ -r.	trac.	c.k.; meas. ^3HHO produced.	68-0209
		9	1×10^9 (rel.)	$k/k_{\text{RNO}} = 0.08$	γ -r.	opt.	c.k.	68-0310
		—	8.0×10^8 (rel.)	$k/k_{\text{BzO}^-} = 0.14$	p.r.	opt.	c.k.; obs.	68-0304
			8.3×10^8 (rel.)	$k/k_{\text{PA}^-} = 0.105$			hydroxycyclohex-	
			8.3×10^8 (rel.)	$k/k_{\text{PNBA}^-} = 0.32$			adienyl radical buildup.	
		~1.2	6.4×10^8 (rel.)	$k/k_{2\text{-PrOH}} = 0.29$	γ -r.	chem.	c.k.	68-0602
		6.98	7.9×10^8 (rel.)	$k/k_{2\text{-PrOH}} = 0.36$	γ -r.	chem.	c.k.; $\mu = 0.1$; ratio = 0.34 at $\mu = 1.1$.	68-0602
		—	9.5×10^8 (rel.)	$k/k_{\text{RNO}} = 0.076$	p.r.	opt.	c.k.	69-0156
		1	1.1×10^9 (rel.)	$k/k_{\text{thym}} = 0.206$	Fenton	esr	c.k.; $k/k_{\text{perox}} = 15.0$.	69-5278
		1	9.8×10^8 (rel.)	$k/k_{\text{thym}} = 0.18$	Ti(III) - H_2O_2	esr	c.k.; $k/k_{\text{perox}} = 12.8$.	69-5278
		9	1.1×10^9 (rel.)	$k/k_{\text{RNO}} = 0.09$	γ -r.	opt.	c.k.; $E_a = -1.9 \pm 0.08$ kcal/mol (7.9 kJ/mol) at -8 to 23°C.	71-0469
		nat.	8.6×10^8 (rel.)	$k/k_{\text{ferro}} = 0.0925$	p.r.	opt.	c.k.	71-0578
		0.82	9.9×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 4.30$	Fenton	chem.	c.k.	71-9132
		—	—	$k_{\text{II}}/k_{\text{I}} = 0.075$	p.r.	opt.	hydroxymethyl radical identified by reaction with TNM, methoxy radical by I^- .	73-0126
		~1	1.3×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.63$	γ -r.	chem.	obs. effect of alcohols on oxid. Sb(III) \rightarrow Sb(IV).	73-0289
		10.4	1.2×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.62$	X-r.	lum.	obs. effect. of alcohols on quenching chemiluminescence from fluorescein.	73-6068
<i>For other ratios see: 3.8, 3.10, 3.12, 3.25, 3.27, 3.41, 3.50, 3.54, 3.58, 3.66, 3.71, 3.80, 3.82, 3.88, 3.90, 3.91, 3.100, 3.102, 3.103, 3.106, 3.107, 3.112, 3.129, 3.131, 3.132, 3.144, 3.225, 3.246, 3.271, 3.304, 3.319, 3.358, 3.368, 3.386, 3.436, 3.446, 3.478, 3.500, 3.503, 3.546, 3.592, 3.593, 3.636, 3.637, 3.669, 3.673, 3.680, 3.698, 3.711, 3.755.</i>								
3.512	methanol- d_3 $\text{OH} + \text{CD}_3\text{OH} \rightarrow \text{HDO} + \text{CD}_2\text{OH}$	6	4.2×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.227$	p.r.	chem.	c.k. with Br^- .	66-0423
3.513	methionine	6-7	8.5×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.77$	p.r.	opt.	c.k.	65-0387
		2-2.2	6.5×10^9 (rel.)	$k/k_{\text{thym}} = 1.2$	γ -r.	opt.	c.k.	65-0388
		5.5-5.7	8.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.74$	p.r.	opt.	c.k.	65-0388
		6.6	6.5×10^9 (rel.)	$k/k_{\text{RNO}} = 0.52$	γ -r.	opt.	c.k.	73-0548
3.514	methoxyacetate ion	9	6.0×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.323$	γ -r.	opt.	c.k. with RNO.	66-0423
3.515	<i>p</i> -methoxybenzoate ion	9	5.0×10^9 (rel.)	$k/k_{\text{EtOH}} = 2.7$	γ -r.	opt.	c.k. with RNO.	66-0441
3.516	2-methoxyethanol	9	1.3×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.7$	γ -r.	opt.	c.k. with RNO.	66-0423

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.517	5-methoxyindole	9.0	$(1.39 \pm 0.04) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) =$ $(1.25 \pm 0.3) \times 10^{10}$.	71-0556
3.517a	1-methoxy-2-methyl-1-phenylpropane	1.7-1.8	8.6×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 3.9$	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.518	<i>o</i> -methoxyphenol	9	1.5×10^{10} (rel.)	$k/k_{\text{RNO}} = 1.18 \pm 0.15$	γ -r.	opt.	c.k.	72-0837
3.519	<i>p</i> -methoxyphenol	9	1.45×10^{10} (rel.)	$k/k_{\text{RNO}} = 1.15 \pm 0.23$	γ -r.	opt.	c.k.	72-0837
3.520	<i>p</i> -methoxyphenyl- β -D-glucopyranoside	—	7.0×10^9 (rel.)	—	p.r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{X}) = 4.4 \times 10^9$ where $\text{X} = \text{phenyl-}\beta\text{-D-glucopyranoside}$.	71-0056
3.521	<i>N</i> -methylacetamide $\text{OH} + \text{CH}_3\text{CONHCH}_3 \rightarrow \text{CH}_3\text{CONHCH}_2 + \text{H}_2\text{O}$	—	1.1×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.04$	p.r.	opt.	c.k.	71-0056
		5.5	1.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.146$	p.r.	opt.	c.k.	70-0098, 71-0645
3.522	methyl acetate	1	2×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.85$	Fenton	chem.	c.k.	49-0002
		6-7	1.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0106$	p.r.	opt.	c.k.	65-0387
		2.0	1.3×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0121$	p.r.	opt.	c.k.	65-0387
		9	1.1×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.0595$	γ -r.	opt.	c.k. with RNO.	66-0423
		12	2.4×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.3$	γ -r.	opt.	c.k. with RNO.	66-0423
3.523	methylamine $\text{OH} + \text{CH}_3\text{NH}_2 \rightarrow \text{H}_2\text{O} + \text{CH}_2\text{NH}_2 + \text{CH}_3\text{NH}$	11.5-	4.7×10^9 (rel.)	$k/k_{\text{NB}} = 1.47$	p.r.	opt.	c.k.	69-0573
		12.5						
		10.5	1.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.16$	p.r.	opt.	c.k.	71-0595
		11.1	3.3×10^9 (rel.)	$k/k_{\text{NB}} = 1.04$	p.r.	opt.	c.k.; at pH 9.7, 12.8 ratio = 0.13 and 1.5, resp.	71-0595
3.524	methylammonium ion	5	1.9×10^7 (rel.)	$k/k_{\text{EtOH}} = 0.0103$	γ -r.	opt.	c.k. with RNO.	66-0423
		6-8	7.5×10^7 (rel.)	$k/k_{\text{NB}} = 0.0234$	p.r.	opt.	c.k.	69-0573
		2	2.8×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.0025$	p.r.	opt.	c.k.	70-0371
		4	3.5×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.0032$	p.r.	opt.	c.k.	70-0371
		7	5.9×10^7 (rel.)	$k/k_{\text{CNS}^-} = 0.0054$	p.r.	opt.	c.k.	70-0371
3.525	methylarabinoside	6.5	2.4×10^9 (rel.)	$k/k_{\text{RNO}} = 0.19 \pm 0.04$	γ -r.	opt.	c.k.	69-0580
3.526	2-methyl-2-butanol	9	1.85×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.0$	γ -r.	opt.	c.k. with RNO.	66-0423
3.527	3-methyl-1-butanol (isoamyl alcohol)	—	3.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.34$	p.r.	opt.	c.k.	73-1077
			3.4×10^9 (rel.)	$k/k_{\text{ferro}} = 0.36$				
3.528	methyl butyrate	6-7	1.7×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.152$	p.r.	opt.	c.k.	65-0387
3.529	2-methylbutyrate ion	9	2.2×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.18$	γ -r.	opt.	c.k. with RNO.	66-0423
3.530	3-methylbutyrate ion	9	2.3×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.27$	γ -r.	opt.	c.k. with RNO.	66-0423
3.531	3-methylbutyric acid	1	7.6×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 3.3$	Fenton	chem.	c.k.	49-0002
3.532	<i>S</i> -methylcysteine	5.4	8.0×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.73$	—	—	c.k.; $pK_a \cong 2.8.8$.	73-0090
		11.0	7.9×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.72$				
3.533	5-methylcytosine	4.2	$(4.7 \pm 0.5) \times 10^9$	—	p.r.	opt.	p.b.k. at 450 nm.	70-3069
		6-7	5.2×10^9 (rel.)	$k/k_{\text{RNO}} = 0.42$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.534	methylene blue $\text{OH} + \text{MB}^+ \rightarrow \text{MB}^{2+} + \text{OH}^-$	—	4.1×10^{10} (rel.)	$k/k_{\text{EtOH}} = 22$	γ -r.	chem.	c.k.; obs. $G(-\text{MB}^+)$.	71-0682
3.535	<i>N</i> -methylformamide	5.5	1.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.11$	p.r.	opt.	c.k.	70-0098
3.536	methylgalactoside	6.5	1.6×10^9 (rel.)	$k/k_{\text{RNO}} = 0.13 \pm 0.01$	γ -r.	opt.	c.k.	69-0580
3.537	methylglucoside	6.5	2.4×10^9 (rel.)	$k/k_{\text{RNO}} = 0.19 \pm 0.02$	γ -r.	opt.	c.k.	69-0580
3.538	<i>O</i> -methylhydroxylamine	4.5	$\leq 4.0 \times 10^8$ (rel.)	$k/k_{\text{CNS}^-} = 0.036$	p.r.	opt.	c.k.	71-0493
		9.1	1.4×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.3$	p.r.	opt.	c.k.	71-0493

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.538a	<i>N</i> -methylimidazole 5.4	5.0×10^9	—	p.r.	opt.	p.b.k.; $pK_a = 7.0$; OH addn.	75-1066
	9.4	8.1×10^9					
3.539	1-methylindole 9.0	$(1.45 \pm 0.01) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.540	2-methylindole 9.0	1.44×10^{10} (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
	9.0	$(3.41 \pm 0.28) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 320 nm.	71-0556
3.541	3-methylindole 9.0	$(1.05 \pm 0.09) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
	9.0	$(3.34 \pm 0.08) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 320 nm.	71-0556
3.542	5-methylindole 9.0	$(1.66 \pm 0.06) \times 10^{10}$ (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{tryptophan}) = 1.25 \times 10^{10}$.	71-0556
3.543	<i>N</i> -methylisobutyramide 5-6	1.9×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.176$	p.r.	opt.	c.k.	71-0414
	methyl mercaptan <i>See</i> methanethiol (3.510).						
3.543a	2-methyl-4-phenyl-2-butanol 1.7-1.8	6.8×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 3.1$	Fenton	chem.	c.k. with cycloheptanol.	74-9006
3.543b	2-methyl-5-phenyl-2-pentanol 1.7-1.8	$< 4.4 \times 10^7$ (rel.)	$k/k_{2-\text{PrOH}} = < 0.02$	Fenton	chem.	c.k. with 3-pentanol.	74-9006
3.543c	2-methyl-1-phenyl-1-propanol 1.7-1.8	1.1×10^{10} (rel.)	$k/k_{2-\text{PrOH}} = 5.0$	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.543d	2-methyl-1-phenyl-1-propanol-1- <i>d</i> 1.7-1.8	9.9×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 4.5$	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.543e	2-methyl-1-phenyl-2-propanol 1.7-1.8	2.0×10^{10} (rel.)	$k/k_{2-\text{PrOH}} = 9.0$	Fenton	chem.	c.k. with cycloheptanol.	74-9006
3.544	<i>N</i> -methylpivalamide 5-6	2.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.218$	p.r.	opt.	c.k.	71-0414
3.545	2-methyl-1-propanol 7	3.3×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.303$	p.r.	opt.	c.k.	65-0387
	9	3.5×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.9$	γ -r.	opt.	c.k. with RNO.	66-0423
	(isobutanol) 2-2.2	3.8×10^9 (rel.)	$k/k_{\text{thym}} = 0.70 \pm 0.05$	γ -r.	opt.	c.k.	67-0461
	OH + $(\text{CH}_3)_2\text{CHCH}_2\text{OH} \rightarrow (\text{CH}_3)_2\text{CHCHOH}$ —	3.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.33$	p.r.	opt.	c.k.	73-1077
	(75%, 69-0522)	2.6×10^9 (rel.)	$k/k_{\text{ferro}} = 0.28$				
	+ H ₂ O + CH ₃ CH(CH ₂)CH ₂ OH, etc.						
3.546	2-methyl-2-propanol (<i>tert</i> -butanol) 1	1.4×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.0065$	Fenton	chem.	c.k.	49-0002
	7	4.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.038$	p.r.	opt.	c.k.	65-0387
	9	4.6×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.25$	γ -r.	opt.	c.k. with RNO.	66-0423
	(I) OH + 2-2.2	7.3×10^8 (rel.)	$k/k_{\text{thym}} = 0.135 \pm 0.015$	γ -r.	opt.	c.k.	67-0461
	(CH ₃) ₃ COH \rightarrow (CH ₃) ₂ COHCH ₂ + H ₂ O nat.	5.2×10^8 (rel.)	$k/k_{\text{ferro}} = 0.056$	p.r.	opt.	c.k.	71-0578
	0.82	4.3×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 1.90$	Fenton	chem.	c.k.	71-9132
	(II) OH + — —	—	$k_{11}/k_1 = 0.045$	p.r.	opt.	detd. % alkoxy radical by reaction with I ⁻ .	73-0126
	(CH ₃) ₃ COH \rightarrow H ₂ O + (CH ₃) ₃ CO —	$\sim 6 \times 10^8$ (rel.)	$k/k_{\text{EtOH}} \cong 0.33$	Fenton	chem.	c.k.	73-9105
	7	5.8×10^8 (rel.)	$k/k_{\text{MeOH}} = 0.65$	Ti(III) + H ₂ O ₂	esr	obs. radical ratios.	74-5144
	1.7-1.8	2.2×10^7 (rel.)	$k/k_{2-\text{PrOH}} = 0.01$	Fenton	chem.	c.k. with 3-pentanol.	74-9006

For other ratios see: 3.41, 3.97, 3.310, 3.343, 3.498.

TABLE 4. Reactions of OH with organic solutes – Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.547	<i>N</i> -methylpropion- amide	5–6	1.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS[−]} = 0.13	p.r.	opt.	c.k.	71–0414
3.548	methyl propionate	6–7	4.5 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{CNS[−]} = 0.041	p.r.	opt.	c.k.	65–0387
3.549	2-methylpropionate ion (isobutyrate ion)	9	1.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 0.68	γ-r.	opt.	c.k. with RNO.	66–0423
3.550	2-methylpyridine	9	2.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.20 ± 0.01	γ-r.	opt.	c.k.	69–0280
3.551	3-methylpyridine	9	2.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.19 ± 0.01	γ-r.	opt.	c.k.	69–0280
3.552	methyl sulfide OH + CH ₃ SCH ₃ → CH ₃ S(OH)CH ₃ → (CH ₃ SCH ₃) ⁺ + OH [−]	—	5.2 x 10 ⁹ (rel.)	—	p.r.	opt.	c.k. with MeOH, 2-PrOH and HCOO [−] ; meas. abs. at 470 nm (CH ₃ SCH ₃) ⁺ .	67–0186
3.553	methylsulfide ion OH + CH ₃ S [−] → OH [−] + CH ₃ S	11	(6.0 ± 0.9) x 10 ⁹	—	p.r.	opt.	p.b.k. at 410–420 nm. (RSSR [−]).	69–0553
3.554	methyl thiogly- colate	5.1 10.6	2.1 x 10 ¹⁰ (rel.) 1.8 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS[−]} = 1.9 <i>k</i> / <i>k</i> _{CNS[−]} = 1.6	p.r.	opt.	c.k.; pK = 7.8	73–0090
	5-methyluracil <i>See</i> thymine (3.711). Metronidazole <i>See</i> 1-(2-hydroxyethyl)-2-methyl-5-nitroimidazole (3.453).							
3.555	1-naphthalene- acetic acid	—	1 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 4.6	γ-r.	chem.	c.k.	72–0541
3.556	1-naphthoate ion	9	7.9 x 10 ⁹	—	p.r.	opt.	p.b.k.	73–0110
3.557	2-naphthoate ion	9	7.6 x 10 ⁹	—	p.r.	opt.	p.b.k.	73–0110
3.558	nicotinamide	9.0	1.5 x 10 ⁹	—	p.r.	opt.	p.b.k.	71–0582
3.559	omitted							
3.560	nicotinate ion	9	1.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.13 ± 0.01	γ-r.	opt.	c.k.	69–0280
		9.1	2.3 x 10 ⁹	—	p.r.	opt.	p.b.k.	71–0582
3.561	nicotinic acid	3.1	2.6 x 10 ⁸	—	p.r.	opt.	p.b.k.	71–0582
3.562	nicotinuric acid	7.5	1.1 x 10 ⁹	—	p.r.	opt.	p.b.k.	71–0582
3.563	nitrilotriacetic acid	~0	—	<i>k</i> / <i>k</i> _{acrylamide} = 0.36	Fenton	pol.	c.k.	72–9162
3.564	5-nitrobarbituric acid	5.9	(9.2 ± 0.9) x 10 ⁹ 7.8 x 10 ⁹	— —	p.r.	opt.	p.b.k. at 420 nm. d.k. at 350 nm.	73–1003
3.565	nitrobenzene OH + C ₆ H ₅ NO ₂ → OHC ₆ H ₅ NO ₂	1 10.5	6.7 x 10 ⁸ (rel.) 2.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{Fe2+} = 2.9 <i>k</i> / <i>k</i> _{BzO[−]} = 0.39	Fenton γ-r.	chem. trac.	c.k. c.k.; meas. ¹⁴ CO ₂ formn.	49–0003 65–0099
		9	3.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 1.8	γ-r.	opt.	c.k. with RNO.	66–0441
		—	(4.7 ± 0.5) x 10 ⁹	—	p.r.	opt.	p.b.k. at 410 nm.	67–0458
		—	3.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS[−]} = 0.318	p.r.	opt.	c.k.	67–0458
		—	2.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 1.55	γ-r.	opt.	c.k. with RNO.	68–0157
		7	(3.2 ± 0.4) x 10 ⁹	—	p.r.	opt.	p.b.k. at 410 nm; cor. for (OH + OH) and (H + aromatic).	68–0304
		7	2.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{BzO[−]} = 0.49	r.	chem.	c.k.; meas. sali- cylate formn.	68–0494
		9	3.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.25 ± 0.01	γ-r.	opt.	c.k.	69–0280
			<i>For other ratios see:</i> 3.12, 3.25, 3.66, 3.155, 3.168, 3.169, 3.191, 3.231, 3.233, 3.358, 3.362, 3.479, 3.506, 3.523, 3.524, 3.645, 3.646.					
3.566	nitrobenzene- <i>d</i> ₅	—	3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 1.64	γ-r.	opt.	c.k. with RNO.	68–0157
3.567	<i>p</i> -nitrobenzoate ion (PNBA [−])	9 6–9.4	2 x 10 ⁹ (rel.) (2.6 ± 0.4) x 10 ⁹	<i>k</i> / <i>k</i> _{EtOH} = 1.06 —	γ-r. p.r.	opt. opt.	c.k. with RNO. p.b.k. at 420 nm; cor. for (OH + OH) and (H + aromatic).	66–0441 68–0304
	OH + NO ₂ C ₆ H ₄ COO [−] → NO ₂ C ₆ H ₄ (OH)COO [−]							
			<i>For other ratios see:</i> 3.133, 3.186, 3.280–1, 3.284, 3.286, 3.358, 3.511.					
3.568	<i>anti</i> -5-nitro-2- furaldoxime (nifuroxime)	7	1.0 x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 500 nm as well as d.k. at 360 nm.	73–1018

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.569	5-nitro-2-furaldehyde	7	5.5 x 10 ⁹	—	p.r.	opt.	p.b.k. at 500 nm as well as d.k. at 360 nm.	73-1018
3.570	5-nitro-2-furaldehyde semicarbazone (nitrofurazone)	7	1.06 x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 500 nm as well as d.k. at 360 nm.	73-1018, 73-3016
3.571	5-nitrofuroate ion	7	5.3 x 10 ⁹	—	p.r.	opt.	p.b.k. at 500 nm as well as d.k. at 300 nm.	73-1018, 73-0114
3.572	5-nitroindole	9.0	(1.25 ± 0.24) x 10 ¹⁰	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> (OH + tryptophan) = (1.25 ± 0.3) x 10 ¹⁰ .	71-0556
3.573	nitromethane	9	3.1 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 0.168	γ-r.	opt.	c.k. with RNO.	66-0423
		—	≤ 8.4 x 10 ⁶ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} ≤ 7.6 x 10 ⁻⁴	p.r.	opt.	c.k.	66-0800
3.574	nitromethane ion OH + CH ₂ =NO ₂ ⁻ → CH ₂ (OH)NO ₂ ⁻	10.5	(8.5 ± 1.5) x 10 ⁹	—	p.r.	opt.	p.b.k. at 280 nm.	68-0342
3.575	5-nitro-6-methyluracil	5.9	(5.3 ± 0.5) x 10 ⁹	—	p.r.	opt.	p.b.k. at 420 nm.	73-1003
3.576	5-nitroorotate ion	5.9	(5.8 ± 0.6) x 10 ⁹	—	p.r.	opt.	p.b.k. at 420 nm.	73-1003
3.577	<i>m</i> -nitrophenol	9	7.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.57 ± 0.05	γ-r.	opt.	c.k.	72-0837
3.577a	<i>o</i> -nitrophenol	9	9.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.74 ± 0.06	γ-r.	opt.	c.k.	72-0837
3.577b	<i>p</i> -nitrophenol OH + HOC ₆ H ₄ NO ₂ → HOC ₆ H ₄ NO ₃ ⁻ + H ⁺	—	(3.8 ± 0.6) x 10 ⁹	—	p.r.	opt.	p.b.k. at 290 nm, d.k. at 400 nm.	68-0303
		9	7.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.61 ± 0.08	γ-r.	opt.	c.k.	72-0837
3.578	<i>o</i> -nitrophenyl-β-D-glucopyranoside	—	3.0 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> (OH + X) = 4.4 x 10 ⁹ where X = phenyl β-D-glucopyranoside.	71-0056
3.579	<i>m</i> -nitrophenyl-β-D-glucopyranoside	—	4.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.44	p.r.	opt.	c.k.	71-0056
		—	3.4 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> (OH + X) = 4.4 x 10 ⁹ where X = phenyl β-D-glucopyranoside.	71-0056
3.580	<i>p</i> -nitrophenyl-β-D-glucopyranoside	—	2.8 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO; rel. to <i>k</i> _X = 4.4 x 10 ⁹ where X = phenyl β-D-glucopyranoside.	71-0056
3.581	nitrosobenzene OH + C ₆ H ₅ NO → C ₆ H ₅ NO ₂ ⁻ + H ⁺	—	4.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.395	p.r.	opt.	c.k.	71-0056
		7.0	1.1 x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 285 nm.	66-0433
		7.0	1.8 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.67	p.r.	opt.	c.k.	67-0688, 66-0433
3.582	<i>p</i> -nitrosodimethylaniline (RNO)	9.0	ca. 1.0 x 10 ¹⁰ (rel.)	—	γ-r.	opt.	c.k. with 18 different compounds; meas. loss of abs. at 440 nm; <i>k</i> / <i>k</i> _{ferro} ≅ <i>k</i> / <i>k</i> _{I⁻} ≅ 1.	65-0356

TABLE 4. Reactions of OH with organic solutes – Continued

No.	Solute and Reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.	
3.582 cont.	—	1.8 x 10 ¹⁰	—	p.r.	opt.	d.k. at 440 nm.	68-0066	
	—	(1.8 ₅ ± 0.15) x 10 ¹⁰ (rel.)	—	p.r.	opt.	c.k. with HCOO ⁻ , I ⁻ , AsO ₂ ⁻ , NO ₂ ⁻	68-0066	
	7, 9.0	(7.1 ± 0.5) x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with Br ⁻ , HCOO ⁻ , EtOH. In N ₂ O saturated solution with concn. of above scavengers for complete OH removal, dye still bleaches.	68-0066	
	7	(1.25 ± 0.2) x 10 ¹⁰ <i>For other ratios see:</i> 3.9, 3.12, 3.23, 3.25, 3.27, 3.35, 3.39, 3.52, 3.54, 3.63, 3.64, 3.66, 3.70, 3.73, 3.82, 3.128, 3.131, 3.146, 3.148, 3.151, 3.155, 3.165, 3.166, 3.172, 3.177, 3.180, 3.181, 3.184, 3.186, 3.191, 3.196, 3.216, 3.217, 3.219, 3.236, 3.244, 3.248, 3.253, 3.254, 3.259, 3.260, 3.265, 3.268, 3.269, 3.291, 3.294-5, 3.318, 3.340, 3.341, 3.345, 3.348, 3.358, 3.367, 3.375, 3.384, 3.388, 3.390, 3.391, 3.392, 3.394, 3.395, 3.399, 3.400, 3.403, 3.405, 3.428, 3.429, 3.442, 3.446, 3.448, 3.449, 3.456, 3.457, 3.461, 3.461a, 3.462, 3.463, 3.464, 3.465, 3.466a, 3.473, 3.474, 3.483a, 3.484, 3.485, 3.492, 3.494, 3.497, 3.498, 3.504, 3.511, 3.513, 3.518, 3.519, 3.525, 3.533, 3.536, 3.537, 3.550, 3.551, 3.560, 3.565, 3.577, 3.583, 3.590, 3.592, 3.607, 3.608, 3.614, 3.615, 3.633, 3.637, 3.648, 3.648a, 3.649, 3.651-5, 3.657a 3.660, 3.663, 3.664, 3.666, 3.669, 3.674, 3.676, 3.677, 3.689, 3.708, 3.711, 3.714, 3.715, 3.716, 3.728, 3.733, 3.737, 3.743, 3.746, 3.749, 3.749a, 3.750, 3.751, 3.753, 3.754a.	—	p.r.	opt.	d.k. at 440 nm.	69-0156	
3.583	<i>p</i> -nitro- <i>o</i> -toluene- sulfonic acid	—	<i>k</i> / <i>k</i> _{RNO} = 0.128	r.	opt.	c.k.	72-0425	
3.584	5-nitrouracil	5.9	(5.4 ± 0.5) x 10 ⁹	—	p.r.	opt.	p.b.k. at 420 nm.	73-1003
		—	7.4 x 10 ⁹ (6.5 ± 1) x 10 ⁹	—	p.r.	opt.	d.k. at 350 nm. p.b.k. as well as d.k.	73-0145
3.585	norleucine	2.2	3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.55 ± 0.06	γ-r.	opt.	c.k.	67-0461
3.586	norpseudopellet- tierine <i>N</i> -oxyl (NPPN)	10.5	6.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{carb} = 18.4	p.r.	opt.	c.k.; cor. for CO ₃ ⁻ + NPPN.	71-0061
3.587	norvaline	7	(4.7 - 4.2) x 10 ⁹	—	p.r.	opt.	d.k. at 242 nm.	71-0061
		2-2.2	1.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.28 ± 0.02	γ-r.	opt.	c.k.	67-0461
3.588	1-octanol	2-2.2	6.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.20 ± 0.15	γ-r.	opt.	c.k.	67-0461
3.589	ornithine	2-2.2	1.7 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.032 ± 0.003	γ-r.	opt.	c.k.	67-0461
3.590	orotate ion	5.2	5.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{ferro} = 0.62	p.r.	opt.	c.k.	70-0567
		11	5.3 x 10 ⁹	—	p.r.	opt.	p.b.k. at 340 nm (OH adduct).	70-0567
		11	5.0 x 10 ⁹	—	p.r.	opt.	d.k. at 280 nm (5,6-double bond); ave. <i>k</i> (pH 5-11) by all methods = 5.2 x 10 ⁹ ; <i>k</i> de- creases at pH < 5.	70-0567
3.591	orotidine	6-7	4.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.36	γ-r.	opt.	c.k.; 17°C.	73-0294
		7	4.0 x 10 ⁹	—	p.r.	opt.	p.b.k. (OH adduct).	70-0567

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.592	oxalate ion OH + C ₂ O ₄ ²⁻ → OH ⁻ + COOCOO ⁻ → CO ₂ + CO ₂ ⁻ + OH ⁻	9.0 9-13 7 7 7 6 —	8.4 x 10 ⁶ (rel.) — 1 x 10 ⁷ (rel.) 1 x 10 ⁷ (rel.) — 7.8 x 10 ⁶ (rel.) 1.6 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 6.7 x 10 ⁻⁴ (± 15%) <i>k</i> / <i>k</i> _{HCOO⁻} = <i>k</i> (O ⁻ + C ₂ O ₄ ²⁻)/ <i>k</i> (O ⁻ + HCOO ⁻) <i>k</i> / <i>k</i> _{EtOH} = 0.00565 <i>k</i> / <i>k</i> _{MeOH} = 0.0112 <i>k</i> / <i>k</i> _{perox} = 0.208 <i>k</i> / <i>k</i> _{CNS⁻} = 0.0007 <i>k</i> / <i>k</i> _{I⁻} = 0.0012	γ-r. γ-r. γ-r. γ-r. γ-r. p.r. p.r.	opt. trac. chem. chem. opt. opt. opt.	c.k. meas. formn. of H ₂ C ₂ O ₄ . c.k. c.k. c.k. c.k. c.k.; obs. I ₂ ⁻ formn.	65-0356 66-0151, 66-0621 67-0131, 66-0621 67-0131 70-1050 71-0041 73-0020
3.593	oxalate ion, hydrogen OH + HC ₂ O ₄ ⁻ → OH ⁻ + CO ₂ + CO ₂ H	2.7 3	5.6 x 10 ⁸ (rel.) 6.9 x 10 ⁸ (rel.) 4.7 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 0.3 <i>k</i> / <i>k</i> _{MeOH} = 0.77 <i>k</i> / <i>k</i> _{CNS⁻} = 0.00425	γ-r. p.r.	chem. opt.	c.k. c.k.	67-0131, 66-0621 71-0041
3.594	oxalic acid OH + (COOH) ₂ → H ₂ O + CO ₂ + CO ₂ H	1.3, 2.7 2.0- 2.2 0.5 1.5	< 10 ⁷ (rel.) 9.2 x 10 ⁶ (rel.) 1.45 x 10 ⁶ (rel.) —	— <i>k</i> / <i>k</i> _{thym} = (1.7 ± 0.7) x 10 ⁻³ <i>k</i> / <i>k</i> _{CNS⁻} = 0.00013 <i>k</i> / <i>k</i> _{Cl⁻} = 1.3	γ-r. γ-r. p.r. X-r.	chem. opt. opt. pol.	c.k. with MeOH and EtOH. c.k. c.k. effect of Cl ⁻ and oxalic acid on reaction of U (VI).	66-0621, 67-0131 67-0461 71-0041 71-0542
For other ratios see: 3.382, 3.427.								
3.594a	oxytocin	~6	(1.3 ± 0.2) x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 330 nm.	74-1102
3.595	papain	6.4	4.7 x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 310- 350 nm.	72-3042
5.596	paraldehyde See 2,4,6-trimethyl-1,3,5-trioxane (3.731). DL-penicillamine	1	5.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.09	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 79.2.	69-5278
3.597	penicillamine disulfide	1	8.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.5	Fenton	esr	c.k.; <i>k</i> / <i>k</i> _{perox} = 110.	69-5278
3.598	pentaerythritol	9	3.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 1.73	γ-r.	opt.	c.k. with RNO	66-0423
3.599	1,4-pentadien-3- ol	7.0	(1.0 ± 0.2) x 10 ¹⁰	—	p.r.	opt.	p.b.k.	73-1070
3.600	pentafluorobenzene	—	4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.36	p.r.	opt.	p.b.k.	73-0054
3.600a	pentamethylbenzene	~7	7.5 x 10 ⁹	—	p.r.	opt.	p.b.k.; OH addn. and H abstr.; rate for H abstr. = 2.4 x 10 ⁹ .	75-1009
3.601	1,5-pentanediol	9	3.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 1.9	γ-r.	opt.	c.k. with RNO.	66-0423
3.602	pentanoate ion See valerate ion (3.752). 1-pentanol	9 2-2.2 5-5.5 —	4.9 x 10 ⁹ (rel.) 5.1 x 10 ⁹ (rel.) 5.5 x 10 ⁹ (rel.) 3.7 x 10 ⁹ (rel.) 3.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.65 <i>k</i> / <i>k</i> _{thym} = 0.95 ± 0.10 <i>k</i> / <i>k</i> _{thym} = 1.02 ± 0.10 <i>k</i> / <i>k</i> _{CNS⁻} = 0.34 <i>k</i> / <i>k</i> _{CNS⁻} = 0.38	γ-r. γ-r. γ-r. p.r.	opt. opt. opt. opt.	c.k. with RNO. c.k. c.k. c.k.	66-0423 67-0461 67-0461 73-1077
3.602a	3-pentanol	1.7-1.8	2.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 1.1	Fenton	chem.	c.k. with cyclo- heptanol.	74-9006
3.603	2-pentanone	6-7	1.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.174	p.r.	opt.	c.k.	65-0387
3.604	3-pentanone	6-7	1.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.123	p.r.	opt.	c.k.	65-0387
pentylamine See amylamine (3.168). pentylammonium ion See amylammonium ion (3.169).								
3.605	phenethyl alcohol	—	7.0 x 10 ⁹ (rel.) 5.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.64 <i>k</i> / <i>k</i> _{CNS⁻} = 0.55	p.r.	opt.	c.k.	73-1077
3.606	phenethylammonium ion	4	9.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.89	p.r.	opt.	c.k.	70-0371

TABLE 4. Reactions of OH with organic solutes – Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.607	phenol	7.0	6.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{BzO⁻} = 1.08	γ-r.	trac.	c.k.; meas. ¹⁴ CO ₂ .	65-0099
	OH + C ₆ H ₅ OH →	6-7	1.8 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.61	p.r.	opt.	c.k.	65-0387
	C ₆ H ₅ (OH) ₂	9	8.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 4.64	γ-r.	opt.	c.k. with RNO.	66-0441
		7.4-7.7	(1.4 ± 0.3) x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 330 nm.	67-0122
		~1.2	9.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 4.4	γ-r.	chem.	c.k.	68-0602
		6.98	1.1 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 5.2	γ-r.	chem.	c.k.	68-0602
		9	8.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.68 ± 0.02	γ-r.	opt.	c.k.	72-0837
3.608	phenoxide ion OH + C ₆ H ₅ O ⁻ → C ₆ H ₅ (OH)O ⁻	10.7	9.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{BzO⁻} = 1.62	γ-r.	trac.	c.k.; meas. ¹⁴ CO ₂ .	65-0099
3.608a	<i>p</i> -phenoxybenzoate ion	—	7.0 x 10 ⁹	—	p.r.	opt.	p.b.k.; OH addn.	75-1001
3.609	phenylacetamide	9	5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.73	γ-r.	opt.	c.k. with RNO.	66-0441
3.610	phenyl acetate	9	5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.73	γ-r.	opt.	c.k. with RNO.	66-0441
3.611	phenylacetate ion (PA ⁻)	9	4.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.36	γ-r.	opt.	c.k.	66-0441
		6-8	(7.9 ± 1.1) x 10 ⁹	—	p.r.	opt.	p.b.k. at 325 nm, cor. for (OH + OH) and (H + aromatic).	68-0304
For other ratios see: 3.199, 3.200, 3.358, 3.511.								
3.612	phenylacetic acid	1	1.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{Fe²⁺} = 4.8	Fenton	chem.	c.k.	49-0003
		—	1.8 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 8.2	γ-r.	chem.	c.k.	72-0541
3.613	phenylalanine, positive ion	2-2.2	5.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.515	p.r.	opt.	c.k.	65-0388
		2-2.2	7.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{thym} = 1.42 ± 0.08	γ-r.	opt.	c.k.	65-0388, 67-0461
3.614	phenylalanine, zwitterion	5.5-6	5.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.53	γ-r.	opt.	c.k.	65-0388, 67-0461
		nat.	6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{ferro} = 0.645	p.r.	opt.	c.k.	71-0578
		nat.	6.6 x 10 ⁹	—	p.r.	opt.	p.b.k. at 300 nm.	71-0578
		6.9	7.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.57	γ-r.	opt.	c.k.	73-0548
	For other ratios see:			3.178, 3.347, 3.697.				
3.615	phenylalanine, negative ion	10.6	8.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{carb} = 23	p.r.	opt.	c.k.	68-0062
		—	1.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.12	p.r.	opt.	c.k.	69-0156
3.615a	1-phenyl-3-butanol	1.7-1.8	2.0 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 9	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.615b	1-phenylethanol	1.7-1.8	1.3 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 5.8	Fenton	chem.	c.k. with cycloheptanol.	74-9006
3.615c	1-phenylethanol-1- <i>d</i>	1.7-1.8	1.3 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 5.9	Fenton	chem.	c.k. with bromo-phenylethanol.	74-9006
3.616	2-phenylethanol	See phenethyl alcohol (3.605).						
	phenyl-β-D-glucopyranoside	6.8	4.4 x 10 ⁹	—	p.r.	opt.	p.b.k. at 300 nm.	71-0055, 71-0056
	OH + C ₆ H ₅ OC ₆ H ₁₁ O ₅ → C ₆ H ₅ (OH)OC ₆ H ₁₁ O ₅	—	5.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.53	p.r.	opt.	c.k.	71-0055, 71-0056
For other ratios see:				3.213, 3.255, 3.256, 3.274, 3.337, 3.458, 3.520, 3.578, 3.579, 3.580, 3.729.				
3.617	phenylhydroxyl-amine	3.7-11.5	1.5 x 10 ¹⁰	—	p.r.	opt.	p.b.k. at 290 nm.	67-0191, 67-0688
	OH + C ₆ H ₅ NHOH → C ₆ H ₅ (OH)NHOH	—	2 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.82	p.r.	opt.	c.k.	67-0688
3.617a	1-phenyl-1-propanol	1.7-1.8	1.2 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 5.5	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.617b	1-phenyl-2-propanol	1.7-1.8	2.4 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 11	Fenton	chem.	c.k. with 1-phenylethanol.	74-9006
3.617c	2-phenyl-2-propanol	1.7-1.8	5.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 2.4	Fenton	chem.	c.k. with cycloheptanol.	74-9006
3.618	phthalate ion	9	5.9 x 10 ⁹	—	p.r.	opt.	p.b.k.	73-0110
		—	3.0 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO assuming <i>k</i> (OH + sulfanilic acid) = 2.93 x 10 ⁹ .	73-0094

