# THE CHEMISTRY OF AMIDOXIMES AND RELATED COMPOUNDS

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#### AMIDOXIMES AND RELATED COMPOUNDS

#### I. Introduction

# A. NOMENCLATURE

The amidoxime function (I) can be considered either as an amide (II) in which the oxygen atom of the car-

bonyl group has been replaced by an isonitroso group, or as an amidine (III) whose hydrogen atom of the imido group has been exchanged for a hydroxy radical. For this reason amidoximes are sometimes named oxamidines. They are also related to the hydroxamic acids (IV).

The lowest homolog (i.e., R = H) was named "isuretin" (V) by Lossen and Schifferdecker (91).

The name "amidoxime" was first used by Tiemann (175), who elucidated the structure of this class of compounds in 1884.

In the German papers published at the end of the nineteenth century, the name of the amidoximes is formed by adding "amidoxime" to the (obsolete) name of the corresponding trivalent radical. In the French literature, the suffix "oxime" is added to the name of the corresponding amide. According to Rule 32 of the International Union of Pure and Applied Chemistry, amidoximes should be named by adding "amidoxime" to the name of the corresponding hydrocarbon or by using the ending "carbonamidoxime." Chemical Abstracts considers the monoderivatives as amides of hydroxamic acids, i.e., hydroxamamides, and the polyderivatives as oximes of the amides.

Examples for some simple compounds are summarized in Table 1.

Imidoximes can be considered as imides of organic dicarboxylic acids in which the oxygen atom of one or both carbonyl groups has been replaced by an isonitroso group. For example:

#### B. MOLECULAR STRUCTURE

Although the first amidoxime was prepared in 1873 by Lossen and Schifferdecker (91) from hydrogen cyanide and hydroxylamine, these authors did not establish the structure of this compound, which they called "isuretin." It was Tiemann (175) who assigned a structural formula to the novel functional group, naming it "amidoxime." He prepared two related compounds (VI and VII) by the addition reaction of hydroxylamine to benzaldehyde cyanohydrin and to benzonitrile.

$$C_6H_6CHOHCN + NH_2OH \rightarrow C_6H_6CHOHC$$
 $NH_2$ 

VI, Mandelamidoxime

NOH
$$C_{6}H_{5}CN + NH_{2}OH \rightarrow C_{6}H_{5}C$$

$$NH_{2}$$
VII, Benzamidoxime

Tiemann proved the structure of the amidoxime function by showing the simultaneous presence of NH<sub>2</sub> and NOH groups (193): benzamidoxime forms salts with metals and with mineral acids just as would oximes and amines, respectively. The isonitroso group is identified by its acidic character and its reaction with nitrous acid with evolution of nitrous oxide.

NOH 
$$C_0H_5C \begin{array}{c} NOH \\ + HNO_2 \rightarrow C_0H_5CONH_2 + N_2O + H_2O \\ NH_2 \end{array}$$

When the sodium salt of benzamidoxime is alkylated with methyl iodide the resulting alkylated compound reacts with nitrous acid to yield nitrogen, thus proving the presence of the amino group.

NONa NOCH<sub>2</sub>

$$C_6H_6C + CH_2I \rightarrow C_6H_6C + NaI$$

$$NH_2 NOCH_3 NOCH_4$$

$$C_6H_6C + HONO \rightarrow C_6H_6C + N_2 + H_2O$$

$$OH$$

Tiemann recognized that the amidoximes may be present in two tautomeric forms (VIIIa and VIIIb), with VIIIa predominant.



The problem of tautomerism has been investigated more recently in the case of aminoglyoximes by Ponzio

TABLE 1 Nomenclature NOH NOH Nomenclature HC NΗ German Methenylamidoxime Ethenylamidoxime French Formamidoxime Acetamidoxime I.U.P.A.C. Methanamidoxime Ethanamidoxime Formhydroxamamide Acetohydroxamamide C.A.HON NOH NOH Nomenclature C<sub>6</sub>H<sub>5</sub>C C(CH<sub>2</sub>)<sub>5</sub>C NH  $H_2N$ NH: German Benzenylamidoxime French Pimelamidoxime Benzamidoxime I.U.P.A.C. Benzamidoxime Heptanediamidoxime Benzohydroxamamide Pimelamide dioxime

and his collaborators (132). Several aminoglyoximes are known under two forms with distinct physical properties. According to Ponzio they should be mere tautomers IXa and IXb.

The latter compounds can be transformed irreversibly into the former. However, the evidence given cannot be considered conclusive and only the more modern methods such as spectroscopy may finally settle the question of tautomerism in aminoglyoximes.

#### C. SCOPE OF THE REVIEW

This report deals with the chemistry of amidoximes. General and particular methods of their preparation, and their physical and chemical properties will be reviewed. The different classes of substitution products on both NOH and NH<sub>2</sub> groups and their derivatives will be examined, with the notable exception of the 1,2,4-oxadiazoles (X) formed by the cyclization of acylated amidoximes.

The greatest part of the work in the field of amidoximes was done by Tiemann and his co-workers (175–195) at the University of Berlin. They prepared most of the amidoximes and related compounds known at present. The more recent papers describe several new substances, but only a few new properties have been discovered.

#### II. SYNTHESIS OF AMIDOXIMES

#### 1. Monoamidoximes

Most of the monoamidoximes which can be found in the literature since 1873 are listed in Table 2. The details of the different methods of preparation (A through L) are given in the following paragraphs.

This is the most used process for the preparation of amidoximes. The experimental procedure recommended by Tiemann and Krüger (175) consists in liberating hydroxylamine from its hydrochloride using sodium carbonate, adding an equivalent amount of nitrile and enough alcohol to obtain a clear solution, and keeping the mixture at 60–80° during a few hours (method A1).

Instead of sodium carbonate, sodium or potassium hydroxide or sodium ethoxide also have been used. In the case of high-molecular-weight amidoximes Eitner and Weitz (35) used twice the theoretical amount of hydroxylamine and kept the reaction mixture for 25 hr. at 80°.

Yields rarely are given. Aromatic amidoximes generally are obtained with better yields than are aliphatic. As far as the lower members are concerned, their very high solubility in water and ethanol renders their isolation tedious.

To avoid the separation of amidoximes from potassium chloride or sodium chloride some authors (3, 12, 48, 61, 93, 148, 169, 173, 201) used a solution of free hydroxylamine in absolute methanol or ethanol (method A2).

Highest yields are obtained when a 15% excess of a solution of hydroxylamine in butanol is used (81a). A solution of the nitrile in the same solvent is introduced and the mixture is left for 48 hr. at 60°. The amidoxime separates as a practically pure crystalline compound.

A procedure described by Schiff (157) and related to the classical method of Tiemann consists in forming hydroxylamine *in situ* by the oxidation of ammonia with an alkaline hypochlorite solution (method A3). This method of forming hydroxylamine gives very poor yields and has never been used since.

# Method B: Action of Hydroxylamine on Amides or Thioamides

Hydroxylamine, as a rule, does not react with amides. This method has only been reported (32, 157) for the preparation of the two amidoximes.

NOH O—N NOH HOOCC and 
$$C_6H_6C$$
 C—C NH2 XI XII

Some aromatic amidoximes have been prepared by the action of hydroxylamine on thioamides (45, 54a, 168, 179).

$$RCSNH_2 + NH_2OH \rightarrow RC(=NOH)NH_2 + H_2S$$

Hydroxylamine is liberated from its hydrochloride by an equivalent amount of aqueous sodium carbonate. The thioamide is then introduced and ethanol is added until the mixture is clear. The solution is refluxed for a few hours and the amidoxime isolated. This procedure is used when the thioamide is more easily available than the corresponding nitrile.