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.619	pinelic acid 2-2.2	3×10^9 (rel.)	$k/k_{\text{thym}} = 0.55 \pm 0.06$	γ -r.	opt.	c.k.	67-0461
3.620	pinacol 1	2.9×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 1.25$	Fenton	chem.	c.k.	49-0002
	9	5.4×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.29$	γ -r.	opt.	c.k. with RNO.	66-0423
3.621	pivalamide 5-6	1.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.132$	p.r.	opt.	c.k.	71-0414
3.622	pivalate ion 9	1.4×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.78$	γ -r.	opt.	c.k. with RNO.	66-0423
3.623	polyacrylate ion —	$(3.2 - 4.5) \times 10^8$ (rel.)	—	p.r.	opt.	c.k. with RNO and CNS ⁻ ; <i>k</i> depends on chain length; at mol. wt. 9×10^3 <i>k</i> = $(1 \rightarrow 3) \times 10^8$ as pH varies 2 \rightarrow 8.	73-1095
3.624	polyadenylic acid (poly A) 4.6	1.6×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.015$	p.r.	opt.	c.k.; rate in terms of nucleotide concn.	68-0845
	5.9	2.8×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.025$				
	6.3	3.6×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.033$				
	7.3	3.8×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.035$				
	7	$(9 \pm 1) \times 10^8$	—	p.r.	opt.	p.b.k.	73-1071
3.625	polyadenylic + -uridylic acid (poly A + U) 7	$(5 \pm 1) \times 10^8$	—	p.r.	opt.	p.b.k.	73-1071
3.626	polycytidylic acid (poly C) 7	$(1.2 \pm 0.1) \times 10^9$	—	p.r.	opt.	p.b.k. at 425 nm; $\epsilon = 780 \pm 80$; mol. wt. $> 10^5$.	73-1071
3.627	polyethylene oxide H abstr. 7.3	$> 2.4 \times 10^8$ (rel.)	$k/k_{\text{cyst}} = 2.86 \times 10^{-2}$	p.r.	opt.	c.k.; <i>k</i> based on monomer unit of mol. wt. 44, and $k_{\text{cyst}} = 8.5 \times 10^9$.	69-0088
	7	$> 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS ⁻ , BzO ⁻ and RNO; <i>k</i> based on monomer unit; varies with chain length and concn.	70-0394
	—	$< 2.0 \times 10^9$ (rel.)	—	r.	visc.	c.k.; $k > 2.0 \times 10^6$; effect of dioxane on crosslinking; rel. to $k(\text{OH} + \text{dioxane}) = 2.35 \times 10^9$.	70-2058
	—	$(2.8-7.6) \times 10^8$ (rel.)	—	p.r.	opt.	c.k.; <i>k</i> depends on concn. and mol. wt. of polymer; rel. to ferro-cyanide or I ⁻ .	73-1046
	—	$\sim 1 \times 10^9$ (rel.)	$k/k_{\text{Br}^-} \cong 0.026$	r.	chem.	c.k.; effect of Br ⁻ on product yields, assume $k(\text{OH} + \text{Br}^-) = 3.7 \times 10^{10}$.	73-2129, 73-2126
3.628	poly(ethylenesulfonate) (poly-anion) —	1.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0106$	p.r.	opt.	c.k.; concn. in monomer units.	68-0352
3.629	polyoxyethylene-(15)nonylphenol (Igepal CO-730) —	1.1×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1$	p.r.	opt.	c.k.; concn. $< 10^{-4} M$; at higher concn. <i>k</i> decreases.	71-0001, 71-0586

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.630	poly(styrenesul- fonate) (polyanion)	— 3.3 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.0303$	p.r.	opt.	c.k.; concn. in monomer units.	68-0352
3.631	polyuridylic acid (poly U)	7 < (3.8 ± 0.4) x 10 ⁹ 7 1.4 x 10 ⁹ (rel.)	— $k/k_{\text{CNS}^-} = 0.13 \pm 0.06$	p.r. p.r.	opt. opt.	p.b.k. c.k.; rate per base unit.	69-0571 69-0571
3.632	polyvinyl- pyrrolidone	7 (1.25 ± 0.05) x 10 ⁹ 7 > 10 ¹⁰ (rel.)	— —	p.r. p.r.	opt. opt.	p.b.k. at 390 nm. c.k. with CNS ⁻ , BzO ⁻ and RNO; <i>k</i> varies with chain length and is per monomer unit.	73-1071 70-0394
3.632a	proflavine	— (1.0 ± 0.2) x 10 ¹⁰	—	p.r.	opt.	d.k. at 444 nm; deduced $k \cong 2 \times 10^9$ for dye bound to DNA.	75-3094
3.633	proline	2-2.2 3.1 x 10 ⁸ (rel.) 6.8 6.5 x 10 ⁸ (rel.)	$k/k_{\text{thym}} = 0.0565$ $k/k_{\text{RNO}} = 0.052$	γ-r. γ-r.	opt. opt.	c.k. c.k.	65-0388 73-0548
3.634	1,2-propanediol (I) OH + C ₃ H ₈ O ₂ → MeCHOHCHOH or MeCOHCH ₂ OH + H ₂ O (II) OH + C ₃ H ₈ O ₂ → CH ₂ CHOHCH ₂ OH + H ₂ O	7 1.7 x 10 ⁹ (rel.) 9 1.7 x 10 ⁹ (rel.) 6 1.6 x 10 ⁹ (rel.) 2-2.2 1.9 x 10 ⁹ (rel.) — — — —	$k/k_{\text{CNS}^-} = 0.153$ $k/k_{\text{EtOH}} = 0.9$ $k/k_{\text{EtOH}} = 0.855$ $k/k_{\text{thym}} = 0.35 \pm 0.03$ $k_{\text{II}}/k_{\text{I}} = 0.26$	p.r. γ-r. γ-r. γ-r. p.r.	opt. opt. chem. opt.	c.k. c.k. with RNO. c.k. with Br ⁻ . c.k. detd. % α-alco- hol radical by reaction with TNM ≤ 0.1% alkoxy radi- cal detd. by reac- tion with I ⁻ .	65-0387 66-0423 66-0423 67-0461 73-0126
3.635	1,3-propanediol	9 2.4 x 10 ⁹ (rel.) 6 2.0 x 10 ⁹ (rel.)	$k/k_{\text{EtOH}} = 1.3$ $k/k_{\text{EtOH}} = 1.1$	γ-r. γ-r.	opt. chem.	c.k. with RNO. c.k. with Br ⁻ .	66-0423 66-0423
3.636	1-propanol (PrOH) (I) OH + PrOH → H ₂ O + MeCH ₂ CHOH (61%, 69-0522) (II) OH + PrOH → H ₂ O + MeCHCH ₂ OH (III) OH + PrOH → H ₂ O + MeCH ₂ CH ₂ O	1 6.0 x 10 ⁸ (rel.) 7, 2.7 x 10 ⁹ (rel.) 10.7 2.5 x 10 ⁹ (rel.) 9 2.8 x 10 ⁹ (rel.) 2-2.2 3.2 x 10 ⁹ (rel.) -5-5.5 3.0 x 10 ⁹ (rel.) nat. 2.7 x 10 ⁹ (rel.) — — — 1.5 x 10 ⁹ (rel.)	$k/k_{\text{Fe}^{2+}} = 2.6$ $k/k_{\text{carb}} = 7.5$ $k/k_{\text{CNS}^-} = 0.223$ $k/k_{\text{EtOH}} = 1.5$ $k/k_{\text{thym}} = 0.60 \pm 0.05$ $k/k_{\text{thym}} = 0.56 \pm 0.06$ $k/k_{\text{ferro}} = 0.29$ $k_{\text{II}}/k_{\text{I}} = 0.86$ $k_{\text{III}}/k_{\text{I}} \leq 0.01$ $k/k_{\text{MeOH}} = 1.65$	Fenton p.r. p.r. γ-r. γ-r. γ-r. p.r. p.r. p.r. Ti(III) + H ₂ O ₂	chem. opt. opt. opt. opt. opt. opt. opt. esr	c.k. c.k. c.k. c.k. with RNO. c.k. c.k. c.k. c.k. c.k. c.k. c.k. with HSO ₄ ⁻ . c.k.	49-0002 65-0190, 65-0387 65-0387 66-0423 67-0461 67-0461 71-0578 73-0126 73-5253
3.637	2-propanol (2-PrOH) (I) OH + 2-PrOH → H ₂ O + (CH ₃) ₂ COH (89%, 69-0522) (II) OH + 2-PrOH → H ₂ O + CH ₃ CHOHCH ₂ (III) OH + 2-PrOH → H ₂ O + CH ₃ CHOCH ₃	1 6.9 x 10 ⁸ (rel.) 7 2.2 x 10 ⁹ (rel.) — 2.1 x 10 ⁹ (rel.) 9.0 2.1 x 10 ⁹ (rel.) 7 6.5 x 10 ⁹ (rel.) 2-2.2 2.1 x 10 ⁹ (rel.) — 3.2 x 10 ⁹ (rel.) 6.8 1.4 x 10 ⁹ (rel.) 6 2.2 x 10 ⁹ (rel.) 9 2.0 x 10 ⁹ (rel.)	$k/k_{\text{Fe}^{2+}} = 3.0$ $k/k_{\text{I}^-} = 0.17 \pm 0.006$ $k/k_{\text{ferro}} = 0.23$ $k/k_{\text{RNO}} = 0.17$ $k/k_{\text{CNS}^-} = 0.591$ $k/k_{\text{thym}} = 0.387$ $k/k_{\text{MeOH}} = 3.6$ $k/k_{\text{ferro}} = 0.15 \pm 0.03$ $k/k_{\text{EtOH}} = 1.2$ $k/k_{\text{EtOH}} = 1.1$	p.r. p.r. phot. γ-r. p.r. γ-r. p.r. X-r. γ-r. γ-r.	opt. opt. — opt. opt. opt. opt. — chem. opt.	c.k. c.k. c.k. c.k. c.k. c.k. c.k. with HSO ₄ ⁻ . c.k. c.k. with Br ⁻ . c.k. with RNO.	49-0002 65-0010, 67-0041 65-0247 65-0356 65-0387 65-0388 66-0019 66-0234 66-0423 66-0423

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.637 cont.	2-propanol-2- <i>d</i> (I) OH + (CH ₃) ₂ CDOH → HDO + (CH ₃) ₂ COH (II) OH + (CH ₃) ₂ CDOH → H ₂ O + CH ₂ CDOHCH ₃	2-2.2	2.3 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 0.42 \pm 0.03$	γ -r.	opt.	c.k.	67-0461
		5-5.5	2.3 x 10 ⁹ (rel.)	$k/k_{\text{thym}} = 0.42 \pm 0.05$	γ -r.	opt.	c.k.	67-0461
		—	—	$k/k_{\text{TCOO}^-} = 0.45$	γ -r.	trac.	c.k.; meas. ³ HHO.	68-0209
		2-10	1.9 x 10 ⁹ (rel.)	$k/k_{\text{CNS}^-} = 0.17$	p.r.	opt.	c.k.	68-0316
		—	1.5 x 10 ⁹ (rel.)	$k/k_{\text{RNO}} = 0.12$	p.r.	opt.	c.k.	69-0156
		—	—	$k_1/k_{\text{II}} = 6.2$	r.	—	c.k. with H ₂ O ₂ .	70-0104
		—	—	$k_1/k_{\text{II}} = 5.2 \pm 0.1$	γ -r.	chem.	c.k. with H ₂ O ₂ ; $k_{\text{H}}/k_{\text{D}}(\text{I}) = 1.38 \pm 0.05$; $k_{\text{H}}/k_{\text{D}}(\text{II}) = 2.08 \pm 0.12$ or 2.15 ± 0.16 .	71-0081
		nat.	2.0 x 10 ⁹ (rel.)	$k/k_{\text{ferro}} = 0.216$	p.r.	opt.	c.k.	71-0578
		0.82	1.3 x 10 ⁹ (rel.) (I)	$k_1/k_{\text{Fe}^{2+}} = 5.73$	Fenton	chem.	c.k.	71-9132
		0.82	2.1 x 10 ⁸ (rel.) (II)	$k_{\text{II}}/k_{\text{Fe}^{2+}} = 0.92$	Fenton	chem.	c.k.	71-9132
		0	—	$k/k_{\text{bisulf}} [\text{HSO}_4^-] = 202 \pm 12 \text{ M}^{-1}$	γ -r.	chem.	c.k.; computer anal.; 4 M H ₂ SO ₄ .	72-0094
		0.8	—	$k/k_{\text{bisulf}} [\text{HSO}_4^-] = (1.1 \pm 0.2) \times 10^4 \text{ M}^{-1}$	γ -r.	chem.	c.k.; computer anal.	72-0094
		0	—	$k_{\text{III}}/k_1 = 1.4 \pm 0.3$	γ -r.	chem.	calcd. by comparing oxid. of Ce(III) in HCOOH and 2-ProOH solns.	72-0094
		—	—	$k_1/k_{\text{II}} = 6.4$ $k_{\text{III}}/k_1 = 0.014$	p.r.	opt.	detd. % of α -alcohol and alkoxy radicals by reaction with TNM and I ⁻ , resp.	73-0126
		10.4	2.3 x 10 ⁹ (rel.)	$k/k_{\text{EtOH}} = 1.23$	X-r.	lum.	c.k.; effect of alcohols on quenching of chemiluminescence from fluorescein.	73-6068
		<i>For other ratios see: 3.12, 3.66, 3.80, 3.107, 3.111, 3.186, 3.198a, 3.201, 3.212a, 3.248a-3.249, 3.274, 3.277-8, 3.337a, 3.358, 3.371, 3.375a, 3.384, 3.385, 3.468, 3.469, 3.471, 3.511, 3.517a, 3.543a-e, 3.546, 3.555, 3.602, 3.607, 3.612, 3.615a-c, 3.617a-c, 3.687, 3.695a.</i>						
3.638	2-propanol-2- <i>d</i> (I) OH + (CH ₃) ₂ CDOH → HDO + (CH ₃) ₂ COH (II) OH + (CH ₃) ₂ CDOH → H ₂ O + CH ₂ CDOHCH ₃	6	1.4 x 10 ⁹ (rel.)	$k/k_{\text{EtOH}} = 0.78$	γ -r.	chem.	c.k. with Br ⁻ .	66-0423
		0.82	7.9 x 10 ⁸ (I) (rel.)	$k_1/k_{\text{Fe}^{2+}} = 3.42$	Fenton	chem.	c.k.	71-9132
			2.1 x 10 ⁸ (II) (rel.)	$k_{\text{II}}/k_{\text{Fe}^{2+}} = 0.91$				
3.639	2-propanol- <i>d</i> ₆ (I) OH + (CD ₃) ₂ CHOH → H ₂ O + (CD ₃) ₂ COH (II) OH + (CD ₃) ₂ CHOH → HDO + CD ₃ CHOHCD ₂	0.82	1.2 x 10 ⁹ (I) (rel.)	$k_1/k_{\text{Fe}^{2+}} = 5.43$	Fenton	chem.	c.k.	71-9132
			5.0 x 10 ⁷ (II) (rel.)	$k_{\text{II}}/k_{\text{Fe}^{2+}} = 0.22$				
3.640	propionamide OH + C ₂ H ₅ CONH → H ₂ O + CH ₃ CHCONH ₂ + CH ₂ CH ₂ CONH ₂ + CH ₃ CH ₂ CONH	5-6	7.0 x 10 ⁸ (rel.)	$k/k_{\text{CNS}^-} = 0.064$	p.r.	opt.	c.k.; 45% CH ₃ CHCONH ₂ formed; anal. of transient spectra.	71-0414, 71-0645

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.641	propionate ion	9	8.0×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.43$	γ -r.	opt.	c.k. with RNO.	66-0423
3.642	propionic acid	1	2.0×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.86$	Fenton	chem.	c.k.	49-0002
		2-2.2	5.2×10^8 (rel.)	$k/k_{\text{thym}} = 0.097 \pm 0.01$	γ -r.	opt.	c.k.	67-0461
3.643	propionitrile	—	1.0×10^8 (rel.)	$k/k_{\text{HCOO}^-} = 0.029$	γ -r.	chem.	c.k.	73-0364
3.644	propyl acetate	6-7	1.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.124$	p.r.	opt.	c.k.	65-0387
3.645	propylamine	—	7.3×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.66$	p.r.	opt.	c.k.	73-0016
			4.8×10^9 (rel.)	$k/k_{\text{NB}} = 1.5$				
	2-propylamine See isopropylamine (3.487).							
3.646	propylammonium ion	2	7.5×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.068$	p.r.	opt.	c.k.	70-0371
		4	1.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.143$	p.r.	opt.	c.k.	70-0371
		—	1.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.13$	p.r.	opt.	c.k.	73-0016
			6.7×10^9 (rel.)	$k/k_{\text{NB}} = 0.21$				
3.647	propylene	—	8.3×10^9 (rel.)	$k/k_{\text{I}^-} = 0.64$	p.r.	opt.	c.k.	67-0041
	propylene oxide	See 1,2-epoxypropane (3.353).						
3.648	propyl gallate	6.5	1.2×10^{10} (rel.)	$k/k_{\text{RNO}} = 0.94 \pm 0.16$	γ -r.	opt.	c.k.	69-0580
3.648a	purine	6-7	3.0×10^8 (rel.)	$k/k_{\text{RNO}} = 0.024$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.649	pyridine	7	$(3.0 \pm 0.6) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = 0.272 \pm 0.054$	p.r.	opt.	c.k.	67-0251
	OH + C ₅ H ₅ N →							
	OHC ₅ H ₅ N +	9	2×10^9 (rel.)	$k/k_{\text{RNO}} = 0.16$	γ -r.	opt.	c.k.	69-0280
	C ₅ H ₅ N-OH	7.0	1.8×10^9	—	p.r.	opt.	p.b.k.	71-0582
3.650	pyridine- <i>d</i> ₅	7	$(2.7 \pm 0.9) \times 10^9$ (rel.)	$k/k_{\text{CNS}^-} = 2.42 \times 10^{-1} (\pm 0.073)$	p.r.	opt.	c.k.	67-0251
3.651	3-pyridine-carboxamide	9	1×10^9 (rel.)	$k/k_{\text{RNO}} = 0.08$	γ -r.	opt.	c.k.	69-0280
3.652	4-pyridine-carboxamide	9	1.5×10^9 (rel.)	$k/k_{\text{RNO}} = 0.12$	γ -r.	opt.	c.k.	69-0280
3.653	3-pyridinecarboxylate ion	9	2.3×10^9 (rel.)	$k/k_{\text{RNO}} = 0.18$	γ -r.	opt.	c.k.	69-0280
3.654	4-pyridinecarboxylate ion	9	2.6×10^9 (rel.)	$k/k_{\text{RNO}} = 0.21$	γ -r.	opt.	c.k.	69-0280
3.655	4-pyridinenitrile	9	7.5×10^8 (rel.)	$k/k_{\text{RNO}} = 0.06$	γ -r.	opt.	c.k.	69-0280
3.656	pyridinium ion	1	4.1×10^7 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.18$	Fenton	chem.	c.k.	49-0002
	OH + C ₅ H ₅ NH ⁺ →	1-2	$(3.3 \pm 0.7) \times 10^7$ (rel.)	$k/k_{\text{CNS}^-} = (3 \pm 0.6) \times 10^{-3}$	p.r.	opt.	c.k.	67-0251
	OHC ₅ H ₅ NH ⁺	2.0	2×10^7	—	p.r.	opt.	p.b.k.	71-0582
3.657	pyridinium ion- <i>d</i> ₅	1-2	3.6×10^7 (rel.)	$k/k_{\text{CNS}^-} = (3.3 \pm 1) \times 10^{-3}$	p.r.	opt.	c.k.	67-0251
3.657a	pyridoxine(PH)	7.2	6.3×10^9	—	p.r.	opt.	p.b.k.	75-1024
	PH ₂ ⁺	3.6	4.3×10^9	—				
	P ⁻	10.5	7.4×10^9	—				
3.657b	pyrimidine	6-7	1.6×10^8 (rel.)	$k/k_{\text{RNO}} = 0.013$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.658	pyrrole	—	1.5×10^{10}	—	p.r.	opt.	p.b.k. at 300 nm.	71-0360
	OH + C ₄ H ₄ NH →							
	(OH)C ₄ H ₄ NH							
3.659	pyrrolidinium ion	6.2	5.7×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.52$	p.r.	opt.	c.k.; see also 70-0006.	75-1016
3.660	pyruvate ion	9	3.1×10^7 (rel.)	$k/k_{\text{RNO}} = 0.0025$	γ -r.	opt.	c.k.; assume $k_{\text{RNO}} = k_{\text{ferro}}$.	67-0555
3.661	rennin	6.4	2.1×10^{10} (rel.)	—	X-r.	biol.	effects of methanol, malonate, glycerol, ethanol, glycylglycine, formate, glucose and adenine on enzyme inactivation.	73-3030
3.662	Rhodamine B (RhB)	—	$\sim 10^{10}$	—	p.r.	opt.	d.k. at 530 nm (RhB) as well as p.b.k. at 460 nm.	67-0239, 67-6053

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.663	ribonuclease	—	2.6×10^{10} (rel.) ($T = 20^\circ\text{C}$) 5.2×10^{10} (rel.) ($T = 60^\circ\text{C}$)	—	p.r.	opt.	c.k. with CNS^- ; mol. wt. 13,683; ref. rate not given.	68-3007
		3.5	$(3.6 \pm 0.5) \times 10^{10}$	—	p.r.	opt.	p.b.k.	72-3079
		5.6	$(1.9 \pm 0.3) \times 10^{10}$	—				
		~7	$(2.4 \pm 0.6) \times 10^{10}$	—				
		6.5	2.5×10^{10} (rel.)	$k/k_{\text{RNO}} = 2$	γ -r.	opt.	c.k.	73-0548
3.664	ribose	9	2.1×10^9 (rel.)	$k/k_{\text{RNO}} = 0.17$	γ -r.	opt.	c.k.; assume $k_{\text{RNO}} = k_{\text{ferro}}$	67-0555
		6.5	4.4×10^8 (rel.)	$k/k_{\text{RNO}} = 0.035 \pm 0.03$	γ -r.	opt.	c.k.	69-0580
		7	1.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.145$	p.r.	opt.	c.k.	73-1071
		—	1.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.11$	p.r.	opt.	c.k.	73-1077
		—	1.0×10^9 (rel.)	$k/k_{\text{ferro}} = 0.101$	p.r.	opt.	c.k.	73-1077
3.665	ribose-5-phosphate	7	1.3×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.12$	p.r.	opt.	c.k.	73-1071
3.666	RNA	6.5	1.9×10^9 (rel.)	$k/k_{\text{RNO}} = 0.15 \pm 0.02$	γ -r.	opt.	c.k.	69-0580
3.667	Safranine T (S.T)	3-5.5	9.3×10^9 (rel.)	$k/k_{\text{pH}} = 1.19$	γ -r.	chem.	c.k.; OH addn.	69-0279
3.668	Safranine T, protonated (S.TH ⁺) $\text{OH} + \text{S.TH}^+ \rightarrow \text{OH} \cdot \text{S.TH}^+$	0.4	3.4×10^{10} (rel.)	$k/k_{\text{pH}} = 4.35$	γ -r.	chem.	c.k.	69-0279
3.668a	salicylaldehyde	9	8.6×10^9 (rel.)	$k/k_{\text{RNO}} = 0.69 \pm 0.16$	γ -r.	opt.	c.k.	72-0837
3.669	salicylate ion $\text{OH} + \text{HOC}_6\text{H}_4\text{COO}^- \rightarrow (\text{HO})_2\text{C}_6\text{H}_4\text{COO}^-$	10.7	5.8×10^9 (rel.)	$k/k_{\text{BzO}^-} = 1.01$	γ -r.	trac.	c.k.; meas. $^{14}\text{CO}_2$	65-0099
		9.0	9.4×10^9 (rel.)	$k/k_{\text{RNO}} = 0.752 \pm 0.038$	γ -r.	opt.	c.k.	65-0356
		7	1.2×10^{10}	—	p.r.	opt.	p.b.k. at 350 nm.	68-0305
		7	2.0×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.8 \pm 0.2$	p.r.	opt.	c.k.	68-0305
		9	8.6×10^9 (rel.)	$k/k_{\text{RNO}} = 0.69 \pm 0.16$	γ -r.	opt.	c.k.	72-0837
3.670	sarcosine anhydride	5.0, 11.0	2.6×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.236$	p.r.	opt.	c.k.	71-0554
3.671	sebacic acid	2-2.2	5.4×10^9 (rel.)	$k/k_{\text{thym}} = 1.00 \pm 0.10$	γ -r.	opt.	c.k.	67-0461
3.672	selenocystine (RSeSeR)	7	1.0×10^{10}	—	p.r.	opt.	p.b.k. at 460 nm (RSe \cdot); c.k. with CNS^- gave $k = 1.7 \times 10^{10}$.	73-1010
3.672a	selenomethionine	7	$\sim 1 \times 10^{10}$	—	p.r.	opt.	p.b.k. at 380 nm; c.k. gave $k/k_{\text{CNS}^-} = 1.2$.	74-1092
3.673	selenourea $\text{OH} + \text{NH}_2\text{CSeNH}_2 \rightarrow \text{H}_2\text{O} + \text{NHCSNH}_2$	6.5	6.9×10^9 5.5×10^9 1.2×10^{10} (rel.) 1.1×10^{10} (rel.) 1.2×10^{10} (rel.)	— — $k/k_{\text{CNS}^-} = 1.09$ $k/k_{\text{EtOH}} = 6.2$ $k/k_{\text{MeOH}} = 13.8$	p.r. p.r. p.r. p.r. p.r.	opt. opt. opt. opt. opt.	d.k. at 250 nm. p.b.k. at 410 nm. c.k. c.k. c.k.	70-0240 70-0240 70-0240 70-0240 70-0240
3.674	serine	2-2.2	2.5×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0228$	p.r.	opt.	c.k.	65-0388
		5.5-6	3.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0288$	p.r.	opt.	c.k.	65-0388
		2-2.2	2.9×10^8 (rel.)	$k/k_{\text{thym}} = 0.0532$	γ -r.	opt.	c.k.	65-0388
		6.6	2.3×10^8 (rel.)	$k/k_{\text{RNO}} = 0.184$	γ -r.	opt.	c.k.	73-0548
3.675	serum albumin, human	—	2.3×10^{10}	—	—	—	—	66-0844
		—	$\sim 6 \times 10^{10}$	—	—	—	calcd.	70-0253
3.676	starch, corn	6.5	2.8×10^7 (rel.)	$k/k_{\text{RNO}} = 0.0023 \pm 0.002$	γ -r.	opt.	c.k.	69-0580

TABLE 4. Reactions of OH with organic solutes — Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.677	starch, waxy	6.5	2.5×10^7 (rel.)	$k/k_{\text{RNO}} = 0.002 \pm 0.003$	γ -r.	opt.	c.k.	69-0580
3.678	styrene	5.5	$(6.0 \pm 0.6) \times 10^9$	—	p.r.	opt.	p.b.k. at 320 nm (66% $\text{C}_6\text{H}_5\text{CHCH}_2\text{OH}$) and 345 nm (33% ring addn.).	74-1138
3.679	suberic acid	2-2.2	4.0×10^9 (rel.)	$k/k_{\text{thym}} = 0.75 \pm 0.07$	γ -r.	opt.	c.k.	67-0461
3.680	succinic acid	1	7×10^6 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.03$	Fenton	chem.	c.k.	49-0002
		2-2.2	1.4×10^8 (rel.)	$k/k_{\text{thym}} = 0.026 \pm 0.002$	γ -r.	opt.	c.k.	67-0461
		1	8.9×10^7 (rel.)	$k/k_{\text{MeOH}} = 0.097$	Fenton	chem.	c.k.; $k_{\text{MeOH}}/k_{\text{Fe}^{2+}} = 4.3$.	73-9341
3.681	succinimide	3.5	5.0×10^8	—	p.r.	opt.	p.b.k.	71-0145
3.682	succinonitrile	—	3.0×10^7 (rel.)	$k/k_{\text{HCOO}^-} = 0.012$	γ -r.	chem.	c.k.	73-0364
3.683	sucrose	2-2.2	2.8×10^9 (rel.)	$k/k_{\text{thym}} = 0.52 \pm 0.05$	γ -r.	opt.	c.k.	67-0461
3.684	sulfacetamide, Na	—	4.7×10^9 (rel.)	—	γ -r.	—	c.k. with RNO.	71-0128
3.685	sulfaguanidine	—	3.1×10^9 (rel.)	—	γ -r.	opt.	c.k. with RNO assuming $k(\text{OH} + \text{sulfanilic acid}) = 2.93 \times 10^9$.	73-0094
3.686	sulfanilamide	—	3.2×10^9 (rel.)	—	γ -r.	—	c.k. with RNO.	71-0128
		—	1.6×10^9 (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{sulfanilic acid}) = 2.93 \times 10^9$.	73-0094
3.687	sulfanilic acid	7-8	1.9×10^9 (rel.)	$k/k_{\text{ferro}} = 0.21$	γ -r.	chem.	c.k.	74-0283
		—	3.4×10^9 (rel.)	—	γ -r.	—	c.k. with RNO.	71-0128
		0.4	2.1×10^9 (rel.)	$k/k_{2-\text{PrOH}} = 0.95$	γ -r.	chem.	c.k.	73-0270
		—	2.93×10^9	—	p.r.	opt.	p.b.k. at 270 nm.	73-0094
<i>For other ratios see: 3.189, 3.191, 3.700, 3.701, 3.702.</i>								
sulfasuccidine <i>See</i> 4-(2-thiazolylsulfamoyl)succanilic acid (3.701).								
sulfathiazole <i>See</i> N'-(2-thiazolyl)sulfanilamide (3.702).								
3.687a	superoxide dismutase	7.2	5.3×10^{10}	—	p.r.	opt.	p.b.k. at 330 nm; rate for bovine enzyme; human enzyme gave $k = 4.6 \times 10^{10}$.	74-3081
TAN <i>See</i> 2,2,6,6-tetramethylpiperidone-N-oxyl (3.697).								
3.688	tartaric acid	2-2.2	5.9×10^8 (rel.)	$k/k_{\text{thym}} = 0.11 \pm 0.01$	γ -r.	opt.	c.k.	67-0461
3.689	tartrate ion	9	6.7×10^8 (rel.)	$k/k_{\text{RNO}} = 0.054$	γ -r.	opt.	c.k.; assume $k_{\text{ferro}} = k_{\text{RNO}}$.	67-0555
		9	7.5×10^8 (rel.)	$k/k_{\text{RNO}} = 0.06$	r.	opt.	c.k.; $E_a = -1.2 \pm 0.3$ kcal/mol (-5 kJ/mol) (-8 to 23°).	71-0469
3.690	terephthalate ion	9	3.2×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.75$	γ -r.	opt.	c.k. with RNO.	66-0441
3.691	tetrachloro-ethylene $\text{OH} + \text{CCl}_2=\text{CCl}_2 \rightarrow \text{CCl}_2\text{CCl}_2\text{OH}$	—	$(2.3 \pm 0.3) \times 10^9$	—	p.r.	condy.	p.b.k. (Cl^-); $(\text{CCl}_2\text{CCl}_2\text{OH} \rightarrow \text{H}^+ + \text{Cl}^- + \text{CCl}_2\text{COCl})$	71-0709
		—	$(1.7 \pm 0.3) \times 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS^- ; reference rate not given.	71-0709
3.692	1,2,3,4-tetrafluorobenzene	—	5×10^9 (rel.)	—	p.r.	opt.	c.k. with CNS^- .	73-0054
3.693	tetrahydrofuran	1	1.4×10^9 (rel.)	$k/k_{\text{Fe}^{2+}} = 6.2$	Fenton	chem.	c.k.	49-0002
		9	2.7×10^5 (rel.)	$k/k_{\text{EtOH}} = 1.46$	γ -r.	opt.	c.k. with RNO.	66-0423
3.694	tetrahydropyran	1	1.0×10^9 (rel.)	$k/k_{\text{Fe}^{2+}} = 4.5$	Fenton	chem.	c.k.	49-0002

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
3.695	tetrahydroxysuc- cinate ion	9	1.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 0.7	γ-r.	opt.	c.k. with RNO.	66-0423
3.695a	α-tetralol	1.7-1.8	8.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{2-PrOH} = 3.7	Fenton	chem.	c.k. with 1- phenylethanol.	74-9006
3.695b	1,2,3,4-tetra- methylbenzene	~7	7.2 x 10 ⁹	—	p.r.	opt.	p.b.k.; OH addn. as well as H abstr.; rate for H abstr. = 1.9 x 10 ⁹ .	75-1009
3.695c	1,2,3,5-tetra- methylbenzene	~7	7.1 x 10 ⁹	—	p.r.	opt.	p.b.k.; OH addn. and H abstr.; rate for H abstr. = 1.8 x 10 ⁹ .	75-1009
3.695d	1,2,4,5-tetra- methylbenzene (Durene)	~7	7.0 x 10 ⁹	—	p.r.	opt.	p.b.k.; OH addn. and H abstr.; rate for H abstr. = 1.8 x 10 ⁹ .	75-1009
3.696	N,N,N',N'-tetra- methyl-1,2-dia- zenedicarboxamide	7.1 7 10.3- 11	7.7 x 10 ⁹ 3 x 10 ⁹ (rel.) 4.8 x 10 ⁹ (rel.)	— <i>k</i> / <i>k</i> _{CNS⁻} = 0.27 <i>k</i> / <i>k</i> _{carb} = 14.1	p.r. p.r. p.r.	opt. opt. opt.	p.b.k. at 400 nm; c.k. c.k.	74-1061 74-1061 74-1061
3.697	2,2,6,6-tetra- methylpiperidone- N-oxyl (TAN)	10.5 7 5-6 nat.	3.8 x 10 ⁹ (rel.) (4.1 ± 0.4) x 10 ⁹ < 10 ⁸ (rel.) (3.3 ± 0.3) x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{carb} = 10.5 — <i>k</i> / <i>k</i> _{ferro} < 10 ⁻² <i>k</i> / <i>k</i> _X = 5.2 ± 0.4	p.r. p.r. p.r. p.r.	opt. opt. opt. opt.	c.k.; cor. for CO ₃ ⁻ + TAN. d.k. at 230 nm. c.k. c.k. with phenyl- alanine (X); <i>k</i> _X = 6.3 x 10 ⁹ ; cor. for H; obs. X abs. at 320 nm.	71-0061 71-0061 71-0618 72-3021
3.698	tetrasulfonated Cu phthalocyanine	10.7	7.3 x 10 ⁹ (rel.) 7.6 x 10 ⁹ (rel.) 7.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{carb} = 20 <i>k</i> / <i>k</i> _{EtOH} = 4.1 <i>k</i> / <i>k</i> _{MeOH} = 8.6	γ-r. γ-r. γ-r.	chem. chem. chem.	c.k. c.k. c.k.	69-0827 69-0827 69-0827
3.699	tetronate ion	7	9.2 x 10 ⁹	—	p.r.	opt.	d.k. at 248 nm.	74-1053
3.700	thalamyd	—	6.3 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO rel. to <i>k</i> (OH + sulfanilic acid) = 2.93 x 10 ⁹ .	73-0094
3.701	4-(2-thiazolyl- sulfamoyl)- succinanic acid (sulfasuccidine)	—	4.6 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO rel. to <i>k</i> (OH + sulfanilic acid) = 2.93 x 10 ⁹ .	73-0094
3.702	N'-(2-thiazolyl)- sulfanilamide (sulfathiazole)	—	7.8 x 10 ⁹ (rel.)	—	γ-r.	opt.	c.k. with RNO rel. to <i>k</i> (OH + sulfanilic acid) = 2.93 x 10 ⁹ .	73-0094
3.703	thiodiglycolic acid	1	6.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.546	p.r.	opt.	c.k.	65-0387
3.704	thioglycolic acid	1	6.2 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{Fe²⁺} = 2.7	Fenton	chem.	c.k.	49-0002
3.705	thioglycolate ion	6.6 11.1	5.9 x 10 ⁹ (rel.) 5.5 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.53 <i>k</i> / <i>k</i> _{CNS⁻} = 0.05	p.r.	opt.	c.k.; p <i>K</i> _a = 3.7, 10.3 for thiogly- colic acid.	73-0090
3.706	thiolactate ion	7.2 10.8	1.7 x 10 ¹⁰ (rel.) 1.6 x 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.55 <i>k</i> / <i>k</i> _{CNS⁻} = 1.45	p.r.	opt.	c.k.; p <i>K</i> _a ≅ 4, 10.7 for thiolactic acid.	73-0090
3.707	thiophene OH + C ₄ H ₄ S → (OH)C ₄ H ₄ S	—	3.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.304	p.r.	opt.	c.k.	71-0360
3.708	threonine	2-2.2 6.6	3.9 x 10 ⁸ (rel.) 5.1 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{thym} = 0.0727 <i>k</i> / <i>k</i> _{RNO} = 0.041	γ-r. γ-r.	opt. opt.	c.k. c.k.	65-0388 73-0548

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.	
3.709	thymidine	2-2.2	4.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.417	p.r.	opt.	c.k.	65-0388	
		5-5.2	5.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.455	p.r.	opt.	c.k.	65-0388	
		7.4-7.6	4.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.417	p.r.	opt.	c.k.	65-0388	
		~7	4.7 x 10 ⁹	—	p.r.	opt.	p.b.k. at 375 (pH = 7) and 400 (pH = 12.4) nm.	68-0312	
		~12.4	2.1 x 10 ⁹	—	p.r.	opt.	p.b.k. at 400 nm.	70-3069	
3.710	thymidylic acid	5.6	(5.3 ± 0.5) x 10 ⁹	—	p.r.	opt.	p.b.k. at 400 nm.	70-3069	
		2-2.2	4.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.394	p.r.	opt.	c.k.; NH ₄ ⁺ salt.	65-0388	
		6.5-7.0	5.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.477	p.r.	opt.	c.k.; NH ₄ ⁺ salt.	65-0388	
3.711	thymine OH addn. to 5,6-double bond	0.7-7	3.7 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.00 ± 0.10	γ-r.	opt.	c.k.	65-0133	
		1	5.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.47	p.r.	opt.	c.k.	65-0387	
		2-2.2	5.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.47	p.r.	opt.	c.k.	65-0388	
		5-5.5	5.0 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.455	p.r.	opt.	c.k.	65-0388	
		7.2-7.4	5.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.485	p.r.	opt.	c.k.	65-0388	
		2	7.8 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{pH} = 1	γ-r.	opt.	c.k.	67-0461	
		9	6.2 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{RNO} = 0.5	γ-r.	opt.	c.k.; assume <i>k</i> _{ferro} = <i>k</i> _{RNO} .	67-0555	
		~7	7.4 x 10 ⁹	—	p.r.	opt.	p.b.k.; obs. transient at 400 and 550 (pH = 12.4) nm.	68-0312	
		~11	3.9 x 10 ⁹	—	p.r.	opt.	p.b.k.; obs. disappearance of 5,6-double bond at 270 nm.	69-0571	
		~12.4	1.1 x 10 ⁹	—	p.r.	opt.	p.b.k.; OH-adduct obs. at 385 nm.	69-0571	
		—	5.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{1⁻} = 0.435	X-r.	opt.	c.k.	68-0359	
		—	4.1 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{MeOH} = 4.55	X-r.	opt.	c.k.	68-0359	
		—	4.9 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.63	X-r.	opt.	c.k.	68-0359	
		7	7.4 x 10 ⁹	—	p.r.	opt.	p.b.k. at 400 nm (adduct).	68-0597	
		7	(7.4 ± 0.5) x 10 ⁹	—	p.r.	opt.	d.k.; obs.	69-0571	
		7	(4.6 ± 0.3) x 10 ⁹	—	p.r.	opt.	p.b.k.; OH-adduct obs. at 385 nm.	69-0571	
		7	7.6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = (0.69 ± 0.15)	p.r.	opt.	c.k.; cor. for incomplete scav. of <i>e</i> _{aq} ⁻ by H ₂ O ₂ .	69-0571	
		1	—	<i>k</i> / <i>k</i> _{perox} = 72.4	Fe(II) + H ₂ O ₂	esr	c.k.	69-5278	
		1	—	<i>k</i> / <i>k</i> _{perox} = 71.5	Ti(III) + H ₂ O ₂	esr	c.k.	69-5278	
		3.712	<i>p</i> -toluate ion OH + CH ₃ C ₆ H ₄ COO ⁻ → CH ₃ (OH)C ₆ H ₄ COO ⁻	9	4.4 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.38	γ-r.	opt.	c.k. with RNO.
9	8 x 10 ⁹			—	p.r.	opt.	p.b.k. at 340 nm.	72-0047	
For other ratios see: 3.25, 3.27, 3.129, 3.131, 3.149, 3.150, 3.154, 3.160, 3.161, 3.162, 3.177, 3.180, 3.181, 3.182, 3.186, 3.225, 3.226, 3.239, 3.247, 3.266, 3.289-90, 3.291-94, 3.343, 3.346, 3.358, 3.361, 3.369, 3.371, 3.374, 3.384, 3.385, 3.394, 3.399, 3.400, 3.401, 3.402, 3.403, 3.404, 3.409, 3.410, 3.411, 3.415, 3.418, 3.419, 3.420, 3.421, 3.422, 3.423, 3.424, 3.425, 3.426, 3.435, 3.441, 3.461, 3.484, 3.491, 3.493, 3.497, 3.501, 3.503, 3.507, 3.511, 3.513, 3.545, 3.546, 3.585, 3.587, 3.588, 3.589, 3.594, 3.596, 3.597, 3.602, 3.613, 3.619, 3.633, 3.634, 3.636, 3.637, 3.642, 3.671, 3.674, 3.679, 3.680, 3.683, 3.688, 3.708, 3.735, 3.741, 3.753.									
nat.	4.7 x 10 ⁹ (rel.)			<i>k</i> / <i>k</i> _{ferro} = 0.505	p.r.	opt.	c.k.	71-0578	
nat.	5.1 x 10 ⁹			—	p.r.	opt.	d.k.	71-0578	
9	5.5 x 10 ⁹			—	p.r.	opt.	p.b.k. at 375 nm.	72-0047	
6-7	4.7 x 10 ⁹ (rel.)			<i>k</i> / <i>k</i> _{RNO} = 0.38	γ-r.	opt.	c.k.; 17°C.	75-0294	

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction pH	k	Ratio	Source	Method	Comment	Ref.
3.713	toluene 3 (I) $\text{OH} + \text{C}_6\text{H}_5\text{CH}_3 \rightarrow$ $\text{C}_6\text{H}_5\text{CH}_2 + \text{H}_2\text{O}$ ~7 (II) $\text{OH} + \text{C}_6\text{H}_5\text{CH}_3 \rightarrow$ $\text{C}_6\text{H}_5(\text{OH})\text{CH}_3$	$(3.0 \pm 0.7) \times 10^9$ 6.8×10^9 4.0×10^8 (I)	— — $k_I/k_{II} = 0.033$	p.r. p.r.	opt. opt.	p.b.k. at 313 and 309 nm. p.b.k. at 258 nm ($\text{C}_6\text{H}_5\text{CH}_2$).	64-0115 73-0089, 75-1009
3.714	<i>o</i> -toluenesulfonate ion —	3.2×10^9 (rel.)	$k/k_{\text{RNO}} = 0.258$	r.	opt.	c.k.	72-0425
3.715	<i>m</i> -toluenesulfonate ion —	3.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.303$	r.	opt.	c.k.	72-0425
3.716	<i>p</i> -toluenesulfonate ion 9	1.8×10^9 (rel.) 3.7×10^9 (rel.)	— $k/k_{\text{RNO}} = 0.294$	— r.	— opt.	c.k. with RNO. c.k. with RNO.	66-0843 72-0425
3.717	<i>o</i> -tolyl- β -D-glucopyranoside —	5.3×10^9 (rel.) 3.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.485$ —	p.r. γ -r.	opt. opt.	c.k. c.k. with RNO; relative to $k(\text{OH} + \text{X}) = 4.4 \times 10^9$, X = phenyl- β -D-glucopyranoside.	71-0056 71-0056
3.718	<i>m</i> -tolyl- β -D-glucopyranoside —	3.0×10^9 (rel.)	—	γ -r.	opt.	c.k. with RNO; relative to $k(\text{OH} + \text{X}) = 4.4 \times 10^9$, X = phenyl- β -D-glucopyranoside.	71-0056
3.719	<i>p</i> -tolyl- β -D-glucopyranoside —	6.2×10^9 (rel.) 2.7×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.56$ —	p.r. γ -r.	opt. opt.	c.k. c.k. with RNO; rel. to $k(\text{OH} + \text{X}) = 4.4 \times 10^9$, X = phenyl- β -D-glucopyranoside.	71-0056 71-0056
3.720	<i>p</i> -tolyl-S- β -D-thioglucopyranoside —	3.6×10^9 (rel.) 8.7×10^9	$k/k_{\text{CNS}^-} = 0.33$ —	p.r. p.r.	opt. opt.	c.k. transient absorbs at 320 nm.	71-0056 70-1056
3.720a	tributyl phosphate 1.2	1.03×10^{10} (rel.)	$k/k_{\text{EtOH}} = 5.5$	γ -r.	chem.	c.k.; $k/k_{\text{HNO}_3} = 77$.	74-0439
3.721	1,1,2-trichloroethylene — $\text{OH} + \text{CHCl}=\text{CCl}_2 \rightarrow$ $\text{CHCl}(\text{OH})\text{CCl}_2$ —	$(4.0 \pm 0.4) \times 10^9$ $(2.6 \pm 0.3) \times 10^9$ (rel.)	— —	p.r. p.r.	condy. opt.	p.b.k. (Cl^-); ($\text{CHClOHCCl}_2 \rightarrow \text{H}^+ + \text{Cl}^- + \text{CCl}_2\text{CHO}$). c.k. with CNS^- ; reference rate not given.	71-0709 71-0709
3.722	2,4,6-trichlorophenyl- β -D-glucopyranoside —	1.9×10^9 (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{X}) = 4.4 \times 10^9$, X = phenyl- β -D-glucopyranoside.	71-0056
3.723	triethylamine — (I) $\text{OH} + \text{Et}_3\text{N} \rightarrow \text{H}_2\text{O} + \text{CH}_3\text{CHNEt}_2$ (II) $\text{OH} + \text{Et}_3\text{N} \rightarrow \text{H}_2\text{O} + \text{CH}_2\text{CH}_2\text{NEt}_2$ 12	1.1×10^{10} (rel.) 3.7×10^9 (I) (rel.)	$k/k_{\text{CNS}^-} = 1$ $k_I/k_{\text{EtOH}} = 2$	p.r. γ -r.	opt. chem.	c.k.; extrapolated value based on $k/k_{\text{CNS}^-} = 0.73$ at pH 11. c.k.; no II obs.; may be O^- reaction.	71-0585 71-0590
3.724	triethylammonium ion 1 3.6 (I) $\text{OH} + \text{Et}_3\text{NH}^+ \rightarrow \text{CH}_3\text{CHNH}^+\text{Et}_2 + \text{H}_2\text{O}$ 1.5, 6.5 (II) $\text{OH} + \text{Et}_3\text{NH}^+ \rightarrow \text{H}_2\text{O} + \text{CH}_2\text{CH}_2\text{NH}^+\text{Et}_2$	1.8×10^8 (rel.) 3.5×10^8 (rel.) 1.3×10^8 (rel.)	$k/k_{\text{Fe}^{2+}} = 0.8$ $k/k_{\text{CNS}^-} = 0.032$ $k_I/k_{\text{EtOH}} = 0.068$	Fenton p.r. γ -r.	chem. opt. chem.	c.k. c.k. c.k.; $k_{II}/k_I = 0.76$.	49-0002 71-0585 71-0590
3.725	trifluoroacetate ion 9	2×10^5 (rel.)	—	—	—	c.k. with RNO.	66-0843
3.725a	1,2,3-trimethoxybenzene —	$(8.0 \pm 0.8) \times 10^9$	—	p.r.	—	—	75-1171

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.725b	1,2,4-trimethoxy- benzene	—	$(8.1 \pm 0.8) \times 10^9$	—	p.r.	—	—	75-1171
3.725c	1,3,5-trimethoxy- benzene	—	$(8.1 \pm 0.8) \times 10^9$	—	p.r.	—	—	75-1171
3.726	trimethylacetate ion trimethylamine $\text{OH} + (\text{CH}_3)_3\text{N} \rightarrow$ $\text{H}_2\text{O} + \text{CH}_2\text{N}(\text{CH}_3)_2$	—	1.3×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.2$	p.r.	opt.	c.k.; extrapolat- ed value based on $k/k_{\text{CNS}^-} = 1.1$ at pH 10.9.	71-0585
3.727	trimethylammonium ion $\text{OH} + (\text{CH}_3)_3\text{NH}^+ \rightarrow$ $\text{H}_2\text{O} +$ $\text{CH}_2\text{NH}^+(\text{CH}_3)_2$	7.5	4×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.0364$	p.r.	opt.	c.k.	71-0585
3.727a	1,2,3-trimethyl- benzene	~7	7.0×10^9	—	p.r.	opt.	p.b.k.; OH addn. as well as H abstr.; rate for H abstr. = 1.3×10^9 .	75-1009
3.727b	1,2,4-trimethyl- benzene	~7	6.2×10^9	—	p.r.	opt.	p.b.k.; OH addn. and H abstr.; rate for H abstr. = 1.15 $\times 10^9$.	75-1009
3.727c	1,3,5-trimethyl- benzene (Mesitylene)	~7	6.4×10^9	—	p.r.	opt.	p.b.k.; OH addn. and H abstr.; rate for H abstr. = 1.2 $\times 10^9$.	75-1009
3.728	2,4,6-trimethyl- 3-hydroxypyri- dine	6.5	2.5×10^9 (rel.)	$k/k_{\text{RNO}} = 0.20 \pm$ 0.03	γ -r.	opt.	c.k.	69-0580
3.729	2,4,5-trimethyl- phenyl- β -D-glucoside	—	3.2×10^9 (rel.)	—	γ -r.	opt.	c.k. with RNO; rel. to $k(\text{OH} + \text{X}) =$ 4.4×10^9 , X = phenyl- β -D-glucoside.	71-0056
3.730	trimethyl phosphate $\text{OH} + (\text{CH}_3\text{O})_3\text{PO} \rightarrow$ $\text{H}_2\text{O} +$ $\text{CH}_2\text{O}(\text{CH}_3\text{O})_2\text{PO}$	—	1.2×10^8 (rel.)	$k/k_{\text{CNS}^-} = 0.011$	p.r.	opt.	c.k.	72-3008
3.731	2,4,6-trimethyl- 1,3,5-trioxane	9	1×10^9 (rel.)	$k/k_{\text{EtOH}} = 0.546$	γ -r.	opt.	c.k. with RNO.	66-0423
3.732	1,3,5-trioxane tryptaflavin <i>See</i> acriflavin (3.141).	9	4.9×10^8 (rel.)	$k/k_{\text{EtOH}} = 0.264$	γ -r.	opt.	c.k. with RNO.	66-0423
3.733	trypsin	—	2.5×10^{10} (rel.)	—	X-r.	biol	effect on enzyme inact. compared with acetone, glycylglycine, glycerol, glucose, ethanol, formate ion.	67-3044
		~7	$(8.2 \pm 1.2) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 330 nm. or c.k. with glu- cose ($k = 1 \times$ 10^9).	71-3069
3.734	trypsinogen	6.3 7.4	3.9×10^{10} (rel.) $(8.5 \pm 0.5) \times 10^{10}$ (rel.)	$k/k_{\text{RNO}} = 3.1$ —	γ -r. p.r.	opt. opt.	c.k. c.k. with glucose ($k = 1 \times 10^9$); obs. 330 nm abs.	73-0548 71-3069
3.735	tryptophan, positive ion	2-2.2 2-2.2	1.1×10^{10} (rel.) 7.7×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.985$ $k/k_{\text{thym}} = 1.42 \pm$ 0.15	p.r. γ -r.	opt. opt.	c.k. c.k.; k from initial slope of competition plot.	65-0388 65-0388, 67-0461

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.736	L-tryptophan, positive ion	1	$(1.25 \pm 0.2) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 560 nm.	69-0459
3.737	tryptophan, zwitterion	6.1-6.3	1.4×10^{10} (rel.)	$k/k_{\text{CNS}^-} = 1.29$	p.r.	opt.	c.k.	65-0388
		6.2	7.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.62$	γ -r.	opt.	c.k.	73-0548
3.738	L-tryptophan, zwitterion	8.8	$(1.2 \pm 0.3) \times 10^{10}$	—	p.r.	opt.	p.b.k. at 310 nm.	69-0459
<i>For other ratios see: 3.211, 3.252, 3.273, 3.333, 3.334, 3.335, 3.454, 3.468, 3.470, 3.472, 3.517, 3.539, 3.540, 3.541, 3.542.</i>								
3.739	omitted							
3.740	tyramine, negative ion	11.2	$(1.5 \pm 0.2) \times 10^{10}$	$k/k_{\text{CNS}^-} = 1.36$	p.r.	opt.	c.k.	73-0003
3.741	tyrosine, positive ion	2-2.2	1×10^{10} (rel.)	$k/k_{\text{thym}} = 1.87$	p.r.	opt.	c.k.	65-0388
3.742	L-(-)-tyrosine, positive ion	4.0	1.5×10^{10} (rel.)	$k/k_{\text{HCOO}^-} = 4.2$	p.r.	opt.	c.k.; meas. transient at 310-320 nm.	68-0062
3.743	tyrosine, negative ion	5.2	$(1.4 \pm 0.3) \times 10^{10}$ (rel.)	$k/k_{\text{CNS}^-} = 1.3$	p.r.	opt.	c.k.	73-0003
		6.5	1.05×10^{10} (rel.)	$k/k_{\text{RNO}} = 0.84$	γ -r.	opt.	c.k.	73-0548
3.744	tyrosine, dinegative ion	11.2	$(1.3 \pm 0.3) \times 10^{10}$ (rel.)	$k/k_{\text{CNS}^-} = 1.2$	p.r.	opt.	c.k.	73-0003
3.745	L-(-)-tyrosine, negative ion	10.6	2×10^{10} (rel.)	$k/k_{\text{carb}} = 53.7$	p.r.	opt.	c.k.	68-0062
3.746	uracil	9.0	6.8×10^9 (rel.)	$k/k_{\text{RNO}} = 0.542 \pm 0.027$	γ -r.	opt.	c.k.	65-0356,
	$\text{OH} + \text{C}_5\text{H}_6\text{N}_2\text{O}_2 \rightarrow$							67-0555
	$\text{C}_5\text{H}_6\text{N}_2\text{O}_2 \cdot \text{OH}$	2-2.2	4.8×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.44$	p.r.	opt.	c.k.	65-0388
		5-5.2	5.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.469$	p.r.	opt.	c.k.	65-0388
		7.3-7.5	5.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.477$	p.r.	opt.	c.k.	65-0388
		7.0	$(7.4 \pm 1.0) \times 10^9$	—	p.r.	opt.	d.k. at 270 nm.	68-0316
		6.5	7.4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.67$	p.r.	opt.	c.k.; cor. for e_{aq}^- not scav. by $10^{-2} M \text{H}_2\text{O}_2$.	68-0316, 69-0571
		7	$(6.0 \pm 0.3) \times 10^9$	—	p.r.	opt.	d.k. at 270 nm.	69-0571
		7	$(6.5 \pm 0.7) \times 10^9$	—	p.r.	opt.	p.b.k. at 385 nm.	69-0571
		5.9	$(5.8 \pm 0.2) \times 10^9$	—	p.r.	opt.	p.b.k. at 400 nm.	70-3069
		nat.	4.2×10^9 (rel.)	$k/k_{\text{ferro}} = 0.452$	p.r.	opt.	c.k.	71-0578
		—	6.0×10^9	—	p.r.	opt.	p.b.k. as well as as d.k.	73-3016
		6-7	4.5×10^9 (rel.)	$k/k_{\text{RNO}} = 0.36$	γ -r.	opt.	c.k.; 17°C.	75-0294
<i>For other ratios see: 3.289, 3.290.</i>								
3.747	uracil dinucleotide (UpU)	7	$(3.8 \pm 0.2) \times 10^9$	—	p.r.	opt.	d.k. at 270 nm.	69-0571
		7	5.3×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.48 \pm 0.08$	p.r.	opt.	c.k.; rates calcd. per nucleotide base.	69-0571
uracil mononucleotides See uridine monophosphate (3.751).								
3.748	uracil oligonucleotide (oligo U)	7	$(4.3 \pm 0.2) \times 10^9$	—	p.r.	opt.	d.k. at 270 nm.	69-0571
		7	4×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.36 \pm 0.07$	p.r.	opt.	c.k.; rate calcd. per nucleotide base.	69-0571
uracil polynucleotide (poly U) See polyuridylic acid (3.631).								
3.749	urea	9.0	$< 1.25 \times 10^6$	$k/k_{\text{RNO}} < 10^{-4}$	γ -r.	opt.	c.k.	65-0356
3.749a	uric acid	6-7	7.2×10^9 (rel.)	$k/k_{\text{RNO}} = 0.58$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.750	uridine	7	4.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.38 \pm 0.08$	p.r.	opt.	c.k.; cor. for incomplete scavenging of e_{aq}^- by H_2O_2 .	68-0316, 69-0571
		7	$(6.5 \pm 0.5) \times 10^9$	—	p.r.	opt.	d.k. at 270 nm.	69-0571
		7	$(4.1 \pm 0.2) \times 10^9$	—	p.r.	opt.	p.b.k.; OH adduct obs. at 385 nm.	69-0571
		6.5	2.4×10^9 (rel.)	$k/k_{\text{RNO}} = 0.19$	γ -r.	opt.	c.k.	69-0580

TABLE 4. Reactions of OH with organic solutes - Continued

No.	Solute and Reaction	pH	k	Ratio	Source	Method	Comment	Ref.
3.750 cont.		5.4	$(4.5 \pm 0.3) \times 10^9$	—	p.r.	opt.	p.b.k. at 400 nm.	70-3069
		7	4.1×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.36$	p.r.	opt.	c.k.; unpubl. data.	73-1071
3.751	uridine monophosphate (uridylic acid)	7	5.2×10^9 (rel.)	$k/k_{\text{CNS}^-} = 0.47 \pm 0.1$	p.r.	opt.	c.k.; cor. for incomplete scav. of e_{aq}^- by H_2O_2 .	68-0316, 69-0571
		7	$(4.6 \pm 0.3) \times 10^9$	—	p.r.	opt.	d.k. at 270 nm.	69-0571
		7	$(4.0 \pm 0.4) \times 10^9$	—	p.r.	opt.	p.b.k.; OH adduct obs. at 385 nm.	69-0571
		6.5	2.5×10^9 (rel.)	$k/k_{\text{RNO}} = 0.2 \pm 0.02$	γ -r.	opt.	c.k.	69-0580
		7.0	$(5.1 \pm 0.3) \times 10^9$	—	p.r.	opt.	p.b.k. at 400 nm.	70-3069
		7	4.5×10^9	—	p.r.	opt.	p.b.k. at 390 nm.	73-1071
3.752	valerate ion	9	2.9×10^9 (rel.)	$k/k_{\text{EtOH}} = 1.55$	γ -r.	opt.	c.k. with RNO.	66-0423
3.753	valine	2-2.2	7.2×10^8 (rel.)	$k/k_{\text{thym}} = 0.134$	γ -r.	opt.	c.k.	65-0388
		6.6	6.6×10^8 (rel.)	$k/k_{\text{RNO}} = 0.053$	γ -r.	opt.	c.k.	73-0548
3.754	vinyl chloride $\text{OH} + \text{CH}_2=\text{CHCl} \rightarrow \text{CH}_2\text{OHCHCl}$ vinyl methyl ketone <i>See</i> 1-butene-3-one (3.229). Vitamin B12 <i>See</i> cyanocobalamin (3.272a). Vitamin B12a <i>See</i> hydroxocobalamin (3.447).	—	$(7.1 \pm 0.5) \times 10^9$ (rel.)	—	p.r.	opt.	c.k. with CNS^- ; reference rate not given.	71-0709
3.754a	xanthine	8.0	8.9×10^9 (rel.)	$k/k_{\text{RNO}} = 0.71$	γ -r.	opt.	c.k.; 17°C.	75-0294
3.755	<i>o</i> -xylene	~7	6.7×10^9	—	p.r.	opt.	p.b.k.; OH addn. and H abstr.; rate for H abstr. = 8.0×10^8 .	75-1009
3.756	<i>m</i> -xylene	~7	7.5×10^9	—	p.r.	opt.	p.b.k.; OH addn. and H abstr.; rate for H abstr. 9.0×10^8 .	75-1009
3.757	<i>p</i> -xylene	~7	7.0×10^9	—	p.r.	opt.	p.b.k.; OH addn. and H abstr.; rate for H abstr. 8.4×10^8 .	75-1009
3.758	xlenol orange	11	2.2×10^{10} (rel.)	$k/k_{\text{MeOH}} = 24.5$	γ -r.	opt.	c.k.	71-0437

TABLE 5. Reactions of O^- with water, transients from water, inorganic solutes, and organic solutes

No.	Solute and reaction	pH	k	Ratio	Source	Method	Comment	Ref.
4.1	H_2O $O^- + H_2O \rightarrow$ $OH^- + OH$	11	2×10^6 (rel.)	$k/k_{EtOH} = (9.6 \pm 1.0) \times 10^{-2}$ $M[H_2O]$	p.r.	opt.	c.k. with CO_3^{2-} ; N_2O and O_2 -satd. solns.	70-0511
		11	1.5×10^6 (rel.)	$k/k_{MeOH} = (1.3 \pm 0.1) \times 10^{-1}$ $M[H_2O]$	p.r.	opt.	c.k. with CO_3^{2-} ; N_2O and O_2 -satd. solns.	70-0511
		8-8.8	$(1.75 \pm 0.4) \times 10^6$ (rel.)	—	p.r.	opt.	c.k.; soln. contains ferrocyanide with methanol or ethanol; assumed $k(O^- + ferro) \leq 3 \times 10^7$, $k(O^- + EtOH) = 9.8 \times 10^8$, $k(O^- + MeOH) = 5.3 \times 10^8$.	71-0137
4.2	$O^- \rightarrow$ 1st order decay	13- 13.7	$4.3 \times 10^4 s^{-1}$ (rel.)	$k/k_{oxy} = (1.2 \pm 0.24) \times 10^{-5} dm^{-3} mol$	f.phot.	opt.	d.k. of O_3^- .	68-7277
		>13	$4.3 \times 10^4 s^{-1}$ (rel.)	$k/k_{oxy} = (1.2 \pm 0.4) \times 10^{-5} dm^{-3} mol$	p.r.	opt.	d.k. of O_3^- .	69-0002
4.3	e_{aq}^- $O^- + e_{aq}^- \rightarrow 2OH^-$	alk.	$\sim 2 \times 10^{10}$	—	—	—	See 1.9, S1.5, NSRDS-NBS 43 and supplement.	73-0030 75-0002
4.4	OH $O^- + OH \rightarrow HO_2^-$	alk.	$< 2.6 \times 10^{10}$	—	—	—	See 3.4 (Table 2)	—
4.5	O^- $O^- + O^- \rightarrow O_2^{2-}$	12- 13	$\sim 1 \times 10^9$	—	p.r.	opt.	curve fitting; $N_2O-Fe(CN)_6^{4-}$ soln.	64-0213
		13	8.3×10^9 (rel.)	$k/k_{oxy} = 2.3$	p.r.	opt.	c.k.; obs. O_3^- .	66-0001
		>12	$\leq 9 \times 10^8$ (rel.)	—	p.r.	opt.	c.k. with $Fe(CN)_6^{4-}$; est. based on numerous assumptions; $pK_a(OH) = 11.9$.	66-0424
4.6	BH_4^- $O^- + BH_4^- + (H_2O) \rightarrow$ $BH_4 + 2OH^-$	11- 12.83	$< 4 \times 10^8$	—	p.r.	opt.	calcd. from p.b.k.; assumed $pK_a(OH) = 11.8$ and $k(OH + BH_4^-) = 1.2 \times 10^{10}$.	70-1046
4.7	Br^- $O^- + Br^- \rightleftharpoons BrO^{2-}$ $BrO^{2-} + (H_2O) \rightleftharpoons$ $Br + 2OH^-$	13	4.5×10^7 (rel.)	$k/k_{2-PrOH} = 0.03$	γ -r.	chem.	c.k.; obs. $G(\text{acetone})$.	68-0602
		6-7	2×10^8 (rel.)	$k/k_{EtOH} = 0.18$ $k/k_{MeOH} = 0.34$	p.r.	opt.	c.k.; soln. contains N_2O ($e_{aq}^- + N_2O \rightarrow N_2 + O^-$).	71-0137
4.8	BrO^- $O^- + BrO^- (+ H_2O) \rightarrow$ $BrO + O^{2-}$ or	11- 13	4.4×10^9 (rel.)	$k/k(OH + CO_3^{2-}) = 11$	p.r.	opt.	c.k.; $pK_a(OH) = 11.9$; $\mu = 0.4$.	68-0153
	$\rightarrow BrO + 2OH^-$	12- 13	$(2.0 \pm 0.4) \times 10^9$ (rel.)	—	f.phot.	opt.	d.k. of O_3^- ; anal. of data is complex.	69-7340

TABLE 5. Reactions of O^- with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction pH	k	Ratio	Source	Method	Comment	Ref.
4.9	BrO_2^- $O^- + BrO_2^- (+ H_2O) \rightarrow BrO_2 + 2OH^-$ 13	1.7×10^9 (rel.)	$k/k(OH + CO_3^{2-}) = 4.5$	p.r.	opt.	c.k.; assume $k(OH + BrO_2^-) = 1.9 \times 10^9$ and $pK_a(OH) = 11.9$; $\mu = 0.4$.	68-0153
	12-13	$(1.1 \pm 0.2) \times 10^9$ (rel.)	—	f.phot.	opt.	d.k. of O_3^- ; data anal. is complex.	69-7340
4.10	BrO_3^- $O^- + BrO_3^- (+ H_2O) \rightarrow BrO_3 + 2OH^-$ 12-13	$(1.2 \pm 0.2) \times 10^6$ (rel.)	—	f.phot.	chem.	c.k. of O_3^- ; more than one rate involved in calcn.; may be up to 30% lower.	69-7340
4.11	BrO_4^- 7	$< 10^7$	—	p.r.	opt.	very slow or no reaction.	73-0106
4.12	CNS^- $O^- + CNS^- (+ H_2O) \rightarrow CNSOH^- + OH^-$ 13.5	1.0×10^9	—	p.r.	opt.	p.b.k.; assume product is CNS.	65-0386
	$CNSOH^- + CNS^- \rightleftharpoons (CNS)_2^- + OH^-$ 6-7	1.6×10^9 (rel.) 1.6×10^9 (rel.)	$k/k_{EtOH} = 1.5$ $k/k_{MeOH} = 2.8$	p.r.	opt.	c.k.	71-0137
	$CNSOH^- \rightleftharpoons CNS + OH^-$ alk.	1.1×10^9	—	p.r.	opt.	p.b.k. at 0.36 M NaOH; $k = 1.3 \times 10^9$ at 1.08 M NaOH.	71-0137
	$CNS + CNS^- \rightleftharpoons (CNS)_2^-$ 13	$(3.7 \pm 0.3) \times 10^9$	—	p.r.	opt.	p.b.k.	72-0126
4.13	CO_3^{2-} $O^- + CO_3^{2-} (+ H_2O) \rightarrow CO_3 + 2OH^-$ —	$\leq 10^7$ $\leq 5 \times 10^5$	—	p.r.	opt.	no details given.	66-0139
4.14	Ce^{3+} $O^- + Ce^{3+} \rightarrow Ce^{4+} + 2OH^-$ 2.3-2.6	6.6×10^8 (rel.)	$k/k_{EtOH} = 0.6 \pm 0.2$	p.r.	opt.	c.k.; assuming $k(O^- + H_2O) = 1.75 \times 10^6$.	71-0137
4.15	ClO^- $O^- + ClO^- \rightarrow ClO + O^{2-}$ 13	$(2.2 \pm 0.1) \times 10^8$ (rel.)	$k/k(OH + CO_3^{2-}) = 0.6$	p.r.	opt.	c.k.	72-0301
4.16	ClO_2^- $O^- + ClO_2^- \rightarrow ClO_2 + O^{2-}$ 13	$(1.7 \pm 0.1) \times 10^8$ (rel.)	$k/k(OH + CO_3^{2-}) = 0.48$	p.r.	opt.	c.k.	72-0301
4.17	ClO_3^- 13	$< 10^6$	—	p.r.	opt.	no effect on CO_3^- formn. in carbonate soln.	72-0301
4.18	Fe^{2+} $O^- + Fe^{2+} \rightarrow Fe^{3+} + 2OH^-$ 4.4-4.8	3.5×10^9 (rel.)	$k/k_{EtOH} = 3.2 \pm 1.2$	p.r.	opt.	c.k.	71-0137
4.19	$Fe(CN)_6^{4-}$ $O^- + Fe(CN)_6^{4-} (+ H_2O) \rightarrow Fe(CN)_6^{3-} + 2OH^-$ 13	5.8×10^8 (rel.)	$k/k_{MeOH} = 0.98$	γ -r.	chem.	c.k.; assuming that at pH = 13 most of OH is present as O^- ; not cor. for OH.	63-0072
	13	1.5×10^9 (rel.)	$k/k_{EtOH} = 1.36$	p.r.	opt.	c.k.; $k_{EtOH}/k_{oxy} = 0.35$; not cor. for OH.	65-0007
	13	9×10^8 (rel.)	$k/k_{HCOO^-} = 0.9 \pm 0.1$	X-r.	chem.	c.k.; assuming $k(O^- + HCOO^-) = 1 \times 10^9$; not cor. for OH.	67-0064
	—	$\leq 3 \times 10^7$	—	p.r.	opt.	estd. from $k_{obs} = 2.57 \times 10^8$ and	71-0137

TABLE 5. Reactions of O^- with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction pH	k	Ratio	Source	Method	Comment	Ref.
4.19 cont.						1.6×10^8 for 0.366 and 1.11 M NaOH, resp.	
4.20	FeO_4^{2-}, FeO_2^- (I) $O^- + FeO_4^{2-} \rightarrow HFeO_4^{2-} + HO_2^-$ (II) $O^- + FeO_2^- \rightarrow FeO_3^{2-}$	14 —	$k_I/k_{II} \cong 0.03$	γ -r.	chem.	c.k.; $K \cong 10^{-4}$ for $Fe(OH)_3 + OH^- \rightleftharpoons Fe(OH)_4^-$ is involved in calcn.	67-0614
4.21	H_2 $O^- + H_2 \rightarrow H + OH^-$	13.3 (8 \pm 4) $\times 10^7$ (rel.)	—	p.r.	opt.	rel. to $2k = 1.1 \times 10^{10}$ for $e_{aq}^- + e_{aq}^- \rightarrow H_2 + 2OH^-$	65-0009
4.22	HO_2^- $O^- + HO_2^- \rightarrow OH^- + O_2^-$	13.0 (1.0 \pm 0.4) $\times 10^9$ (rel.)	$k/k_{oxy} = 0.28 \pm 0.12$	p.r.	opt.	c.k.; k is a composite of $O^- + HO_2^-$, $O^- + H_2O_2$ and $OH^- + HO_2^-$ (69-0002).	67-0132
		alk. 3.9 $\times 10^8$ (rel.)	—	p.r.	opt.	p.b.k. at 260 nm; anal. of data is complex.	68-0298
		13-13.7 7 $\times 10^8$ (rel.)	$k/k_{oxy} = 0.2$	f.phot.	opt.	c.k.; obs. O_3^- at 430 nm.	68-7277
		11-13 (7.2 \pm 0.8) $\times 10^8$ (rel.)	$k/k(OH + CO_3^{2-}) = 1.98$	p.r.	opt.	c.k.; $\mu = 0.4$; cor. for OH and HCO_3^- .	69-0379
4.23	H_2O_2 $O^- + H_2O_2 \rightarrow H_2O + O_2^-$	alk. $\cong 5 \times 10^7$	—	p.r.	opt.	p.b.k. at 260 nm; more than one rate constant is involved in calcn.	68-0298
		11 $k + 1.4 k(OH + HO_2^-) = (8 \pm 0.8) \times 10^9$ (rel.)	—	p.r.	opt.	c.k. with CO_3^{2-} ; rel. to $k(OH + CO_3^{2-}) = (4 \pm 0.2) \times 10^8$ and $pK_a(OH) = 11.9$; $\mu = 0.4$.	69-0379
4.24	I^- $O^- + I^- (+ H_2O) \rightarrow 2OH^- + I$	13 (8.6 \pm 1) $\times 10^8$ (rel.)	$k/k(OH + CO_3^{2-}) = 2.3$	p.r.	opt.	c.k. with CO_3^{2-} ; cor. for HCO_3^- and OH present.	69-0379
		13 2.8 $\times 10^9$ (rel.)	$k/k_{2-PrOH} = 1.82$	γ -r.	chem.	c.k.; obs. $G(\text{acetone})$; $\mu = 0.1$; ratio increases with μ .	68-0602
		14 2.6 $\times 10^9$ (rel.)	$k/k_{2-PrOH} = 1.70$				
		6-7 2.6 $\times 10^9$ (rel.) 2.5 $\times 10^9$ (rel.)	$k/k_{EtOH} = 2.35$ $k/k_{MeOH} = 4.35$	p.r.	opt.	c.k.	71-0137
		alk. 2.0 $\times 10^9$	—	p.r.	opt.	p.b.k. at 0.58 M NaOH; $k = 1.9 \times 10^9$ at 1.1 M NaOH.	71-0137

TABLE 5. Reactions of O⁻ with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
4.24 cont.	alk.	2.2 x 10 ⁹ (rel.) 2.3 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EtOH} = 2.04 <i>k</i> / <i>k</i> _{MeOH} = 3.85	p.r. p.r.	opt. opt.	c.k. c.k.	71-0137 71-0137
4.25	IO ⁻ O ⁻ + IO ⁻ (+ H ₂ O) → 2OH ⁻ + IO	13.6 6 x 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{oxy} = 1.84	f.phot.	opt.	c.k.; effect of IO ⁻ on decay of O ₃ ⁻ .	70-0018
4.26	IO ₃ ⁻ O ⁻ + IO ₃ ⁻ (+ H ₂ O) → IO ₃ + 2OH ⁻ or → IO ₄ ²⁻	12.4 2.9 x 10 ⁸ (rel.) 12.6 (3 ± 0.5) x 10 ⁹	<i>k</i> / <i>k</i> _{oxy} = 0.08 —	f.phot. p.r.	opt. opt.	c.k.; effect of IO ₃ ⁻ on decay of O ₃ ⁻ . p.b.k. at 360 nm (IO ₄ ²⁻); cor. for OH reaction.	70-0018 72-0017
		12.05 1.6 x 10 ⁹	—	p.r.	opt.	p.b.k. at 360 nm (IO ₃).	73-0027
4.27	NO ₂ ⁻ O ⁻ + NO ₂ ⁻ (+ H ₂ O) → 2OH ⁻ + NO ₂	13 (2.4 ± 0.3) x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> (OH + CO ₃ ²⁻) = 0.67	p.r.	opt.	c.k. with CO ₃ ²⁻ ; cor. for OH and HCO ₃ ⁻ .	69-0379
		12 3.6 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{oxy} ≅ 10 ⁻¹	f.phot.	opt.	c.k.; obs. O ₃ ⁻ at 430 nm; based on <i>k</i> (OH + NO ₂ ⁻)/ <i>k</i> _{oxy} = 4.0 ± 0.4.	70-7264
4.28	Ni(dimethylglyoxime) ²⁺	>13 <i>For other ratios see: 4.32.</i> (2.5 ± 0.3) x 10 ⁹	—	p.r.	opt.	p.b.k. at 440 nm; incl. oxid. of free ligand.	72-0584
4.29	O ₂ (oxy) O ⁻ + O ₂ → O ₃ ⁻	13 alk. 4 x 10 ⁹	— —	p.r. —	opt. —	p.b.k. at 430 nm. unpubl. data cited.	66-0001 66-0424
		~11 3.6 x 10 ⁹ <i>For other ratios see: 4.2, 4.5, 4.22, 4.25-7, 4.33, 4.48, 4.65, 4.80, 4.89, 4.103.</i>	—	p.r.	opt.	p.b.k. at 430 nm.	69-0379
4.30	O ₃ ⁻ O ⁻ + O ₃ ⁻ → O ₄ ²⁻ or → O ₂ + O ₂ ²⁻	13- 13.7 (8 ± 2) x 10 ⁸ (rel.)	—	f.phot.	opt.	d.k. at 430 nm; complex anal. uses other rate constants.	68-7277
4.31	HPO ₄ ²⁻	>13 ~ 5 x 10 ⁸	—	p.r.	opt.	d.k.; <i>k</i> estd.	69-0002
4.32	RuO ₄ ²⁻ O ⁻ + RuO ₄ ²⁻ (+ H ₂ O) → RuO ₄ ⁻ + 2OH ⁻	12.35 2.7 x 10 ⁶ (rel.) >13 —	<i>k</i> / <i>k</i> _{MeOH} = 0.0046 <i>k</i> / <i>k</i> _{nitrite} = 7.6	p.r. p.r. γ-r.	opt. — chem.	c.k.; μ ≅ 0.75.	73-1049 68-0063
4.33	SO ₃ ²⁻ O ⁻ + SO ₃ ²⁻ → SO ₃ ⁻ + 2OH ⁻	14 3 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{oxy} = 0.083	r.	opt.	c.k.; obs. O ₃ ⁻ at 430 nm.	71-0461
4.34	acetate ion O ⁻ + CH ₃ COO ⁻ → OH ⁻ + CH ₂ COO ⁻	14 5 x 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{3HX} = 0.077	p.r.	opt.	c.k.	75-1003
4.35	acetonitrile	14 2.2 x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{3HX} = 0.34	p.r.	opt.	c.k.	75-1003
4.36	acetylenedicarboxylate ion	14 ≤ 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{3HX} = 0.063	p.r.	opt.	c.k.; cor. for OH reactions; <i>k</i> _{obs} = 4 x 10 ⁷ .	75-1003
4.37	aconitate ion	14 ~ 1.5 x 10 ⁸	—	p.r.	opt.	p.b.k. (allylic radicals from H abstr.).	75-1003
4.38	acrylamide	~12 (6.4 ± 0.8) x 10 ⁸ (rel.)	<i>k</i> / <i>k</i> (OH + CO ₃ ²⁻) = 1.75	p.r.	opt.	c.k. with CO ₃ ²⁻ ; μ = 0.4; assume p <i>K</i> _a (OH) = 11.9.	70-0052