# Method C: Reduction of Nitrosolic and Nitrolic Acids

Amidoximes can be obtained by reduction of nitro-

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TABLE 2

R = H  H— 104-105 A1 Ethanc  114-115 A1 Water  135 A1 Ethanc  135-135.5 A2 Butanc  CH <sub>1</sub> CH <sub>2</sub> — 55-58 A2 Butanc  (CH <sub>1</sub> ) <sub>2</sub> CH— 58 A1 Ethanc  CH <sub>1</sub> (CH <sub>2</sub> ) <sub>10</sub> — 92-92.5 A1 Ethanc  CH <sub>1</sub> (CH <sub>2</sub> ) <sub>11</sub> — 97 A1 Ethanc  CH <sub>1</sub> (CH <sub>2</sub> ) <sub>12</sub> — 101.5-102 A1 Ethanc  CH <sub>2</sub> (CH <sub>2</sub> ) <sub>14</sub> — 101.5-102 A1 Ethanc  CH <sub>3</sub> (CH <sub>2</sub> ) <sub>16</sub> — 106-106.5 A1 Ethanc	ol-water ol ol ol ol ol-water ol-water ol-water	°C.  40 0-5 30-40 40 60 Reflux 75 75 75	48 48 60-80 48 36 30 13-25 13-25 13-25	Ethanol Ethyl acetate Ethanol—ether Butyl ether  Benzene Benzene Benzene	60 80 20–25 90 75	91 110 112, 121, 194 36c 112 36a 55
CH₂ —     114-115     A1     Water       CH₂ —     135     A1     Ethan       135-135.5     A2     Butand       CH₃ CH₂ —     55-58     A2     Butand       (CH₃)₂ CH —     58     A1     Ethand       CH₃ (CH₂)₂ —     92-92.5     A1     Ethand       CH₃ (CH₂)₂ —     97     A1     Ethand       CH₃ (CH₂)₂ —     101.5-102     A1     Ethand       CH₃ (CH₂)₂ —     106-106.5     A1     Ethand	ol-water ol ol ol ol ol-water ol-water ol-water	0-5 30-40 40 60 Reflux 75 75 75	48 60-80 48 36 30 13-25 13-25	Ethyl acetate Ethanol-ether Butyl ether  Benzene Benzene	80 20–25 90	110 112, 121, 194 36c 112 36a
CH₂ —     114-115     A1     Water       CH₂ —     135     A1     Ethan       135-135.5     A2     Butand       CH₃ CH₂ —     55-58     A2     Butand       (CH₃)₂ CH —     58     A1     Ethand       CH₃ (CH₂)₂ —     92-92.5     A1     Ethand       CH₃ (CH₂)₂ —     97     A1     Ethand       CH₃ (CH₂)₂ —     101.5-102     A1     Ethand       CH₃ (CH₂)₂ —     106-106.5     A1     Ethand	ol-water ol ol ol ol ol-water ol-water ol-water	0-5 30-40 40 60 Reflux 75 75 75	48 60-80 48 36 30 13-25 13-25	Ethyl acetate Ethanol-ether Butyl ether  Benzene Benzene	80 20–25 90	110 112, 121, 194 36c 112 36a
135-135.5   A2   Butano	ol ol ol-water ol-water ol-water ol-water	40 60 Reflux 75 75 75	48 36 30 13–25 13–25	Butyl ether Benzene Benzene	90	36c 112 36a
CH₁CH₂→       A1         (CH₁)₂CH→       55-58       A2       Butano         (CH₁)₂CHCH₂CH₂→       58       A1       Ethano         CH₃(CH₂)10→       92-92.5       A1       Ethano         CH₃(CH₂)12→       97       A1       Ethano         CH₃(CH₂)14→       101.5-102       A1       Ethano         CH₃(CH₂)16→       106-106.5       A1       Ethano	ol ol ol-water ol-water ol-water	60 Reflux 75 75 75	36 30 13–25 13–25	Benzene Benzene		112 36a
(CH <sub>2</sub> ) <sub>2</sub> CH— 55–58 A2 Butano (CH <sub>3</sub> ) <sub>4</sub> CHCH <sub>2</sub> CH <sub>2</sub> — 58 A1 Ethano CH <sub>3</sub> (CH <sub>2</sub> ) <sub>10</sub> — 92–92.5 A1 Ethano CH <sub>3</sub> (CH <sub>2</sub> ) <sub>12</sub> — 97 A1 Ethano CH <sub>3</sub> (CH <sub>2</sub> ) <sub>14</sub> — 101.5–102 A1 Ethano CH <sub>3</sub> (CH <sub>2</sub> ) <sub>16</sub> — 106–106.5 A1 Ethano	ol ol-water ol-water ol-water ol-water	Reflux 75 75 75	30 13–25 13–25	Benzene	75	36a
(CH <sub>2</sub> ) <sub>2</sub> CHCH <sub>2</sub> CH <sub>2</sub> —     58     A1     Ethanc       CH <sub>4</sub> (CH <sub>2</sub> ) <sub>10</sub> —     92-92.5     A1     Ethanc       CH <sub>4</sub> (CH <sub>2</sub> ) <sub>12</sub> —     97     A1     Ethanc       CH <sub>4</sub> (CH <sub>2</sub> ) <sub>14</sub> —     101.5-102     A1     Ethanc       CH <sub>4</sub> (CH <sub>2</sub> ) <sub>16</sub> —     106-106.5     A1     Ethanc	ol ol-water ol-water ol-water ol-water	Reflux 75 75 75	30 13–25 13–25	Benzene	70	
CH4(CH2)10—     92-92.5     A1     Ethanc       CH4(CH2)12—     97     A1     Ethanc       CH2(CH2)14—     101.5-102     A1     Ethanc       CH4(CH2)16—     106-106.5     A1     Ethanc	ol-water ol-water ol-water ol-water	75 75 75	13-25 $13-25$	Benzene		
CH <sub>4</sub> (CH <sub>2</sub> ) <sub>12</sub> — 97 A1 Ethanc CH <sub>4</sub> (CH <sub>2</sub> ) <sub>14</sub> — 101.5-102 A1 Ethanc CH <sub>4</sub> (CH <sub>2</sub> ) <sub>16</sub> — 106-106.5 A1 Ethanc	bl-water bl-water bl-water	75 75	13-25			35
CH <sub>4</sub> (CH <sub>2</sub> ) <sub>16</sub> — 106-106.5 Al Ethano	ol-water		13-25	Denzene		35
		75		Benzene		35
(CHa)um	ol-water		13–25	Benzene		35
COLDIZ 69 AI EURING		Reflux	24	Ethanol		19
CH2=CH(CH2)s 69 A2 Ethano	ol			Benzene		61
C <sub>9</sub> H <sub>15</sub> — 101 A2 Ethano		Reflux	Days			48
• •	ol-water	60-80	Hours	Water	100	175. 193
* * * * * * * * * * * * * * * * * * *	ol-water ol-water	80 Reflux	18 15–18			63 179
C Water	T-waret	Cold	10-19			179 212
D Ether		0014				207
E Ethano	ol	0		Ether		82
76–77 F						88
80 G Water	•	20	Days	Ethanol-water		119
77-78 H Ethano CH <sub>i</sub> C <sub>5</sub> H <sub>4</sub> 149.5 Al Ethano		175 Reflux	8 6	Water		89 162
	l-water	80	1.5	Ethanol	75–90	22
	l-water	80-90	6	Water	10 00	161
C <sub>6</sub> H <sub>5</sub> CH <sub>2</sub> — 67 A1 Ethano	l-water	40-50	36-48	Ethanol-water	40-50	57
CH <sub>3</sub> 178 A1 Ethano	l-water	Reflux	5–6	Ethanol		114
	l-water	60-70	Days	Ethanol	25	217
93 A2 Methat		0	72	Water		148
• •	l-water	Reflux 80-90	10-12	Water		36. 151 150
3-Naphthyl 150 Al Ethano	l-water	pressure) 80-90	10-12	Ethanol-water		36, 151
• •	l-water	80-90	10-12	Ethanol-water		150
		pressure)				
R # H	eterocyclic ra	adical				
·	l-water	70	21	_		79
R-Benzofuryl 190-191 Al <sup>a</sup> Ethano		70 70	24	Benzene		79 70
Henzofuryl 106-107 A1° Ethano H-Pyridyl 116-117 A1 Ethano		70 70	30 5	Benzene Benzen <del>e</del>		79 79
• •	l-water	70	5	Water	80	27
		Reflux	1.5	Benzene	75–90	22
128 Al Ethano	l-water	70	8	Chloroform	57	27
128 Al Water		70	8	Chloroform		99
-Pyridyl 197-198 Al Ethano		70 70	5 5	Water Water	38	79 27
	l-water l-water :	Reflux	5 24	Water Ethanol		27 79
		Reflux	36	Ethanol		79
-Quinolyl 195 Al <sup>a</sup> Ethano	1	70	24	Benz. ethanol		79
		Reflux	0.0	Alcohol		13
R-Thienyl 91-92 A1° Ethano R-Thienyl 91-92 A1	l-water	Reflux	30	Benzene Benz. pet. ether		79, 98 98
C <sub>6</sub> H <sub>6</sub> NC 208-210 A1 Ethano	l-water	50-70	Hours	Ethanol		15
C						
CH <sub>1</sub> * Excess of NH <sub>2</sub> OH used						

<sup>&</sup>lt;sup>a</sup> Excess of NH<sub>2</sub>OH used.

TABLE 2 (Continued)

	Melting			Reaction	Reaction			
	point		Reaction	temp.,	time,	Crystallized	Yield,	
R	°C.	Method	medium	°C.	hr.	from:	%	Ref.
0-N		·					<del></del>	
/		_						
HC C—	115	ь						81a
o—n								
7.0		ъ.	36.0 1	2.		T. 1		0.0
6H6C C—	158	В	Methanol	65		Ethanol		3 <b>2</b>
`n´								
		R =	substituted hydro	carbon radic	al .			
CH:	91-92	A1	Water	30	0.25	Benzene	70	169, 171
2CH—	103-104	Al	Water	0	Min.	Benzene	60	169, 171
ıC—	128-129	Al	Water	0	Min.	Benzene		169, 171
·CH <sub>2</sub> —	95-96	A2	Methanol	8-0		Methanol		169, 173
rsCH—	120	A2	Methanol	0		Toluene		169, 173
C—	126	A2	Methanol	0	Hours	Ethanol		169, 173
CH <sub>2</sub> —	123-124	A2	Methanol	20	24	Ethanol	37.5	169, 171
ClC <sub>6</sub> H <sub>4</sub> —	117	D	Ethanol			Water		206
ClC <sub>6</sub> H <sub>4</sub> —	134-135	Al	Ethanol-water	Reflux	20	Ethanol	61	3
BrC6H4—	146-147	Al	Ethanol-water	Reflux	1.5	Ethanol	75–90	22
	144-145	Al	Ethanol-water	Reflux	20		72	3
H <sub>4</sub> CH(OH)—	115-116	Al	Water			Ethyl acetate	50	157
CH <sub>1</sub> ) <sub>1</sub> C(OH)—	52	A2	Ethanol	Reflux	2	Ether	60-70	70
)II)(C(OI)—	55–60	A3	Water	Itenua	-	Ether	33	157
MIL CHOU CHION								
CH <sub>1</sub> ) <sub>2</sub> CHCH <sub>2</sub> CH(OH)—	176.5	Al	Water	~~	70	Ethanol-water	90-95	157
Hich(OH)—	141	Al	Water	<b>7</b> 5	72	Ethanol-water	90-95	157
H₅CH(OH)—	140	A1	Ethanol-water	20	Days			175
	158-159	Al	Ethanol-water	25-30	Days	Ethanol	25	50, 51
	163-164	K		0		Ethanol-chlorof.		24
₅H₅CH≕CHCH(OH)—	136	Al	Ethanol-water	7	12		50	17
l <sub>s</sub> CCH(OH)—	156-157	Al	Water	20	Hours	Water	90-95	152, 157
HOC <sub>5</sub> H <sub>4</sub> —	98-99	В	Ethanol-water	Reflux	4	Water		100, 168
-HOC₀H₄—	71	Al	Ethanol-water	Reflux	6	Water		23
				Cold	5-6 days			23
HOC₀H₄—	153	Al	Ethanol-water	70	10	Water	80	62
CH <sub>3</sub>								
<b>(</b> )-	123-124	В	Ethanol-water	Reflux	3-4	Benzene		45
OH								
	100 7		TO 1 0	00				110
CH OH	126.5	Al	Ethanol-water	60	15			116
CH <sub>3</sub> —OH								
но 🍆	150	A 1	Ethanol	D.4	1 5			110
CH <sub>3</sub>	152	Al	Ethanol-water	Reflux	15			116
Br								
		-	The bound of the state of			Title - 1		100
Br OH	180	В	Ethanol-water			Ethanol-water		168
c								
но	166	A2	Ethanol	20	6-8 days	Ethanol-water		93
/_OH					-			
~~ ^~	_			D -		_		
·CH <sub>3</sub> OC <sub>6</sub> H <sub>4</sub> —	123	Al	Ethanol-water	Reflux	1.5	Benzene	75–90	22
	123	Al	Ethanol-water	90	6–8	Benzene	100	100
-CH2OC6H4—	122-123	Al	Ethanol-water	90	6-8	Benzene	81	100
H0(	100	A2	Ethanol	80	15	Water		93
• •								
-CH <sub>2</sub> (CH <sub>2</sub> ) <sub>5</sub> OC <sub>6</sub> H <sub>4</sub> —	110	Al	Ethanol-water	70	24	Ethanol-water	50	115
			Tu -1	60				00
<u></u>	151	A2	Ethanol	60				93
<b>6</b>	104 105		Ditarial	D. Ø	0.4	Eth and	**	0
CH,	164–165	Al	Ethanol-water	Reflux	24	Ethanol	55	3
CH <sub>3</sub> O CH <sub>2</sub>	137	Al	Ethanol-water	Reflux	8	Ethanol		69
J1130\ /CH2								