TABLE 5. Reactions of O^- with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction pH	k	Ratio	Source	Method	Comment	Ref.
4.39	acrylate ion 14	1.5×10^8 (rel.)	$k/k_{3HX} = 0.307$	p.r.	opt.	c.k.; cor. for OH + acrylate ion.	75-1003
4.40	adipate ion 14	4.5×10^8 (rel.)	$k/k_{3HX} = 0.69$	p.r.	opt.	c.k.	75-1003
4.41	allyl alcohol 14.0	$(2.9 \pm 0.5) \times 10^9$	—	p.r.	opt.	p.b.k.; H abstr.	73-1070
	14	2.2×10^9 (rel.)	$k/k_{3HX} = 3.4$	p.r.	opt.	c.k. with ethanol	75-1003
4.42	allylbenzene 14	5×10^8	—	p.r.	opt.	$k_{3HX}/k_{EtOH} = 0.53$. p.b.k. (allylic radicals).	75-1003
4.43	allyl cyanide 14	1.05×10^9 (rel.)	$k/k_{3HX} = 1.61$	p.r.	opt.	c.k.	75-1003
4.44	amylamine —	1.6×10^{10} (rel.)	$k/k_{ferro} = 1.7$	p.r.	opt.	c.k.; calcd.	73-0016
		1.42×10^{10} (rel.)	$k/k_{CNS^-} = 1.3$			from obs. val-	
		9.0×10^9 (rel.)	$k/k_{NB} = 2.8$			ues at pH 8-13.1 assuming equal OH and O^- rates for ferro, CNS^- and NB.	
4.45	aniline 13.3	$(3.1 \pm 0.6) \times 10^9$	—	p.r.	opt.	p.b.k. at 300 and 400 nm.	72-0289
	$O^- + C_6H_5NH_2 \rightarrow OH^- + C_6H_5NH$ 14	1.6×10^9	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{obs} = 1.7 \times 10^9$.	75-1002
4.46	9-anthroate ion 14	4.8×10^8	—	p.r.	opt.	p.b.k.; cor. for OH.	73-0110
4.47	benzene 13	7.5×10^7 (rel.)	$k/k_{2-PrOH} = 0.05$	γ -r.	chem.	c.k.	68-0602
4.48	benzoate ion >13	$< 8 \times 10^6$ (rel.)	$k/k_{oxy} < 0.0024$	p.r.	opt.	c.k.; obs	69-0002
	$O^- + C_6H_5COO^- \rightarrow OHC_6H_5COO^- + OH^-$					O_3^- at 430 nm; $pK_a(OH) = 11.8 \pm 2$; assume $k(OH + C_6H_5COO^-) = 6 \times 10^9$.	
	14	4×10^7	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{obs} = 8.5 \times 10^7$.	72-0047
4.49	benzonitrile 14	7×10^7 (rel.)	$k/k_{3HX} = 0.154$	p.r.	opt.	c.k.; cor. for OH contribution	75-1003, 75-1002
	$O^- + C_6H_5CN (+ H_2O) \rightarrow OH^- + C_6H_5(OH)CN$					$k_{obs} = 1.0 \times 10^8$.	
4.50	4-biphenylcarboxylate ion 14	7.0×10^7	—	p.r.	opt.	p.b.k.; cor. for OH.	73-0110
4.51	2,2'-biphenyldicarboxylate ion (diphenate ion) 14	$\leq 2.9 \times 10^7$	—	p.r.	opt.	p.b.k.; cor. for OH.	73-0110
4.52	4,4'-biphenyldicarboxylate ion 14	$\leq 2.8 \times 10^7$	—	p.r.	opt.	p.b.k.; cor. for OH.	73-0110
4.53	2-butene-1,4-diol 14	2.3×10^9 (rel.)	$k/k_{3HX} = 3.54$	p.r.	opt.	c.k. with ethanol, $k_{3HX}/k_{EtOH} = 0.53$.	75-1003
	2-butenenitrile See crotononitrile (4.60).						
	3-butenenitrile See allyl cyanide (4.43).						
	2-butenate ion See crotonate ion (4.59).						
4.54	3-butenate ion 14	7.2×10^8 (rel.)	$k/k_{3HX} = 1.1$	p.r.	opt.	c.k.	75-1003
4.55	butylamine —	1.3×10^{10} (rel.)	$k/k_{ferro} = 1.4$	p.r.	opt.	c.k.; k calcd.	73-0016
		1.34×10^{10} (rel.)	$k/k_{CNS^-} = 1.2$			frm obs. values at pH = 8-13.1 assuming $k_{O^-} = k_{OH}$ for ferro, CNS^- and NB.	
		7.7×10^9 (rel.)	$k/k_{NB} = 2.4$				
4.56	butyrate ion 14	6.5×10^8 (rel.)	$k/k_{3HX} = 1.0$	p.r.	opt.	c.k.; H abstr.	75-1003
4.57	citrate ion 14	4.2×10^7 (rel.)	$k/k_{3HX} = 0.0645$	p.r.	opt.	c.k.	75-1003
4.58	<i>o</i> -cresol See <i>o</i> -methylphenoxide ion (4.82a)						

TABLE 5. Reactions of O^- with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction	pH	k	Ratio	Source	Method	Comment	Ref.
4.59	crotonate ion	14	9.0×10^8	—	p.r.	opt.	p.b.k. at 250 nm (allylic radical); also c.k. with ethanol and 3HX.	75-1003
4.60	crotononitrile	14	9.9×10^8 (rel.)	$k/k_{3HX} = 1.53$	p.r.	opt.	c.k.	75-1003
4.61	cyanoacetate ion	14	4.1×10^8 (rel.)	$k/k_{3HX} = 0.63$	p.r.	opt.	c.k.	75-1003
4.62	<i>p</i> -cyanophenoxide ion	14	6.2×10^8	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{obs} = 6.8 \times 10^8$.	75-1002
	$O^- + CNC_6H_4O^- (+ H_2O) \rightarrow 2OH^- + CNC_6H_4O$							
4.63	<i>p</i> -cyanotoluene	See <i>p</i> -tolunitrile (4.112).						
4.64	diphenylacetate ion	14	6×10^7	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{obs} = 9 \times 10^7$.	72-0047
	$O^- + (C_6H_5)_2CHCOO^- \rightarrow (C_6H_5)_2CCOO^- + OH^-$							
4.65	ethanol	>13	1.2×10^9 (rel.)	$k/k_{oxy} = 0.35$	p.r.	opt.	c.k.	65-0007
	$O^- + C_2H_5OH \rightarrow$	>13	1.2×10^9 (rel.)	$k/k_{oxy} = 0.337 \pm 0.028$	p.r.	opt.	c.k.	69-0002
	$OH^- + \cdot C_2H_4OH \rightleftharpoons \cdot C_2H_4O^- + H^+$	13	1.1×10^9 (rel.)	$k/k_{oxy} = 0.324$	f.phot.	opt.	c.k.; soln. contains NO_3^- .	69-7218
		13.92	$(11.3 \pm 1.7) \times 10^8$	—	p.r.	opt.	p.b.k. at 360 nm ($\cdot C_2H_4O^-$).	70-0080
		11	9.5×10^8 (rel.)	$k/k(OH + CO_3^{2-}) = 2.6$	p.r.	opt.	c.k.	70-0511
		14	4.5×10^8 (rel.)	$k/k(OH + EtOH) \cong 0.24$	X-r.	lum.	obs. effect of quenching chemiluminescence from fluorescein at pH 10.4 and 14.	73-6068
		14	1.22×10^9 (rel.)	$k/k_{3HX} = 1.89$	p.r.	opt.	c.k.; obs. reduction in allylic radical formn. from 3HX by addn. of EtOH.	75-1003
	For other ratios see: 4.1, 4.7, 4.12, 4.14, 4.18, 4.19, 4.24, 4.95, 4.114.							
4.66	ethylamine	—	5.8×10^9 (rel.) 8.9×10^9 (rel.)	$k/k_{NB} = 1.8$ $k/k_{CNS^-} = 8.1$	p.r.	opt.	c.k.; calcd. k from obs. values at pH 8-13.1 assuming equal OH and O^- rates for NB and CNS^- .	73-0016
4.67	ethyl ether	13	1.2×10^9 (rel.)	$k/k_{2-PrOH} = 0.79$	γ -r.	chem.	c.k.	68-0602
4.68	formate ion	—	—	$k/k_{oxalate} = 410$	γ -r.	chem.	c.k.	66-0621, 66-0151
		13	9×10^8 (rel.)	$k/k_{2-PrOH} = 0.60$	γ -r.	chem.	c.k.	68-0602
		14	1.0×10^9 (rel.)	$k/k_{2-PrOH} = 0.68$				
		11-13	1.3×10^9 (rel.)	$k/k(OH + CO_3^{2-}) = 3.5$	p.r.	opt.	c.k.; $\mu = 0.4$; assume $pK_a(OH) = 11.9$.	69-0379
	For other ratios see: 4.19.							
4.69	fumarate ion	14	$\leq 10^7$ (rel.)	$k/k_{3HX} = 0.063$	p.r.	opt.	c.k.; cor. for OH, $k_{obs} = 4 \times 10^7$.	75-1003
4.70	glutaconate ion	14	3.0×10^8	—	p.r.	opt.	p.b.k. at 250-270 nm (allylic radical); also c.k. with EtOH.	75-1003

TABLE 5. Reactions of O⁻ with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
4.71	glycine, negative ion 14	5.6 x 10 ⁸ (rel.)	$k/k_{3HX} = 0.865$	p.r.	opt.	c.k.	75-1003
4.72	2,4-hexadien-1-ol 14.0	(4.3 ± 0.8) x 10 ⁹	—	p.r.	opt.	p.b.k.; H abstr.	73-1070
4.72a	hexamethylbenzene ~13 O ⁻ + C ₆ (CH ₃) ₆ → OH ⁻ + C ₆ (CH ₃) ₅ (CH ₂)	~ 2.5 x 10 ⁹	—	p.r.	opt.	p.b.k.	75-1009
4.73	hexanoate ion 14	1.44 x 10 ⁹ (rel.)	$k/k_{3HX} = 2.2$	p.r.	opt.	c.k.	75-1003
4.74	2-hexene-1,6-dioate ion 14	6.9 x 10 ⁸	—	p.r.	opt.	p.b.k. at 250-270 nm (allylic radical); also c.k. with EtOH.	75-1003
4.75	3-hexene-1,6-dioate ion (3HX) 14 O ⁻ + ⁻ O ₂ CCH ₂ CH=CHCH ₂ CO ₂ ⁻ → OH ⁻ + ⁻ O ₂ CCH ₂ CHCHCHCO ₂ ⁻	(6.5 ± 0.3) x 10 ⁸	—	p.r.	opt.	p.b.k. at 266 nm (allylic radicals); cor. for background reactions; $k_{obs} = 6.3 \times 10^8$.	75-1003
For other ratios see: 4.34, 4.35, 4.36, 4.39, 4.40, 4.41, 4.43, 4.49, 4.53, 4.54, 4.56, 4.57, 4.60, 4.61, 4.65, 4.69, 4.71, 4.73, 4.76, 4.77, 4.78, 4.79, 4.83, 4.84, 4.94, 4.95, 4.96, 4.97, 4.102.							
<i>o</i> -hydroxybenzaldehyde See salicylaldehyde (4.100).							
<i>o</i> -hydroxybenzoate ion See salicylate ion (4.101).							
4.76	maleate ion 14	~ 3 x 10 ⁷ (rel.)	$k/k_{3HX} = 0.123$	p.r.	opt.	c.k.; cor. for OH, $k_{obs} = 8 \times 10^7$.	75-1003
4.77	malonate ion 14	2.1 x 10 ⁷ (rel.)	$k/k_{3HX} = 0.0323$	p.r.	opt.	c.k.	75-1003
4.78	methacrylonitrile 14	1.76 x 10 ⁹ (rel.)	$k/k_{3HX} = 2.7$	p.r.	opt.	c.k.	75-1003
4.79	methacrylate ion 14	4.8 x 10 ⁸ (rel.)	$k/k_{3HX} = 0.74$	p.r.	opt.	c.k.	75-1003
4.80	methanol >13 (I) O ⁻ + CH ₃ OH → OH ⁻ + •CH ₂ OH 13.92 ⇌ •CH ₂ O ⁻ + H ⁺ (II) O ⁻ + CH ₃ OH → >13 OH ⁻ + CH ₃ O•	7 x 10 ⁸ (rel.) (5.8 ± 0.8) x 10 ⁸	$k/k_{oxy} = 0.209 \pm 0.014$ — $k_{II}/k_I = 0.075$	p.r. p.r.	opt. opt.	c.k. p.b.k. at 360 nm (•CH ₂ O ⁻). detd. % of α-alcohol and alkoxy radicals by reactions with TNM and I ⁻ , resp.	69-0002 70-0080 73-0126
For other ratios see: 4.1, 4.7, 4.12, 4.18, 4.24, 4.31, 4.95.							
4.81	<i>o</i> -methoxyphenoxide ion 13	7 x 10 ⁸ (rel.)	$k/k_{2-PrOH} = 0.46 \pm 0.09$	γ-r.	chem.	c.k.	72-0837
4.82	methylamine 13.1	7.5 x 10 ⁹ (rel.)	$k/k_{CNS^-} = 0.71$	p.r.	opt.	c.k. assuming $k_{CNS^-} = 1.1 \times 10^{10}$.	71-0595
4.82a	<i>o</i> -methylphenoxide ion 13	5 x 10 ⁸ (rel.)	$k/k_{2-PrOH} = 0.33 \pm 0.03$	γ-r.	chem.	c.k.	72-0837
4.82b	<i>p</i> -methylphenoxide ion 14 (I + II) (I) O ⁻ + CH ₃ C ₆ H ₄ O ⁻ → OH ⁻ + CH ₂ C ₆ H ₄ O ⁻ (II) O ⁻ + CH ₃ C ₆ H ₄ O ⁻ (+ H ₂ O) → 2OH ⁻ + CH ₃ C ₆ H ₅ O	1.6 x 10 ⁹ (I + II) 1.0 x 10 ⁹ (I)	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{obs} = 1.65 \times 10^9$.	75-1002
4.83	2-methyl-2-propanol (<i>tert</i> -butanol) 14	3.3 x 10 ⁸ (rel.)	$k/k_{3HX} = 0.51$	p.r.	opt.	c.k.	75-1003
4.84	muconate ion 14	~ 2 x 10 ⁹ (rel.)	$k/k_{3HX} \cong 3.1$	p.r.	opt.	c.k.	75-1003
4.85	1-naphthoate ion 14	1.2 x 10 ⁸	—	p.r.	opt.	p.b.k.; cor. for OH.	73-0110
4.86	2-naphthoate ion 14	1.3 x 10 ⁸	—	p.r.	opt.	p.b.k.; cor. for OH.	73-0110

TABLE 5. Reactions of O⁻ with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
4.87	nitrobenzene O ⁻ + C ₆ H ₅ NO ₂ (+ H ₂ O) → C ₆ H ₅ (OH)NO ₂ + OH ⁻	14	< 7 × 10 ⁷	—	p.r.	opt.	p.b.k.; cor. for OH; <i>k</i> _{obs} ≤ 1 × 10 ⁸ .	75-1002
4.88	<i>p</i> -nitrotoluene O ⁻ + CH ₃ C ₆ H ₄ NO ₂ → OH ⁻ + CH ₂ C ₆ H ₄ NO ₂	14	7.6 × 10 ⁸	—	p.r.	opt.	p.b.k.; cor. for OH; <i>k</i> _{obs} = 8 × 10 ⁸ ; <i>k</i> for abstr. from methyl group = 7 × 10 ⁸ .	75-1002
4.89	oxalate ion O ⁻ + C ₂ O ₄ ²⁻ (+ H ₂ O) → CO ₂ + 2OH ⁻ + CO ₂ .	13	2.6 × 10 ⁷ (rel.)	<i>k</i> / <i>k</i> _{oxy} = 7.2 × 10 ⁻³	γ-r.	chem.	c.k.; ratio in D ₂ O = 9.4 × 10 ⁻³ .	66-0068, 66-0621 68-0015
<i>For other ratios see: 4.68.</i>								
4.89a	1,4-pentadien-3-ol	14.0	(2.4 ± 0.5) × 10 ⁹	—	p.r.	opt.	p.b.k.	
4.89b	pentamethylbenzene O ⁻ + C ₆ H(CH ₃) ₅ → OH ⁻ + C ₆ H(CH ₃) ₄ (CH ₂)	~13	2.6 × 10 ⁹	—	p.r.	opt.	p.b.k.	75-1009
4.90	phenoxide ion O ⁻ + C ₆ H ₅ O ⁻ (+ H ₂ O) → 2OH ⁻ + C ₆ H ₅ O	13 14	1.1 × 10 ⁹ (rel.) 6.5 × 10 ⁸	<i>k</i> / <i>k</i> _{2-PrOH} = 0.75 —	γ-r. p.r.	chem. opt.	c.k. p.b.k. at 402 nm (phenoxyl radical); cor. for OH addn.; <i>k</i> _{obs} = 7.1 × 10 ⁸ .	68-0602 75-1001, 75-1002
4.91	phenoxybenzoate ion	14	1.6 × 10 ⁸	—	p.r.	opt.	p.b.k. at 337 nm (hydroxycyclohexadienyl radical); cor. for OH; <i>k</i> _{obs} = 2.1 × 10 ⁸ .	75-1001, 75-1002
4.92	phenylacetate ion O ⁻ + C ₆ H ₅ CH ₂ COO ⁻ (+ H ₂ O) → OH ⁻ + HOC ₆ H ₅ CH ₂ COO ⁻	14	(2 ± 0.6) × 10 ⁸	—	p.r.	opt.	p.b.k. at 290 nm; <i>k</i> _{obs} = 2.2 × 10 ⁸ ; assume OH contribution is 6.2 × 10 ⁷ .	72-0047
4.93	phthalate ion, dianion	14	≤ 1.8 × 10 ⁷	—	p.r.	opt.	p.b.k.; cor. for OH.	73-0110
4.94	1-propanol	14	1.51 × 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{3HX} = 2.32	p.r.	opt.	c.k.	75-1003
4.95	2-propanol	13	1.7 × 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EiOH} = 1.56	γ-r.	chem.	c.k.	68-0602
	(I) O ⁻ +	13	1.5 × 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{MeOH} = 2.56	γ-r.	chem.	c.k.	68-0602
	(CH ₃) ₂ CHOH →	14	1.6 × 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{EiOH} = 1.43	γ-r.	chem.	c.k.	68-0602
	OH ⁻ + (CH ₃) ₂ COH	14	1.2 × 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{MeOH} = 2.13	γ-r.	chem.	c.k.	68-0602
	(II) O ⁻ +	13.5	—	<i>k</i> _I / <i>k</i> _{II} = 5.6 ± 0.3	γ-r.	chem.	c.k.; <i>k</i> _H / <i>k</i> _D (I) = 1.35 ± 0.10 and <i>k</i> _H / <i>k</i> _D (II) = 3.26 ± 0.23.	72-0167
	(CH ₃) ₂ CHOH → OH ⁻ + CH ₂ (CH ₃)CHOH	14	1.22 × 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{3HX} = 1.88	p.r.	opt.	c.k.	75-1003
<i>For other ratios see: 4.7, 4.23, 4.34, 4.40-2, 4.48.</i>								
4.96	propionate ion	14	3.3 × 10 ⁸ (rel.)	<i>k</i> / <i>k</i> _{3HX} = 0.51	p.r.	opt.	c.k.	75-1003
4.97	propionitrile	14	1.0 × 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{3HX} = 1.54	p.r.	opt.	c.k.	75-1003
4.98	propylamine	—	1.02 × 10 ¹⁰ (rel.) 6.4 × 10 ⁹ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 0.93 <i>k</i> / <i>k</i> _{NB} = 2.0	p.r.	opt.	c.k., calcd. from obs. values at pH 8-13.1 assuming equal O ⁻ and OH rates for CNS ⁻ and NB.	73-0016
4.99	pyridine O ⁻ + C ₅ H ₅ N (+ H ₂ O) → OH ⁻ + C ₅ H ₅ N(OH)	14	< 7 × 10 ⁷	—	p.r.	opt.	p.b.k.; cor. for OH; <i>k</i> _{obs} ≤ 1 × 10 ⁸ .	75-1002
4.99a	pyrrolidine	13.2	2.1 × 10 ¹⁰ (rel.)	<i>k</i> / <i>k</i> _{CNS⁻} = 1.9	p.r.	opt.	c.k. assuming <i>k</i> _{CNS⁻} = 1.1 × 10 ¹⁰ .	75-1016

TABLE 5. Reactions of O⁻ with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
4.100	salicylaldehyde, 13 anion	4.0 x 10 ⁸ (rel.)	$k/k_{2-\text{PrOH}} = 0.27 \pm 0.06$	γ-r.	chem.	c.k.	72-0837
4.101	salicylate ion 13 O ⁻ + ⁻ O ₂ CC ₆ H ₄ O ⁻ (+ H ₂ O) → 2OH ⁻ + 14 ⁻ O ₂ CC ₆ H ₄ O	4.8 x 10 ⁸ (rel.) 4.5 x 10 ⁸	$k/k_{2-\text{PrOH}} = 0.32 \pm 0.05$ —	γ-r. p.r.	chem. opt.	c.k. p.b.k.; cor. for OH; $k_{\text{obs}} = 5.1 \times 10^8$.	72-0837 75-1002
4.102	succinate ion 14	1.35 x 10 ⁸ (rel.)	$k/k_{3\text{HX}} = 0.207$	p.r.	opt.	c.k.	75-1003
4.103	1,2,3,4-tetra- ~13 methylbenzene (prehnitine) O ⁻ + C ₆ H ₂ (CH ₃) ₄ → OH ⁻ + C ₆ H ₂ (CH ₃) ₃ CH ₂	2.4 x 10 ⁹	—	p.r.	opt.	p.b.k.	75-1009
4.104	1,2,3,5-tetra- ~13 methylbenzene (isodurene)	2.6 x 10 ⁹	—	p.r.	opt.	p.b.k.	75-1009
4.105	1,2,4,5-tetra- ~13 methylbenzene (durene)	2.3 x 10 ⁹	—	p.r.	opt.	p.b.k.	75-1009
4.106	2,2,6,6-tetra- 13 methyl-4-piperidone N-oxyl (TAN)	1.6 x 10 ⁹ (rel.)	$k/k_{\text{oxy}} = 0.46$	p.r.	opt.	c.k.	71-0618
4.107	thymine >13	4 x 10 ⁸	—	p.r.	opt.	p.b.k.	72-0047
4.108	<i>o</i> -toluate ion 14 O ⁻ + CH ₃ C ₆ H ₄ CO ₂ ⁻ → OH ⁻ + CH ₂ C ₆ H ₄ CO ₂ ⁻	3.4 x 10 ⁸	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{\text{obs}} = 3.8 \times 10^8$; $k_{\text{abstr}} = 3 \times 10^8$.	75-1002
4.109	<i>m</i> -toluate ion 14	7.5 x 10 ⁸	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{\text{obs}} = 7.9 \times 10^8$; $k_{\text{abstr}} = 7 \times 10^8$.	75-1002
4.110	<i>p</i> -toluate ion 14	5 x 10 ⁸	—	p.r.	opt.	p.b.k. at 280 nm; contribution of OH reaction < 10%.	72-0047
	14	8.2 x 10 ⁸	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{\text{obs}} = 8.6 \times 10^8$; $k_{\text{abstr}} = 8 \times 10^8$.	75-1002
4.111	toluene ~13 O ⁻ + C ₆ H ₅ CH ₃ → C ₆ H ₅ CH ₂ + OH ⁻	(2.1 ± 0.3) x 10 ⁹	—	p.r.	opt.	p.b.k.	73-0089, 75-1009
4.112	<i>p</i> -tolunitrile 14 O ⁻ + CH ₃ C ₆ H ₄ CN → OH ⁻ + CH ₂ C ₆ H ₄ CN	8.8 x 10 ⁸	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{\text{obs}} = 9.2 \times 10^8$; $k_{\text{abstr}} = 8 \times 10^8$.	75-1002
4.113	<i>p</i> -toluidine 14 (I) O ⁻ + CH ₃ C ₆ H ₄ NH ₂ → OH ⁻ + CH ₂ C ₆ H ₄ NH ₂ (II) O ⁻ + CH ₃ C ₆ H ₄ NH ₂ → OH ⁻ + CH ₃ C ₆ H ₄ NH	3.0 x 10 ⁹ (I + II) 1.5 x 10 ⁹ (I)	—	p.r.	opt.	p.b.k.; cor. for OH; $k_{\text{obs}} = 3.1 \times 10^9$.	75-1002
4.114	triethylamine 12	2.4 x 10 ⁹ (rel.)	$k/k_{\text{EtOH}} = 2$	γ-r.	chem.	c.k.; may be OH reaction.	71-0590
4.115	1,2,3-trimethyl- ~13 benzene O ⁻ + C ₆ H ₃ (CH ₃) ₃ → OH ⁻ + C ₆ H ₃ (CH ₃) ₂ (CH ₂)	2.1 x 10 ⁹	—	p.r.	opt.	p.b.k.	75-1009

TABLE 5. Reactions of O⁻ with water, transients from water, inorganic solutes, and organic solutes - Continued

No.	Solute and reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
4.116	1,2,4-trimethyl benzene	~13	2.1×10^9	—	p.r.	opt.	p.b.k.	75-1009
4.117	1,3,5-trimethyl- benzene	~13	2.4×10^9	—	p.r.	opt.	p.b.k.	75-1009
4.118	uracil	13.5	4.1×10^9 (rel.)	$k/k(\text{OH} + \text{CNS}^-)$ = 0.374	p.r.	opt.	c.k.; authors doubtful about value.	68-0316, 69-0571
		12	1.8×10^9	—	p.r.	opt.	d.k.; double bond bleaching; value from graph.	69-0571
4.119	<i>o</i> -xylene $\text{O}^- + \text{C}_6\text{H}_4(\text{CH}_3)_2 \rightarrow$ $\text{OH}^- +$ $\text{C}_6\text{H}_4(\text{CH}_3)(\text{CH}_2)$	~13	1.8×10^9	—	p.r.	opt.	p.b.k.	75-1009
4.120	<i>m</i> -xylene	~13	2.2×10^9	—	p.r.	opt.	p.b.k.	75-1009
4.121	<i>p</i> -xylene	~13	1.8×10^9	—	p.r.	opt.	p.b.k.	75-1009

TABLE 6. Reactions of HO₂ (H₂O₂⁺ and O₂⁻) with transients from water, inorganic solutes, and organic solutes

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
<i>H₂O₂⁺ Reactions</i>							
5.1	OH H ₂ O ₂ ⁺ + OH → H ₃ O ⁺ + O ₂	—	—	—	—	See 3.6, Table 2.	—
<i>HO₂ Reactions</i>							
5.2	H HO ₂ + H → H ₂ O ₂	—	—	—	—	See 2.4, NSRDS-NBS 51.	75-0001
5.3	OH HO ₂ + OH → H ₂ O + O ₂ or → H ₂ O ₃	—	—	—	—	See 3.5, Table 2.	—
5.4	HO ₂ HO ₂ + HO ₂ → H ₂ O ₂ + O ₂	nat. (3.4 ± 2.5) × 10 ⁶ nat. 3.1 × 10 ⁶	— —	phot. γ-r.	chem. chem.	<i>k</i> at 25°C. <i>k</i> at 0°C; no pH effects discussed; rates are probab- ly for O ₂ ⁻ + O ₂ ⁻ .	53-0014 53-0014
		2.7 2.5 × 10 ⁶	—	f.phot.	opt.	d.k.; ε(254 nm) = 350 mol ⁻¹ cm ² .	62-0050
		0.5- 1.55 (2.4 ± 0.4) × 10 ⁶	—	therm.	esr	d.k.; flow syst- em; Ce ⁴⁺ + H ₂ O ₂ soln.; <i>E</i> _a = 5.9 ± 0.4 kcal/mol(25kJ/ mol).	62-0054
		2 2.3 × 10 ⁶ , 2.2 × 10 ⁶ (rel.)	—	e-r.	chem.	c.k.; obs. reaction of HO ₂ with tetranitro- methane.	63-0075
		1.7- 3.0 2.7 × 10 ⁶	—	p.r.	opt.	d.k.; ε(253.7 nm) = 830 ± 125 dm ³ mol ⁻¹ cm ⁻¹ .	64-0064
		1 ~ 2 × 10 ⁶	—	γ-r.	chem.	c.k.; rotating sector method; H ₂ O ₂ soln.	65-0046
		2 (2.5 ± 0.5) × 10 ⁶ 2.8- 2.9 2.5 × 10 ⁶	— —	p.r. e-r.	opt. chem.	d.k. c.k.; also from Ce(IV) + H ₂ O ₂ ; <i>k</i> decreases below pH 2 and increases above pH 3.	66-0001 66-0614
		— 2.65 × 10 ⁶	—	therm.	esr	d.k.; from Ce(IV) + H ₂ O ₂ ; <i>E</i> _a = 4.7 kcal/mol (19.6 kJ/ mol).	68-9083
		0.3-2 0.7 × 10 ⁶	—	p.r.	opt.	d.k.; ε(254 nm) = 540 dm ³ mol ⁻¹ cm ⁻¹ ; more than one rate constant is involv- ed in calcn.; <i>k</i> = 2.8 × 10 ⁹ exp (-4900/RT).	68-0382
		2-5 6.7 × 10 ⁵	—	p.r.	opt.	d.k.; ε(240 nm) = 1150 dm ³ mol ⁻¹ cm ⁻¹ ; p <i>K</i> _a (HO ₂) = 4.8.	69-0418
		0-7.7 7.6 × 10 ⁵	—	p.r.	opt.	d.k.; p <i>K</i> _a (HO ₂) = 4.88.	70-0304
		0 — 5.5 —	<i>k</i> _H / <i>k</i> _D = 7 <i>k</i> _H / <i>k</i> _D = 3	—	—	<i>E</i> _a in D ₂ O = 7.1 ± 0.4 kcal/mol (29.7 kJ/mol); unpubl. data.	70-0642

TABLE 6. Reactions of HO₂ (H₂O₂⁺ and O₂⁻) with transients from water, inorganic solutes, and organic solutes — Continued

No.	Solute and reaction	pH	k	Ratio	Source	Method	Comment	Ref.
5.5	O_2^- $\text{HO}_2 + \text{O}_2^- \rightarrow \text{HO}_2^- + \text{O}_2$	0	$(9 \pm 1) \times 10^5$	—	Ce(IV)	opt.	$\epsilon(230 \text{ nm}) = 1100 \text{ dm}^3 \text{ mol}^{-1} \text{ cm}^{-1}$.	70-0920,
		1.1	2.8×10^6	—	+ H_2O_2			69-9139
		0-2	$(1.35 \pm 0.3) \times 10^6$	—	f.phot.	opt.	d.k.; $\epsilon(240 \text{ nm}) = 10^3 \text{ dm}^3 \text{ mol}^{-1} \text{ cm}^{-1}$.	70-0920
		2.6-7	$10^6 - 10^7$	—	$e-r.$	chem.	obs. HO_2 reaction with tetra-nitromethane in formate soln.	72-0308
		7	$\sim 2 \times 10^8$	—	f.phot.	opt.	c.k.	62-0050
		7	1×10^7	—	f.phot.	opt.	d.k.; data of 53-0014.	62-0050
		—	$< 7 \times 10^7$	—	$e-r.$	chem.	flow technique; pH effects, c.k.	63-0075
		7	5.3×10^7	—	p.r.	opt.	d.k.	66-0001
		0-5	3×10^7	—	p.r.	opt.	d.k.; more than one rate constant is involved in calcn.	68-0382
		2-9.7	7.9×10^7	—	p.r.	opt.	d.k.; $\text{p}K_a(\text{HO}_2) = 4.8$.	69-0418
5.6	Br_2 $\text{HO}_2 + \text{Br}_2 \rightarrow \text{H}^+ + \text{Br}^- + \text{Br} + \text{O}_2$	0-7.7	8.5×10^7	—	p.r.	opt.	d.k.; $\text{p}K_a(\text{HO}_2) = 4.88$.	70-0304
		~ 1	$(1.5^{+1.5}_{-0.8}) \times 10^8$ (rel.)	—	p.r.	opt.	c.k.; indirect estimation; more than one rate constant is involved.	65-0382
		2	$< 4 \times 10^6$	—	p.r.	opt.	p.b.k. and d.k.; mechanistic anal. of data.	65-0383
		2-7	$(1.1^{+0.6}_{-0.4}) \times 10^8$ (rel.)	—	$e-r$	chem.	c.k. in formate- Br_2 soln.; rel. to $k(\text{O}_2^- + \text{C}(\text{NO}_2)_4) = 2 \times 10^9$.	72-0308
5.7	Br_2^- $\text{HO}_2 + \text{Br}_2^- \rightarrow \text{Br}_2 + \text{HO}_2^-$	2		$k/k_X = 4 \times 10^{-4}$	$\gamma-r.$	chem.	c.k.; $k_X = k(\text{HO}_2 + \text{Br}_2) \times k(\text{Br}_2^- + \text{Br}_2^- \rightarrow \text{Br}_3^- + \text{Br}^-)^{1/2}$.	65-0055
		2	$(3.8 \pm 0.9) \times 10^9$ (rel.)	—	p.r.	opt.	d.k.; $k/\epsilon(\text{Br}_2^-) = (4.6 \pm 0.4) \times 10^5 \text{ cm/s}$; more than one rate constant is involved in calcn.	65-0382
		2	$(1.6 \pm 0.5) \times 10^9$ (rel.)	—	p.r.	opt.	c.k.; obs. decay of $\text{Br}_2^- + \text{Br}_2^- \rightarrow \text{Br}_3^- + \text{Br}^-$ at 360 nm; data fitting.	65-0383
5.8	Br_3^- $\text{HO}_2 + \text{Br}_3^- \rightarrow \text{H}^+ + \text{Br}_2^- + \text{Br}^- + \text{O}_2^-$	2	$(1 \pm 0.5) \times 10^8$ (rel.)	—	p.r.	opt.	c.k.; mechanistic anal.	65-0383
		2-7	$< 10^7$ (rel.)	—	$e-r.$	chem.	c.k. in formate- Br_2 soln.; rel. to $k(\text{O}_2^- + \text{C}(\text{NO}_2)_4) = 2 \times 10^9$.	72-0308

TABLE 6. Reactions of HO₂ (H₂O₂[•] and O₂^{•-}) with transients from water, inorganic solutes, and organic solutes—Continued

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
5.9	CNS HO ₂ + CNS → CNS ⁻ 1 + H ⁺ + O ₂	1.6 × 10 ⁹ (rel.)	—	p.r.	opt.	c.k.; pH effect on decay CNS + CNS → (CNS) ₂ .	65-0386
5.10	Ce ³⁺ HO ₂ + Ce ³⁺ (+ H ⁺) → Ce ⁴⁺ + H ₂ O ₂	0.4 (2.1 ± 0.2) × 10 ⁵	—	p.r.	opt.	p.b.k. at 320 nm, Ce(IV).	74-1107
5.11	Ce ⁴⁺ HO ₂ + Ce ⁴⁺ → Ce ³⁺ + H ⁺ + O ₂	0.4 0.4	<i>k</i> / <i>k</i> _{Ce3+} = 7.7 <i>k</i> / <i>k</i> _{Ce3+} = 13 ± 2	therm. therm.	chem. chem.	0°C. d.k.; flow technique; Ce(IV) + H ₂ O ₂ .	57-9009 63-9017
5.12	Cu ⁺ (I) HO ₂ + Cu ⁺ (+H ₂ O) → Cu ²⁺ + H ₂ O ₂ + OH ⁻ (II) H ₂ O ₂ [•] + Cu ⁺ → Cu ²⁺ + H ₂ O ₂ (III) H ₂ O ₂ + Cu ⁺ → Cu ²⁺ + OH + OH ⁻	2.3 6 × 10 ⁸ (I) 0.8-2 2.3	— <i>k</i> _I / <i>k</i> _{II} = 2.4 — <i>k</i> _I / <i>k</i> _{III} = 0.015 × <i>k</i> (HO ₂ + Cu ²⁺) Ms	phot. p.r. phot.	opt. opt. opt.	rotating sector; μ = 0.1; soln. contains Cu ²⁺ and 4.5 M H ₂ O ₂ ; see also 73-7514. d.k. at 245 nm; Cu ²⁺ soln. rotating sector; assume <i>k</i> _{III} = 4.7 × 10 ³ ; <i>k</i> (HO ₂ + Cu ²⁺) = 3.4 × 10 ⁷ .	69-7082, 69-7083 73-0112 73-7514
5.13	Cu ²⁺ (I) HO ₂ + Cu ²⁺ → Cu ⁺ + H ⁺ + O ₂ (II) O ₂ ⁻ + Cu ²⁺ → Cu ⁺ + O ₂	1.35- 2.65 ~2 2.3 0.4	<i>k</i> _I / <i>k</i> _{Fe2+} = 3.5 - 103 <i>k</i> _I / <i>k</i> _{Fe2+} = 55 — <i>k</i> _I / <i>k</i> _{Fe2+} = 0.4	Fenton γ-r. f.phot. γ-r.	chem. chem. opt. chem.	c.k.; pH dependent; 0°C. c.k.; 0.01 M H ⁺ d.k. at 254 nm. c.k.	51-9004 55-0039 62-0050 66-0334, 68-0355
		2.3 0.8-2 >2.5	— — <i>k</i> _I / <i>k</i> _{II} = 0.024	phot. p.r. phot.	opt. opt. opt.	rotating sector. d.k. at 245 nm. rotating sector.	69-7083 73-0112 73-7514
5.14	Fe ²⁺ HO ₂ + Fe ²⁺ → Fe ³⁺ •HO ₂ ⁻ or Fe ²⁺ + HO ₂ ⁻ (+ H ⁺) → Fe(OH) ₂ ⁺ + H ₂ O ₂	1.35- 2.65 2.7 ~2 0.3 0-2.1 0.38- 2 1	<i>k</i> / <i>k</i> _{Fe3+} = 1.0-7 <i>k</i> / <i>k</i> _{Fe3+} = 3.3 <i>k</i> / <i>k</i> _{Fe3+} = 160- 190 — <i>k</i> / <i>k</i> _{Fe3+} = 30[H ⁺] <i>M</i> —	Fenton γ-r. γ-r. γ-r. p.r. p.r. γ-r. p.r.	chem. chem. chem. chem. opt. opt. chem. opt.	c.k.; pH dependent; at 0°C ratio = 1.1-8. c.k.; at pH 2.0 ratio = 9, at pH ~ 0.5 ratio > 100. c.k.; at pH 0.8 ratio is 300. p.b.k. at 305 nm; several reactions are involved in analysis. p.b.k.; obs. (Fe ³⁺ •HO ₂ ⁻); supercedes value in 64-0090. c.k. p.b.k. at 250 nm; μ = 1.0; <i>k</i> = 9.1 × 10 ⁵ at 20°C; <i>E</i> _s = 10.0 ± 1.0 kcal/mol (42 kJ/mol).	51-9004 57-0010 58-0004 60-0102 64-0090 69-0434 69-0642 73-0038

For other ratios see: 5.13, 5.28, 5.44.

TABLE 6. Reactions of HO₂ (H₂O₂⁺ and O₂⁻) with transients from water, inorganic solutes, and organic solutes—Continued

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
5.15	Fe(CN) ₆ ⁴⁻ HO ₂ + Fe(CN) ₆ ⁴⁻ → HO ₂ ⁻ + Fe(CN) ₆ ³⁻	~2 1.64 × 10 ⁵ 0.46–(3.0 ± 1.5) × 10 ⁴ 4.37	—	p.r.	opt.	p.b.k. at 420 nm (ferricyanide).	65-0007
	HO ₂ + HFe(CN) ₆ ³⁻ HO ₂ + H ₂ Fe(CN) ₆ ²⁻ HO ₂ + KFe(CN) ₆ ³⁻	(1.4 ± 0.1) × 10 ⁵ (1.0 ± 0.3) × 10 ⁴ (3.0 ± 1.5) × 10 ⁴	—	p.r.	opt.	p.b.k. at 420–460 nm; pH effects obs.	72-0431
5.16	Fe ³⁺ HO ₂ + Fe ³⁺ → Fe ²⁺ + H ⁺ + O ₂	1	<i>k</i> / <i>k</i> _{H⁺} = 1.20	γ-r.	chem.	c.k.	69-0642
		<i>For other ratios see:</i>	5.14, 5.43.				
5.17	H ⁺ HO ₂ + H ⁺ ⇌ H ₂ O ₂ ⁺	<i>For ratio see:</i> 5.16.					
5.18	H ₂ O ₂ HO ₂ + H ₂ O ₂ → H ₂ O + O ₂ + OH	— 530	—	γ-r.	chem.	no pH effects considered; termination rate <i>k</i> (2HO ₂ + H ₂ O ₂) = 2.7 × 10 ¹⁰ dm ⁶ mol ⁻² s ⁻¹ .	52-0018
		nat. 3.7 ± 1.6	—	phot.	chem.	propagation step in chain reaction; <i>k</i> at 25°C; no pH effects considered; see 5.46.	53-0014
		nat. 1.1	—	γ-r.	chem.	<i>k</i> at 0°C. no pH effects considered; probably for O ₂ ⁻ + H ₂ O ₂ .	53-0014
		1 1 × 10 ⁻²	—	γ-r.	chem.	mechanistic fit; <i>k</i> at 10°C; concn. H ₂ O ₂ ~ 1–35 <i>M</i> .	65-0046
		0.8–1.5 0.20 ± 0.01 (rel.)	<i>k</i> / <i>k</i> (HO ₂ + HO ₂) = 1.8 × 10 ⁻⁷	γ-r.	chem.	c.k.; obs. <i>G</i> (-H ₂ O ₂); includes <i>k</i> (H ₂ O ₂ + H ₂ O ₂ → H ₃ O ⁺ + O ₂ + OH); <i>k</i> (HO ₂ + HO ₂) = 1.1 × 10 ⁶ .	69-0643
5.19	MnO ₄ ⁻ HO ₂ + MnO ₄ ⁻ → H ⁺ + O ₂ + MnO ₄ ²⁻	2 8 × 10 ⁶	—	p.r.	opt.	d.k.	65-0385
5.20	OsO ₄ HO ₂ + OsO ₄ → OsO ₄ ⁻ + H ⁺ + O ₂	<1 5.7 × 10 ⁵ (rel.)	<i>k</i> / <i>k</i> (HO ₂ + HO ₂) = 0.24	γ-r.	chem.	c.k.; obs. <i>G</i> (H ₂ O ₂); <i>k</i> (HO ₂ + HO ₂) = 2.35 × 10 ⁶ ; dose rate 9.7 × 10 ¹⁸ eV cm ⁻³ h ⁻¹ .	64-0050
5.21	Te(IV) HO ₂ + Te(IV) → Te(VI) + OH	0.4 ~ 7.5 × 10 ³ (rel.)	<i>k</i> / <i>k</i> (HO ₂ + HO ₂) ≅ 3 × 10 ⁻³	γ-r.	chem.	c.k.; preliminary value; assume <i>k</i> (HO ₂ + HO ₂) = 2.5 × 10 ⁶ .	67-0553
		0.4 > 50 (rel.)	<i>k</i> / <i>k</i> (HO ₂ + HO ₂) ≧ 2 × 10 ⁻⁵	γ-r.	chem.	c.k.; more than one rate involved in calcn.; <i>k</i> (HO ₂ + HO ₂) = 2.5 × 10 ⁶ .	68-0356

TABLE 6. Reactions of HO₂ (H₂O₂⁺ and O₂⁻) with transients from water, inorganic solutes, and organic solutes—Continued

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
5.22	Th(IV) (I) HO ₂ + Th(IV) \rightleftharpoons Th(IV)-HO ₂ (II) HO ₂ + Th(IV)-HO ₂ \rightarrow Th(IV) + H ₂ O ₂ + O ₂ (III) 2Th(IV)-HO ₂ \rightarrow 2Th(IV) + H ₂ O ₂ + O ₂	~1 $\geq 5 \times 10^6$ (I) (8.0 \pm 2.0) $\times 10^5$ (II) (5 \pm 2) $\times 10^2$ (III) (1.8 \pm 0.2) $\times 10^6$ (I)	— — —	therm. p.r.	esr opt.	$K_1 = (1.7 \pm 0.4) \times 10^5 M^{-1}$; d.k. as well as p.b.k.; Ce ⁴⁺ + H ₂ O ₂ soln. p.b.k.; $K_1 = (4 \pm 1) \times 10^4 M^{-1}$.	73-9071 74-1107
5.23	Tl ²⁺ HO ₂ + Tl ²⁺ \rightarrow Tl ⁺ + H ⁺ + O ₂	1 (2.5 \pm 1) $\times 10^9$ (rel.)	—	p.r.	opt.	d.k. (Tl ²⁺); rel. to $k(Tl^{2+} + Tl^{2+}) = 2.3 \times 10^9$.	66-0097
5.24	UO ₂ ²⁺ (I) HO ₂ + UO ₂ ²⁺ \rightleftharpoons U(VI)-HO ₂ (II) U(VI)-HO ₂ + HO ₂ \rightarrow U(VI) + H ₂ O ₂ + O ₂ (III) 2U(VI)-HO ₂ \rightarrow 2U(VI) + H ₂ O ₂ + O ₂	~1 $\geq 1 \times 10^5$ (I) (9.0 \pm 1.5) $\times 10^5$ (II) 1 (1.5 \pm 0.1) $\times 10^5$ (I) (5 \pm 1) $\times 10^5$ (II) (8 \pm 2) $\times 10^4$ (III)	— — —	therm. p.r.	esr opt.	d.k. as well as p.b.k.; Ce ⁴⁺ + H ₂ O ₂ ; $K_1 = (2.7 \pm 0.4) 10^3 M^{-1}$. p.b.k. and d.k.; $K_1 = (1.7 \pm 0.3) \times 10^3 M^{-1}$.	73-9071 74-1107
5.25	VO(O ₂) ⁺ HO ₂ + VO(O ₂) ⁺ \rightarrow complex	— (9.4 \pm 1) $\times 10^4$ (rel.)	$k/k(\text{HO}_2 + \text{HO}_2) = 0.1$	therm.	esr	flow technique; Ce ⁴⁺ + H ₂ O ₂ ; assume $k(\text{HO}_2 + \text{HO}_2) = 9 \times 10^5$.	70-9058
5.26	cyclohexaneperoxy radical (RO ₂) HO ₂ + RO ₂ \rightarrow O ₂ + RO ₂ H	— 2.26 $\times 10^6$	—	γ -r.	chem.	detd. H ₂ O ₂ and RO ₂ H yields; assume $k(\text{RO}_2 + \text{RO}_2) = 2.7 \times 10^6$; see also 5.49.	67-0737
5.26a	cytochrome C (ferro)	5.3 5 $\times 10^5$ - 5 $\times 10^6$	—	p.r.	opt.	d.k. at 550 nm.	75-3093
5.27	cytochrome C HO ₂ + Fe ³⁺ -cyt \rightarrow no reaction	1.84 1.2-6.2 —	— — —	p.r. p.r.	opt. opt.	no reaction obs. no reaction.	71-0327 75-3093
5.28	ethylene HO ₂ + C ₂ H ₄ C ₂ H ₄ OOH	—	$k/k_{\text{Fe}^{2+}} = 0.167$	γ -r.	chem.	c.k.	67-0037
5.28a	horseradish peroxidase Compound I	— 2.2 $\times 10^8$	—	p.r.	opt.	d.k.; detd. <i>k</i> at pH 3.8 to 8.8.	74-1148
5.29	indigodisulfonate HO ₂ + dye \rightarrow decoloration	0.4 8.5 $\times 10^3$ (rel.)	$k/k(\text{HO}_2 + \text{HO}_2) = 3.9 \times 10^{-3}$	γ -r.	opt.	c.k.; assume $k(\text{HO}_2 + \text{HO}_2) = 2.2 \times 10^6$; $G(\text{HO}_2) = 3.6$.	68-0059
5.30	indigotrisulfonate HO ₂ + dye \rightarrow decoloration	0.4 4.5 $\times 10^3$ (rel.)	$k/k(\text{HO}_2 + \text{HO}_2) = 2 \times 10^{-4}$	γ -r.	opt.	c.k.; assume $k(\text{HO}_2 + \text{HO}_2) = 2.2 \times 10^6$; $G(\text{HO}_2) = 3.6$.	68-0059
5.31	indigotetrasulfonate HO ₂ + dye \rightarrow decoloration	0.4 7.7 $\times 10^2$ (rel.)	$k/k(\text{HO}_2 + \text{HO}_2) = 3.5 \times 10^{-4}$	γ -r.	opt.	c.k.; assume $k(\text{HO}_2 + \text{HO}_2) = 2.2 \times 10^6$; $G(\text{HO}_2) = 3.6$.	68-0059
5.31a	NADH-lactate dehydrogenase	— ~1.2 $\times 10^6$	—	p.r.	opt.	d.k.; detd. from k_{obs} at pH 4.4 to 9; see also 5.61a.	74-1159

TABLE 6. Reactions of HO₂ (H₂O₂⁺ and O₂⁻) with transients from water, inorganic solutes, and organic solutes—Continued

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
5.32	tetranitromethane HO ₂ + C(NO ₂) ₄ → C(NO ₂) ₃ ⁻ + NO ₂ + H ⁺ + O ₂	0-6 < 10 ⁵	—	p.r.	opt.	p.b.k.	65-0183
<i>O₂⁻ Reactions</i>							
5.33	<i>e</i> _{aq} ⁻ O ₂ ⁻ + <i>e</i> _{aq} ⁻ → O ₂ ²⁻	— 1.3 x 10 ¹⁰	—	—	—	See 1.10, NSRDS-NBS 43.	73-0030
5.34	OH O ₂ ⁻ + OH → OH ⁻ + O ₂	— 1 x 10 ¹⁰	—	—	—	See 3.7, Table 2.	—
5.35	O ₂ ⁻ O ₂ ⁻ + O ₂ ⁻ (+ H ₂ O) → HO ₂ ⁻ + O ₂ + OH ⁻	5.5 1.45 x 10 ⁷	—	p.r.	condy.	d.k.	60-0101 60-0121 62-0050
		11 2.7 x 10 ⁵	—	f.phot.	opt.	d.k.	62-0050
		— 1.5 x 10 ⁷	—	e-r.	chem.	c.k.; pH effects, flow techniques.	63-0075
		5-7 1.7 x 10 ⁷	—	p.r.	opt.	d.k.; ε(253.7 nm) = 980 ± 140 dm ³ mol ⁻¹ cm ⁻¹ .	64-0064
		7 5 x 10 ⁷	—	γ-r.	chem.	rotating sector method; H ₂ O ₂ soln.	65-0046
		5.5 1.45 x 10 ⁷	—	p.r.	condy.	d.k.	67-0213
		5.5 2.2 x 10 ⁷	—	f.phot.	opt.	d.k.; ε(260 nm) = 700 dm ³ mol ⁻¹ cm ⁻¹ .	67-7012
		5.7 5.8 x 10 ⁷	—	f.phot.	opt.	d.k.; ε(250 nm) = 1030 dm ³ mol ⁻¹ cm ⁻¹ .	67-7012
		12.8 1.4 x 10 ⁸	—	f.phot.	opt.	d.k.; ε(260 nm) = 900 dm ³ mol ⁻¹ cm ⁻¹ .	67-7012
		10.5 5.8 x 10 ⁷	—	f.phot.	opt.	d.k.; ε(260 nm) = 900 dm ³ mol ⁻¹ cm ⁻¹ .	67-7012
		5 1.2 x 10 ⁷	—	p.r.	opt.	d.k.; ε(241 nm) = 890 dm ³ mol ⁻¹ cm ⁻¹ . <i>k</i> = 4 x 10 ⁸ exp(-2100/RT); more than one rate constant is involved in calcn.	68-0382
		~5- 9.7 < 10 ⁵	—	p.r.	opt.	d.k.; ε(240 nm) = 1970 dm ³ mol ⁻¹ cm ⁻¹ ; <i>k</i> probably < 10 ³ .	69-0418
		13 < 10 ²	—	p.r.	opt.	d.k.; p <i>K</i> _a (HO ₂) = 4.88.	70-0304
		— < 10 ²	—	—	—	reaction with cytochrome C.	71-0327
	alk. 1.0 x 10 ⁷	—	—	f.phot.	condy.	d.k. (20°C); <i>k</i> = 1.63 x 10 ⁷ at 50°C; 9 x 10 ⁻⁵ M NaOH.	72-0404
5.36	Br ₂ O ₂ ⁻ + Br ₂ → O ₂ + Br ₂ ⁻	2-7 (5.6 ± 0.7) x 10 ⁹ (rel.)	—	e-r.	chem.	c.k. in formate- Br ₂ soln.; rel. to <i>k</i> (O ₂ ⁻ + C(NO ₂) ₄) = 2 x 10 ⁹ .	72-0308
5.37	Br ₃ ⁻ O ₂ ⁻ + Br ₃ ⁻ → O ₂ + Br ⁻ + Br ₂ ⁻	2-7 (3.8 ± 0.7) x 10 ⁹ (rel.)	—	e-r.	chem.	c.k. in formate- Br ₂ soln.; rel. to <i>k</i> (O ₂ ⁻ + C(NO ₂) ₄) = 2 x 10 ⁹ ; soln. contains 0.2 M Br ⁻ .	72-0308

TABLE 6. Reactions of HO₂ (H₂O₂[•] and O₂⁻) with transients from water, inorganic solutes, and organic solutes—Continued

No.	Solute and reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
5.38	HOBr O ₂ ⁻ + HOBr → OH ⁻ + O ₂ + Br	2-7	(9.5 ± 0.8) × 10 ⁸ (rel.)	—	<i>e</i> -r.	chem.	c.k. in formate- Br ₂ soln.; rel. to <i>k</i> (O ₂ ⁻ + C(NO ₂) ₄) = 2 × 10 ⁹ .	72-0308
5.39	CO ₃ ⁻ O ₂ ⁻ + CO ₃ ⁻ → CO ₅ ²⁻ or → O ₂ + CO ₃ ²⁻	~13	1.5 × 10 ⁹	—	p.r.	opt.	d.k. at 600 as well as 260 nm; ε(600 nm) for CO ₃ ⁻ = 1.8 × 10 ³ , ε(260 nm) for O ₂ ⁻ = 1.22 × 10 ³ dm ³ mol ⁻¹ cm ⁻¹ .	66-0001
		12.8	1.3 × 10 ⁸	—	f.phot.	opt.	d.k.	67-7012
		—	(4 ± 1) × 10 ⁸	—	f.phot.	opt.	d.k. at 260 nm and 600 nm; at 260 nm ε(CO ₅ ²⁻) = 410 and ε(O ₂ ⁻) = 1850 dm ³ mol ⁻¹ cm ⁻¹ .	70-0247
5.40	HCO ₃ ⁻ O ₂ ⁻ + HCO ₃ ⁻ → CO ₃ ⁻ + HO ₂ ⁻	5.5	1-2 × 10 ⁶	—	p.r.	condy.	d.k. (rotating sector); CO ₂ soln.	72-0404.
5.41	ClO ₂ O ₂ ⁻ + ClO ₂ → O ₂ + ClO ₂ ⁻	5-7		$k/k(\text{O}_2^- + \text{O}_2)^{0.5}$ = 1.7 ± 0.6 <i>M</i> ^{-0.5} s ^{-0.5}	γ-r.	chem.	c.k.	67-0028
5.42	Cu ⁺ O ₂ ⁻ + Cu ⁺ + (2H ₂ O) → Cu ²⁺ + H ₂ O ₂ + 2OH ⁻	~3-6.5	10 ¹⁰	—	p.r.	opt.	d.k. at 245 nm in Cu ²⁺ soln.	73-0112
5.43	Cu ²⁺ O ₂ ⁻ + Cu ²⁺ → Cu ⁺ + O ₂	1.35- 2.65 ~3-6.5	8 × 10 ⁹	<i>k</i> / <i>k</i> _{Fe³⁺} = 25 — For values of <i>k</i> (HO ₂ + Cu ²⁺)/ <i>k</i> (O ₂ ⁻ + Cu ²⁺) See 5.13.	Fenton p.r.	chem. opt.	c.k. d.k. at 245 nm.	51-9002 51-9005 73-0112
5.44	Fe ³⁺ O ₂ ⁻ + Fe ³⁺ → Fe ²⁺ + O ₂	<3 —		$k K(\text{HO}_2)/k(\text{HO}_2$ + Fe ²⁺) = 3.6 × 10 ⁻³ $k K(\text{HO}_2)/k(\text{HO}_2$ + Fe ²⁺) = 7 × 10 ⁻³	γ-r. Fenton	chem. chem.	c.k. c.k.	63-0004 51-9004
5.45	Fe(CN) ₆ ³⁻ O ₂ ⁻ + Fe(CN) ₆ ³⁻ → O ₂ + Fe(CN) ₆ ⁴⁻ O ₂ ⁻ + KFe(CN) ₆ ²⁻ → O ₂ + KFe(CN) ₆ ³⁻	9.5- 9.7	(2.7 ± 0.9) × 10 ² (6.2 ± 0.6) × 10 ³	—	p.r.	opt.	p.b.k. at 420-440 nm; μ = 0.	72-0431
5.46	H ₂ O ₂ O ₂ ⁻ + H ₂ O ₂ → OH + OH ⁻ + O ₂	~7	16.0 ± 3.3 (rel.)	$k/k(\text{O}_2^- + \text{O}_2)$ = 9.5 × 10 ⁻⁷	phot.	chem.	obs. rate of H ₂ O ₂ decompn.; assumed <i>k</i> (O ₂ ⁻ + O ₂) = 1.7 × 10 ⁷ ; recalcd. from 53-0014 <i>k</i> = 9.0, and from 62-0163 <i>k</i> = 12.0.	74-7351
5.47	benzoquinone O ₂ ⁻ + O=C ₆ H ₄ =O → O=C ₆ H ₄ O ⁻ + O ₂	~7 6.9 7.0	9.6 × 10 ⁸ (9.0 ± 0.9) × 10 ⁸ 9.8 × 10 ⁸	— — —	p.r. p.r. p.r.	opt. opt. opt.	p.b.k. at 430 nm. p.b.k. at 430 nm. p.b.k. at 430 nm.	71-0619 73-0049 73-0068
5.48	cyanocobalamin (Vitamin B ₁₂)	—	—	—	p.r.	—	no reaction	73-0116

TABLE 6. Reactions of HO₂ (H₂O₂[•] and O₂⁻) with transients from water, inorganic solutes, and organic solutes—Continued

No.	Solute and reaction pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
5.49	cyclohexaneperoxy radical (RO ₂) O ₂ ⁻ + RO ₂ (+H ⁺) → RO ₂ H + O ₂	— 2.54 x 10 ⁸	—	γ-r.	chem.	pH dependence of H ₂ O ₂ and RO ₂ H yields; assume $k(\text{RO}_2 + \text{RO}_2) = 2.7 \times 10^6$; see 5.26.	67-0737
5.50	cysteine O ₂ ⁻ + RSH(+H ⁺) → RS + H ₂ O ₂	7 > 5 x 10 ⁴ 3-5.1 ~ 1.8 x 10 ⁴	—	γ-r.	chem.	obs. G(H ₂ O ₂) as function of dose.	70-0882
5.50a	cytochrome C (ferro)	—	—	p.r.	opt.	obs. increase in G(H ₂ O ₂) with pH. no reaction.	74-0188 75-3093
5.51	cytochrome C O ₂ ⁻ + Fe ³⁺ -cyt C → Fe ²⁺ -cyt C	8.4 1.6 x 10 ⁵	—	therm.	esr	d.k.; O ₂ ⁻ from tetraacetylriboflavin + O ₂ .	69-9128
		8.5 1.1 x 10 ⁵	—	p.r.	opt.	p.b.k. at 550 nm.	71-0327
		10.4 8 x 10 ³	—	p.r.	opt.	p.b.k. at 550 nm.	71-0327
		7 2.4 x 10 ⁶	—	p.r.	opt.	p.b.k.	75-1012
		9.3 1.5 x 10 ⁵	—	p.r.	opt.	p.b.k.	75-1012
		4.7-6.7 (1.4 ± 0.15) x 10 ⁶	—	p.r.	opt.	p.b.k. at 550 nm; from pH effect pK _a (cyt C) = 7.45, 9.2; $k = (3.0 \pm 0.4) \times 10^5$ for the form present above pH 7.45; the form present above pH 9.2 does not react; E _a = 21.2 at pH 6.75 and 19.9 kJ/mol at pH 8.6.	75-3093
5.52	2,5-dichloro- <i>p</i> -benzoquinone	7.0 1.1 x 10 ⁹	—	p.r.	opt.	p.b.k. at 430 nm.	73-0068
	1,2-dihydroxybenzene-3,5-disulfonate ion	<i>See tiron (5.64).</i>					
5.52a	2,3-dimethylbenzoquinone	7 (4.5 ± 1) x 10 ⁸	—	p.r.	opt.	p.b.k.	73-0125
5.53	2,5-dimethyl- <i>p</i> -benzoquinone	7.0 7.5 x 10 ⁸ 7 (3.6 ± 1) x 10 ⁸	—	p.r.	opt.	p.b.k. at 430 nm.	73-0068
5.53a	2,6-dimethylbenzoquinone	7 (5.8 ± 1) x 10 ⁸	—	p.r.	opt.	p.b.k.	73-0125
5.54	4,4'-dimethyl-1,1'-bipyridylum chloride (Paraquat radical)	— 6.5 x 10 ⁸	—	p.r.	opt.	calcd. from d.k.; $k(\text{O}_2 + \text{PQ}^+) = 7.7 \times 10^8$.	73-1074
5.54a	2,3-dimethylnaphthoquinone	7 4 x 10 ⁶	—	p.r.	opt.	detd. from equil. const. and d.k. of semiquinone.	73-0125
5.55	diphenoquinone	7.0 (1.4 ± 0.14) x 10 ⁹	—	p.r.	opt.	p.b.k. at 400 nm	73-0068
5.55a	DNA	6.2, < 5 x 10 ⁶ 9.2	—	p.r.	—	—	75-3051
5.55b	duroquinone	7 5 x 10 ⁶	—	p.r.	opt.	detd. from equil. const. and d.k. of semiquinone.	73-0125
5.55c	horseradish peroxidase Compound I	7-8.8 1.6 x 10 ⁶	—	p.r.	opt.	d.k. as well as p.b.k., detd. <i>k</i> at pH 3.8 to 8.8.	74-1148, 74-3069
5.56	hydroquinone O ₂ ⁻ + OHC ₆ H ₄ OH → HO ₂ ⁻ + OHC ₆ H ₄ O [•] → O=C ₆ H ₄ O ⁻ + H ⁺	7.0 (1.6 ± 0.1) x 10 ⁷	—	p.r.	opt.	p.b.k. at 430 nm.	73-0068

TABLE 6. Reactions of HO₂ (H₂O₂⁺ and O₂⁻) with transients from water, inorganic solutes, and organic solutes. — Continued

No.	Solute and reaction	pH	<i>k</i>	Ratio	Source	Method	Comment	Ref.
5.57	2-methyl- <i>p</i> -benzoquinone	7.0 7	8.0 × 10 ⁸ (7.6 ± 1) × 10 ⁸	— —	p.r. p.r.	opt. opt.	p.b.k. at 430 nm. p.b.k. (semiqui- none).	73-0068 73-0125
5.58	1,2-naphthoquinone	7.0	7.2 × 10 ⁸	—	p.r.	opt.	p.b.k. at 365 nm.	73-0068
5.59	1,2-naphthoquinone 4-sulfonate ion	7.0	8.4 × 10 ⁸	—	p.r.	opt.	p.b.k. at 365 nm.	73-0068
5.60	1,4-naphthoquinone 2-sulfonate ion	7.0	6.6 × 10 ⁸	—	p.r.	opt.	p.b.k. at 400 nm.	73-0068
5.61	nicotinamide-adenine dinucleotide, reduced (NADH) O ₂ ⁻ + NADH (+ H ⁺) → H ₂ O ₂ + NAD•	8.6	<<27	—	X-r.	biol.	upper limit estd. for soln. contg. KBr and O ₂ .	71-0158
5.61a	NADH-lactate dehydrogenase	7-9	3.6 × 10 ⁴	—	p.r.	opt.	d.k.	74-1159
5.62	superoxide dismutase (E) Dismutation of O ₂ ⁻ (see 5.35) is catalyzed by E.	5.3- 9.5	1.8 × 10 ⁹	—	p.r.	opt.	d.k. at 250 nm (O ₂ ⁻) as well as 650 nm (Cu); enz- yme from bovine blood.	72-3066
		7	(1.4 ± 0.2) × 10 ⁹	—	p.r.	opt.	d.k. at 245 nm (O ₂ ⁻); enzyme from bovine blood.	72-1007, 72-3078
		7.5	(1.2 ± 0.2) × 10 ⁹	—	p.r.	opt.	d.k. at 650 nm (E); soln. con- tains Na formate and EDTA; enzyme from bovine blood.	73-0109
		5.0- 9.5	~ 2 × 10 ⁹ (rel.)	—	chem., biol.	opt.	c.k. (bovine Cu-Zn enzyme); assume <i>k</i> (O ₂ ⁻ + cyt C) = 1.1 × 10 ⁵ and <i>k</i> (O ₂ ⁻ + C(NO ₂) ₄) = 1.9 × 10 ⁹ ; rel. rates at pH 6.0- 10.2 also detd. for <i>E.coli</i> Mn and Fe enzymes and chicken liver mitochondria Mn enzyme.	73-3052
		5.7- 10.5	1.5 × 10 ⁹	—	p.r.	opt.	d.k. at 690 nm; Cu enzyme from human blood.	73-3132
		9.0- 9.9	(2.37 ± 0.18) × 10 ⁹	—	p.r.	opt.	d.k. at 250 nm; bovine Cu-Zn enzyme; super- sedes 72-3066.	74-3017
		7.9	(1.3 ± 0.15) × 10 ⁹	—	p.r.	opt.	d.k. at 248 nm; <i>E.coli</i> Mn enzyme.	74-3059
		9-10.2	2.3 × 10 ⁹	—	elec.	pol.	obs. increased O ₂ formn. with enzyme addn.	74-3132
	Paraquat radical See 4,4-dimethyl-1,1'-bipyridylum chloride (5.54). 2,3,5,6-tetramethylbenzoquinone See duroquinone (5.55a).							
5.63	tetranitromethane	—	(2.0 ± 0.4) × 10 ⁹	—	p.r.	opt.	p.b.k.	64-0133
	O ₂ ⁻ + C(NO ₂) ₄ → + C(NO ₂) ₃ + NO ₂	5.6- 6.2	(1.9 ± 0.4) × 10 ⁹	—	p.r.	opt.	p.b.k.	65-0183
5.64	tiron (1,2-dihydroxy- benzene-3,5-disul- fonate ion)	7 7	5 × 10 ⁸ 1.5 × 10 ⁸ (rel.)	— <i>k</i> / <i>k</i> _{benzoquinone} = 0.17	p.r. p.r.	opt. opt.	p.b.k. at 400 nm. c.k.; assume <i>k</i> _{benzoquinone} = 9 × 10 ⁸ .	75-1087 75-1087
5.65	Vitamin K ₁	7	< 2 × 10 ⁵	—	p.r.	opt.	detd. from equil. const. and d.k. of semiquinone.	73-0125

Formula index

The following formula list refers to entry numbers, not only in the preceding tables, but also in the tables of rates of hydrated electron and hydrogen atom reactions published as part I (and Supplemental data) and part II. The first digit of the entry number identifies the section of the tables where the entry can be found.

1. Part I. Hydrated electron 73-0030 (NSRDS-NBS 43)
- S1. Hydrated electron, Supplemental data 75-0002 (NSRDS-NBS 43-Supp)
2. Part II. Hydrogen Atom 75-0001 (NSRDS-NBS 51)
3. Part III. Hydroxyl radical (this work, tables 2-4)
4. Part III. Oxide ion (this work, table 5)
5. Part III. Perhydroxyl radical and superoxide ion (this work, table 6).

Thus, there are entries for Ag^+ in the tables of hydrated electron reactions (1.11), hydrogen atom reactions (2.5) and hydroxyl radical reactions (3.8), while BrO_4^- entries are found in the supplemental tables for hydrated electrons (S1.6), hydroxyl radical reactions (3.17), and oxide ion reactions (4.11).

- | | |
|---|---|
| Ag^+ Silver(I) ion, 1.11, 2.5, 3.8 | CH_2Cl_2 Dichloromethane (Methylene chloride), S1.266, 2.212 |
| AgH_6N_2^+ Diamminesilver(I) ion, 1.12 | CH_2I_2 Diiodomethane (Methylene iodide), S1.322 |
| Al^{3+} Aluminum(III) ion, 1.16 | CH_2O Formaldehyde, 1.432, 2.238, 3.382 |
| AlH_4O_4^- Aluminate ion, 1.17 | CH_2O_2 Formic acid, 1.435, S1.246, 2.241, 3.385 |
| AsF_6^- Hexafluoroarsenate(V) ion, 1.24, 2.7 | CH_3Cl Chloromethane, 1.367a, S1.174, 2.184 |
| AsHO_4^{2-} Arsenate ion, 1.23 | CH_3DO Methanol- <i>d</i> , 2.297 |
| AsO_2^- Arsenite ion, 1.22, 2.6, 3.9 | CH_3I Iodomethane, 1.495, 2.275, 3.478 |
| AuCl_4^- Tetrachloroaurate(III) ion, 2.8 | CH_3NO Formamide, 1.433, S1.245, 3.383 |
| BF_4^- Tetrafluoroborate ion, 1.26 | CH_3NO_2 Nitromethane, 1.553, 1.554, S1.348, 2.311, 3.573 |
| BH_4^- Tetrahydroborate ion, 3.11, 4.6 | CH_3NO_2^- Nitromethane anion, S1.349, 3.574 |
| $\text{B}_4\text{O}_7^{2-}$ Tetraborate ion 2.9 | CH_3O^- Methoxide ion, 2.299 |
| Br^- Bromide ion, 2.10a, 3.12, 3.13, 4.7 | CH_3S^- Methyl sulfide ion, 3.553 |
| $\text{BrCoH}_{15}\text{N}_5^{2+}$ Bromopentaamminecobalt(III) ion, 1.67 | CH_4 Methane, 1.519, 2.294, 3.509 |
| BrHO Hypobromous acid, 5.38 | $\text{CH}_4\text{N}_2\text{O}$ Urea, 1.650, 2.386, 3.749 |
| $\text{BrH}_{15}\text{N}_5\text{Ru}^{2+}$ Bromopentaammineruthenium(III) ion, S1.56 | $\text{CH}_4\text{N}_2\text{S}$ Thiourea, 1.624, 2.370 |
| BrO^- Hypobromite ion, 1.28, 3.14, 4.8 | $\text{CH}_4\text{N}_2\text{Se}$ Selenourea, 1.609, 2.352, 3.673 |
| BrO_2^- Bromite ion, 1.29, 3.15, 4.9 | CH_4O Methanol, 1.521, 2.296, 3.511, 4.80 |
| BrO_3^- Bromate ion, 1.30, 2.11, 3.16, 4.10 | $\text{CH}_4\text{O}_3\text{S}$ Methanesulfonic acid, 3.509a |
| BrO_4^- Perbromate ion, S1.6, 3.17, 4.11 | CH_4S Methanethiol, 1.520, 2.295, 3.510 |
| Br_2 Bromine, 2.10, 5.6, 5.36 | CH_5N Methylamine, 3.523, 4.82 |
| Br_2^- , 1.27, 5.7 | CH_5NO <i>N</i> -Methylhydroxylamine, S1.328; <i>O</i> -Methylhydroxylamine, S1.329, 3.538 |
| Br_3^- , 5.8, 5.37 | CH_5N_3 Guanidine, 1.463, 1.464, 2.259 |
| CBrF_3 Bromotrifluoromethane, 1.347a | CH_6N^+ Methylammonium ion, 1.524, 2.301, 3.524 |
| CClF_3 Chlorotrifluoromethane, 1.378, 2.189, 3.261 | CH_6NO^+ <i>N</i> -Methylhydroxylammonium ion, S1.330; <i>O</i> -Methylhydroxylammonium ion, S1.331 |
| CCl_2F_2 Dichlorodifluoromethane, 1.399a, 2.211 | CH_6N_2 Methylhydrazine, S1.325 |
| CCl_3F Trichlorofluoromethane, 1.635, 2.377 | CH_7N_2^+ Methylhydrazinium ion, S1.326 |
| CCl_4 Carbon tetrachloride, 1.355, 2.177 | $\text{CH}_{14}\text{CoN}_5\text{O}^{2+}$ Cyanoaquotetraamminecobalt(III) ion, 1.71 |
| CDO_2^- <i>d</i> -Formate ion, 2.240 | $\text{CH}_{15}\text{CoN}_6^{2+}$ Cyanopentaamminecobalt(III) ion, 1.68, 2.29 |
| CF_3I Trifluoriodomethane, 1.638a | $\text{CH}_{15}\text{CoN}_6\text{S}^{2+}$ Thiocyanatopentaamminecobalt(III) ion, 1.69, 2.30 |
| CF_4 Tetrafluoromethane, 2.359 | CN^- Cyanide ion, 1.35, 2.16, 3.23 |
| CHCl_3 Chloroform, 1.367, 2.183, 3.251 | CNO^- Cyanate ion, 1.36 |
| CHDO_2^- <i>d</i> -Formic acid, 2.242 | CNS , 5.9 |
| CHD_3O Methanol- <i>d</i> ₃ , 2.298, 3.512 | |
| CHN Hydrogen cyanide, S1.7, 2.15, 3.24 | |
| CHO_2^- Formate ion, 1.434, 2.239, 3.384, 4.68 | |
| CHO_3^- Bicarbonate ion, 1.33, 2.14, 3.20, 5.40 | |