Formed by the	action of formic	acid on oxamidediox	ime. CThese an	idoximes have be	en prepared	by the method $A_1$ b	ut no data ar	e given (20a):
R'	R = H	R' = Cl	R = Br	R' = Br		ъ	R = C1	R' = Cl
<b>⟨</b> »	{ C1	Cl	H	I	and	110 <sup>1</sup>	H	Br
R/—OH	\ н	Br	I	I		поу	Br	Br
						R	T	Т

TABLE 2 (Continued)

			TABLE 2 (CO	nisnaea)				
	Melting			Reaction	Reaction			
			Reaction		time.	Crystallized	Yield,	
R	point, °C.	Method		temp., °C.	hr.	from:	1 161a, %	Ref.
11	<u> </u>	- Internou	mediam			11041.	/0	1001.
$C_6H_5CH[O(C_{12}H_{21}O_{10})]$ —	135-140	Al	Ethanol-water	20	72	Ethanol-water		156
C <sub>6</sub> H <sub>6</sub> CO—	129-131	D	Ether	20	•-		56	76
H00C-	158	ď	20201	-0			50	52, 53
HOOCCH2							50	
	144							103
HOOC(CH <sub>2</sub> ) <sub>4</sub> —	156-158	C	Acetic acid	25		Methanol	83	44
m-HOOCC <sub>0</sub> H <sub>4</sub> —	200	Al	Ethanol-water	80-100	12			107
$p ext{-}HOOC_6H_4$ —	330	Al	Ethanol-water	20	18	Ethanol-water		107
m-C2H2OCOC6H4-	118	Al	Ethanol-water	80-100	Hours	Water		107
p-C2H5OCOC6H4—	135	Al	Ethanol-water	60-100	8			106
NO <sub>2</sub> CH <sub>2</sub> —	108	Al	Water	20	24	Water		172
						W WIEL		
m-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	174	Al	Water	100	5			159
p-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub>	169	Al	Ethanol-water	90	4-5	Water		201
	170	Al	Ethanol-water	Reflux	1.5	Water	75–90	22
	180	A2	Methanol	Reflux	1	Ethanol	66	12
$C_6H_5CH(NO_2)$ —	125	Al	Water	20		Ether-pet. eth.		170
CH <sub>3</sub> ( )—	161	A2	Ethanol					201
$\sim$ NO <sub>2</sub>								
0-NH2C6H4-	84-85	A1	Ethanol-water	Reflux	8	Benzene	65	121s
m·NH2CoH4—	01 00	î.	Linanoi water	1tenua	0	Donzene	00	160
p-NH2C6H4—	160-174	f				Water		
p-N119C6H4—	100-174	•				water		201
CH <sub>3</sub>								
	166	f					Poor	201
$\sim$ NH <sub>2</sub>								
$_{DL}$ - $C_{6}H_{5}CH_{2}CHNH_{2}$ -	117-118	A2	Methanol			Water	52	118
CI								
$\sim$ NH <sub>2</sub>	128-130	Al	Ethanol-water	Reflux	8	Benzene	60	20b
$H_2NC$ —— $C$ —								
N N	189-190	ø				Water		83
, O , 1								
•								
$\sim$					_			
CH <sub>2</sub> O	292-294	Al	Pyr. water-	100	1		54	47
🔷 🗸			ethanol					
$NH_2$								
∧ .N. ∧								
		Al	Pyr. water-	100	2			47
			ethanol	200	-			21
NHC <sub>6</sub> H <sub>5</sub>			evilation					
· •								
NH <sub>2</sub> COCH <sub>2</sub> —	149	A2	Ethanol	80		Ethanol		74
$NH_2CO(CH_2)_{11}$ —	157-158	A1	Ethanol	60	20	Ethyl acetate		75
DL-C6H6CH2CH(NHCOCH1)—	156-158	λ				Water	66	118
L-	167-169	ħ				Water	74	118
DL-C6H6CH2CH(NHCOC6H6)—	200-202	h				Methanol-water	62	118
		h					44	
L-	200-203					Methanol-water	44	118
						butanol		
NCH2CH2—	185	A1	Dimethylform-	60	24		40	36a
11022012			amide-water					
¥ C0								
/ co~\								
DL-C <sub>6</sub> H <sub>5</sub> CH <sub>2</sub> CH(NCCC)	198-204	Al	Methanol	Reflux	3	Methanol	71	118
1 .co.//								
L-	164-171	Al	Methanol	Reflux	3	Methanol-water	73	118
C <sub>1</sub> H <sub>1</sub> CONHCH <sub>2</sub> —	123-126	***	17000000	20044	v	and of the state o		31
	124-127	4.1	Ethanol-water	20	Hours	Ethanol		
CNCH.		Al						158
CN(CH <sub>2</sub> )	103	Al	Ethanol-water	60-70	10	Water		14, 42
$CN(CH_2)_4$ —	8991	A2	Butanol	50	18		76	219a
CN(CH <sub>2</sub> ) <sub>5</sub> —	71	A2	Butanol	50	18			219a
CN(CH <sub>2</sub> ) <sub>11</sub> —	87-88	Al	Ethanol	60	20	Methanol		<b>7</b> 5
CN(CH2)15-	98	$\mathbf{A}1$	Ethanol	60	20	Ethanol		75
p-CNC <sub>6</sub> H <sub>4</sub> CH <sub>2</sub> —	168		Ethanol-water					154
HONHCOCH:-	152		Ethanol-water	40	3-4			102, 103
	148-152		Water	60	~ -	Water		144, 169, 171
CH(:NOH)—				00		Benzene		
CCIV. MOTIV	152		Ethanol	a =	M:		10	142
CCl(: NOH)—	109		Water	65	Min.	Eth. petr. ether	19	169, 171
CH <sub>1</sub> C(:NOH)—	183-184		Water	20	-	T-1		146
	183-184		Water	20	Hours	Ether		145
CH <sub>2</sub> C(:NOCH <sub>2</sub> )—	99		Methanol	20		Pet. ether		6
CH(:NOCOC6H6)—	157-158	D	Ether-water	20		Ethanol-water		142, 144
β CH(:NOH)C(:NOH)—	147-148		Methanol	20				85
C <sub>6</sub> H <sub>5</sub> C(:NOH)—	α 154	L						140, 141
	β 195		Ethanol	20	Hours	Ethanol		130, 138, 140

<sup>&</sup>lt;sup>d</sup> Formed by partial hydrolysis of the diamidoxime NH<sub>2</sub>C(:NOH)C(:NOH)NH<sub>2</sub>. <sup>e</sup> Formed by partial hydrolysis of the hydroxamic acid HONHCOCH<sub>2</sub>C-(:NOH)NH<sub>2</sub>. <sup>f</sup> Prepared by reduction of the corresponding nitro compound. <sup>g</sup> Formed by hydrolysis of the acetyl derivative. <sup>h</sup> Formed by partial hydrolysis of the diacylated aminoamidoximes. <sup>i</sup> From two moles of NH<sub>2</sub>OH on C<sub>2</sub>H<sub>2</sub>OCOCH<sub>2</sub>CN. <sup>j</sup> From a large excess of hydroxylamine on Cl<sub>2</sub>CHCN. <sup>k</sup> From two moles of NH<sub>2</sub>OH on Cl<sub>2</sub>CCN.

TABLE 2 (Continued)

R	Melting point, °C.	Method	Reaction medium	Reaction temp., °C.	Reaction time, hr.	Crystallized from:	Yield,	Ref.
C <sub>6</sub> H <sub>6</sub> COC(; NOH)—	α 127	L	Water	20	24			143
Conscion(. Non)	β 187	~i	11 2001	20				143
p-CH <sub>4</sub> C <sub>6</sub> H <sub>4</sub> COC(:NOH)—	α 114	L	Water	0				143
CoHoC(:NOCOCoHo)—	171-172	D	Ether-water	20		Ethanol		129, 131
	171-172	L	Ether-water	20		Ethanol		131
p-CH <sub>2</sub> C <sub>2</sub> H <sub>4</sub> C(:NOCOC <sub>6</sub> H <sub>5</sub> )—	α 178-179	L	Ether-water	20		Ethanol		131
C <sub>5</sub> H <sub>5</sub> NHC(:NOH)—	180	Al	Ethanol	75-80		Water		180, 221
p-CH <sub>3</sub> C <sub>6</sub> H <sub>4</sub> NHC(:NOH)—	175	Al	Ethanol	75-80		Water		199
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> NHC(:NH)—	147-148	Al	Ethanol	75-80		Benzene		199
$C_6H_6-N=N-$	125-126	Al	Water	20		Water		86
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —N=N—	164-165	Al	Water	20		Water		86
p-CH <sub>5</sub> SC <sub>6</sub> H <sub>4</sub> —	130	A1	Ethanol-water	Reflux	12	Water	100	20
p-C <sub>2</sub> H <sub>5</sub> SC <sub>6</sub> H <sub>4</sub> —	120	Al	Ethanol-water	Reflux	12	Water		20
p-C <sub>6</sub> H <sub>6</sub> SC <sub>6</sub> H <sub>4</sub> —	125	Al	Ethanol-water	Reflux	40	Ethanol		78, 79
p,p'-ClC6H4SC6H4—	159-161	Al	Ethanol-water	Reflux	40	Ethanol		78, 79
p, p'-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> SC <sub>6</sub> H <sub>4</sub> —	162-164	Al	Ethanol-water	Reflux	40	Benzene		78, 79
p-HO2SC6H6—	236	Al	Ethanol-water	Reflux	3		25	3
HO'S OH	250	m						168
p-CH₁SO₂C₅H₄—	177	Al	Ethanol-water	Reflux	12	Ethanol-water		20
p-011,50204114	188	Al	Ethanol-water	Reflux	1.75	Water	96	3
$p$ -C <sub>6</sub> $H_6$ SO <sub>2</sub> C <sub>6</sub> $H_4$ —	175–176	Al	Ethanol-water	Reflux	40	Ethanol	00	78, 79
p, p'-ClC <sub>0</sub> H <sub>4</sub> SO <sub>2</sub> C <sub>0</sub> H <sub>4</sub> —	201-202	Al	Ethanol-water	Reflux	40	Ethanol		78, 79
p, p'-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> SO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	201-202	Al	Ethanol-water	Reflux	40	Ethanol		78, 79
p-NH <sub>2</sub> SO <sub>2</sub> C <sub>5</sub> H <sub>4</sub> —	-01 -0-	A2	Methanol	Reflux	2	Water	80	12
p=1111200206114		A1	Ethanol-water	Reflux	1.5	,, 400	70-82	3
p-HONHSO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	152-153	Al	Ethanol	20	1.0	Water	51	3
NH <sub>2</sub> SO <sub>2</sub> CH <sub>3</sub>	183–184	Al	Ethanol-water	Reflux	3	Water		3
p-(C2H1)2NSO2C6H4-	123-124	A1	Ethanol-water	Reflux	16		55	3
p-(HOC2H4)2NSO2C6H4—	183-184	Al	Ethanol-water	Reflux	16	Ethanol	40	3
p-CH <sub>2</sub> ONHSO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	198 <sup>n</sup>	Al	Ethanol-water	Reflux	16	Dil. HCl	56	3
p-C <sub>2</sub> H <sub>5</sub> ONHSO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	$151-152^n$	A1	Ethanol-water	Reflux	16	Dil. HCl	57	3
p-C <sub>4</sub> H <sub>7</sub> ONHSO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	145-150 <sup>n</sup>	Al	Ethanol-water	Reflux	16	Dil. HCl	55	3
p-C <sub>6</sub> H <sub>9</sub> ONHSO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	130-140 <sup>n</sup>	Al	Ethanol-water	Reflux	16	Dil. HCl	76	3
p-\bigcolon NSO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> -	196–197	A2	Ethanol			Ethanol	93.5	3
2-0_NSO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> -	158–159	A1	Ethanol-water	Reflux	16	Ethanol	92.5	3
p,p'-(CH <sub>3</sub> )2NSO2C6H4NHSO2C6H4—	130	A2	Ethanol	Reflux	16	Water		3

I Formed by heating the α form in dilute acetic acid. The Formed by sulfonation of α-OHC<sub>6</sub>H<sub>4</sub>C(:NOH)NH<sub>1</sub>. Isolated as hydrochloride.