- CNS^- Thiocyanate ion, 1.37, S1.9, 2.18, 3.25, 4.12
 CN_3O_6^- Trinitromethyl ion, 1.642
 CN_4O_8 Tetranitromethane, 1.618, 2.364, 5.63
 CO Carbon monoxide, 1.31, 2.12, 3.18
 CO_2 Carbon dioxide, 1.32, 2.13, 3.19
 CO_3^- , 5.39
 CO_3^{2-} Carbonate ion, 1.34, 3.21, 4.13
 CS_2 Carbon disulfide, 1.354, 2.176, 3.240
 C_2AgN_2^- Dicyanoargentate(I) ion, 1.13
 C_2AuN_2^- Dicyanoaurate(I) ion, 1.25, 3.10
 $\text{C}_2\text{Cl}_3\text{O}_2^-$ Trichloroacetate ion, 1.634
 C_2Cl_4 Tetrachloroethylene, 1.633a, 3.691
 $\text{C}_2\text{D}_3\text{O}_2^-$ Acetate ion- d_3 , 2.108
 $\text{C}_2\text{F}_3\text{O}_2^-$ Trifluoroacetate ion, 1.637, 3.725
 $\text{C}_2\text{HCl}_2\text{O}_2^-$ Dichloroacetate ion, 1.396a
 C_2HCl_3 1,1,2-Trichloroethylene, 1.634a, 3.721
 $\text{C}_2\text{HD}_5\text{O}$ Ethanol- d_5 , 3.359
 C_2HO_3^- Glyoxylate ion, 2.257
 C_2HO_4^- Oxalate ion, hydrogen, 1.570, 2.316, 3.593
 C_2H_2 Acetylene, 1.295, 2.114, 3.137
 $\text{C}_2\text{H}_2\text{BrO}_2^-$ Bromoacetate ion, 1.335, 2.158, 3.208
 $\text{C}_2\text{H}_2\text{ClO}_2^-$ Chloroacetate ion, 1.358, 2.178, 3.246
 $\text{C}_2\text{H}_2\text{Cl}_2$ 1,1-Dichloroethylene, 1.399b, 3.302;
 1,2-Dichloroethylene, 1.399c, 3.303
 $\text{C}_2\text{H}_2\text{FO}_2^-$ Fluoroacetate ion, 1.423, 2.236, 3.378
 $\text{C}_2\text{H}_2\text{IO}_2^-$ Iodoacetate ion, 1.488
 $\text{C}_2\text{H}_2\text{NO}_3^-$ Oxamate ion, 1.572, S1.359
 $\text{C}_2\text{H}_2\text{O}_2$ Glyoxal, 2.256, 3.427
 $\text{C}_2\text{H}_2\text{O}_3$ Glyoxylic acid, S1.273, 2.258
 $\text{C}_2\text{H}_2\text{O}_4$ Oxalic acid, 1.571, 2.316-7, 3.594
 $\text{C}_2\text{H}_3\text{BrO}_2$ Bromoacetic acid, 2.159
 $\text{C}_2\text{H}_3\text{Cl}$ Vinyl chloride, 1.658a, 3.754
 $\text{C}_2\text{H}_3\text{ClO}_2$ Chloroacetic acid, 1.359, 2.179, 3.247
 $\text{C}_2\text{H}_3\text{Cl}_3\text{O}_2$ Chloral hydrate, S1.171, 3.245
 $\text{C}_2\text{H}_3\text{IO}_2$ Iodoacetic acid, 3.473a
 $\text{C}_2\text{H}_3\text{N}$ Acetonitrile, 1.292, 2.111, 3.133, 4.35
 $\text{C}_2\text{H}_3\text{O}_2^-$ Acetate ion, 1.287, S1.77, 2.107, 3.128, 4.34
 $\text{C}_2\text{H}_3\text{O}_2\text{S}^-$ Thioglycolate ion, 1.621, S1.416, 3.705
 $\text{C}_2\text{H}_3\text{O}_3^-$ Glycolate ion, 2.252, 3.408
 C_2H_4 Ethylene, 1.419, 2.229, 3.365, 5.28
 $\text{C}_2\text{H}_4\text{CdNO}_2^+$ Glycinatocadmium(II) ion, 1.43
 $\text{C}_2\text{H}_4\text{D}_2\text{O}$ Ethanol- d_2 , 2.225
 $\text{C}_2\text{H}_4\text{INO}$ Iodoacetamide, S1.292
 $\text{C}_2\text{H}_4\text{NNiO}_2^+$ Glycinatonickel(II) ion, 1.196
 $\text{C}_2\text{H}_4\text{N}_2$ Aminoacetonitrile, 2.125
 $\text{C}_2\text{H}_4\text{N}_2\text{O}_2$ Oxamide, S1.360
 $\text{C}_2\text{H}_4\text{O}$ Acetaldehyde, 1.284, 2.104, 3.123;
 Ethylene oxide, 3.370
 $\text{C}_2\text{H}_4\text{O}_2$ Acetic acid, 1.288, S1.78, 2.109, 3.129
 $\text{C}_2\text{H}_4\text{O}_2\text{S}$ Thioglycolic acid, 2.367, 3.704
 $\text{C}_2\text{H}_4\text{O}_3$ Glycolic acid, S1.260, 2.253, 3.409
 $\text{C}_2\text{H}_5\text{Br}$ 1-Bromoethane, 1.339, 2.160
 $\text{C}_2\text{H}_5\text{BrO}$ 2-Bromoethanol, 1.340, 2.161, 3.210
 $\text{C}_2\text{H}_5\text{Cl}$ Chloroethane, 2.181
 $\text{C}_2\text{H}_5\text{ClO}$ 2-Chloroethanol, 1.366, S1.173, 2.182, 3.250
 $\text{C}_2\text{H}_5\text{I}$ Iodoethane, 1.494
 $\text{C}_2\text{H}_5\text{NO}$ Acetaldoxime, 1.285; Acetamide, 1.286, S1.75,
 2.105, 3.124; *N*-Methylformamide, 1.530,
 S1.323, 3.535
 $\text{C}_2\text{H}_5\text{NO}_2$ Glycine, 1.443-5, S1.256-8, 2.250-1,
 3.404-3.406, 4.71; Glycine, copper salt,
 1.116a; Hydroxyacetamide, S1.284, 3.447a;
 Nitroethane, 2.310
 $\text{C}_2\text{H}_5\text{NS}$ Thioacetamide, 2.365
 $\text{C}_2\text{H}_5\text{N}_3\text{O}_2$ Biuret, S1.154
 $\text{C}_2\text{H}_5\text{O}^-$ Ethoxide ion, 2.226
 $\text{C}_2\text{H}_5\text{OS}^-$ 2-Hydroxyethylsulfide ion, 3.454
 $\text{C}_2\text{H}_5\text{O}_3\text{S}^-$ Ethanesulfonate ion, S1.234, 3.357
 C_2H_6 Ethane, 2.223, 3.356
 $\text{C}_2\text{H}_6\text{N}_2\text{O}$ 2-Aminoacetamide(Glycine amide), S1.115
 $\text{C}_2\text{H}_6\text{O}$ Ethanol, 1.411, 2.224, 3.358, 4.65
 $\text{C}_2\text{H}_6\text{OS}$ Dimethyl sulfoxide, 1.405, S1.227, 3.342;
 2-Mercaptoethanol, 1.514, S1.304, 2.292,
 3.506
 $\text{C}_2\text{H}_6\text{O}_2$ Ethylene glycol, 2.231, 3.369
 $\text{C}_2\text{H}_6\text{O}_2\text{S}$ Dimethyl sulfone, 3.341a
 $\text{C}_2\text{H}_6\text{O}_4\text{P}^-$ Dimethyl phosphate ion, 3.338
 $\text{C}_2\text{H}_6\text{S}$ Methyl sulfide, 1.404, 3.552
 $\text{C}_2\text{H}_6\text{S}_2$ Dimethyl disulfide, 3.327a
 $\text{C}_2\text{H}_7\text{N}$ Ethylamine, S1.236, 3.362, 4.66
 $\text{C}_2\text{H}_7\text{NO}$ 2-Aminoethanol, 2.126
 $\text{C}_2\text{H}_7\text{NS}$ Cysteamine, 1.389, S1.193, 2.204, 3.289
 $\text{C}_2\text{H}_7\text{O}_4\text{P}$ Ethyldihydrogen phosphate, 2.233
 $\text{C}_2\text{H}_8\text{CdN}_2^{2+}$ Ethylenediaminecadmium(II) ion, 1.48
 $\text{C}_2\text{H}_8\text{N}^+$ Dimethylammonium ion, 3.324; Ethylammonium ion,
 1.417, S1.237, 3.363
 $\text{C}_2\text{H}_8\text{N}_2$ 1,1-Dimethylhydrazine, S1.222, 3.329;
 1,2-Dimethylhydrazine, S1.223, 3.330;
 Ethylenediamine, 3.366
 $\text{C}_2\text{H}_8\text{N}_2\text{Ni}^{2+}$ Ethylenediaminenickel(II) ion, 1.202
 $\text{C}_2\text{H}_9\text{N}_2^+$ 1,1-Dimethylhydrazinium ion, S1.220, 3.331;
 1,2-Dimethylhydrazinium ion, S1.221, 3.332
 $\text{C}_2\text{H}_{10}\text{TI}^+$ Diethylthallium ion, 1.401
 $\text{C}_2\text{H}_{15}\text{CoF}_3\text{N}_5^{2+}$ Trifluoroacetatopentaamminecobalt(III) ion,
 2.38
 $\text{C}_2\text{H}_{18}\text{CoN}_5\text{O}_2^{2+}$ Acetatopentaamminecobalt(III) ion, 1.72,
 2.36
 C_2N_2 Cyanogen, S1.8, 2.17, 3.22
 $\text{C}_2\text{O}_4^{2-}$ Oxalate ion, 1.569, 2.315, 3.592, 4.89
 $\text{C}_3\text{D}_6\text{O}$ Acetone- d_6 , 3.132
 $\text{C}_3\text{HD}_7\text{O}$ 2-Propanol- d_7 , 2.336
 $\text{C}_3\text{H}_2\text{D}_6\text{O}$ 2-Propanol- d_6 , 3.639
 $\text{C}_3\text{H}_2\text{NO}_2^-$ Cyanoacetate ion, 1.382, 3.272, 4.61
 $\text{C}_3\text{H}_2\text{N}_2$ Malononitrile, 2.291
 $\text{C}_3\text{H}_2\text{O}_4^{2-}$ Malonate ion, S1.302, 3.502, 4.77
 $\text{C}_3\text{H}_3\text{F}_3\text{O}$ α,α,α -Trifluoroacetone, 1.638
 $\text{C}_3\text{H}_3\text{F}_3\text{O}_2$ Methyl trifluoroacetate, 1.537
 $\text{C}_3\text{H}_3\text{N}$ Acrylonitrile, S1.102, 3.145
 $\text{C}_3\text{H}_3\text{NS}$ Thiazole, 1.619
 $\text{C}_3\text{H}_3\text{NO}_2$ Cyanoacetic acid, 2.192
 $\text{C}_3\text{H}_3\text{O}_2^-$ Acrylate ion, S1.100, 4.39
 $\text{C}_3\text{H}_3\text{O}_3^-$ Pyruvate ion, 1.601, 3.660
 $\text{C}_3\text{H}_3\text{O}_4^-$ Hydrogen malonate ion, 1.513, S1.250
 $\text{C}_3\text{H}_4\text{BrO}_2^-$ 2-Bromopropionate ion, 1.346, 2.164, 3.214;
 3-Bromopropionate ion, 1.347, 2.165, 3.215
 $\text{C}_3\text{H}_4\text{ClO}_2^-$ 2-Chloropropionate ion, 1.375, 2.185, 3.257;
 3-Chloropropionate ion, 1.376, 2.186,
 3.258
 $\text{C}_3\text{H}_4\text{IO}_2^-$ 2-Iodopropionate ion, 1.497; 3-Iodopropionate
 ion, S1.293
 $\text{C}_3\text{H}_4\text{N}_2$ Imidazole, 1.484, 2.273, 3.467
 $\text{C}_3\text{H}_4\text{N}_2\text{O}_3$ Barbituric acid, S1.130
 $\text{C}_3\text{H}_4\text{O}$ Acrolein, 3.142
 $\text{C}_3\text{H}_4\text{O}_2$ Acrylic acid, S1.101, 3.144
 $\text{C}_3\text{H}_4\text{O}_4$ Malonic acid, S1.303, 2.290, 3.503
 $\text{C}_3\text{H}_4\text{O}_5$ Tartronic acid, 2.358

$C_3H_5BrO_2$ 2-Bromopropionic acid, 2.166; 3-Bromopropionic acid, 2.167
 $C_3H_5ClO_2$ 2-Chloropropionic acid, 2.187; 3-Chloropropionic acid, 2.188
 C_3H_5FO Fluoroacetone, 1.424
 $C_3H_5FO_2$ Methyl fluoroacetate, 1.529
 $C_3H_5IO_2$ 3-Iodopropionic acid, 3.479
 C_3H_5N Propionitrile, 1.593, 2.339, 3.643, 4.97
 C_3H_5NO Acrylamide, 1.299, S1.99, 2.118, 3.143, 4.38
 $C_3H_5NO_3$ *N*-Formylglycine, S1.247
 $C_3H_5N_2^+$ Imidazolium ion, 1.485
 $C_3H_5O_2^-$ Propionate ion, 2.337, 3.641, 4.96
 $C_3H_5O_2S^-$ 2-Mercaptopropionate ion (Thiolactate ion), S1.304a, 3.706; 3-Mercaptopropionate ion, S1.305, 3.508
 $C_3H_5O_3^-$ Lactate ion, 1.501, S1.296, 3.490; Methoxyacetate ion, 3.514
 C_3H_6 Cyclopropane, 2.202; Propylene, 2.340, 3.647
 C_3H_6ClNO 2-Chloropropionamide, 1.373; 3-Chloropropionamide, 1.374
 $C_3H_6N_2O_2$ Malonamide, S1.300
 C_3H_6O Acetone, 1.289, S1.80, 2.110, 3.131; Allyl alcohol, 1.309, 2.124, 3.156, 4.41; 1,2-Epoxypropane, 3.353
 $C_3H_6O_2$ 2,3-Epoxypropanol, 3.354; Ethyl formate, 3.372; Methyl acetate, 2.300, 3.522; Propionic acid, S1.386, 2.338, 3.642
 $C_3H_6O_2S$ 2-Mercaptopropionic acid, S1.306, 2.292a; 3-Mercaptopropionic acid, S1.307, 2.292b; Methyl thioglycolate, S1.336, 3.554
 $C_3H_6O_3$ Lactic acid, S1.297, 2.285, 3.491; Methyl 2-hydroxyacetate, 1.532, S1.327; 1,3,5,-Trioxane, S1.431, 3.732
 C_3H_7Br 1-Bromopropane, 1.345
 C_3H_7Cl 1-Chloropropane, 1.372
 C_3H_7DO 2-Propanol-2-*d*, 2.335, 3.638
 C_3H_7I 1-Iodopropane, 1.496
 C_3H_7NO Acetone oxime, 1.291, S1.81; *N,N*-Dimethylformamide, 1.403, S1.218, 3.328; *N*-Methylacetamide, 3.521; Propionamide, 1.592, S1.385, 3.640
 $C_3H_7NO_2$ Alanine, 1.303-4, S1.110, 2.122, 3.150-3.152; β -Alanine, 1.305, S1.111, 2.123; 2-Hydroxypropionamide, 3.461a; Methyl 2-aminoacetate (Glycine methyl ester), 1.523, S1.318; 1-Nitropropane, 2.312; Sarcosine, 1.608, 2.351
 $C_3H_7NO_2S$ Cysteine, 1.390-2, S1.194, 2.205-6, 3.290, 5.50
 $C_3H_7NO_3$ Serine, 1.610, 2.353, 3.674
 C_3H_8 Propane, 2.332
 $C_3H_8N^+$ Allylammonium ion, 3.157
 C_3H_8O 1-Propanol, 2.333, 3.636, 4.94; 2-Propanol, 2.334, 3.637, 4.95
 $C_3H_8O_2$ Dimethoxymethane, 3.322; 2-Methoxyethanol, 3.516; 1,2-Propanediol, 3.634; 1,3-Propanediol, 2.331, 3.635
 $C_3H_8O_3$ Glycerol, 2.249, 3.403
 C_3H_9N Isopropylamine, 1.500a, 3.487; Propylamine, S1.387, 3.645, 4.98; Trimethylamine, 3.726
 $C_3H_9N_3S$ Mercaptoethylguanidine, 1.515
 $C_3H_9O_4P$ Trimethyl phosphate, S1.430, 3.730
 $C_3H_{10}N^+$ Isopropylammonium ion, 3.488; Propylammonium ion, 1.593a, S1.388, 3.646; Trimethylammonium ion, 3.727
 $C_3H_{10}N_2$ Trimethylhydrazine, S1.428
 $C_3H_{11}N_2^+$ Trimethylhydrazinium ion, S1.429
 $C_3O_5^{2-}$ Oxomalonate ion, S1.362
 $C_4CdN_4^{2-}$ Tetracyanocadmiate(II) ion, 1.42, 2.20
 $C_4CuN_4^{2-}$ Tetracyanocuprate(II) ion, 1.122
 $C_4H_2BrO_3^-$ α -Bromotetronate ion, 3.218
 $C_4H_2O_4$ Acetylenedicarboxylic acid, 2.115
 $C_4H_2O_4^{2-}$ Fumarate ion, 1.436, 4.69; Maleate ion, 1.512, S1.299, 4.76
 $C_4H_3BrN_2O_2$ 5-Bromouracil, 1.348, S1.159, 2.168, 3.219
 $C_4H_3ClN_2O_2$ 5-Chlorouracil, S1.178, 2.190, 3.262
 $C_4H_3FN_2O_2$ 5-Fluorouracil, S1.244, 2.237, 3.381
 $C_4H_3IN_2O_2$ Iodouracil, 1.499
 $C_4H_3N_2O_3^-$ Barbiturate ion, 2.140
 $C_4H_3N_3O_4$ 5-Nitrouracil, S1.355, 3.584
 $C_4H_3N_3O_5$ 5-Nitrobarbituric acid, S1.340, 3.564
 $C_4H_3O_3^-$ Tetronate ion, 3.699
 $C_4H_3O_4^-$ Hydrogen fumarate ion, 2.243; Hydrogen maleate ion, 1.511, S1.299; α -Hydroxytetronate ion, 3.466
 $C_4H_3O_5^-$ Oxalacetate ion, 1.568, S1.358
 $C_4H_4CrO_{10}^-$ Dioxalatodiaquochromate(III) ion, 1.111
 $C_4H_4N_2$ Pyrazine, 2.343; Pyridazine, 2.344; Pyrimidine, 2.347, 3.657b; Succinonitrile, 2.682
 $C_4H_4N_2O_2$ Uracil, 1.647-8, S1.437, 2.385, 3.746, 4.118
 $C_4H_4N_2O_2S$ Thiobarbituric acid, 1.620
 $C_4H_4N_2O_3$ Barbituric acid, 2.141
 C_4H_4O Furan, 1.437, 3.390
 $C_4H_4O_4$ Fumaric acid, 2.244, 3.386; Maleic acid, 2.288, 3.500
 $C_4H_4O_4^{2-}$ Succinate ion, 1.614, S1.401, 2.354, 4.102
 $C_4H_4O_4S^{2-}$ Thiodiacetate ion, S1.414
 $C_4H_4O_4S_2^{2-}$ 2,2'-Dithiobisacetate ion, S1.229
 $C_4H_4O_5$ Oxalacetic acid, 2.314
 $C_4H_4O_5^{2-}$ Malate ion, 1.510, 3.499
 $C_4H_4O_6$ 2,3-Dihydroxyfumaric acid, 2.216
 $C_4H_4O_6^{2-}$ Tartrate ion, 3.689
 $C_4H_4O_8^{2-}$ Tetrahydroxysuccinate ion, 3.695
 C_4H_4S Thiophene, 1.622, 3.707
 C_4H_5N 3-Butenenitrile (Allyl cyanide), 1.351, 4.43; Crotononitrile, 4.60; Methacrylonitrile, 4.78; Pyrrole, 1.597, 3.658
 $C_4H_5NO_2$ Methyl cyanoacetate, 1.526; Succinimide, 1.615, S1.403, 3.681
 $C_4H_5NO_4^{2-}$ Aspartate ion, 1.322
 $C_4H_5N_3$ 2-Aminopyrimidine, 1.313, 3.167; 4-Aminopyrimidine, 1.313a
 $C_4H_5N_3O$ Cytosine, 1.396, S1.204, 2.209, 3.295
 $C_4H_5O_2^-$ 3-Butenoate ion, 4.54; Crotonate ion, S1.188, 4.59; Methacrylate ion, 1.518, S1.309, 4.79
 $C_4H_5O_3^-$ Acetoacetate ion, S1.79
 $C_4H_5O_4^-$ Succinate ion, 1.613, S1.401
 C_4H_6 1,3-Butadiene, 1.349, 2.168a, 3.220
 $C_4H_6NO_3$ *N*-Acetyl glycine, 2.117, 3.139
 $C_4H_6NO_4^-$ Aspartate ion, 1.321, 2.138
 $C_4H_6N_2$ *N*-Methylimidazole, 3.538a
 $C_4H_6N_2O_2$ Glycine anhydride, S1.259, 3.407; Hydrouacil, 1.474, 3.318
 $C_4H_6N_3O_4P$ Cytidine-5'-phosphate, 3.290
 C_4H_6O 1-Butene-3-one, 3.229; Crotonaldehyde, 3.270
 $C_4H_6O_2$ Biacetyl, 2.155, 3.202; 2,3-Butanedione, 1.350; Crotonic acid, S1.189, 3.271; Cyclopropanecarboxylic acid, 2.203; Methacrylic

- acid, S1.310; Methyl acrylate, S1.315;
Vinyl acetate, S1.439
- $C_4H_6O_4$ Succinic acid, S1.402, 2.355, 3.680
- $C_4H_6O_4S$ Thiodiglycolic acid, 2.366, 3.703; Thiomalic acid, 2.368
- $C_4H_6O_4S_2$ Dithiodiglycolic acid, 2.219
- $C_4H_6O_5$ Malic acid, 2.289, 3.501
- $C_4H_6O_6$ Tartaric acid, 2.357, 3.688
- C_4H_7N Isobutyronitrile, 2.281
- C_4H_7NO Methacrylamide, S1.308; 2-Pyrrolidone, 1.600
- $C_4H_7NO_2$ Diacetamide, S1.208
- $C_4H_7NO_3$ *N*-Acetylglycine, 1.296, S1.87
- $C_4H_7NO_4$ Aspartic acid, S1.128, 2.139, 3.181; Iminodi-acetic acid, 2.274
- $C_4H_7O_2^-$ Butyrate ion, 2.174, 3.238, 4.56; 2-Methylpropio-nate ion (Isobutyrate ion), 2.279, 3.549
- C_4H_8 1-Butene, 2.173, 3.228; Isobutylene, 2.278, 3.482
- $C_4H_8CdN_2O_4$ Bis(glycinato)cadmium(II), 1.44
- $C_4H_8CuN_2O_4^{2+}$ Bis(glycinato)copper(II) ion, 3.43
- $C_4H_8NO_2^-$ 4-Aminobutyrate ion, 1.312
- $C_4H_8N_2NiO_2$ Dimethylglyoximenickelate(II) ion, 4.28
- $C_4H_8N_2NiO_4$ Bis(glycinato)nickel(II), 1.197
- $C_4H_8N_2O_2$ *N*-Acetylglycine amide, S1.88; Succinamide, S1.400
- $C_4H_8N_2O_3$ Asparagine, 1.319–20, 2.137, 3.180; Glycylgly-cine, 1.450–2, S1.261–2, 2.254, 3.411–13
- C_4H_8O 2-Butanone, 3.227; Butyraldehyde, 3.237; 1,2-Epoxy-butane, 3.352; Tetrahydrofuran, 2.360, 3.693
- $C_4H_8O_2$ Acetoin, 3.130; 2-Butene-1,4-diol, 4.53; Butyric acid, 2.175, 3.239; Dioxane, 2.218, 3.343; Ethyl acetate, 1.415, 2.227, 3.361; 3-Hydroxy-2-butanone, 1.480; Isobutyric acid, 2.280; Methyl propionate, 1.536, 3.548
- $C_4H_8O_3$ 2-Hydroxybutyric acid, 3.450, 2-Hydroxyethyl acetate, 3.451
- C_4H_9Br 1-Bromobutane, 1.338
- C_4H_9Cl 1-Chlorobutane, 1.364; 2-Chlorobutane, 1.365; 1-Chloro-2-methylpropane, 1.368
- C_4H_9I 1-Iodobutane, 1.493
- C_4H_9N Pyrrolidine, 1.598–9, 3.659, 4.99a
- C_4H_9NO *N,N*-Dimethylacetamide, S1.215, 3.323; *N*-Ethyl-acetamide, 1.414; Isobutyramide, 3.483; *N*-Methylpropionamide, 3.547
- C_4H_9NOS *N*-Acetylcysteamine, S1.85
- $C_4H_9NO_2$ 2-Aminobutyric acid, 2.125a, 3.160; 3-Amino-butyric acid, 3.161; 4-Aminobutyric acid, 3.162; 2-Amino-2-methylpropionic acid, 2.127; Ethyl 2-aminoacetate, 1.416
- $C_4H_9NO_2S$ Cysteine, methyl ester, S1.195; *S*-Methylcysteine, S1.319, 3.532
- $C_4H_9NO_3$ 2-Methyl-2-nitro-1-propanol, 1.556; Threonine, 1.625, 2.371, 3.708
- $C_4H_9NO_4$ 2-Methyl-2-nitro-1,3-propanediol, 1.555
- $C_4H_9N_3O$ Acetone semicarbazone, 1.290
- $C_4H_9N_3O_2$ Creatine, 1.381; Glycylglycine amide, S1.263, 3.414
- C_4H_{10} Butane, 2.169; Isobutane, 2.276
- $C_4H_{10}N^+$ Pyrrolidinium ion, 1.599, 3.659
- $C_4H_{10}O$ 1-Butanol, 2.170, 3.225; 2-Butanol, 2.171, 3.226; Ethyl ether, 1.421, 2.232, 3.371, 4.67; 2-Methyl-1-propanol (Isobutanol), 2.277, 3.546, 4.83; 2-Methyl-2-propanol (*tert*-Butanol), 1.352, 2.172, 3.545
- $C_4H_{10}O_2$ 1,2-Butanediol, 3.221; 1,3-Butanediol, 3.222; 1,4-Butanediol, 3.223; 2,3-Butanediol, 3.224; 1,2-Dimethoxyethane, 3.321; 2-Ethoxyethanol, 3.360
- $C_4H_{10}O_2S_2$ Dithiothreitol, 3.347
- $C_4H_{10}O_3$ Diethyleneglycol, 3.308
- $C_4H_{10}O_4$ Erythritol, 3.355
- $C_4H_{10}S$ *tert*-Butylmercaptan, 1.353, 3.235
- $C_4H_{10}S_2$ Diethyl disulfide, 3.307a
- $C_4H_{10}Tl^+$ Diethylthallium ion, 1.401
- $C_4H_{11}N$ Butylamine, S1.161, 3.231, 4.55; *tert*-Butylamine, 3.232
- $C_4H_{11}NO$ *N,N*-Diethylhydroxylamine, S1.212
- $C_4H_{12}N^+$ Butylammonium ion, S1.162, 3.233; *tert*-Butylam-monium ion, 1.352a, 3.234; Diethylammonium ion, 3.307; Isobutylammonium ion, 3.481
- $C_4H_{12}NO^+$ *N,N*-Diethylhydroxylammonium ion, S1.211
- $C_4H_{12}N_2S$ 2,2'-Dithiobis(ethylamine), 3.346
- $C_4H_{12}N_2S_2$ Cystamine, 1.388, S1.192
- $C_4H_{16}CdN_4^{2+}$ Bis(ethylenediamine)cadmium(II) ion, 1.49
- $C_4H_{16}Cl_2CoN_4^+$ Dichlorobis(ethylenediamine)cobalt(III) ion, 1.87, 2.43
- $C_4H_{16}Cl_2CrN_4^+$ Dichlorobis(ethylenediamine)chromium(III) ion, 1.107
- $C_4H_{16}CoF_2N_4^+$ Difluorobis(ethylenediamine)cobalt(III) ion, 1.86, 2.42
- $C_4H_{16}CuN_4^{2+}$ Bis(ethylenediamine)copper(II) ion, 3.42
- $C_4H_{16}Ni_4^{2+}$ Bis(ethylenediamine)nickel(II) ion, 1.203
- $C_4H_{18}ClCoN_5^{2+}$ Chloroamminebis(ethylenediamine)cobalt(III) ion, 1.89
- $C_4H_{18}CoFN_4O^{2+}$ Fluoroaquo-bis(ethylenediamine)cobalt(III) ion, 1.91
- $C_4H_{18}CoN_5O_4^+$ Fumaratopentaamminecobalt(III) ion, 1.73, 2.32
- $C_4H_{18}CoN_6O_2^{2+}$ Nitroamminebis(ethylenediamine)cobalt(III) ion, 1.90
- $C_4H_{20}CoN_4O_2^{3+}$ Diaquo-bis(ethylenediamine)cobalt(III) ion, 2.41
- $C_4HgN_4^{2-}$ Tetracyanomercurate(II) ion, 1.150
- $C_4N_4Ni^{2-}$ Tetracyanonickelate(II) ion, 1.195, 3.85
- $C_4N_4Pd^{2-}$ Tetracyanopalladate(II) ion, 1.221
- $C_4N_4Pt^{2-}$ Tetracyanoplatinate(II) ion, 1.226, S1.53, 2.93, 3.100
- $C_4N_4Zn^{2-}$ Tetracyanozincate(II) ion, 1.279
- $C_4O_4^{2-}$ Acetylenedicarboxylate ion, 4.36
- $C_4O_9Ti^{2-}$ Bisoxalatoxo-titanate(IV) ion, 3.117a
- $C_5ClCoN_5^{3-}$ Chloropentacyanocobaltate(III) ion, 1.77
- $C_5CoIN_5^{3-}$ Iodopentacyanocobaltate(III) ion, S1.20
- $C_5CoN_5^{3-}$ Pentacyanocobaltate(II) ion, 1.59, S1.14
- $C_5CoN_6O^{3-}$ Nitrosylpentacyanocobaltate ion, S1.21, 3.35
- $C_5CoN_6O_2^{3-}$ Nitropentacyanocobaltate(III) ion, 1.80
- $C_5CoN_8^{3-}$ Azidopentacyanocobaltate(III) ion, 1.79
- $C_5CrN_6O^{3-}$ Nitrosylpentacyanochromate ion, S1.23, 3.39
- C_5D_5N Pyridine-*d*₅, 3.650
- $C_5FeN_6O^{2-}$ Nitrosylpentacyanoferrate(III) ion, 1.138, 3.57
- $C_5HCoN_5^{3-}$ Hydridopentacyanocobaltate(III) ion, S1.16
- $C_5HCoN_5O^{3-}$ Hydroxypentacyanocobaltate(III) ion, 1.78, S1.19
- $C_5HD_5N^+$ Pyridinium ion-*d*₅, 3.657
- $C_5H_2BrN_2O_4^{5-}$ Bromoorotate ion, S1.155
- $C_5H_2CoN_5O^{2-}$ Aquopentacyanocobaltate(III) ion, S1.18
- $C_5H_2NO_5^{5-}$ Nitrofuroate ion, S1.347, 3.571
- $C_5H_2N_3O_6^{5-}$ Nitrooorotate ion, 3.576
- $C_5H_3BrN_2O_4$ 5-Bromoorotic acid, S1.156, 2.162, 3.212

$C_5H_3FeN_6^{3-}$ Pentacyanoammineferrate(II) ion, 1.135
 $C_5H_3NO_4$ 5-Nitro-2-furaldehyde, S1.343, 3.569
 $C_5H_3N_2O_4^-$ Isoorotate ion, 1.500c, 3.485; Orotate ion, 1.567, 3.590
 $C_5H_3N_3O_6$ 5-Nitroorotic acid, S1.351
 $C_5H_3O_3^-$ 2-Furoate ion, 3.392
 C_5H_4BrN 2-Bromopyridine, 3.216; 3-Bromopyridine, 3.217
 C_5H_4ClN 2-Chloropyridine, 3.259; 3-Chloropyridine, 3.260
 $C_5H_4NO^-$ 2-Hydroxypyridine, anion, 3.462; 3-Hydroxypyridine, anion, 3.464; 4-Hydroxypyridine, anion, 3.465
 $C_5H_4N_2O_4$ Isoorotic acid, 2.283; *Anti*-5-Nitro-2-furaldoxime, S1.346, 2.310a, 3.568; Orotic acid, 2.313
 $C_5H_4N_4$ Purine, 1.595, S1.389, 2.342, 3.648a
 $C_5H_4N_4O$ Hydroxypurine, 3.461b; Hypoxanthine, 1.483, 3.466a
 $C_5H_4N_4O_2$ Xanthine, 3.754a
 $C_5H_4N_4O_3$ Uric acid, 1.651, 3.749a
 $C_5H_4O_2$ 2-Furaldehyde, 3.388
 $C_5H_4O_4^{2-}$ Glutaconate ion, 4.70
 $C_5H_4O_5^{2-}$ 2-Oxoglutarate ion, 1.573, S1.361
 C_5H_5N Pyridine, 1.596, S1.391, 2.345, 3.649, 4.99
 C_5H_5NO 3-Hydroxypyridine, 3.463
 $C_5H_5NO_3$ *N*-Acetylalanine, 3.135
 $C_5H_5N_2O_4^-$ Hydroorotate ion, 1.472a, 3.316
 $C_5H_5N_3O_4$ 5-Nitro-6-methyluracil, S1.350, 3.575
 $C_5H_5N_5$ Adenine, 1.300, S1.105, 2.119, 3.146
 $C_5H_5N_5O$ Guanine, S1.274, 3.428; Isoguanine, 3.483a
 $C_5H_5O_4^-$ Methyl fumarate ion, S1.324
 C_5H_6I Iodobenzene, 3.474
 $C_5H_6N^+$ Pyridinium ion, 2.346, 3.656
 $C_5H_6N_2$ 2-Aminopyridine, 3.165; 4-Aminopyridine, 3.166
 $C_5H_6N_2O_2$ 6-Methyluracil, 1.539, 2.303; Thymine, 1.627, S1.418, 2.374, 3.711, 4.107
 $C_5H_6O_2$ Furfuryl alcohol, 3.391
 $C_5H_7NO_2$ Ethyl cyanoacetate, 1.418; *N*-Methylsuccinimide, S1.335
 $C_5H_7N_3O$ 1-Methylcytosine, S1.320; 5-Methylcytosine, 1.527, 3.533
 $C_5H_7O_2^-$ Cyclobutanecarboxylate ion, 3.276
 C_5H_8 Cyclopentene, 3.288
 $C_5H_8NO_4^-$ Glutamate ion, 1.440, 2.246
 $C_5H_8N_2O_2$ Dihydro-6-methyluracil, 3.315; 5,6-Dihydrothymine, 1.473a, 2.217, 3.317
 C_5H_8O 1,4-Pentadien-3-ol, 3.599, 4.89a
 $C_5H_8O_2$ Acetylacetone, 2.113; Cyclobutanecarboxylic acid, 2.193; Ethyl acrylate, S1.235; Methyl methacrylate, S1.332
 $C_5H_8O_4$ Glutaric acid, 3.401
 C_5H_9N Trimethylacetoneitrile, 2.379
 $C_5H_9NO_2$ Proline, 1.590-1, 2.330, 3.633
 $C_5H_9NO_3$ *N*-Acetylalanine, 1.293, 1.294, 2.116; *N*-Acetylglycine, methyl ester, S1.89; *N*-Acetylsarcosine, S1.96; Hydroxyproline, 1.482, 2.272, 3.461
 $C_5H_9NO_3S$ *N*-Acetylcysteine, S1.86
 $C_5H_9NO_4$ Glutamic acid, 2.247, 3.399
 $C_5H_9O_2^-$ 2-Methylbutyrate ion, 3.529; 3-Methylbutyrate ion (Isovalerate ion), 2.284, 3.530; Pentanoate ion (Valerate ion), 2.319, 3.752; Trimethylacetate ion (Pivalate ion), 2.378, 3.622
 $C_5H_9O_9P$ Ribose-5-phosphate, 3.665
 C_5H_{10} Cyclopentane, 2.200, 3.286
 $C_5H_{10}N_2O_3$ Alanine, 1.307, 3.154; Glutamine, 3.400; Glycylalanine, 1.447, 3.410; Glycylsarcosine, S1.272; Sarcosylglycine, S1.395
 $C_5H_{10}N_2O_4$ Glycylserine, 3.424
 $C_5H_{10}O$ 2-Pentanone, 3.603; 3-Pentanone, 3.604; Tetrahydropyran, 3.694
 $C_5H_{10}O_2$ Ethyl propionate, 3.376; Isopropylacetate, 3.486; Methyl butyrate, 3.528; 2-Methylbutyric acid, 2.302; 3-Methylbutyric acid (Isovaleric acid), 3.531; Propyl acetate, 3.644; Trimethylacetic acid (Pivalic acid), 1.588
 $C_5H_{10}O_4$ Deoxyribose, S1.205, 2.210, 3.299
 $C_5H_{10}O_5$ Arabinose, 1.315, 2.133; Ribose, 1.605, 2.349, 3.664; Xylose, 1.661
 $C_5H_{11}NO$ *N*-Methylisobutyramide, 3.543; Pivalamide, S1.380, 3.621
 $C_5H_{11}NO_2$ Norvaline, 2.312a, 3.587; Valine, 1.657-8, 2.387-8, 3.753
 $C_5H_{11}NO_2S$ 3-Mercaptovaline (Penicillamine), 1.517, S1.365, 2.317a, 3.596; Methionine, 1.522, S1.314, 2.298a, 3.513
 $C_5H_{11}NO_2Se$ Selenomethionine, 3.672a
 C_5H_{12} Pentane, 2.318
 $C_5H_{12}NO^+$ Betaine, 2.154
 $C_5H_{12}N_2O_2$ Ornithine, 3.589
 $C_5H_{12}O$ 2-Methyl-2-butanol, 3.526; 3-Methyl-1-butanol (Isobutanol), 3.527; Neopentyl alcohol, 2.305; 1-Pentanol, 3.602; 3-Pentanol, 3.602a
 $C_5H_{12}O_2$ Diethoxymethane, 3.306; 1,5-Pentanediol, 3.601
 $C_5H_{12}O_4$ Pentaerythritol, 3.598
 $C_5H_{13}N$ Amylamine, 1.313b, S1.121, 3.168, 4.44; Isoamylamine, 1.499a
 $C_5H_{14}N^+$ Amylammonium ion, S1.122, 3.169; Isoamylammonium ion, 3.480
 $C_5H_{16}CoN_4O_3^+$ Carbonatobis(ethylenediamine)cobalt(III) ion, 1.88, 2.44
 $C_5H_{20}CoN_6^{3+}$ Pentaamminepyridinecobalt (III) ion, 3.34a
 $C_5MnN_6O^{3-}$ Nitrosylpentacyanomanganate ion, 3.70
 $C_6CoN_6^{3-}$ Hexacyanocobaltate(III) ion, 1.76, S1.15, 2.35
 $C_6CoN_6S^{3-}$ Pentacyanothiocyanoatocobaltate(III) ion, S1.17
 $C_6CoO_3^{3-}$ Trioxalatocobaltate(III) ion, S1.17
 $C_6CrN_6^{3-}$ Hexacyanochromate(III) ion, 1.105
 $C_6CrN_6^{4-}$ Hexacyanochromate(II) ion, 1.100
 $C_6CrO_{12}^{3-}$ Trioxalatochromate(III) ion, S1.24, 2.51
 $C_6D_5NO_2$ Nitrobenzene-*d*₅, 3.566
 C_6D_6 Benzene-*d*₆, 3.187
 C_6F_6 Hexafluorobenzene, 1.465c, S1.279, 3.438
 $C_6FeN_6^{3-}$ Hexacyanoferrate(III) ion, 1.137, S1.30, 2.63, 5.45
 $C_6FeN_6^{4-}$ Hexacyanoferrate(II) ion, 1.134, 3.54, 3.55, 4.19, 5.15
 $C_6FeO_{12}^{3-}$ Trioxalatoferrate(III) ion, 2.64
 C_6HF_5 Pentafluorobenzene, 1.573a, S1.367, 3.600
 $C_6HFeN_6^{3-}$ Hydrogen hexacyanoferrate(II) ion, S1.29
 $C_6H_2Cl_2O_2$ 2,5-Dichloro-*p*-benzoquinone, 5.52
 $C_6H_2F_4$ 1,2,3,4-Tetrafluorobenzene, 1.633b, S1.411, 3.692
 $C_6H_2N_3O_7^-$ Picrate ion, 1.587
 $C_6H_3O_6^{3-}$ Aconitate ion, 1.297, 4.37
 $C_6H_4BrO^-$ *o*-Bromophenoxide ion, 1.341; *m*-Bromophenoxide ion, 1.342; *p*-Bromophenoxide ion, 1.344, S1.158
 $C_6H_4ClO^-$ *o*-Chlorophenoxide ion, 1.369; *m*-Chlorophenoxide ion, 1.370; *p*-Chlorophenoxide ion, 1.371