Recently, an apparently general method was described (44) for the preparation of the monoamidoximes of dicarboxylic acids, by catalytic reduction of nitrolic acids (XIV).

For example, adipomonoamidoxime was prepared by the reduction of adipomononitrolic acid; the latter was obtained from cyclohexanol and cold nitric acid.

> Method D: Action of Ammonia on Hydroximic Acid Chlorides (Chloroximes)

Hydroximic acid chlorides (XV) are formed by

direct chlorination of aldoximes. These compounds react easily with ammonia to yield amidoximes. This procedure was used by Werner (206, 207) to prepare benzamidoxime, o-chlorobenzamidoxime (206, 207), and terephthalamidoxime (36a).

Method E: Reduction of Oxyamidoximes

Hydroximic acid chlorides also react with hydroxylamine to form oxyamidoximes (XVI). This reaction was used by Ley and Ulrich (82) to prepare benzoxyamidoxime.

NOH NOH NOH C<sub>6</sub>H<sub>6</sub>C + NH<sub>2</sub>OH 
$$\rightarrow$$
 C<sub>6</sub>H<sub>5</sub>C + HCl NHOH XVI

This compound then is reduced with sulfur dioxide to the corresponding amidoxime.

Method F: Action of Hydroxylamine on Iminoethers

This reaction was reported by Pinner (119) and Lossen (88), who obtained benzamidoxime by treating ethyl iminobenzoate (XVII) with hydroxylamine.

Since benzonitrile is the starting material for the synthesis of the iminoether, this reaction is not a practical method for the synthesis of amidoximes (compare with the more direct method A).

# Method G: Action of Hydroxylamine on Amidine Hydrochlorides

Pinner (119) prepared benzamidoxime by treating benzamidine hydrochloride (XVIII) with hydroxylamine.

This reaction has no practical interest, since amidines generally are obtained from nitriles, thioamides, or iminoethers (compare with methods A, B, and F).

# Method H: Action of Ammonia on Oximinoethers

When heated in a pressure bottle for 8 hr. at 175°, an alcoholic solution of ammonia and ethyl benzhydroxamic acid (XIX) yields benzamidoxime (89).

NOH
$$C_{\delta}H_{\delta}C + NH_{\bullet} \rightarrow C_{\delta}H_{\delta}C + C_{2}H_{\delta}OH$$

$$OC_{2}H_{\delta}$$

$$XIX$$

$$NOH$$

$$+ C_{2}H_{\delta}OH$$

$$NH_{2}$$

This reaction has not found general application.

# Method K: Action of Formamidoxime on Aromatic Aldehydes

According to Conduché (24) formamidoxime reacts with aromatic aldehydes, leading to an aldol condensation

$$C_{\bullet}H_{\circ}C$$
  $O$  NOH NOH NOH NOH NOH NOH NH<sub>2</sub>

Mandelamidoxime is the only compound that has been prepared by this method.

Method L: Action of Ammonia on Glyoxime Peroxides

Dehydrogenation of substituted glyoximes yields "oxides" (XX), the structure of which has not yet been established with certainty. Depending on the relative configuration of the NOH groups (syn, anti, or amphi) and on the nature of the substituents, the "oxides" are considered as nitrile oxides (XXa), furazane oxides (XXb,c), or peroxides (XXd).

On treatment with ammonium hydroxide phenylglyoxime peroxide yields an oxime containing amidoxime (XXI) (140, 141, 215).

$$C_0H_5(C_2N_2O_2)H \xrightarrow{NH_4OH} C_0H_5C - C$$

NOH NH<sub>2</sub>

XXI

The reaction is analogous to that of the *O*-benzoylated nitrile-oxide XXII (131).

Another example is the reaction of dibenzoylglyoxime peroxide with ammonium hydroxide (143).

# 2. Diamidoximes

Diamidoximes, *i.e.*, compounds containing two amidoxime groups simultaneously can be prepared by reactions analogous to those described in section 1. The diamidoximes heretofore prepared and their methods of preparation are summarized in Table 3.

$$\begin{array}{c} \text{TABLE 3} \\ \text{HON} \\ \text{NOH} \\ \text{C--R--C} \\ \text{H}_2\text{N} \\ \text{NH}_2 \end{array}$$

R	Melting point, °C.	Method	Reaction medium	Reaction temp., °C.	Reaction time, hours	Crystallized from:	Yield, %	Ref.
—(oxamidedioxime)	196–200	Al	Water	0		Water	40	40
	195	Al	Ethanol	20				180
	196-200	Al	Ethanol	Reflux		Water		199, 221
	212	Al	Water	0-5		Water	53	36a
	198	В	Ethanol	Reflux		Ethanol		37
	200	D	Ethyl acetate	20		Water		54
—CH₂—	163-167	Al	Ethanol-water	20	24	Ethanol-water	80	158
-CH2CH2-	188	A1	Ethanol-water	20	4 days	Water		52, 166
	188	Al	No solvent	60-65				81 c
	188	A1	Ethanol	70	8	Water		42
-CH <sub>2</sub> CH <sub>2</sub> CH <sub>2</sub> -	233	Al	Ethanol-water	60-70	10	Water		14, 42
—(CH <sub>2</sub> ) <sub>4</sub> —	226	A2	Butanol	50	48	Butanol	100	36a, 81a
(+).	168-170	A2	Butanol	70			47	219a
—(CH <sub>2</sub> ) <sub>5</sub> —	142-144	A1ª	Ethanol	60	20-30	Ethanol		75
—(CH <sub>2</sub> ) <sub>6</sub> —		A2	Butanol	50	40			36a, 81a
—(CH <sub>2</sub> );—	156	A1ª	Ethanol	60	20-30	Ethanol		75
—(CH <sub>2</sub> ) <sub>8</sub> —	152	A2	Butanol	70				219a
—(CH <sub>2</sub> ) <sub>8</sub> —	167	A1a	Ethanol	60	20-30	Methanol		75
—(CH2)10—	184186	A1a	Ethanol	60	20-30	Ethanol		75
—(CH <sub>2</sub> ) <sub>11</sub> —	166	A1ª	Ethanol	60	20-30	Methanol		75
—(CH2)11—	170	A1ª	Ethanol	60	20-30	Ethanol		75
-CH=CH-	212	Al	Water	20	15	Water	16	92
-C(:NOH)-	154	A2	Ethanol	50-60	36	Water	100	83
-C(:NOH)-	181-182	A1	Water	50-60	00	***************************************	100	83
-CC(CH <sub>2</sub> ) <sub>2</sub> N=NC(CH <sub>2</sub> ) <sub>2</sub> C-	151-152	Al	Ethanol-water	20	24	Ethanol-pet. ether	82	174a
m-CsH4—	193	A2	Ethanol	Reflux	24	Ethanol	02	46
	180	D	Ethanor	Itenux		Ethanoi	100	36a, 81a
p-C <sub>6</sub> H <sub>4</sub> —	192	A1	Ethanol-water	20	24	Water	100	30a, 31a 154
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —		$A1^b$	Ethanol	60	20-30	Ethanol		75
p,p'-C6H4-C6H4-	245	A1	Butanol				100	
	290°	$^{ m A2}_{ m A1^b}$		45	48	Acetone-water	100	36a, 81a
p, p'-C <sub>6</sub> H <sub>4</sub> CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	245		Ethanol	60	20-30	Ethanol		75
p,p'-C6H4CH2CH2C6H4-	243	$A1^b$	Ethanol	60	20-30	Ethanol		75
p,p'-C6H4CH=CHC6H4-	>320	$A1^b$	Ethanol	60	20-30	Ethanol		75
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> CH <sub>2</sub> —	250	A2	Butanol	45	48	Butanol	85	36a, 81a
p,p'-C6H4O(CH2)1OC6H4-			<b>.</b>					7
p, p'-C <sub>6</sub> H <sub>4</sub> O(CH <sub>2</sub> ) <sub>5</sub> OC <sub>6</sub> H <sub>4</sub> —	190	A2	Butanol	45	48	<b>~</b> .	100	36a, 81a
p,p'-C <sub>6</sub> H <sub>4</sub> SC <sub>6</sub> H <sub>4</sub> —	200	Al	Ethanol-water	Reflux	40	Dioxane-water		78, 79
p, p'-C <sub>6</sub> H <sub>4</sub> SO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	214–220	Al	Ethanol-water	Reflux	40	Dioxane-water Dimethylformamide-		78, 79
$p, p'$ -(— $C_6H_4CH_2S$ ) <sub>2</sub>	195	A2	Butanol	45	72	water	85	36a, 81a

<sup>&</sup>lt;sup>a</sup> Twice the theoretical amount of hydroxylamine used. <sup>b</sup> Excess of hydroxylamine used. <sup>c</sup> Analyzed as the dihydrochloride.

The most usual synthesis is still the Tiemann method (dinitrile + hydroxylamine). Highest yields are obtained if free hydroxylamine in butanol is used (81a). This procedure also facilitates the isolation of the product. With high molecular weight dinitriles an excess of hydroxylamine is needed to obtain good yields.

In the case of oxamidedioxime a variant of method A1 has been described (180, 199, 221). Instead of cyanogen, its addition compound with aniline, diphenyloxamidine (XXIII) is treated with hydroxylamine hydrochloride.

$$(CN)_{2} \xrightarrow{C_{\theta}H_{1}NH_{2}} C_{\theta}H_{\delta}NHC(:NH)C(:NH)NHC_{\theta}H_{\delta}$$

$$\xrightarrow{XXIII}$$

$$HON \qquad NOH$$

$$C-C \qquad + 2C_{\theta}H_{\delta}NH_{2}\cdot HC$$

$$H_{2}N \qquad NH_{2}$$

Two by-products also are formed during the reaction

HON NOH HON NH
$$C-C \qquad \text{and} \qquad C-C$$

$$H_2N \qquad NHC_{\mathfrak{b}}H_{\mathfrak{b}} \qquad H_2N \qquad NHC_{\mathfrak{b}}H_{\mathfrak{b}}$$

Other amines such as benzylamine and p-toluidine can be used.

It appears that best yields and the purest product are obtained if gaseous cyanogen is led directly into an aqueous hydroxylamine solution at 0° (36a).

#### 3. Imidoximes

The formation of diamidoximes often is accompanied by side reactions which are influenced by such factors as the proportion of reagents, temperature, and the choice of solvents. When intramolecular cyclization can occur, mono and dioximes of the corresponding imides are formed. These reactions take place, for instance, with the dinitriles of succinic, glutaric, and o-phthalic acids and with o-cyanomethylbenzonitrile.

 $\mathrm{CH_{2}CN}$ 

NH<sub>2</sub>OH

When one mole of succinonitrile reacts with two moles of hydroxylamine in aqueous ethanol at room temperature, a small amount of succinimidedioxime (XXV) accompanies the major product, succinamidedioxime (XXIV). At 60-70° the yield in XXV increases (166). The latter is formed by splitting off ammonia from XXIV.

HON NOH CH<sub>2</sub>—C NH

$$CCH_2CH_2C$$
 NH

 $NH_2$  CH<sub>2</sub>—C NOH

 $CH_2$ —C NOH

 $XXIV$  XXV

If the molar ratio of reactants is one, either an open chain diamidoxime or a cyclic imidoxime is formed (XXVI), depending on the reaction conditions: in absolute ethanol, at room temperature, the former is the main reaction product, while at 60–70° or in the presence of water the cyclic compound (XXVI) is formed (42). Probably an intermediate monoamidoxime first cyclizes into an imide which then hydrolyzes readily into an imidoxime.

The same end product XXVI can be obtained through two other distinct methods. The first consists in treating succinimidedioxime (XXV) with an equivalent amount of nitrous acid.

NOH NOH

$$CH_{2}C$$

$$NH + HNO_{2} \rightarrow \qquad NH + N_{2}O + H_{2}O$$

$$CH_{2}C$$

$$NOH$$

$$O$$

In the second,  $\beta$ -cyanopropionamide (XXVII) reacts with hydroxylamine.

$$\begin{array}{c} \text{NOH} \\ \text{CH}_2\text{CN} \\ \text{CH}_2\text{CONH}_2 \end{array} \xrightarrow{\text{NH}_4\text{OH}} \begin{array}{c} \text{CH}_2\text{C} \\ \text{NH}_2 \\ \text{NH}_2 \end{array} \xrightarrow{-\text{NH}_4} \begin{array}{c} \text{CH}_2\text{C} \\ \text{NH} \\ \text{CH}_2\text{C} \end{array}$$

The behavior of glutaronitrile is very similar (14). Imidoximes are still able to react with nitrous acid, yielding the corresponding imides.

$$(CH_2)_{2,3} \xrightarrow{\text{NOH}} (CH_2)_{2,3} \xrightarrow{\text{NOH$$

However, these cyclic imides cannot be converted into the oximes by hydroxylamine; while glutarimide fails to react, succinimide yields disuccinimide-dihydroxamic acid (XXVIII) (42).

Imidoximes also were prepared from certain aromatic nitriles with hydroxylamine. For instance, o-cyanomethylbenzonitrile and ethyl o-cyanobenzoate, respectively, yield the imidoximes (34, 188, 107).

$$\begin{array}{c|c} CH_2CN & \stackrel{NH_2OH}{\longrightarrow} & C=NOH \\ \hline CN & \stackrel{H_2OH}{\longrightarrow} & C \\ \hline C=NOH \\ \hline CN & NH \\ \hline COOC_2H_5 & \stackrel{NH_2OH}{\longrightarrow} & C \\ \hline NH \\ \hline \end{array}$$

The latter compound is hydrolyzed easily to phthalimide, which does not react further with hydroxylamine.

Phthalonitrile yields phthalimidedioxime (22).

$$\begin{array}{c}
\text{CN} & \xrightarrow{\text{NH}_2\text{OH}} & \text{CN} & \text{NOH} \\
\text{CN} & \xrightarrow{\text{NOH}} & \text{CN} & \text{NOH}
\end{array}$$

Table 4 lists all imidoximes and imidedioximes described in the literature.

# 4. Polyamidoximes

Alkylenediaminetetraacetamidoximes recently have been synthesized and used as fungicides (11a)

TABLE 4
Imidoximes and Imidedioximes

Melting

	point,	
Compound	°C.	Ref.
NOH CHr-C		
NH CH2-C  NOH  CH2-C	197	42
CH <sub>2</sub> —C NH CH <sub>2</sub> —C	196	42
NH	250	107
CH <sub>2</sub> C=NOH NH NOH	158	34, 188
NH CH2—C NOH NOH CH3—C	198	42, 166
CH <sub>2</sub> —C NOH NOH	193	14, 42
NOH	271	22

Polymers containing the amidoxime function have been obtained by Schouteden (160a, b and c) by treating a polyacrylonitrile of low molecular weight with a slight excess of hydroxylamine. The reaction was carried out in dimethylformamide as solvent, and its kinetics have been studied.

$$\begin{bmatrix} -\text{CHCH}_2 - \\ -\text{CN} \end{bmatrix}_n \xrightarrow{+\text{NH}_2\text{OH}} \begin{bmatrix} -\text{CHCH}_2 - \\ -\text{NOH} \\ -\text{CN} \end{bmatrix}$$

# III. PROPERTIES AND REACTIONS OF AMIDOXIMES

#### 1. Physical Properties

The amidoximes are crystalline, colorless compounds which generally decompose when heated over their melting point. The melting points of amidoximes are listed in Tables 2, 3, and 4. Aryl amidoximes are more stable than the aliphatic amidoximes (193).

The first members of the aliphatic series are soluble in water but their solubility decreases with increasing molecular weight. Aryl amidoximes are less or not soluble in water but soluble in alcohol and in most organic solvents.

The infrared spectra of the amidoximes show two well defined absorption bands (36a): The first is a doublet at  $2.87-2.93~\mu$  and  $2.96-3.03~\mu$ , assigned to the two NH<sub>2</sub> stretching modes; the second between 5.95 and 6.08  $\mu$  to the C=N stretching (see Table 5). The OH stretching band of the NOH group is very broad and has its maximum at approximately  $3.2~\mu$ .

	—NH <sub>2</sub> st	$C = N$ stretching, $\mu$	
Formamidoxime	2.93	3.03	5.95
Acetamidoxime	2.87	2.98	6.03
Benzamidoxime	2.87	2.96	6.08
Oxamidedioxime	2.88	2.98	6.05
Adipodiamidoxime	2.88	2.98	6.05

Crystallographic data about amidoximes are very scarce: only those of benzamidoxime (178) and succinamidedioxime (166) have been published.

#### 2. Chemical Properties

# (a) Salt Formation

The amidoximes are amphoteric substances, soluble in dilute mineral acids as well as in aqueous alkaline solutions (193).

The amino group in the molecule confers basic properties to the amidoximes. Salts of amidoximes with mineral or organic acids are known; they are crystallized easily and have well defined melting points.

On the other hand, the hydrogen atom of the NOH group can be substituted, as in the case of oximes, by a metal. Many sodium and silver salts have been described.

Amidoximes form colored crystalline compounds with the salts of some metals (176). Werner (204) prepared a great number of such compounds with different amidoximes and proved that they are internal complexes in which the metal atom is linked to the oxime group as well as to the amino group.

These amidoximes have been used as analytical reagents for various cations (197, 198): formamidoxime (68), hydroxyisobutyramidoxime (70), benzamidoxime (30, 117), hippuramidoxime (31), phenylacetamidoxime (71, 117), o- and p-toluamidoxime (117), homoveratramidoxime (69), 2- and 4-pyridinamidoxime (117), oxamidoxime (74), oxalhydroxamamidoxime (72), oxamidedioxime (30, 113, 117), malonamidedioxime (73, 117), and succinamidedioxime (117).

Oxamidedioxime ("Niccolox"), which forms complex salts with Ni<sup>++</sup>, Cu<sup>++</sup>, Ag<sup>+</sup>, Co<sup>++</sup>, has been applied in quantitative analysis (21, 65, 66, 67).

Nicotinamidoxime can be used for the spectrophotometric determination of uranium (196).

Finally, analytical applications of amidoximes were studied recently by Pearse (117), who proposed a colorimetric method for the determination of cobalt based upon the benzamidoxime complex and a spectrophotometric method for the determination of cobalt and nickel using a single reagent, oxamidedioxime.

# (b) Organic Complexes

Amidoximes form with chloral bimolecular complexes, which are insoluble in water and soluble in organic solvents. They have sharp melting points and may be used for the identification of amidoximes (39, 55, 114, 175).

N-Phenylbenzamidoxime also forms a complex with chloral (109). However, oxamidedioxime is reported to react with chloral to give a product C<sub>8</sub>H<sub>6</sub>N<sub>4</sub>O<sub>4</sub>Cl<sub>2</sub> whose structure has not been elucidated (199).

Trichloroacetic acid yields with an aqueous solution of adipamidedioxime a crystalline precipitate, soluble in alcohol. Its formula corresponds to the addition compound of 2 molecules of trichloroacetic acid to 1 molecule of adipamidedioxime (36a). It has not been established definitely whether this substance is a salt or a molecular complex.

#### (c) Thermal Decomposition

Generally, amidoximes are decomposed when heated in the neighborhood of their melting point. Benzamidoxime, melting at 80°, is stable up to 170°. At this temperature it decomposes, yielding several products which were identified (81) as nitrogen, nitrous oxide, ammonia, water, benzonitrile, benzamide, diphenyl1,2,4-oxadiazole, diphenyl-1,2,4-triazole and triphenyl-1,3,5-triazine.

The thermal decomposition of *p*-phenylsulfonylbenzamidoxime yields *p*-phenylsulfonylbenzamide and 3,5-diphenylsulfone-1,2,4-oxadiazole.

NOH 
$$C_6H_5SO_2C_6H_4C \longrightarrow NH_2$$
 
$$N-O$$
 
$$C_6H_5SO_2C_6H_4C \longrightarrow CC_6H_4SO_2C_6H_5 \text{ and } C_6H_5SO_2C_6H_4CONH_2$$

Many amidoximes, which at room temperature form soluble salts with dilute mineral acids and alkalies, are hydrolyzed completely when heated in the same media (176). Amides and hydroxylamine are formed, and under drastic conditions the amides are hydrolyzed into the corresponding acids:

(d) Hydrolysis

At 200°, a solution of ammonium hydroxide hydrolyzes benzamidoxime into benzamide and ammonium benzoate (90). Oxamidedioxime is hydrolyzed by concentrated hydrochloric acid into oxalic acid, ammonia, and hydroxylamine (52, 53).