- $C_6H_4Cl_2$ *o*-Dichlorobenzene, 1.397; *m*-Dichlorobenzene, 1.398; *p*-Dichlorobenzene, 1.399
 $C_6H_4FO^-$ *o*-Fluorophenoxide ion, 1.429; *m*-Fluorophenoxide ion, 1.430; *p*-Fluorophenoxide ion, 1.431
 $C_6H_4F_2$ *o*-Difluorobenzene, S1.213, 3.313; *p*-Difluorobenzene, S1.214, 3.314
 $C_6H_4NO_2^-$ Isonicotinate ion, 1.500b; Nicotinate ion, 1.549, 3.560; Picolinate ion (2-Pyridinecarboxylate ion), 1.586a; 3-Pyridinecarboxylate ion, 3.653; 4-Pyridinecarboxylate ion, 3.654
 $C_6H_4NO_3^-$ *o*-Nitrophenoxide ion, 1.557; *m*-Nitrophenoxide ion, 1.558; *p*-Nitrophenoxide ion, 1.560
 $C_6H_4N_2$ 3-Pyridinenitrile, 3.655
 $C_6H_4O_2$ Benzoquinone, 1.330, S1.138, 2.152, 3.195, 5.47
 $C_6H_4O_4^{2-}$ Muconate ion, 4.84
 $C_6H_4O_8S_2^{2-}$ Tiron (1,2-Dihydroxybenzene-3,5-disulfonate ion), 5.64
 C_6H_5Br Bromobenzene, 1.336
 C_6H_5BrO *p*-Bromophenol, 1.343, S1.157, 2.163
 C_6H_5Cl Chlorobenzene, 1.360, 3.248
 C_6H_5ClO *m*-Chlorophenol, 3.253; *o*-Chlorophenol, 3.254
 C_6H_5F Fluorobenzene, 1.425, S1.243, 3.379
 C_6H_5I Iodobenzene, 1.489
 C_6H_5NO Nitrosobenzene, 1.563, 3.581
 $C_6H_5NO_2$ Nicotinic acid, 2.306, 3.561; Nitrobenzene, 1.551, S1.341, 2.308, 3.565, 4.87
 $C_6H_5NO_2^-$ Nitrobenzene anion, S1.342
 $C_6H_5NO_3$ *m*-Nitrophenol, 3.577; *o*-Nitrophenol, 3.577a; *p*-Nitrophenol, 1.559, S1.352, 3.577b
 $C_6H_5O^-$ Phenoxide ion, 1.576, S1.370, 3.608, 4.90
 $C_6H_5O_2^-$ *p*-Hydroxyphenoxide ion, 1.473
 $C_6H_5O_3S^-$ Benzenesulfonate ion, 1.326, 2.145, 3.189
 $C_6H_5O_7^{3-}$ Citrate ion, 4.57
 $C_6H_5S^-$ Thiophenoxide ion, 1.623
 C_6H_6 Benzene, 1.324, S1.132, 2.144, 3.186, 4.47
 $C_6H_6AlNO_6$ Nitrilotriacetatoaluminum(III), 1.19
 $C_6H_6AsO_3^-$ Phenylarsonate(V) ion, 1.581
 $C_6H_6NNiO_6^-$ Nitrilotriacetatonickelate(II) ion, 1.199
 $C_6H_6NO_3S^-$ Sulfanilate ion, 1.616, S1.406
 $C_6H_6NO_6^{3-}$ Nitrilotriacetate ion, 1.550
 $C_6H_6NO_6Zn^-$ Nitrilotriacetatozincate(II) ion, 1.281
 $C_6H_6N_2O$ Isonicotinamide, 1.500a; Nicotinamide, 1.546a, 3.558; 3-Pyridinecarboxamide, 3.651; 4-Pyridinecarboxamide, 3.652
 $C_6H_6N_4O_4$ 5-Nitro-2-furaldehyde semicarbazone, S1.345, 3.570
 C_6H_6O Phenol, 1.575, S1.369, 2.320, 3.607
 $C_6H_6O_2$ Hydroquinone, 3.446, 5.56; *m*-Hydroxyphenol, 3.456; *o*-Hydroxyphenol, 3.457
 $C_6H_6O_3S$ Benzenesulfonic acid, 3.190
 $C_6H_6O_4^{2-}$ 2-Hexene-1,6-dioate ion, 4.74; 3-Hexene-1,6-dioate ion, 4.75
 C_6H_6S Thiophenol, 2.369
 C_6H_7N Aniline, 1.314, S1.123, 2.128, 3.170, 4.45
 C_6H_7NO Phenylhydroxylamine, 1.582, 3.617
 $C_6H_7NO_2$ *N*-Ethylmaleimide, 1.421a, S1.239, 2.234, 3.374
 $C_6H_7NO_2S$ Benzenesulfonamide, 1.325, S1.133, 3.188
 $C_6H_7NO_3S$ Sulfanilic acid, 1.616a, S1.407, 3.687
 $C_6H_7N_3$ 2-Methyladenine, S1.316; 7-Methyladenine, S1.317
 $C_6H_7O_2^-$ Sorbate ion, S1.397
 $C_6H_7O_6^-$ Ascorbate ion, S1.127, 2.135, 3.178
 $C_6H_7O_7^-$ Citrate ion, 1.380; Isocitrate ion, 1.500
 C_6H_8 1,3-Cyclohexadiene, 1.384, 2.195, 3.280; 1,4-Cyclohexadiene, 1.385, 2.196, 3.281
 $C_6H_8CoN_2O_8^-$ Dioxalatoethylenediaminecobaltate(III) ion, 2.46
 C_6H_8N 2-Methylpyridine, 3.550; 3-Methylpyridine, 3.551
 $C_6H_8N^+$ Anilinium ion, 2.129, 3.171
 $C_6H_8NO_3^-$ *N*-Ethylmaleamate ion, S1.238
 $C_6H_8N_2$ *o*-Phenylenediamine, 2.325; *m*-Phenylenediamine, 2.326; *p*-Phenylenediamine, 2.327
 $C_6H_8N_2O_2$ 1,3-Dimethyluracil, 1.406; 1,6-Dimethyluracil, 1.407; 3,6-Dimethyluracil, 1.408; 4-Ethoxyuracil, 1.413
 $C_6H_8N_2O_2S$ Sulfanilamide, 1.615b, S1.405, 3.686
 $C_6H_8O_2$ Sorbic acid, S1.398
 $C_6H_8O_4$ Dimethyl fumarate, S1.219, 3.217a; Dimethyl maleate, S1.224
 $C_6H_8O_4^{2-}$ Adipate ion, 4.40
 $C_6H_8O_4S^{2-}$ 3,3'-Thiodipropionate ion, S1.415
 $C_6H_8O_4S_2^{2-}$ 2,2'-Dithiobispropionate ion, S1.230
 $C_6H_8O_6$ Ascorbic acid, 2.136, 3.179; *D*-Glucuronolactone, 3.398
 $C_6H_8O_7$ Citric acid, 2.191, 3.266
 C_6H_9NO *N*-Vinylpyrrolidone, S1.442
 $C_6H_9NO_3$ *N*-Ethylmaleamic acid, 2.233a, 3.373
 $C_6H_9NO_6$ Nitrilotriacetic acid, 2.307, 3.563
 $C_6H_9N_2O_4$ *N*-Acetylglycylglycine, S1.90, 3.140
 $C_6H_9N_3O_2$ Histidine, 1.466-8, 2.269, 3.442
 $C_6H_9N_3O_3$ 1-(2-Hydroxyethyl)-2-methyl-5-nitroimidazole, 3.453
 $C_6H_9O_2^-$ Cyclopentanecarboxylate ion, 3.287
 $C_6H_9O_7^-$ *D*-Glucuronate ion, 1.439a, S1.252, 3.396
 C_6H_{10} Cyclohexene, 1.387, 2.198, 3.284
 $C_6H_{10}N_2O_2$ Alanine anhydride, S1.112, 3.153; Sarcosine anhydride, S1.394, 3.670
 $C_6H_{10}N_3O_6$ Glycylasparagine, 1.448-9
 $C_6H_{10}O$ Cyclohexanone, 1.386; 2,4-Hexadien-1-ol, 3.437, 4.72
 $C_6H_{10}O_2$ 1-Cyclopentanecarboxylic acid, 2.201
 $C_6H_{10}O_3$ Ethyl acetoacetate, 2.228
 $C_6H_{10}O_4$ Adipic acid, 3.149
 $C_6H_{10}O_7$ Glucuronic acid, 3.397
 $C_6H_{11}N_3O_3$ *N*-Acetylglycylglycine amide, S1.191
 $C_6H_{11}N_3O_4$ Glycylglycylglycine, 1.453-1.455, S1.266-7, 2.255, 3.415-7
 $C_6H_{11}O_2^-$ Hexanoate ion, 2.264, 3.440, 4.73; 3,3-Dimethylbutyrate ion, 3.327
 C_6H_{12} Cyclohexane, 2.197
 $C_6H_{12}AlN_3O_6$ Tris(glycinato)aluminum(III), 1.18
 $C_6H_{12}CdN_3O_6^-$ Tris(glycinato)cadmate(II) ion, 1.45
 $C_6H_{12}CuN_2O_4^{2+}$ Bis(2-aminopropionato)copper(II) ion, 3.44; Bis(3-aminopropionato)copper(II) ion, 3.45
 $C_6H_{12}CuN_3O_6^-$ Tris(glycinato)cuprate(II) ion, 1.117
 $C_6H_{12}HgN_3O_6^-$ Tris(glycinato)mercurate(II) ion, 1.151
 $C_6H_{12}MnN_3O_6^-$ Tris(glycinato)manganate(II) ion, 1.171
 $C_6H_{12}N_2O_3$ Alanylalanine, 1.306; Glycylglycine, ethyl ester, S1.264
 $C_6H_{12}N_2O_4S_2$ Cystine, 1.393, 1.394, S1.196-7, 2.207, 3.291
 $C_6H_{12}N_2O_4Se_2$ Selenocystine, S1.396, 3.672
 $C_6H_{12}N_3NiO_6^-$ Tris(glycinato)nickelate(II) ion, 1.198
 $C_6H_{12}N_3O_6Pb^-$ Tris(glycinato)plumbate(II) ion, 1.216
 $C_6H_{12}N_3O_6Zn^-$ Tris(glycinato)zincate(II) ion, 1.283
 $C_6H_{12}N_4O_2$ *N,N,N',N'*-Tetramethyl-1,2-diazenedicarboxamide ('Diamide'), 3.696

$C_6H_{12}N_4O_3$ Glycylglycylglycine amide, S1.268
 $C_6H_{12}O$ Vinyl isobutyl ether, S1.441
 $C_6H_{12}O_2$ Cyclohexaneperoxy radical, 5.26, 5.49; Ethyl butyrate, 3.364; Hexanoic acid, 2.265; Methyl trimethylacetate (Methyl pivalate), 1.538
 $C_6H_{12}O_3$ 2,4,6-Trimethyl-1,3,5-trioxane, 3.731
 $C_6H_{12}O_5$ Methylarabinoside, 3.525
 $C_6H_{12}O_6$ Glucose, 1.439, 2.245, 3.394; Inositol, 3.473
 $C_6H_{13}N$ Cyclohexylamine, 1.387a; Hexamethyleneimine, 2.261
 $C_6H_{13}NO$ *N-tert*-Butylacetamide, S1.160, 3.230; *N,N*-Diethylacetamide, S1.210; *N*-Methylpivalamide, 3.544
 $C_6H_{13}NO_2$ Isoleucine, 2.282, 3.484; Leucine, 1.502, 2.286, 3.493-5; Norleucine, 1.566, 3.585
 $C_6H_{13}NO_5$ 2-Amino-2-deoxy- β -galactose, S1.118, 3.163; Glucosamine, 1.438
 $C_6H_{13}NO_8S$ 2-Deoxy-2-sulfoamino- β -glucose, S1.206, 3.300
 $C_6H_{13}O_9P$ Glucosephosphate, 3.395
 C_6H_{14} Hexane, 2.262
 $C_6H_{14}N^+$ Cyclohexylammonium ion, 3.285
 $C_6H_{14}N_2O_2$ Lysine, 1.508, 2.287, 3.497
 $C_6H_{14}N_4O_2$ Arginine, 1.316-8, 2.134, 3.177
 $C_6H_{14}O$ 1-Hexanol, 2.266, 3.441
 $C_6H_{14}O_2$ 1,1-Diethoxyethane, 3.305; 1,6-Hexanediol, 3.439; Pinacol, 3.620
 $C_6H_{14}O_6$ Sorbitol, 1.611
 $C_6H_{14}S_2$ Di-2-propyl disulfide, 3.345a
 $C_6H_{15}N$ Triethylamine, 3.723, 4.114
 $C_6H_{16}CoN_4O_4^+$ Oxalatobis(ethylenediamine)cobalt(III) ion, 2.45
 $C_6H_{16}CoN_6S_2^+$ Dithiocyanatobis(ethylenediamine)cobalt(III) ion, 1.92
 $C_6H_{16}CrN_6S_2^+$ Dithiocyanatobis(ethylenediamine)chromium(III) ion, 1.108
 $C_6H_{16}N^+$ 1-Hexylammonium ion, 2.267; Triethylammonium ion, 3.724
 $C_6H_{16}N_2$ 1,6-Hexanediamine, 2.263
 $C_6H_{16}N_6S_2$ Bis(2-guanidinoethyl)disulfide, 1.516
 $C_6H_{24}CdN_6^{2+}$ Tris(ethylenediamine)cadmium(II) ion, 1.50
 $C_6H_{24}CoN_6^{3+}$ Tris(ethylenediamine)cobalt(III) ion, 1.85, 2.40
 $C_6H_{24}CrN_6^{3+}$ Tris(ethylenediamine)chromium(III) ion, 1.106
 $C_6H_{24}CuN_6^{2+}$ Tris(ethylenediamine)copper(II) ion, 1.121
 $C_6H_{24}HgN_6^{2+}$ Tris(ethylenediamine)mercury(II) ion, 1.149
 $C_6H_{24}Ni_6^{2+}$ Tris(ethylenediamine)nickel(II) ion, 1.204
 $C_6H_{24}Pb_6^{2+}$ Tris(ethylenediamine)lead(II) ion, 1.219
 $C_6H_{24}Zn_6^{2+}$ Tris(ethylenediamine)zinc(II) ion, 1.278
 $C_6MnN_6^{4-}$ Hexacyanomanganate(II) ion, 1.174
 C_6N_4 Tetracyanoethylene, 1.617
 $C_6N_6Os^{4-}$ Hexacyanoosmate(II) ion, 1.207
 $C_6N_6Ru^{4-}$ Hexacyanoruthenate(II) ion, 1.231
 $C_7H_4BrO_2^-$ *p*-Bromobenzoate ion, 1.337, 3.209
 $C_7H_4ClO_2^-$ *o*-Chlorobenzoate ion, 1.361, 3.248a; *m*-Chlorobenzoate ion, 1.362, 3.248b; *p*-Chlorobenzoate ion, 1.363, 3.249
 $C_7H_4FO_2^-$ *o*-Fluorobenzoate ion, 1.426; *m*-Fluorobenzoate ion, 1.427; *p*-Fluorobenzoate ion, 1.428, 3.380
 $C_7H_4IO_2^-$ *o*-Iodobenzoate ion, 1.490, 3.475; *m*-Iodobenzoate ion, 1.491, 3.476; *p*-Iodobenzoate ion, 1.492, 3.477
 $C_7H_4NO^-$ *p*-Cyanophenoxide ion, 4.62
 $C_7H_4NO_4^-$ *p*-Nitrobenzoate ion, 3.567
 $C_7H_4O_3^{2-}$ Salicylate ion, dianion, 4.101
 $C_7H_5ClO_2$ *p*-Chlorobenzoic acid, 2.180
 $C_7H_5Cl_3$ α,α,α -Trichlorotoluene, 1.636
 $C_7H_5F_3$ α,α,α -Trifluorotoluene, 1.639
 C_7H_5N Benzonitrile, 1.328, 2.150, 3.193, 4.49
 C_7H_5NO *o*-Hydroxybenzonitrile, 1.477; *m*-Hydroxybenzonitrile, 1.478; *p*-Hydroxybenzonitrile, 1.479
 $C_7H_5NO_4$ *p*-Nitrobenzoic acid, 2.309
 $C_7H_5O_2^-$ Benzoate ion, 1.327, 2.148, 3.191, 4.48; Salicylaldehyde, anion, 4.100
 $C_7H_5O_3^-$ *m*-Hydroxybenzoate ion, 1.475; *p*-Hydroxybenzoate ion, 1.476, 3.449; Salicylate ion, 1.607, 2.350, 3.669
 $C_7H_6NO_2^-$ *p*-Aminobenzoate ion, 1.310, S1.116, 3.158
 $C_7H_6N_2$ *o*-Aminobenzonitrile, 1.311
 $C_7H_6N_4O_5$ Furamazone, S1.250, 3.389
 C_7H_6O Benzaldehyde, 2.142, 3.184
 $C_7H_6O_2$ Benzoic acid, 1.327a, S1.135, 2.149, 3.192; *p*-Hydroxybenzaldehyde, 3.448; Salicylaldehyde, 3.668a; 2-Methyl-*p*-benzoquinone, 5.57
 $C_7H_6O_3$ *p*-Hydroxybenzoic acid, 2.270
 $C_7H_7^+$ Tropylium ion, 2.381
 C_7H_7Br Benzyl bromide, S1.145
 C_7H_7Cl Benzyl chloride, 1.332, S1.146; *p*-Chlorotoluene, 1.377
 C_7H_7I *p*-Iodotoluene, 1.498
 C_7H_7N Vinylpyridine, 1.659
 C_7H_7NO Benzamide, 1.323, S1.131, 2.143, 3.185
 $C_7H_7NO_2$ *p*-Aminobenzoic acid, 3.159; Anthranilic acid (*o*-Aminobenzoic acid), 3.173; *p*-Nitrotoluene, 1.565, 4.88
 $C_7H_7NO_5S$ *p*-Nitro-*o*-toluenesulfonic acid, 3.583
 $C_7H_7O^-$ *o*-Methylphenoxide ion, 4.82a; *p*-Methylphenoxide ion, 4.82b
 $C_7H_7O_2^-$ *p*-Methoxyphenoxide ion, 4.81
 $C_7H_7O_3S^-$ *o*-Toluenesulfonate ion, 3.714; *m*-Toluenesulfonate ion, 3.715; *p*-Toluenesulfonate ion, 1.632, S1.420, 3.716
 C_7H_8 Cycloheptatriene, S1.191, 2.194, 3.279; Toluene, 1.631, 2.375, 3.713, 4.111
 $C_7H_8N^+$ Vinylpyridinium ion, 1.660
 C_7H_8O Anisole, 2.130, 3.172; Benzyl alcohol, 1.330, S1.144, 2.153, 3.196; *o*-Cresol, 3.268; *p*-Cresol, S1.186, 3.269; Hydroxycycloheptatriene, S1.285
 $C_7H_8O_2$ *o*-Methoxyphenol, 3.518; *p*-Methoxyphenol, 3.519
 C_7H_8S Benzyl mercaptan, S1.149
 C_7H_9N Benzylamine, 1.331a; *p*-Toluidine, S1.421, 4.113
 $C_7H_9N_2O$ 1-Methylnicotinamide, 1.535
 $C_7H_{10}N$ 2,4-Dimethylpyridine, 3.340; 2,6-Dimethylpyridine, 3.341
 $C_7H_{10}N^+$ Benzylammonium ion, 3.197
 $C_7H_{10}N_2O_2$ 4-Ethoxy-1-methyluracil, 1.412; 1,3,5-Trimethyluracil, 1.641
 $C_7H_{10}N_4O_2S$ Sulfaguanidine, S1.404, 3.685
 $C_7H_{11}O_2^-$ Cyclohexanecarboxylate ion, 3.282
 $C_7H_{12}N_2O_3$ Glycylproline, 1.459, 3.423
 $C_7H_{12}N_2O_4S_2^{2-}$ Djenkolate ion(3,3'-Methylenedithiobis(2-aminopropionate ion)), 1.409
 $C_7H_{12}O_4$ Diethyl malonate, 3.311; Pimelic acid, 3.619
 $C_7H_{13}N_3O_4$ β -Alanylglycylglycine, S1.113; Glycylglycyl- β -alanine, S1.265
 $C_7H_{14}N_2O_3$ Glycylvaline, 1.462, 3.426
 $C_7H_{14}N_2O_3S$ Glycylmethionine, 3.421
 $C_7H_{14}O$ Cycloheptanol, 3.277-8
 $C_7H_{14}O_6$ Methylgalactoside, 3.536; Methylglucoside, 3.537
 $C_7H_{15}NO$ *N,N*-Dimethylpivalamide, S1.226, 3.339

- $C_7H_{16}O$ 1-Heptanol, 3.435
 $C_7H_{20}CoN_5O_2^{2+}$ Benzoatopentaamminecobalt(III) ion, 2.39, 3.34
 $C_8H_4NO_2^-$ *p*-Cyanobenzoate ion, 1.383
 $C_8H_4N_2$ *o*-Dicyanobenzene, 2.213; *m*-Dicyanobenzene, 2.214; *p*-Dicyanobenzene, S1.209, 2.215, 3.304
 $C_8H_4O_4^{2-}$ *o*-Phthalate ion, 1.584, 3.618, 4.93; *m*-Phthalate ion, 1.585; *p*-Phthalate ion, 1.586, 3.690
 $C_8H_5O_4^-$ *o*-Phthalate ion, 1.583
 C_8H_6BrN 5-Bromoindole, 3.211
 C_8H_6ClN 5-Chloroindole, 3.252
 $C_8H_6ClO_2^-$ 2-Chloro-2-phenylacetate ion, S1.175
 $C_8H_6NO_4^-$ *p*-Nitrophenylacetate ion, 1.561
 $C_8H_6N_2O_2$ 5-Nitroindole, 3.572
 $C_8H_6N_4O_5$ Furadantin, S1.249, 3.387
 $C_8H_6O_4$ Phthalic acid, S1.379
 C_8H_7N Indole, 1.487, S1.289, 2.274a, 3.468; *p*-Tolunitrile, 1.633, 4.112
 C_8H_7NO 5-Hydroxyindole, 3.455
 C_8H_7NS Benzylthiocyanate, S1.153
 $C_8H_7N_3O_2$ Luminol, 3.496
 $C_8H_7O_2^-$ Phenylacetate ion, 1.577, S1.372, 2.322, 3.611, 4.92; *o*-Toluate ion, 1.628, 4.108; *m*-Toluate ion, 1.629, 4.109; *p*-Toluate ion, 1.630, 3.712, 4.110
 $C_8H_7O_3^-$ *p*-Methoxybenzoate ion, 3.515
 C_8H_8 Styrene, 1.612, 3.678
 $C_8H_8INO_3$ Iodotyrosine, S1.294
 $C_8H_8N_2$ 5-Aminoindole, 3.164
 $C_8H_8N_2O_3$ Nicotinuric acid, 1.549a, 3.562
 C_8H_8O Acetophenone, S1.82, 2.112, 3.134
 $C_8H_8O_2$ Benzyl formate, S1.147; 2,3-Dimethylbenzoquinone, 5.52a; 2,5-Dimethyl-*p*-benzoquinone, 5.53; 2,6-Dimethylbenzoquinone, 5.53a; Phenyl acetate, 2.321, 3.610; Phenylacetic acid, 2.323, 3.612
 C_8H_9BrO 1-(*p*-Bromophenyl)-1-ethanol, 3.212a
 C_8H_9Cl 1-Chloro-2-phenylethane, S1.176
 C_8H_9N Indoline, 3.472
 C_8H_9NO Acetanilide, 2.106, 3.127; Phenylacetamide, 3.609
 $C_8H_9NO_2$ Phenylglycine, S1.377
 C_8H_{10} *o*-Xylene, 3.755, 4.119; *m*-Xylene, 3.756, 4.120; *p*-Xylene, 3.757, 4.121
 $C_8H_{10}N_2O$ *p*-Nitrosodimethylaniline, 1.564, 3.582
 $C_8H_{10}N_2O_3S$ Sulfacetamide, 1.615a, 3.684
 $C_8H_{10}O$ Benzyl methyl ether, 3.198a; Phenethyl alcohol, 3.605; 1-Phenylethanol, 3.615b-c
 $C_8H_{10}O_2$ 1,2-Dimethoxybenzene, 3.320a; 1,3-Dimethoxybenzene, 3.320b; 1,4-Dimethoxybenzene, 3.320c
 $C_8H_{10}O_4$ *cis*-4-Cyclohexene-1,2-dicarboxylic acid, 2.199
 $C_8H_{11}N$ *N,N*-Dimethylaniline, 3.325; Phenethylamine, 1.574a
 $C_8H_{11}NO$ 4-Ethyl-5-hydroxy-2-methylpyridine, 3.375; 2,4,6-Trimethyl-3-hydroxypyridine, 3.728; Tyramine, S1.435, 3.740
 $C_8H_{11}NO_3$ Pyridoxine, 3.657a
 $C_8H_{12}N^+$ *N,N*-Dimethylanilinium ion, 3.326; Phenethylammonium ion, 3.606
 $C_8H_{12}NO_2$ Norpseudopelletierine *N*-oxyl, S1.356, 3.586
 $C_8H_{12}N_2O_2$ 2,4-Diethoxypyrimidine, 1.400
 $C_8H_{12}N_2O_3S$ 6-Aminopenicillanic acid, S1.119
 $C_8H_{13}N_3O_5$ *N*-Acetylglycylglycylglycine, S1.92
 $C_8H_{13}O_2S_2^-$ Lipoate ion, 1.507, S1.298
 $C_8H_{14}N_4O_5$ Glycylglycylglycylglycine, 3.418
 $C_8H_{14}O_2$ 2,5-Dihydroxy-2,5-dimethyl-3-hexyne, 3.319
 $C_8H_{14}O_4$ Diethylsuccinate, 3.312; Suberic acid, 3.679
 $C_8H_{15}NO_6$ 2-Acetamido-2-deoxy-*D*-galactose, S1.76, 3.125; 2-Acetamido-2-deoxy-*D*-glucose, 3.126; *N*-Acetylglucosamine, 3.138
 $C_8H_{15}N_5O_4$ Glycylglycylglycylglycine amide, S1.269
 $C_8H_{16}CuN_2O_4^{2+}$ Bis(2-aminobutyrate)copper(II) ion, 3.46; Bis(3-aminobutyrate)copper(II) ion, 3.47; Bis(4-aminobutyrate)copper(II) ion, 3.48; Bis(2-amino-2-methylpropionate)copper(II) ion, 3.49
 $C_8H_{16}N_2O_3$ Glycylisoleucine, 3.419; Glycylleucine, 1.456-7, 3.420; Leucylglycine, 1.504
 $C_8H_{16}N_2O_4S_2$ Cystine, dimethylester, S1.198; Homocystine, 1.470
 $C_8H_{18}O$ 1-Octanol, 3.588
 $C_8H_{18}O_3$ Diethyleneglycol, diethyl ether, 3.309
 $C_8H_{18}S_2$ Di-*tert*-butyl disulfide, 3.301a
 $C_8H_{19}CoN_5O_4^+$ Terephthatopentaamminecobalt(III) ion, 1.74
 $C_8H_{26}CoN_6^{3+}$ Bis(diethylenetriamine)cobalt(III) ion, 1.93
 $C_8H_{34}Co_2N_9O_2^{4+}$ Tetrakis(ethylenediamine)- μ -amidoperoxodico-balt(III) ion, 1.94
 $C_8MoN_8^{4-}$ Octacyanomolybdate(IV) ion, 1.176
 $C_9H_3O_6^{3-}$ Trimesate ion, 1.640
 $C_9H_6NO_2^-$ Indole-2-carboxylate ion, 1.487a; Indole-3-carboxylate ion, 1.487b; Indole-5-carboxylate ion, 1.487c
 $C_9H_6N_2$ 5-Cyanoindole, 3.273
 $C_9H_6O_6$ 1,3,5-Benzenetricarboxylic acid, 2.146
 $C_9H_7O_2^-$ Cinnamate ion, 1.379
 C_9H_8NO 1-(*p*-Cyanophenyl)-1-ethanol, 3.274
 $C_9H_8O_2$ Vinyl benzoate, S1.440
 $C_9H_8O_3^-$ *p*-Hydroxyphenylpropionate ion, 3.459
 C_9H_9N 1-Methylindole, 3.539; 2-Methylindole, 1.533, 3.540; 3-Methylindole, 1.534, 3.541; 5-Methylindole, 3.542
 C_9H_9NO Cinnamamide, S1.183; 5-Methoxyindole, 3.517
 $C_9H_9NO_3$ Hippuric acid, 2.268
 $C_9H_9NO_7$ 2-Nitro-2-furaldehyde, diacetate, S1.344
 $C_9H_9N_3O_2S_2$ Sulfathiazole, S1.409, 3.702
 $C_9H_9O_2^-$ Hydrocinnamate ion, 1.471
 $C_9H_9O_3^-$ *p*-Hydroxyphenylpropionate ion, 1.481, S1.287, 2.270a
 C_9H_{10} Allylbenzene, 4.42
 $C_9H_{10}N_2$ 5,6-Dimethylbenzimidazole, S1.216
 $C_9H_{10}O$ Phenylacetone, S1.373
 $C_9H_{10}O_2$ Benzyl acetate, S1.143; Hydrocinnamic acid, 1.472
 $C_9H_{10}O_3$ *p*-Hydroxyphenylpropionic acid, 2.271, 3.460
 $C_9H_{11}NO_2$ Phenylalanine, 1.578, 1.579, S1.374, 2.324, 3.613-5
 $C_9H_{11}NO_3$ Tyrosine, 1.645, 1.646, S1.436, 2.384, 3.741-5
 $C_9H_{11}NO_4$ 3-(3,4-Dihydroxyphenyl)alanine, 1.402
 $C_9H_{11}N_3O_7P^-$ Cytidine 2',3'-cyclicphosphate ion, S1.201
 C_9H_{12} 1,2,3-Trimethylbenzene (Hemimellitene), 3.727a, 4.115; 1,2,4-Trimethylbenzene (Pseudocumene), 3.727b, 4.116; 1,3,5-Trimethylbenzene (Mesitylene), 2.293, 3.727c, 4.117
 $C_9H_{12}N_2O$ Phenylalanine amide, S1.375
 $C_9H_{12}N_2O_6$ Uridine, 1.652-3, 3.750
 $C_9H_{12}N_3O_8P$ Cytidine 5'-phosphate(5'-Cytidylic acid), S1.200, 3.293
 $C_9H_{12}O$ 1-Phenyl-1-propanol, 3.617a; 1-Phenyl-2-propanol, 3.617b; 2-Phenyl-2-propanol, 3.617c
 $C_9H_{12}O_3$ 1,2,3-Trimethoxybenzene, 3.725a; 1,2,4-Trimethoxybenzene, 3.725b; 1,3,5-Trimethoxybenzene, 3.725c

- $C_9H_{13}N_2O_9P$ Uridine monophosphate(Uridylic acid), 1.654–6, 3.751
- $C_9H_{13}N_3O_5$ Cytidine, 1.395, S1.99, 3.292
- $C_9H_{14}N^+$ Trimethylanilinium ion, 2.380
- $C_9H_{14}N_3O_7P$ Deoxycytidylic acid, 3.297
- $C_9H_{16}NO_2$ 2,2,6,6-Tetramethyl-4-piperidone *N*-oxyl (TAN), S1.412, 2.363, 3.697, 4.106
- $C_9H_{16}O_4$ Azelaic acid, 3.182
- $C_9H_{18}N_2O_3$ Alanulleucine, 1.308; Leucylalanine, 1.503
- $C_{10}Co_2N_{10}O_2^{5-}$ Decacyano- μ -peroxodicobalt(III) ion, 1.95
- $C_{10}H_5O_5S^-$ 1,2-Naphthoquinone-2-sulfonate ion, S1.338, 5.59; 1,4-Naphthoquinone-2-sulfonate ion, S1.339, 2.304, 5.60
- $C_{10}H_6NO_2^-$ Quinoline-2-carboxylate ion, 1.602a
- $C_{10}H_6O_2$ 1,2-Naphthoquinone, 5.58
- $C_{10}H_7O^-$ 1-Naphthyl oxide ion, 1.543; 2-Naphthyl oxide ion, 1.544
- $C_{10}H_8$ Naphthalene, 1.540
- $C_{10}H_8NO_2^-$ Indole-3-acetate ion, S1.290
- $C_{10}H_8N_2$ 2,2'-Bipyridine, 1.334, 2.156, 3.206; 4,4'-Bipyridine, 1.334a, 2.157, 3.207
- $C_{10}H_8O_8S_2$ 4,5-Dihydroxy-2,7-naphthalenedisulfonic acid, 3.320
- $C_{10}H_9NO_2$ Indole-3-acetic acid, 2.274b, 3.469; Indole-5-acetic acid, 3.470
- $C_{10}H_9N_3$ Dipyrindylamine, 1.408c
- $C_{10}H_{11}N$ 1,2-Dimethylindole, 3.333; 1,3-Dimethylindole, 3.334; 2,3-Dimethylindole, 3.335
- $C_{10}H_{11}NO_3$ *N*-Acetylphenylglycine, S1.95
- $C_{10}H_{12}AgN_2O_8^{3-}$ Ethylenediaminetetraacetatoargentate(I) ion, 1.15
- $C_{10}H_{12}AlN_2O_8^-$ Ethylenediaminetetraacetatoaluminate(III) ion, 1.21
- $C_{10}H_{12}CdN_2O_8^{2-}$ Ethylenediaminetetraacetatocadmuate(II) ion, 1.47
- $C_{10}H_{12}CeN_2O_8^-$ Ethylenediaminetetraacetatocerate(III) ion, 1.52
- $C_{10}H_{12}CoN_2O_8^-$ Ethylenediaminetetraacetatocobaltate(III) ion, 1.84
- $C_{10}H_{12}CoN_2O_8^{2-}$ Ethylenediaminetetraacetatocobaltate(II) ion, 1.60
- $C_{10}H_{12}CrN_2O_8^-$ Ethylenediaminetetraacetatochromate(III) ion, 1.109
- $C_{10}H_{12}CuN_2O_8^{2-}$ Ethylenediaminetetraacetatocuprate(II) ion, 1.119, 3.50
- $C_{10}H_{12}DyN_2O_8^-$ Ethylenediaminetetraacetatodysprosate(III) ion, 1.124
- $C_{10}H_{12}ErN_2O_8^-$ Ethylenediaminetetraacetatoerbate(III) ion, 1.126
- $C_{10}H_{12}EuN_2O_8^-$ Ethylenediaminetetraacetatoeuropate(III) ion, 1.128
- $C_{10}H_{12}FeN_2O_8^{2-}$ Ethylenediaminetetraacetatoferrate(II) ion, 1.133
- $C_{10}H_{12}FeN_2O_8^-$ Ethylenediaminetetraacetatoferrate(III) ion, 1.139, 3.58
- $C_{10}H_{12}GaN_2O_8^-$ Ethylenediaminetetraacetatogallate(III) ion, 1.140
- $C_{10}H_{12}GdN_2O_8^-$ Ethylenediaminetetraacetatogadolinate(III) ion, 1.142
- $C_{10}H_{12}HgN_2O_8^{2-}$ Ethylenediaminetetraacetatomercurate(II) ion, 1.153
- $C_{10}H_{12}HoN_2O_8^-$ Ethylenediaminetetraacetatoholmate(III) ion, 1.155
- $C_{10}H_{12}InN_2O_8^-$ Ethylenediaminetetraacetatoindate(III) ion, 1.161
- $C_{10}H_{12}LaN_2O_8^-$ Ethylenediaminetetraacetatolanthanate(III) ion, 1.167
- $C_{10}H_{12}LuN_2O_8^-$ Ethylenediaminetetraacetatolutetate(III) ion, 1.169
- $C_{10}H_{12}MnN_2O_8^{2-}$ Ethylenediaminetetraacetatomanganate(II) ion, 1.173
- $C_{10}H_{12}N_2NdO_8^-$ Ethylenediaminetetraacetatoneodymate(III) ion, 1.192
- $C_{10}H_{12}N_2NiO_8^{2-}$ Ethylenediaminetetraacetatonickelate(II) ion, 1.201, 3.88
- $C_{10}H_{12}N_2O_4$ Thymine dimer, S1.419
- $C_{10}H_{12}N_2O_5S$ 7-Aminocephalosporanic acid, S1.117
- $C_{10}H_{12}N_2O_8$ Orotidine, 1.567b, 3.591
- $C_{10}H_{12}N_2O_8^{4-}$ Ethylenediaminetetraacetate ion, 1.420, 3.367
- $C_{10}H_{12}N_2O_8Pb^{2-}$ Ethylenediaminetetraacetatoplumbate(II) ion, 1.218
- $C_{10}H_{12}N_2O_8Pr^-$ Ethylenediaminetetraacetatopraseodymate(III) ion, 1.224
- $C_{10}H_{12}N_2O_8Sc^-$ Ethylenediaminetetraacetatoscandate(III) ion, 1.244
- $C_{10}H_{12}N_2O_8Sm^-$ Ethylenediaminetetraacetatosamarate(III) ion, 1.251
- $C_{10}H_{12}N_2O_8Sn^{2-}$ Ethylenediaminetetraacetatostannate(II) ion, 1.255
- $C_{10}H_{12}N_2O_8Tb^-$ Ethylenediaminetetraacetatoterbate(III) ion, 1.259
- $C_{10}H_{12}N_2O_8Ti^-$ Ethylenediaminetetraacetatotitanate(III) ion, 1.262
- $C_{10}H_{12}N_2O_8Tm^-$ Ethylenediaminetetraacetatothuliate(III) ion, 1.267
- $C_{10}H_{12}N_2O_8Y^-$ Ethylenediaminetetraacetatoyttrate(III) ion, 1.271
- $C_{10}H_{12}N_2O_8Yb^-$ Ethylenediaminetetraacetatoytterbate(III) ion, 1.273
- $C_{10}H_{12}N_2O_8Zn^{2-}$ Ethylenediaminetetraacetatozincate(II) ion, 1.280
- $C_{10}H_{12}O$ α -Tetralol, 3.695a
- $C_{10}H_{12}O_2$ Duroquinone, 5.55b
- $C_{10}H_{12}O_5$ Propylgallate, 3.648
- $C_{10}H_{13}N_5O_4$ Adenosine, 1.301, S1.108, 2.120, 3.147
- $C_{10}H_{13}N_5O_5$ Guanosine, 3.429
- $C_{10}H_{14}$ 1,2,3,4-Tetramethylbenzene (Prehnitine), 3.695a, 4.103; 1,2,3,5-Tetramethylbenzene (Isodurene), 3.695b, 4.104; 1,2,4,5-Tetramethylbenzene (Durene), 3.695c, 4.105
- $C_{10}H_{14}N_2O_4S$ Methylpenicillin, S1.334
- $C_{10}H_{14}N_2O_5$ Thymidine, 2.373, 3.709
- $C_{10}H_{14}N_5O_6P$ Deoxyadenylic acid, 3.296
- $C_{10}H_{14}N_5O_7P$ Deoxyguanylic acid, 3.298; Adenosine-5'-phosphate(Adenylic acid), 1.302, S1.109, 2.121, 3.148
- $C_{10}H_{14}N_5O_8P$ Guanylic acid, 3.430
- $C_{10}H_{14}O$ *p*-(*tert*-Butyl)phenol, 3.236; 1-(*p*-Ethylphenyl)-1-ethanol, 3.375a; 2-Methyl-1-phenyl-1-propanol, 3.453c-d; 2-Methyl-1-phenyl-2-propanol, 3.543e; 1-Phenyl-3-butanol, 3.615a
- $C_{10}H_{15}N_2O_8P$ Thymidylic acid, 1.626, 2.372, 3.710
- $C_{10}H_{16}N^+$ Benzyltrimethylammonium ion, 3.201
- $C_{10}H_{16}N_2$ *N, N, N', N'*-Tetramethyl-*p*-phenylenediamine, 2.362
- $C_{10}H_{16}N_2O_8$ Ethylenediaminetetraacetic acid, 2.230, 3.368
- $C_{10}H_{17}N_3O_6S$ Glutathione, reduced, 1.441, S1.254, 2.248, 3.402