Polyacrylamidoximes of low molecular weight have been hydrolyzed in aqueous solution at different pH values (160c, 160d). The reaction can be limited at will to the NH<sub>2</sub> groups or extended to both NH<sub>2</sub> and NOH groups. A polymer finally is obtained which contains simultaneously hydroxamic acid, amide, and carboxylic groups.

# (e) Reduction

The reduction of benzamidoxime with sodium amalgam (193, 195) produces ammonia and benzaldoxime with a yield of only 10 to 12%; most of the amidoxime remains unchanged.

$$C_6H_5C$$
 $NH_6$ 
 $NH_6$ 
 $NOH$ 
 $NA_6(Hg)$ 
 $N_6(Hg)$ 
 $NH_6(Hg)$ 
 $NH_6(Hg)$ 
 $NH_6(Hg)$ 
 $NH_6(Hg)$ 

Phenylglyoxalamidoxime (XXIX) has been reduced to phenylethanolamine on a palladium charcoal catalyst under 10 to 20 atmospheres of hydrogen (76).

$$\begin{array}{ccc} \text{NOH} & & & \\ \text{C}_{6}\text{H}_{5}\text{COC} & \xrightarrow{\text{H}_{2}(\text{Pd})} & \text{C}_{6}\text{H}_{5}\text{CHOHCH}_{2}\text{NH}_{2} \\ & & \text{XXIX} & & & \end{array}$$

When N-substituted amidoximes are reduced under the same conditions only 3 moles of hydrogen are taken up; there is evidence that both nitrogen atoms are still present in the reaction product but the structure of these substances has not yet been established.

Amidines can be prepared by reduction of the corresponding amidoximes (7) (20a) in the presence of Raney nickel at 30 atm. and 60-80°.

$$\begin{array}{ccc} \text{NOH} & \text{NH} \\ \text{RC} & \xrightarrow{\text{H}_2(\text{Ni})} & \text{RC} \\ & \text{NH}_2 & & \text{NH} \end{array}$$

The electrolytic reduction of benzamidoxime in 3.3% HCl also yields benzamidine (7). The same compound is formed when benzamidoxime is acetylated with thioacetic acid. The hydrogen sulfide produced during the reaction reduces the amidoxime *in statu nascendi* into benzamidine (36a).

Zinc in hydrochloric acid does not reduce amidoximes. Therefore this reagent reduces *p*-nitrobenzamidoxime to *p*-aminobenzamidoxime (201).

#### (f) Oxidation

Oxidizing agents such as potassium ferricyanide, chlorine, or bromine in acetic acid, and iodine in aqueous bicarbonate react with benzamidoxime to yield a product C<sub>14</sub>H<sub>13</sub>N<sub>3</sub>O which corresponds to an amino-dihydroöxadiazole XXX (64).

Compound XXX also is obtained together with nitrous oxide and aminodiazobenzene when benzenediazonium chloride or sulfonate reacts with benzamidoxime (174). The mechanism of formation of XXX is not known.

Iodine in sodium hydroxide yields benzonitrile with benzamidoxime.

The transformation of benzamidoxime into 3,5-diphenyl-1,2,4-oxadiazole when heated with a carboxylic acid also has been interpreted as an oxidation-reduction disproportionation to XXX with subsequent loss of ammonia (see page 175) (64, 164).

The oxidation of oxamidedioxime, studied by Holleman (52), failed to yield any definite product.

# (g) O-Alkylation

As mentioned before amidoximes and their N-substituted derivatives exhibit acidic properties and form salts with metals. Sodium salts, readily obtained from sodium alcoholate, yield O-alkyl ethers (XXXI) when treated with aliphatic halogen compounds (41a, 63, 108, 193, 208a)

Instead of sodium ethoxide, potassium or sodium hydroxide in aqueous alcoholic solution can be used (202). O-Methyl derivatives have been prepared from amidoximes and methyl sulfate (6, 18).

The O-alkyl derivatives of aliphatic amidoximes are oily unstable compounds which have not yet been obtained in a pure state. Those of aromatic amidoximes are low melting stable compounds, soluble in common organic solvents, and can be prepared in good yields (63, 112, 194).

The reaction of benzamidoxime with  $\alpha,\omega$ -dihalides yielded the expected O,O'-dialkylene derivatives (7, 8, 39). The ethylene di-O,O'-benzamidoxime (XXXV, n=2) on treatment with nitrite in the presence of hydrochloric acid (see page 169) gave the corresponding chlorobenzaldoxime derivative. The latter with sodium ethoxide furnished the ethylene ester of an ethylbenzhydroxamic acid which was hydrolyzed to ethylenedioxydiamine (208a).

NOH
$$C_{6}H_{5}C + Br(CH_{2})_{n}Br \rightarrow NH_{2}$$

$$NO(CH_{2})_{n}ON$$

$$C_{6}H_{5}C \xrightarrow{NH_{2}} CC_{6}H_{5} \xrightarrow{HNO_{2}} HCl$$

$$NH_{2} H_{2}N$$

$$XXXV$$

$$NO(CH_{2})_{n}ON$$

$$C_{6}H_{5}C \xrightarrow{NaOC_{2}H_{5}} Cl$$

$$Cl Cl$$

$$NO(CH_{2})_{n}ON$$

$$C_{6}H_{5}C \xrightarrow{NaOC_{2}H_{5}} HCl$$

$$CC_{6}H_{5} \xrightarrow{HCl} CC_{6}H_{5} \xrightarrow{HCl} HCl$$

$$CC_{6}H_{5}C \xrightarrow{NaOC_{2}H_{5}} CC_{6}H_{5} \xrightarrow{HCl} CC_{6}H_{5}COC_{2}H_{5}$$

$$CC_{6}H_{5}C \xrightarrow{HCl} CC_{6}H_{5}COC_{2}H$$

However this method could not be applied to the preparation of other alkylenedioxydiamines, since all attempts to convert hexamethylene and dodecamethylene di-O,O'-benzamidoxime (XXXV, n=6, 12) into the chloride oximes failed completely (7).

The similar reaction of benzamidoxime with an  $\alpha$ -halogenated carboxylic ester has been used for the synthesis of  $\alpha$ -hydroxylaminocarboxylic acids (209).

NOH
$$C_{6}H_{5}C \xrightarrow{\text{NOH}_{2}} + \text{ClCH}_{2}\text{COOC}_{2}H_{5} \xrightarrow{\text{OH}^{-}} \\ \text{NH}_{2} \xrightarrow{\text{NOCH}_{2}\text{COOH}} \\ C_{6}H_{5}C \xrightarrow{\text{NH}_{2}} + \text{H}_{2}\text{NOCH}_{2}\text{COOH}$$

The  $\alpha$ -hydroxylamine derivatives of acetic, propionic, butyric, and isobutyric acids have been prepared by this method (205, 208).

It should be mentioned that the acid formed in the first step is always accompanied by its corresponding lactam, 3-phenyl-5,6-dihydro-1,2,4-oxadiazine-5-one

The 6-substituted homologs also have been obtained (59, 205, 208) with ethyl chlorofumarate and benzamidoxime or  $\alpha$ -phenylacetamidoxime; only the cyclic product is isolated (216).

NOH Cl
$$C_6H_6C + C = CHCOOC_2H_6 \rightarrow NH_2 C_2H_6OOC$$

$$N = O$$

$$C_6H_6C NH = O$$

$$C = CHCOOC_2H_6$$

$$NH = O$$

O-Alkyl substituted sulfamidobenzamidoximes have been obtained by an altogether different method (3). An alcoholic solution of sulfamidobenziminoethyl ether hydrochloride (XXXII) is treated with an O-alkylhydroxylamine, at 37° for a fortnight in a pressure bottle. NH<sub>4</sub>Cl separates from the reaction mixture and two products can be isolated from the solution. The first, insoluble in dilute hydrochloric acid, is the dialkyl ether of sulfamidobenzoxyamidoxime (XXXIV). The second, soluble in diluted HCl, is the O-alkyl derivative of the amidoxime (XXXIII). For example, O-methyl hydroxylamine reacts

$$\begin{array}{c} p\text{-}\mathrm{NH_2SO_2C_6H_4C} \\ \\ XXXII \\ \mathrm{NH \cdot HCl} \\ \\ p\text{-}\mathrm{NH_2SO_2C_6H_4C} \\ \\ XXXIII \\ \mathrm{NH_2 \cdot HCl} \\ \\ \\ P\text{-}\mathrm{NH_2SO_2C_6H_4C} \\ \\ XXXIII \\ \mathrm{NH_2 \cdot HCl} \\ \\ \\ P\text{-}\mathrm{NH_2SO_2C_6H_4C} \\ \\ \\ XXXIV \\ \mathrm{NHOCH_3} \\ \\ \\ \end{array}$$

The O-alkyl amidoximes are listed in Table 6.