- $C_{10}H_{18}N_2O_7$ 2-Hydroxyethylethylenediaminetriacetic acid, 3.452
 $C_{10}H_{18}O_4$ Sebacic acid, 3.671
 $C_{10}H_{19}N_3O_4$ Leucylglycylglycine, 1.505, 1.506
 $C_{10}H_{20}N_2O_4S_2$ Penicillamine disulfide, S1.366, 3.597
 $C_{11}H_7N$ 1-Naphthonitrile, 1.545; 2-Naphthonitrile, 1.546
 $C_{11}H_7O_2^-$ 1-Naphthoate ion, 1.541, 3.556, 4.85; 2-Naphthoate ion, 1.542, 3.557, 4.86
 $C_{11}H_8O_2$ 2-Methyl-1,4-naphthoquinone, (Menaquinone), S1.333, 3.505
 $C_{11}H_{10}NO_2^-$ Indole-3-propionate ion, S1.291
 $C_{11}H_{11}NO_2$ Indole-3-propionic acid, 2.274c, 3.471
 $C_{11}H_{12}ClNO_3$ *N*-(2-Chloroacetyl)phenylalanine, S1.172
 $C_{11}H_{12}N_2O_2$ Tryptophan, 1.643, 1.644, S1.434, 2.382-3, 3.735-8
 $C_{11}H_{13}NO_3$ *N*-Acetylphenylalanine, S1.93
 $C_{11}H_{14}N_2O_2$ *N*-Acetylphenylalanine amide, S1.94
 $C_{11}H_{14}N_2O_3$ Glycylphenylalanine, 1.458, S1.270, 3.422
 $C_{11}H_{14}N_2O_4$ Glycyltyrosine, 1.461, 3.425
 $C_{11}H_{16}$ Pentamethylbenzene, 3.600a, 4.89b
 $C_{11}H_{16}O$ 2,2-Dimethyl-1-phenyl-1-propanol, 3.337a; 1-Methoxy-2-methyl-1-phenylpropane, 3.517a; 2-Methyl-4-phenyl-2-butanol, 3.543a
 $C_{11}H_{19}N_3O_5$ *N*-Acetylalanylalanylalanine, S1.83, 3.136; *N*-Acetylsarcosylsarcosylsarcosine, S1.97
 $C_{12}H_8N_2$ 1,10-Phenanthroline, 1.574
 $C_{12}H_8O_2$ Diphenoquinone, 5.55
 $C_{12}H_9NO$ 2-Benzoylpyridine, S1.140, 2.152a; 3-Benzoylpyridine, S1.141, 2.152b; 4-Benzoylpyridine, S1.142, 2.152c
 $C_{12}H_{10}$ Biphenyl, 3.202a
 $C_{12}H_{10}O_2$ 2,3-Dimethylnaphthoquinone, 5.54a; 1-Naphthaleneacetic acid, 2.303a, 3.555
 $C_{12}H_{11}N$ Diphenylamine, 3.345
 $C_{12}H_{12}AgN_2O_{12}^{5-}$ Bis(nitritotriacetato)argentate(I) ion, 1.14
 $C_{12}H_{12}AlN_2O_{12}^{3-}$ Bis(nitritotriacetato)aluminate(III) ion, 1.20
 $C_{12}H_{12}CdN_2O_{12}^{4-}$ Bis(nitritotriacetato)cadmate(II) ion, 1.46
 $C_{12}H_{12}CoN_2O_{12}^{4-}$ Bis(nitritotriacetato)cobaltate(II) ion, 1.59a
 $C_{12}H_{12}CuN_2O_{12}^{4-}$ Bis(nitritotriacetato)cuprate(II) ion, 1.118
 $C_{12}H_{12}HgN_2O_{12}^{4-}$ Bis(nitritotriacetato)mercurate(II) ion, 1.152
 $C_{12}H_{12}MnN_2O_{12}^{4-}$ Bis(nitritotriacetato)manganate(II) ion, 1.172
 $C_{12}H_{12}N_2NiO_{12}^{4-}$ Bis(nitritotriacetato)nickelate(II) ion, 1.200
 $C_{12}H_{12}N_2O_2S$ Sulfanilamide, 1.615b
 $C_{12}H_{12}N_2O_{12}Pb^{4-}$ Bis(nitritotriacetato)plumbate(II) ion, 1.217
 $C_{12}H_{12}N_2O_{12}Zn^{4-}$ Bis(nitritotriacetato)zincate(II) ion, 1.282
 $C_{12}H_{13}Cl_3O_6$ 2,4,6-Trichlorophenyl- β -D-glucopyranoside, 3.722
 $C_{12}H_{14}Cl_2N_2$ 4,4'-Dimethyl-1,1'-bipyridylum chloride, S1.217, 5.53
 $C_{12}H_{15}BrO_6$ *m*-Bromophenyl- β -D-glucopyranoside, 3.213
 $C_{12}H_{15}ClO_6$ *m*-Chlorophenyl- β -D-glucopyranoside, 3.255; *p*-Chlorophenyl- β -D-glucopyranoside, S1.177, 3.256
 $C_{12}H_{15}NO_8$ *o*-Nitrophenyl- β -D-glucopyranoside, S1.353, 3.578; *m*-Nitrophenyl- β -D-glucopyranoside, 3.579; *p*-Nitrophenyl- β -D-glucopyranoside, S1.354, 3.580
 $C_{12}H_{16}N_6O_3$ Histidylhistidine, 1.469, 3.443
 $C_{12}H_{16}O_6$ Phenyl- β -D-glucopyranoside, S1.378, 2.328, 3.616
 $C_{12}H_{16}O_7$ *p*-Hydroxyphenyl- β -D-glucopyranoside, S1.286, 3.458
 $C_{12}H_{18}$ Hexamethylbenzene, 3.438a, 4.72a
 $C_{12}H_{18}O$ 2-Methyl-5-phenyl-2-pentanol, 3.543b
 $C_{12}H_{22}O_{11}$ Cellobiose, 3.244; Lactose, 3.492; Melibiose, 3.504; Sucrose, 2.356, 3.683
 $C_{12}H_{24}N_2O_3$ Leucylleucine, 1.507
 $C_{12}H_{25}NaO_4S$ Dodecyl sodium sulfate, 1.409a, S1.232, 2.221, 3.349
 $C_{12}H_{27}O_4P$ Tributyl phosphate, 3.720a
 $C_{12}H_{33}ClN_3Pd^+$ Chloro-1,1,7,7-tetraethyldiethylenetriamine-palladium(II) ion, 1.222
 $C_{12}H_{33}ClN_3Pt^+$ Chloro-1,1,7,7-tetraethyldiethylenetriamine-platinum(II) ion, 1.227
 $C_{13}H_8O$ Fluorenone, S1.241, 2.235
 $C_{13}H_9O_2^-$ Biphenyl-4-carboxylate ion, 1.333a, 3.203, 4.50
 $C_{13}H_9O_3^-$ *p*-Phenoxybenzoate ion, 3.608a, 4.91
 $C_{13}H_{10}O$ Benzophenone, 1.329, S1.137, 2.151, 3.194
 $C_{13}H_{12}NO^+$ 3-Benzoyl-*N*-methylpyridinium ion, S1.139
 $C_{13}H_{12}N_3^+$ Proflavine, 3.632a
 $C_{13}H_{13}N_3O_5S_2$ Sulfasuccidine, S1.408, 3.701
 $C_{13}H_{15}NO_6$ *p*-Cyanophenyl- β -D-glucopyranoside, 3.275
 $C_{13}H_{15}N_3O_3$ Glycyltryptophan, 1.460
 $C_{13}H_{17}N_3O_4$ Glycylphenylalanylglycine, S1.271; Phenylalanylglycylglycine, S1.376
 $C_{13}H_{18}O_5S$ *p*-Tolyl-*S*- β -D-thioglycopyranoside, S1.425, 3.720
 $C_{13}H_{18}O_6$ β -Benzylglucoside, S1.148, 3.198; *o*-Tolyl- β -D-glucopyranoside, S1.422, 3.717; *m*-Tolyl- β -D-glucopyranoside, S1.423, 3.718; *p*-Tolyl- β -D-glycopyranoside, S1.424, 2.376, 3.719
 $C_{13}H_{18}O_7$ *p*-Methoxyphenyl- β -D-glucopyranoside, 3.520
 $C_{14}H_7O_5S^-$ 9,10-Anthraquinone-1-sulfonate ion, S1.124, 3.174; 9,10-Anthraquinone-2-sulfonate ion, S1.125, 2.132, 3.175
 $C_{14}H_8O_4^{2-}$ 2,2'-Biphenyldicarboxylate ion (Diphenate ion), 1.408a, 3.204, 4.51; 4,4'-Biphenyldicarboxylate ion, 1.408a, 3.205, 4.52
 $C_{14}H_{10}$ Anthracene, 2.131
 $C_{14}H_{10}O$ Anthrone, S1.126
 $C_{14}H_{10}O_2$ Benzil, S1.134, 2.147
 $C_{14}H_{11}O_2^-$ Diphenylacetate ion, 3.344, 4.64
 $C_{14}H_{12}O_2$ Benzoin, S1.136
 $C_{14}H_{14}ClN_3$ Acriflavin, 1.298a, S1.98, 3.141
 $C_{14}H_{20}O_6$ 2,3-Dimethylphenyl- β -D-glucopyranoside, S1.225, 3.336; 3,4-Dimethylphenyl- β -D-glucopyranoside, 3.337
 $C_{14}H_{22}O_8$ *trans*-1,2-Cyclohexanediaminetetraacetic acid, 3.283
 $C_{14}H_{23}N_3O_{10}$ Diethylenetriaminepentaacetic acid, 3.310
 $C_{15}H_9O_2^-$ 9-Anthroate ion, 3.176, 4.46
 $C_{15}H_{14}N_2O_6S_2$ Cephalothin, S1.169
 $C_{15}H_{14}O$ 1,3-Diphenylacetone, S1.228
 $C_{15}H_{20}N_4O_6$ Riboflavin, 1.603
 $C_{15}H_{22}O_6$ 2,4,5-Trimethylphenyl- β -D-glucopyranoside, 3.729
 $C_{15}H_{23}N_3O_{10}$ Glutamylglutamylglutamic acid, S1.253
 $C_{15}H_{24}CoO_6^{3+}$ Tris(acetylacetonato)cobalt(III) ion, 1.98, 2.48, 3.36

- $C_{16}H_6N_2O_{14}S_4^{4-}$ Indigotetrasulfonate ion, 1.486, 5.31
 $C_{16}H_7N_2O_{11}S_3^{3-}$ Indigotrisulfonate ion, 5.30
 $C_{16}H_8N_2O_8S_2^{2-}$ Indigodisulfonate ion, 5.29
 $C_{16}H_{10}$ Pyrene, S1.390
 $C_{16}H_{14}N_2O_6S$ Thalamyd, S1.413, 3.700
 $C_{16}H_{18}ClN_3S$ Methylene blue, 1.528, 3.534
 $C_{16}H_{18}N_2O_4S$ Benzylpenicillin, S1.150, 3.199
 $C_{16}H_{18}N_2O_5S$ Phenoxymethylpenicillin, S1.371
 $C_{16}H_{19}N_3O_4S$ Ampicillin, S1.120
 $C_{16}H_{20}N_2O_5S$ Benzylpenicilloic acid, S1.152, 3.200
 $C_{16}H_{21}N_3O_8S$ Cephalosporin C, S1.168
 $C_{17}H_{18}N_2O_6S$ Carbenicillin, S1.163
 $C_{17}H_{20}ClN_3$ Acridine orange, 1.298
 $C_{17}H_{20}N_2O_4S$ Benzylpenicillin, methyl ester, S1.151
 $C_{17}H_{20}N_2O_6S$ Methicillin, S1.313
 $C_{18}H_{11}N_5O_9S$ *p*-Sulfodiphenylpicrylhydrazyl, S1.410
 $C_{18}H_{16}N_3O_4S_2$ Cephaloridine, S1.167
 $C_{18}H_{20}N_2O_3$ Phenylalanylphenylalanine, 1.580
 $C_{18}H_{22}N_2O_4S$ Phenethicillin, S1.368
 $C_{18}H_{31}O_2$ Oleate ion, S1.357
 $C_{18}H_{35}O_2$ Stearate ion, S1.399
 $C_{19}H_{18}ClN_3O_5S$ Cloxacillin, S1.184
 $C_{19}H_{22}N_2O_6S$ Penamceillin, S1.364
 $C_{19}H_{42}BrN$ Hexadecyltrimethylammonium bromide, 1.465a, S1.278, 2.260, 3.436
 $C_{20}H_6Br_4O_5^{2-}$ Eosin (Tetrabromofluorescein), 1.410, 3.351
 $C_{20}H_6I_4O_5^{2-}$ Erythrosin (Tetraiodofluorescein), S1.233, 2.361
 $C_{20}H_{11}O_5^-$ Fluorescein (anion), 1.422
 $C_{20}H_{12}O_5$ Fluorescein, S1.242, 3.377
 $C_{20}H_{19}ClN_4$ Safranin T, 1.577, 3.667, 3.668
 $C_{20}H_{32}N_6O_{12}S_2$ Glutathione, oxidized (disulfide), 1.442, S1.255
 $C_{20}H_{34}N_6O_8$ *N*-Acetylalanylalanylalanylalanylalanylalanine, S1.84
 $C_{21}H_{18}O_5S$ Cresol red, S1.187
 $C_{21}H_{27}FO_6$ Triamcinolone, S1.426
 $C_{21}H_{28}N_7O_{10}P_2$ Nicotinamide-adenine dinucleotide, 1.547, 1.548, 5.61
 $C_{21}H_{30}O_5$ Hydrocortisone, S1.282
 $C_{21}H_{38}ClN$ Hexadecylpyridinium chloride, 1.465b
 $C_{23}H_{32}O_6$ Hydrocortisone acetate, S1.283
 $C_{24}H_{30}F_2O_6$ Fluocinolone acetate, S1.240
 $C_{24}H_{31}FO_6$ Triamcinolone acetonide, S1.427
 $C_{26}H_{35}FO_6$ β -Methazone valerate, S1.311
 $C_{28}H_{31}ClN_2O_3$ Rhodamine B, S1.392, 3.662
 $C_{30}H_{24}CoN_6^{3+}$ Tris(2,2'-bipyridine)cobalt(III) ion, 1.96
 $C_{30}H_{24}FeN_6^{3+}$ Tris(2,2'-bipyridine)iron(III) ion, 2.65
 $C_{30}H_{24}N_6Rh^{3+}$ Tris(2,2'-bipyridine)rhodium(III) ion, 1.230
 $C_{30}H_{24}N_6Ru^{2+}$ Tris(2,2'-bipyridine)ruthenium(II) ion, S1.54
 $C_{30}H_{24}N_6Ru^{3+}$ Tris(2,2'-bipyridine)ruthenium(III) ion, S1.60, 2.93a
 $C_{30}H_{32}N_2O_{10}S$ Xylenol orange, S1.443, 3.758
 $C_{31}H_{46}O_2$ Vitamin K₁, 5.65
 $C_{32}H_{16}CuN_8O_{12}S_4$ Tetrasulfonated Cu phthalocyanine, 3.698
 $C_{34}H_{32}ClFeN_4O_4$ Hemin, 1.464a, 3.431
 $C_{36}H_{24}CoN_6^{3+}$ Tris(1,10-phenanthroline)cobalt(III) ion, 1.97
 $C_{36}H_{24}FeN_6^{3+}$ Tris(1,10-phenanthroline)iron(III) ion, 2.66
 $C_{45}H_{33}CoN_9^{3+}$ Tris(2,2',6',2''-terpyridine)cobalt(III) ion, S1.22
 $C_{63}H_{90}CoN_{14}O_{14}P$ Cyanocobalamin, S1.190, 3.272a, 5.48
 Cd^{+} 3.25b
 Cd^{2+} Cadmium (II) ion, 1.38, S1.10, 2.19, 3.25a
 $CdClH_6O_3^{+}$ Chlorotriaquocadmium(II) ion, 1.40
 $CdH_6IO_3^{+}$ Iodotriaquocadmium(II) ion, 1.41
 $CdH_{12}N_4^{2+}$ Tetraamminecadmium(II) ion, 1.39
 Ce^{3+} Cerium(III) ion, 1.51, 3.26, 4.14, 5.10
 Ce^{4+} Cerium(IV) ion, 2.21, 5.11
 Cl^- Chloride ion, 1.53, 2.21a, 3.27, 3.28
 $ClCoH_{15}N_5^{2+}$ Chloropentaamminecobalt(III) ion, 1.66
 $ClCrH_{15}N_5^{2+}$ Chloropentaamminechromium(III) ion, 1.103, 2.49
 $ClFe^{2+}$ Chloroiron(III) ion, 2.61
 $ClH_{15}N_5Ru^{2+}$ Chloropentaammineruthenium(III) ion, 1.233, S1.57
 $ClHg$ Mercury(I) chloride, 3.65
 ClO^- Hypochlorite ion, 1.54, S1.11, 3.29, 4.15
 ClO_2 Chlorine dioxide, 3.32, 5.41
 ClO_2^- Chlorite ion, S1.12, 3.30, 4.16
 ClO_3^- Chlorate ion, 1.55, S1.13, 3.31, 4.17
 ClO_4^- Perchlorate ion, 1.56
 $Cl_2CrH_8O_4^{+}$ Dichlorotetraaquochromium(III) ion, 2.50
 Cl_2Fe^{+} Dichloroiron(III) ion, 2.62
 Cl_2Hg Mercury(II) chloride, S1.33
 Cl_4Pd^{2-} Tetrachloropalladate(II) ion, 1.220, 3.97
 Cl_4Pt^{2-} Tetrachloroplatinate(II) ion, 1.225, S1.52, 3.99
 Cl_6Ir^{2-} Hexachloroiridate(IV) ion, 1.164
 Cl_6Ir^{3-} Hexachloroiridate(III) ion, 1.162
 Cl_6Pt^{2-} Hexachloroplatinate(IV) ion, 1.228
 Co^{2+} Cobalt(II) ion, 1.57, 3.32a
 $CoBrH_{15}N_5^{2+}$ Bromopentaamminecobalt(III) ion, 2.27
 $CoClH_{15}N_5^{2+}$ Chloropentaamminecobalt(III) ion, 2.26
 $CoFH_{15}N_5^{2+}$ Fluoropentaamminecobalt(III) ion, 1.65, 2.25
 $CoH_9N_6O_6$ Trinitrotrisamminecobalt(III), 2.34
 $CoH_{15}IN_5^{2+}$ Iodopentaamminecobalt(III) ion, 2.28
 $CoH_{15}N_5O_4P$ Phosphatopentaamminecobalt(III), 2.37
 $CoH_{15}N_6O_2^{2+}$ Nitropentaamminecobalt(III) ion, 2.33
 $CoH_{15}N_8^{2+}$ Azidopentaamminecobalt(III) ion, 1.70, 2.31
 $CoH_{16}N_4O_3^{3+}$ Diaquotetraamminecobalt(III) ion, 1.63
 $CoH_{16}N_5O_2^{2+}$ Hydroxopentaamminecobalt(III) ion, 1.64, 2.24
 $CoH_{17}N_5O_3^{3+}$ Aquopentaamminecobalt(III) ion, 1.62, 2.23
 $CoH_{18}N_6^{3+}$ Hexaamminecobalt(III) ion, 1.61, 2.22, 3.33
 $CoN_6O_{12}^{3-}$ Hexanitrocobaltate(III) ion, 1.81
 CoO_2^{2-} Cobaltate(II) ion, 1.58
 $Co_2H_{30}N_{10}O_2^{5+}$ Decaammine- μ -dioxodicobalt(III) ion, 1.75
 Cr^{2+} Chromium(II) ion, 1.99, 3.37
 Cr^{3+} Chromium(III) ion, 1.102, 3.38
 $Cr(V)$, 3.40
 CrF_6^{3-} Hexafluorochromate(III) ion, 1.104
 CrF_6^{4-} Hexafluorochromate(II) ion, 1.101
 CrO_4^{2-} Chromate(VI) ion, 1.112, 2.52
 $Cr_2O_7^{2-}$ Dichromate(VI) ion, 1.113, 2.53
 $Cr_4O_{12}^{3-}$ Trichromatochromate(III) ion, 1.114
 Cu^{+} , 5.12, 5.42
 Cu^{2+} Copper(II) ion, 1.115, S1.25, 2.54, 3.41, 5.13, 5.43
 $CuH_4O_4^{2-}$ Tetrahydroocuprate(II) ion, 1.116
 $CuH_{12}N_4^{2+}$ Tetraamminecopper(II) ion, 1.120
 D Deuterium atom, 1.6, S1.4; see also part II (75-0001)
 D^{+} Deuteron, 1.144
 DO , 1.8; see also part III, tables 2-4.
 D_2 Deuterium, 2.68, 3.60, 3.61
 D_2O Deuterium oxide, 1.2
 D_2O_2 Deuterium peroxide, 1.147
 D_2S Deuterium sulfide, 1.235, 2.93c
 Dy^{3+} Dysprosium(III) ion, 1.123
 Er^{3+} Erbium(III) ion, 1.125
 Eu^{2+} Europium(II) ion, 3.51
 Eu^{3+} Europium(III) ion, 1.127, S1.26

- F^- Fluoride ion, 1.129, 2.55
 FFe^{2+} Fluoroiron(III) ion, 1.230
 FH Hydrofluoric acid, 1.130
 $FH_6NiO_3^+$ Fluorotriaquonickel(II) ion, 1.194
 F_2Fe^+ Difluoroiron(III) ion, 2.60
 F_2H^- , 1.131
 F_3Sn^- Trifluorostannate(II) ion, 1.253, 2.97
 F_6Fe^{3-} Hexafluoroferrate(III) ion, 1.136
 F_6S Sulfur hexafluoride, 1.237, S1.62
 F_6Si^{2-} Hexafluorosilicate(IV) ion, 1.249
 F_6Sn^{2-} Hexafluorostannate(IV) ion, 1.257, 2.99
 F_6Ti^{2-} Hexafluorotitanate(IV) ion, 1.264
 Fe^{2+} Iron(II) ion, 1.132, 2.56, 3.52, 3.53, 4.18, 5.14
 Fe^{3+} Iron(III) ion, S1.27, 2.57, 3.56, 5.16, 5.44
 FeH^{2+} Hydroiron(III) ion, 2.59
 $FeHO^{2+}$ Hydroxoiron(III) ion, 2.58
 FeO_2^- , 4.20
 FeO_4^{2-} , 4.20
 FeO_4S^+ Sulfatoiron(III) ion, S1.28
 Gd^{3+} Gadolinium(III) ion, 1.141
 H Hydrogen atom, 1.5, S1.3; see also part II (75-0001)
 H^+ , 1.143, S1.31, 2.67, 5.17
 HNO_2 Nitrous acid, 2.84
 HNO_3 Nitric acid, 3.84
 $HNO_7S_2^-$ Hydroxylaminedisulfonate ion, 1.185, 3.78
 HO Hydroxyl radical, 1.7, 2.3; see also part III, tables 2-4.
 HO^- Hydroxide ion, 2.90, 3.62
 $HOZn^+$ Hydroxozinc(II) ion, 1.275
 HO_2 Perhydroxyl radical, 2.4, 3.5; see also part III, table 6.
 HO_2^- Hydroperoxide ion, 1.148, 3.63, 4.22
 HO_3P^{3-} Hydrogenphosphite ion, 2.91
 HO_3S^- Bisulfite ion, S1.64, 3.104
 HO_4P^{2-} Hydrogenphosphate ion, S1.49, 2.92, 3.92, 4.31
 HO_4S^- Bisulfate ion, 3.106
 HO_5S^- Peroxysulfate ion, 1.241, 3.108
 $HO_7P_2^{3-}$ Pyrophosphate ion, S1.50
 HS^- Bisulfide ion, 1.236, S1.61, 3.103
 HSe^- Hydroselenide ion, 1.246, 3.111
 H_2 , 1.145, 3.59, 4.21
 H_2N Amide radical, 3.72
 $H_2NO_3S^-$ Sulfamate ion, 1.183
 H_2O Water, 1.1, 4.1
 H_2O_2 Hydrogen peroxide, 1.146, S1.32, 2.69, 3.64, 4.23, 5.18, 5.46
 $H_2O_2^+$, 3.6
 $H_2O_2P^-$ Hypophosphite(III) ion, 1.209, 3.95
 $H_2O_3P^-$ Phosphite ion, 1.210
 H_2O_3Te Telluric(IV) acid, 3.115, 5.21
 $H_2O_4P^-$ Dihydrogenphosphate ion, 1.211, S1.48, 2.91b, 3.91
 H_2O_5S Peroxysulfuric acid, 2.95
 H_2S Hydrogen sulfide, 1.234, 3.102
 H_2Se Hydrogen selenide, 1.245, 3.110
 H_3N Ammonia, 2.80, 3.71
 H_3NO Hydroxylamine, 1.181, S1.40, 3.74
 H_3O_4P Phosphoric acid, 2.91a, 3.90
 H_4N^+ Ammonium ion, 1.178, 2.78
 H_4NO^+ Hydroxylammonium ion, 1.182, S1.41, 3.75
 H_4N_2 Hydrazine, 1.179, S1.38, 3.76
 $H_4O_4Zn^{2-}$ Tetrahydroxozincate(II) ion, 1.276
 H_5N^+ Hydrazinium ion, 1.180, S1.39, 2.79, 3.77
 $H_{12}N_4Zn^{2+}$ Tetraamminezinc(II) ion, 1.277
 $H_{15}IN_5Ru^{2+}$ Iodopentaammineruthenium(III) ion, S1.58
 $H_{15}N_7Ru^{2+}$ Pentaamminenitrogenoruthenium(III) ion, 1.231a, S1.55, 3.301
 $H_{16}N_5ORu^{2+}$ Hydroxopentaammineruthenium(III) ion, S1.59
 $H_{18}IrN_6^{3+}$ Hexaammineiridium(III) ion, 1.163
 $H_{18}N_6Os^{3+}$ Hexaammineosmium(III) ion, 1.208
 $H_{18}N_6Rh^{3+}$ Hexaamminerhodium(III) ion, 1.229
 $H_{18}N_6Ru^{3+}$ Hexaammineruthenium(III) ion, 1.232
 Hg^{2+} , 2.70
 Hg_2^{2+} , 2.70a
 Ho^{2+} Holmium(III) ion, 1.154
 I^- Iodide ion, S1.34, 2.72, 3.66, 4.24
 IO^- Hypoiodite ion, 4.25
 IO_3^- Iodate ion, 1.158, 2.75, 3.67, 4.26
 IO_4^- Periodate ion, 1.159, 3.68
 I_2 Iodine, 1.156, S1.35, 2.71
 I_2^- , 2.73
 I_3^- , 1.157, 2.74
 In^{3+} Indium(III) ion, 1.160
 K^+ Potassium(I) ion, 1.165, S1.36
 La^{3+} Lanthanum(III) ion, 1.166
 Lu^{3+} Lutetium(III) ion, 1.168
 Mn^{2+} Manganese(II) ion, 1.170, 2.76, 3.69
 MnO_4^- Permanganate ion, 1.175, 2.77, 5.19
 NO Nitric oxide, 1.187, 2.82, 3.80
 NO_2 Nitrogen dioxide, 3.81
 NO_2^- Nitrite ion, 1.188, S1.43, 2.85, 3.82, 4.27
 NO_3^- Nitrate ion, 1.189, S1.44, 2.86, 3.83
 $NO_7S_2^{2-}$ Nitrosyldisulfonate ion (Freymy's salt), 1.184, 2.87, 3.79
 N_2O Nitrous oxide, 1.186, S1.42, 2.83
 N_3^- Azide ion, 1.177, 2.81, 3.73
 Na^+ Sodium(I) ion, 1.190
 Nd^{3+} Neodymium(III) ion, 1.191
 Ni^{2+} Nickel(II) ion, 1.193, S1.45, 2.88, 3.84a
 O^- , 1.9, S1.5, 3.4; see also part III, table 5.
 OV^{2+} Oxyvanadium(IV) ion, 3.121
 O_2 Oxygen, 1.205-6, S1.47, 2.89, 4.29
 O_2^- , 1.10, 3.7; see also part III, table 6.
 O_2Pb^{2-} Plumbate(II) ion, 1.215
 O_2Sn^{2-} Stannate(II) ion, 1.252
 O_2U^{2+} Uranyl(VI) ion, 1.268, 2.102
 O_3^- Ozonide ion, 4.30
 O_3P^{3-} Phosphite ion, 3.96
 O_3S^{2-} Sulfite ion, 1.238, S1.63, 3.105, 4.33
 $O_3S_2^{2-}$ Thiosulfate ion, 1.240, S1.65, 3.107
 O_3Sb^- Antimonate(V) ion, 1.243
 O_3Se^{2-} Selenite(IV) ion, 3.112
 O_3Sn^{2-} Stannate(IV) ion, 1.256
 O_3Te^{2-} Tellurate(IV) ion, 1.260, 3.116
 O_3Ti^{2-} Titanate(IV) ion, 1.263
 O_3V^- Vanadate(V) ion, 1.269
 O_3V^+ Oxyvanadium(VI) ion, 5.25
 O_4Os Osmium tetroxide, 5.20
 O_4P^{3-} Phosphate ion, 3.93
 O_4Ru^{2-} Ruthenate(VI) ion, 4.32
 O_4S^{2-} Sulfate ion, 1.239
 O_4Se^{2-} Selenate ion, 1.248
 O_4Te^{2-} Tellurate(VI) ion, 1.261
 $O_6S_2^{2-}$ Dithionate ion, S1.67
 $O_6S_3^{2-}$ Trithionate ion, S1.69
 $O_6S_4^{2-}$ Tetrathionate ion, S1.70
 $O_7P_2^{2-}$ Pyrophosphate ion, 1.212, 3.94
 $O_8S_2^{2-}$ Peroxydisulfate ion, 1.242, S1.68, 2.94, 3.109
 Pb^{2+} Lead(II) ion, 1.214

Pr(III), 1.223, S1.51, 2.92a, 3.98
Sm²⁺ Samarium(II) ion, 3.113
Sm³⁺ Samarium(III) ion, 1.250, S1.71
Sn(II), 2.96, 3.114
Sn(IV), 2.98
Tb³⁺ Terbium(III) ion, 1.258
Te(VI), 2.100
Th(IV), 5.22
Ti³⁺ Titanium(III) ion, 3.117

Tl⁺ Thallium(I) ion, 1.265, S1.72, 2.101, 3.118, 5.23
Tm(II), 3.119
Tm³⁺ Thulium(III) ion, 1.266
Y³⁺ Yttrium(III) ion, 1.270
U(IV), 3.120
U(VI), 5.24
Yb²⁺ Ytterbium(II) ion, 3.122
Yb³⁺ Ytterbium(III) ion, 1.272, S1.73
Zn²⁺ Zinc(II) ion, 1.274, S1.74, 2.103, 3.122a

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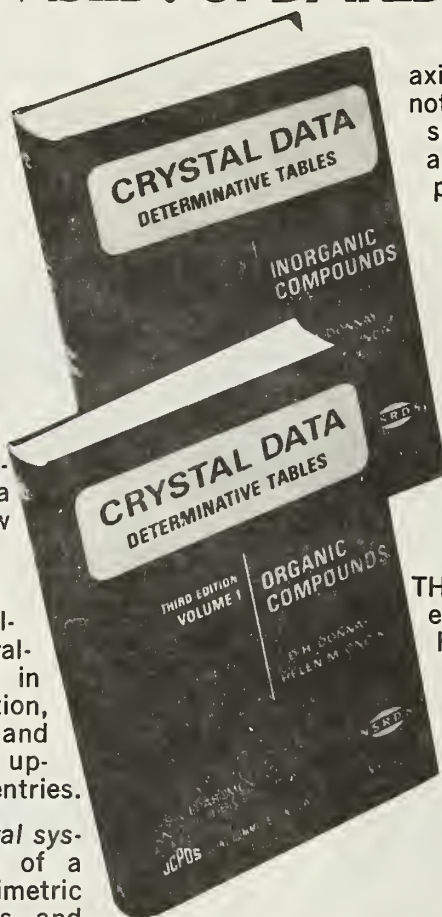
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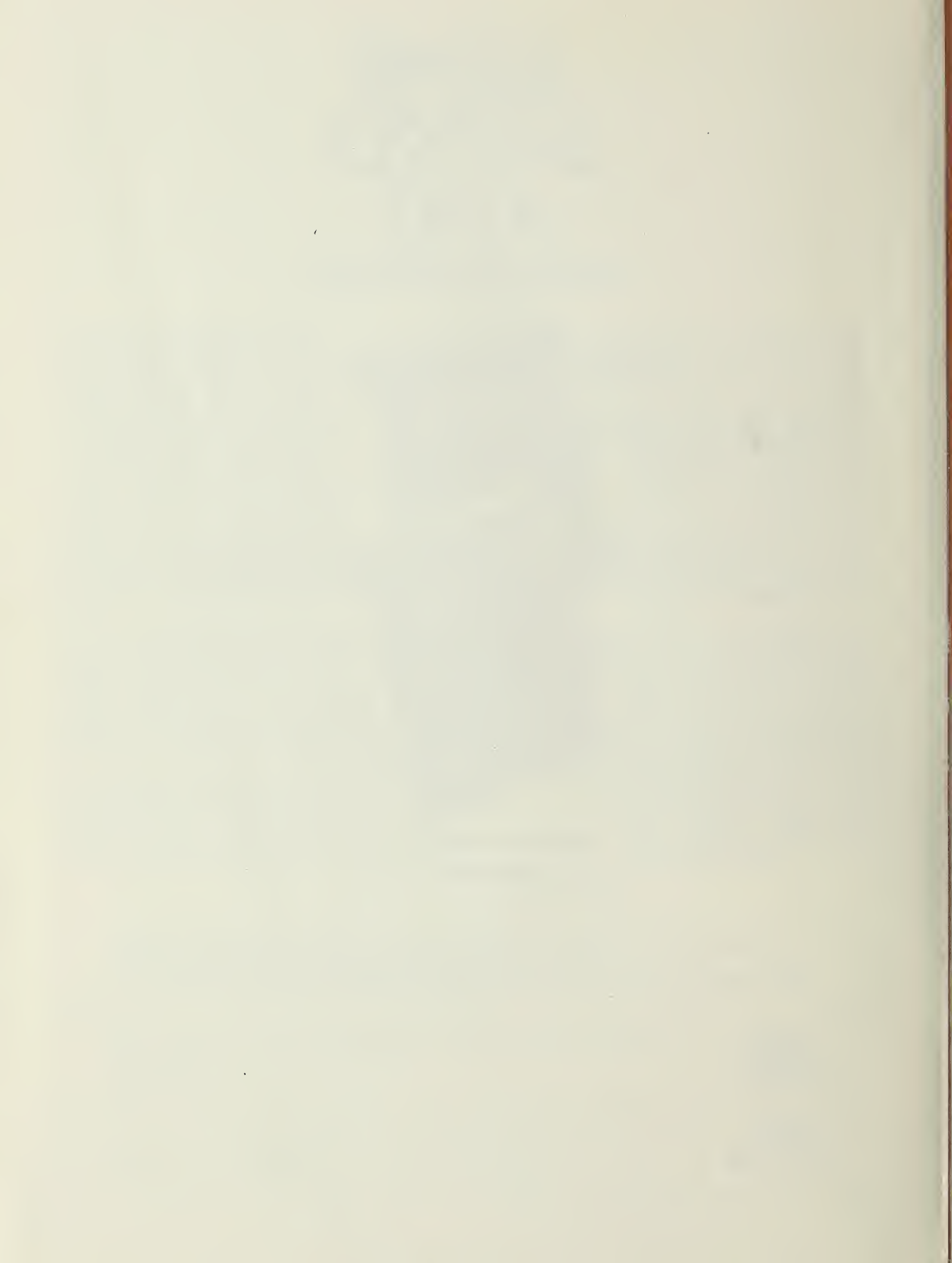
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