	TABLE 6	NO B	
O-Alkulated	Amidoximes R—C	NO—R′	
0 12g	12	NIII	
		$ m NH_2$	
R	R'	M. p., °C.	Ref.
CH.	—CH;	Unst. oil	112
CH-	−C₂H₅	Unst. oil	112
CH₃— (CH₃)₂CHCH₂—	$-CH_2C_6H_6$ $-C_2H_5$	Unst. oil 35	112 55
CH <sub>2</sub> C(: NOCH <sub>2</sub> )—	-CH <sub>2</sub>	B.p. 192	6
C <sub>6</sub> H <sub>5</sub> —	-CH.	57 B.p. 230	
C6H6—	—C <sub>2</sub> H <sub>5</sub>	67	63, 194
C <sub>6</sub> H <sub>6</sub> —	—(CH₂)₅CH₃	49-50	41a
C <sub>6</sub> H <sub>6</sub> — C <sub>6</sub> H <sub>6</sub> —	—CH <sub>2</sub> CH <sub>2</sub> — —(CH <sub>2</sub> ) <sub>6</sub> —	155–156 106	39, 208a 41a
C <sub>6</sub> H <sub>5</sub> —	—(CH <sub>2</sub> ) <sub>12</sub> —	105	4la
C <sub>6</sub> H <sub>5</sub> —	—CH <sub>2</sub> C <sub>6</sub> H <sub>5</sub>	90.5	63
$C_6H_5$ —	$-CH_2C_6H_4NO_2-p$	105-106	202
C <sub>6</sub> H <sub>5</sub> —	—CH <sub>2</sub> COOH	123-124	59
C <sub>5</sub> H <sub>5</sub> —	—СН(СН <sub>1</sub> )СООН	168ª	209
C <sub>6</sub> H <sub>5</sub> — C <sub>6</sub> H <sub>5</sub> —	-CH(C2H5)COOH-CH(C2H5)COOC2H1	81-82 57	208 208
C <sub>6</sub> H <sub>5</sub> —	—C(CH <sub>1</sub> ) <sub>2</sub> COOH	182–185°	205
СеНе—	-C(CH <sub>2</sub> ) <sub>2</sub> COOC <sub>2</sub> H <sub>3</sub>	37-38	205
o-CH <sub>1</sub> C <sub>6</sub> H <sub>4</sub> —	$-C_2H_5$	140	162
p-CH <sub>1</sub> C <sub>6</sub> H <sub>4</sub> —	—СH.	85	161
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	—C₂H₅	64	161
C <sub>6</sub> H <sub>5</sub> CH <sub>2</sub> — C <sub>6</sub> H <sub>6</sub> CH <sub>2</sub> —	—C₂H₃ —CH₂C₅H₅	58 55	57 57
Oin On	-011206116	00	01
CH3—	—C <sub>2</sub> H <sub>5</sub>	172	114
℃H₃			
C₀H₃CH≕CH—	—СН.	98	217
C6H6CH=CH-	-C <sub>2</sub> H <sub>6</sub>	83	217
	—C <sub>2</sub> H <sub>5</sub>	74–75	150, 151
N	—CH₂C₃H₃	80	99
	$-C_2H_5$	85	13
0-HOC6H4—	—C <sub>2</sub> H <sub>5</sub>	B.p. 220	168
C <sub>6</sub> H <sub>6</sub> CHOH—	—C <sub>2</sub> H <sub>5</sub>	(15 cm.) 89	50
C <sub>6</sub> H <sub>6</sub> CHOH—	—CH2C6H5	102-103	50
p-CH <sub>2</sub> OC <sub>6</sub> H <sub>6</sub> —	—C <sub>2</sub> H <sub>5</sub>	51-52	100
0-C2H6OC6H4—	—C <sub>2</sub> H <sub>5</sub>	B.p. 195	168
		(18 cm.)	
m-C <sub>1</sub> H <sub>5</sub> OC <sub>5</sub> H <sub>4</sub> —	—C <sub>2</sub> H <sub>5</sub>	109	23
p-C2H5OC6H4— m-NO2C5H4—	—C <sub>2</sub> H <sub>5</sub> —CH <sub>3</sub>	84 75	62 18
m-NO <sub>2</sub> C <sub>5</sub> H <sub>4</sub> —	—C <sub>2</sub> H <sub>5</sub>	Oil	159
m-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	—CH₂C6H4	58	159
p-NO₂C6H4—	$-C_2H_5$	<b>59–6</b> 0	201
p-NH <sub>2</sub> SO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	—CH:	214-215a	3
p-NH <sub>2</sub> SO <sub>2</sub> C <sub>1</sub> H <sub>4</sub> —	—C₂H₅	221-2224	3
p-NH₁SO₂C₅H₄—	—(CH <sub>2</sub> ) <sub>3</sub> CH <sub>3</sub>	127-128 136-137ª	3
p-NH <sub>2</sub> SO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> — p-CH <sub>3</sub> C <sub>6</sub> H <sub>4</sub> NHC(:NH)—	—(CH <sub>2</sub> )₄CH₄ —C <sub>2</sub> H <sub>5</sub>	132–133	3 199
p-CH <sub>4</sub> C <sub>6</sub> H <sub>4</sub> NHC(:NH)—	—CH2C6H5	165	199
R'—			
	C—R—C		
_			
	H <sub>2</sub> N NH <sub>1</sub>	147	
— (oxamidedioxime)	—CH <sub>1</sub>	144	6
- (oxamidedioxime)	—C₃H₅	114-115	221
—CH₂CH₂—	—C2H3	119	166

a Isolated as hydrochloride.

#### (h) Action of Nitrous Acid

The reaction of nitrous acid on benzamidoxime has been one of the methods which helped to elucidate the structure of the amidoxime group.

Tiemann and Krüger (193) identified the presence of the NOH group by the fact that benzamidoxime, as hydroxylamine itself, evolves nitrous oxide when treated with an equivalent amount of nitrous acid. Under these conditions, benzamidoxime is transformed into benzamide. Acetamidoxime reacts similarly yielding acetamide and nitrous oxide (112).

However, an O-alkylated benzamidoxime treated with nitrous acid evolves nitrogen, which proves that an amino group is present in the molecule (112). In this reaction the corresponding hydroximic acid (XXXVI) is not isolated, but in the presence of an excess hydrochloric acid, the hydroximic acid chloride (XXXVII) is formed in almost quantitative yields (18, 57, 63, 162, 168, 193, 194, 202, 218).

O-Alkyl hydroximic acid chlorides are very stable oils which can be steam distilled without decomposition. They are soluble in most organic solvents.

Hydroximic acid bromides and fluorides are prepared in a similar way if hydrobromic or hydrofluoric acid is used instead of hydrochloric acid in the preceding reaction (162, 189, 203).

Nitrites of the type

(189, 201, 203, 218) are formed when O-alkyl amidoximes are treated with nitrous acid in the presence of sulfuric acid. These very unstable substances explode on heating.

In the presence of acetic acid, the formation of the acetate

is reported by Tiemann (189) but it could not be isolated in a pure state.

When treated with an alcoholic solution of ammonia,

6 to 8 hours at 160–180°, O-alkylated hydroximic acid halides are converted back to the corresponding amidoximes (194).

Unsubstituted hydroximic acid chlorides generally are prepared by direct chlorination of of aldoximes

The action of ammonia on the latter compounds already has been mentioned (p. 161). With NaN<sub>3</sub>, azidoximes (azides of hydroximic acids) are formed.

NOH NOH NOH RC + NaN<sub>3</sub> 
$$\rightarrow$$
 RC + NaCl N<sub>4</sub>

The azide structure of these compounds, heretofore erroneously formulated as hydroxytetrazoles, has been proved recently (36d).

The exceptional action of nitrous acid on the oxime of N-phenyloxamic acid ethyl ester XXXVIII was particularly studied by Jovitschitsch (56) and the structure of the oxidation product formed was proved twenty-five years later by Semper and Lichtenstadt (167) to be 6-nitro-4,1,3-benzoxadiazine.

# (i) N-Substitution

One of the methods outlined below may be used to replace one or both hydrogens of the amino group by an alkyl or aryl radical.

a. Action of an amine on hydroximic acid halides is the usual way of preparing N-substituted amidoximes. Primary and secondary, aliphatic and aromatic amines have been used.

NOH NOH RC + 2 NH<sub>2</sub>R' 
$$\rightarrow$$
 RC + R'NH<sub>2</sub>·HCl NHR'

The reaction is carried out in absolute ethanol or in ether at room temperature and an excess of amine is used to neutralize the hydrochloric acid formed.

b. Action of hydroxylamine on N-substituted thioamides

$$\begin{array}{c} S \\ \parallel \\ RCNHR' + NH_2OH \rightarrow RC \\ NHR' \end{array} + H_2S$$

TABLE 7

NOH

N-Monoalkylated Amidoximes R—C

R	R'	M.p., °C.	Method	Yield, %	Ref.
H—	—C <sub>6</sub> H <sub>5</sub>	138	d	62	110
			b		109
			8.		110
CH.	$-C_5H_5$	120-121	d		112
			b		109
C <sub>2</sub> H <sub>5</sub> OCO—	$-C_6H_6$	109	8.		56
CH <sub>3</sub> C(:NOH)—	$-C_{\mathfrak{b}}\mathbf{H}_{\mathfrak{b}}$				139
C <sub>6</sub> H <sub>6</sub> —	$-CH(CH_1)_2$	Viscous oil	a	90	76
$C_6H_6$ —	CH(CH3)CH2CH2N(C2H5)3		a		7
C <sub>6</sub> H <sub>5</sub> —	$-C_6H_5$	136	b		108
C6H6-	$-C_0H_4CH_{FO}$	147	b	Poor	174
C <sub>6</sub> H <sub>6</sub> —	$-C_6H_4CH_{4-}p$	176	b	Poor	109
		161-162	8.		149
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	$-C_0H_4CH_2-p$	134-135	a		149
$p-(CH_1)_2CHC_5H_4$	—C₀H₀	145-146	a		149
( <sub>o</sub> )	$-C_6H_6$	126–127	8.		149
o-ClCoH-	—С <sub>6</sub> Н <sub>6</sub>	140	8.		206
o-ClC <sub>6</sub> H₄—	-CoH4CH3-0	173	a.		206
p-ClC <sub>6</sub> H <sub>4</sub> —	$-C_0H_4CH_{1}-p$	169.5-170	8.		149
O CH <sub>2</sub> -O	—С <sub>6</sub> Н <sub>4</sub> СН <sub>7</sub> -р	150–151	a		149
C <sub>6</sub> H <sub>6</sub> CO—	—CH <sub>1</sub>	13 <b>2</b> –133°	8.	80	76
C <sub>6</sub> H <sub>6</sub> CO—	—C <sub>2</sub> H <sub>5</sub>	Viscous oil	8.	79	76
C <sub>6</sub> H <sub>6</sub> CO—	-CH2CH2CH1	Viscous oil	a.	68	76
C <sub>6</sub> H <sub>5</sub> CO—	—(CH <sub>2</sub> ) <sub>2</sub> CH <sub>3</sub>	120-125a	a.	85	76
C <sub>6</sub> H <sub>6</sub> CO—	—(CH <sub>2</sub> ) <sub>4</sub> CH <sub>4</sub>	$123-125^a$	a.	70	76
C <sub>6</sub> H <sub>5</sub> CO—	—C <sub>6</sub> H <sub>4</sub>	142	a.	62	76
C <sub>6</sub> H <sub>6</sub> CO—	—CH₂C₀H₅	Viscous oil	a,	82	76
C <sub>6</sub> H <sub>6</sub> CO—		178–179	a	93	76
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> CO—	—СH <sub>1</sub>	124	a	75	76
p-CH <sub>3</sub> C <sub>6</sub> H <sub>4</sub> CO—	—C <sub>6</sub> H <sub>6</sub>	163-164	a	95	76
C <sub>6</sub> H <sub>5</sub> C(:NOH)—	—СН.	α 178	c		87
		β 159-160	a		87
$C_6H_6C(:NOH)$ —	—C <sub>6</sub> H <sub>5</sub>	α 187–188	c		139, 141
,		β 124	c, a		139
		191–192	a,		129
C <sub>6</sub> H <sub>5</sub> C(:NOH)—	$-CH_2C_0H_5$	α 174	c		87
		β 158–159	a.		87
C6H5C(:NOCOCH3)—	—CH <sub>1</sub>	160	"в		87
C <sub>6</sub> H <sub>5</sub> NHC(:NOH)—	—C <sub>6</sub> H <sub>5</sub>	218	a.c		128

<sup>&</sup>lt;sup>a</sup> Isolated as hydrochloride. <sup>b</sup> Formed by the action of acetic anhydride on C<sub>0</sub>H<sub>0</sub>C(:NOH)C(:NOH)NH<sub>1</sub>. <sup>c</sup> Prepared from chlorobromoglyoxime.

The yields are less than in the case of unsubstituted thioamides (108, 109, 174).

c. Action of amines on glyoxime peroxides yields N-substituted aminoglyoximes (XXXIX) (87, 139, 141) (see p. 162).

$$R(C_2N_2O_2)H + R'NH_2 \rightarrow RC$$
——CNHR'

NOH NOH

(XXXIX)

Different isomers of aminodioximes (XXXIX) have been isolated but their configuration is not definitively established (87).

d. Action of Aniline on Amidoximes.—Aniline is reported to react directly on acetamidoxime and formamidoxime hydrochloride; at 80-90° ammonia is

evolved with formation of the corresponding N-substituted amidoximes (110, 112).

NOH NOH RC + 
$$C_6H_6NH_2 \rightarrow RC + NH_5$$

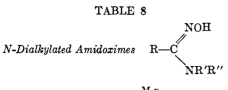
$$NH_2 \qquad NHC_6H_5$$

No other examples of the direct action of an amine on an amidoxime are known.

A series of N-substituted amidoximes with their melting points, methods of preparation, and references are listed in Tables 7 and 8.

#### (j) Action of Isocyanates and Isothiocyanates

According to Tiemann (193) cyanic acid and phenyl isocyanate react with benzamidoxime to yield ureide or



R	NR'R"	М.р., °С	Method	Yield	Ref.
C <sub>6</sub> H <sub>5</sub> — o-ClC <sub>6</sub> H <sub>4</sub> —	Piperidyl Piperidyl	165	a a		7 206
OCH <sub>2</sub> -O	Piperidyl	181–182	а		149
m-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	-N(CH <sub>2</sub> ) <sub>2</sub>	160	a	57	18
$m-NO_2C_0H_4$ —	-N(CH <sub>3</sub> )(C <sub>2</sub> H <sub>5</sub> )	123	8.		18
$C_6H_6CO$ —	Morpholinyl	98	a,	2	76
C <sub>6</sub> H <sub>5</sub> C(:NOH)—	-N(CH <sub>2</sub> CH <sub>2</sub> CH <sub>3</sub> ) <sub>2</sub>	62-66	a		207

phenylureide oximes. However, no proof was ever given for the structure of these compounds which could be as well the isomeric carbamates (176, 177)

In the case of O-alkyl amidoximes, isocyanates can evidently yield only ureide oximes (51, 58, 218).

NOR'
$$+ C_6H_6NCO \rightarrow RC$$

$$NH_2$$
NHCONHC6H6

Benzanilidoxime hydrochloride is reported to react with potassium cyanate (108). The formula proposed for the reaction product is that of an ureide oxime (XLI) but the compound is completely insoluble in alkali and therefore it is more probable that the correct structure is that of a carbamate (XLII).

$$C_6H_5C \xrightarrow{NOH} C_6H_5 \cdot HCl \xrightarrow{KOCN} \begin{cases} NOH \\ C_6H_5C & XLI \\ N-C_6H_5 \\ CONH_2 \\ Or \\ NOCONH_2 \\ C_6H_5C & XLII \\ NHC_6H_5 \end{cases}$$

An example in which the isonitroso group is reported to form a carbamate when treated with phenyl isocyanate is given by Longo (84), who transformed phenylaminoglyoxime into a trisubstituted compound XLIII.

$$\begin{array}{c} \text{NOH} \\ C_6H_5C(:\text{NOH})C & \xrightarrow{C_6H_5NCO} \\ \\ \text{NH}_2 & \text{NOCONHC}_6H_5 \\ \\ C_6H_5C(:\text{NOCONHC}_6H_5)C & \\ \\ \text{NHCONHC}_6H_5 \\ \\ \text{XLIII} \end{array}$$

The formation of carbamates also was observed by Ponzio (132), who claimed that the  $\alpha$  and  $\beta$  forms of the arylaminoglyoximes are structural isomers containing respectively the hydroxamino and the isonitroso groups. For example,  $\alpha$ -phenylaminoglyoxime should have the structure XLIV and the  $\beta$  isomer the structure XLV (see pp. 156 and 176).

According to Ponzio, the β-form treated with cyanic acid gives a ureide oxime XLVI stable in boiling water and ethanol, while the α-form gives a carbamate which in boiling ethanol easily loses CO<sub>2</sub> and NH<sub>3</sub> to yield the phenylaminofurazan XLVII (132)

$$\begin{array}{c} \text{NHOH} \\ \text{Cl}_{\theta}\text{H}_{\delta}\text{C-C} \\ \text{NOH} \\ \text{NH} \\ \text{NOH} \\ \text{NO$$

When phenyl isothiocyanate and benzamidoxime react in equimolecular amounts and the reaction is carried out at room temperature, benzoylphenylthiourea oxime (XLVIII) is formed (60, 63).

$$\begin{array}{c} \text{NOH} \\ \text{C}_6\text{H}_5\text{C} \\ \text{NH}_2 \\ \\ \text{NOH} \\ \text{C}_6\text{H}_5\text{C} \\ \\ \text{NHCSNHC}_6\text{H}_5 \\ \\ \text{XLVIII} \\ \end{array} \begin{array}{c} \text{N-S} \\ \text{C}_6\text{H}_5\text{C} \\ \\ \text{N} \\ \text{XLIX} \\ \end{array}$$

When two moles of isothiocyanate is used for one mole of benzamidoxime and the reaction is carried out in boiling chloroform hydrogen sulfide is evolved and 3-phenyl-5-anilino-1,2,4-thiadiazole (XLIX) is formed.

On the contrary, under similar conditions, p-tolyl isothiocyanate forms the corresponding thiourea oxime, but by cyclization 3-phenyl-5-p-toluidino-1,2,4-oxa-diazole (L) is produced (60).

Finally, with allyl isothiocyanate the thiourea derivative is isolated but no cyclization occurs (60).

NOH 
$$C_{6}H_{5}C + CH_{2}\!\!=\!\!CHNCS \rightarrow \\ NH_{2} \\ NOH \\ C_{6}H_{5}C \\ NHCSNHCH_{2}CH\!\!=\!\!CH_{2}$$

Gheorghiu and Barbos (43) observed that when benzamidoxime reacts with an excess of phenyl isothiocyanate, free sulfur readily is produced, which shows that autoxidative phenomena probably take place.

Most of the known ureidoximes and thioureidoximes are listed in Tables 9 and 10.

TABLE 9

NO—R"

O-Alkyl Derivatives of Ureide Oximes RC

NH—CO—NH—R

R R' R" \*C. Ref.

R	R'	R"	м.р., •С.	Ref.
C <sub>6</sub> H <sub>6</sub> CH <sub>5</sub> —	—С <sub>в</sub> Н <sub>5</sub>	—С2Н5	148	58
C <sub>1</sub> H <sub>5</sub> CH=CH—	—С <sub>с</sub> Н <sub>5</sub>	—С2Н5	155-156	218
C <sub>6</sub> H <sub>6</sub> CHOH—	—С <sub>с</sub> Н <sub>5</sub>	—С2Н5	119	51

# (k) Acylation

O-Acylated Amidoximes.—Amidoximes can be acylated readily at room temperature by acid chlorides or anhydrides.

That the reaction occurs on the isonitroso group is proved by the fact that the acyl derivatives still show basic properties whereas the acidic behavior of the amidoximes has disappeared completely (175). Also the infrared spectra of the acyl derivatives show the presence of the NH<sub>2</sub> and —O—CO-groups and the absence of the broad OH-absorption band at about  $3.2~\mu$  (36c).

Tables 11, 12, 13, 14 list most of the O-acylated amidoximes, diamidoximes, and imidoximes described in current literature. The methods of acylation are all

Т	ABLE 10		
NOH		NOCO	NHR'
Ureide Oximes RC	or Carbamates	$\mathrm{RC}^{/\!\!/}$	
NHCO	NHR'	$ m NH_2$	
		M.p.,	
R	R'	°C.	Ref.
HC(:NOH)—	—н	157	144
HC(: NOCONHC H.)	—C₅H₅	172	84
CH <sub>5</sub> C(: NOCONHC <sub>5</sub> H <sub>5</sub> )— C <sub>6</sub> H <sub>5</sub> —	—С <b>₅Н₅</b> —Н	191-192	84
CoHs—	— ⊓ —C₄H₅	115 115	39 63
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	—H	170	162
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	—C₀H₅	155	162
	V024V	100	102
CH <sub>3</sub>	—н	155	114
CH <sub>3</sub> CH <sub>3</sub>	$-C_{\mathfrak{b}}H_{\mathfrak{b}}$	138	114
CeHeCH2—	$-C_6H_5$	123	57
C <sub>6</sub> H <sub>5</sub> CH=CH−	—н	158-159	218
$C_6H_5CH=CH-$	$-C_6H_5$	158-159	218
C <sub>6</sub> H <sub>6</sub> CHOH—	—н	127	51
$C_6H_6CHOH$ —	$-C_6H_5$	155	51
o-HOC₅H←	<b>—</b> н	148	168
o-HOC₀H₄—	$-\underline{\mathbf{C}}_{6}\mathbf{H}_{5}$	119	168
$p-NH_2SO_2C_6H_4$	— <u>н</u>	202	3
C6H6C(:NOH)—	— <u>н</u>	185	132
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> C(:NOH)—	—H	195–196	132
C <sub>6</sub> H <sub>6</sub> C(: NOCONHC <sub>6</sub> H <sub>6</sub> )—	$-C_{5}H_{5}$	190	84
	—C <sub>5</sub> H <sub>6</sub>	167	99
	—н	164.5	13
HON I	мон		
non	101		
C-R-C		or	
R'HN—CO—HN	NH-CO-NHR'		
	R'NHCOON	NOCO	NHR'
		-R $-$ C	
	/		
	H <sub>2</sub> N	NH2	
— (oxamidedioxime)	—н	191-192	221
—CH₂CH₂—	—н	163.5	166
	NOH		
Thioureide Oximes	R—C		
	NH—C	S-NHR'	
CH			20
C.H.	—CH <sub>2</sub> CH=CH <sub>2</sub> —C <sub>6</sub> H <sub>6</sub>	71 172	60 60. 6 <b>3</b>
C6H6— C6H6—	$-C_6H_6CH_{2^*}p$	67	60
p-CH;C <sub>6</sub> H,—	$-C_6H_6$	190	16 <b>2</b>
$^{\mathrm{CH_{3}}}$ $^{\mathrm{CH_{3}}}$	—С₅Н₅	150	114

based on well-known classical procedures using either an acid chloride, or an anhydride. However, it is noteworthy that formamidoxime yields only dibenzhydroxamic acid when treated with benzoyl chloride (91). The expected O-benzoyl derivative is obtained when benzoic anhydride is used as an acylating agent (36c).

Instead of acid anhydrides and chlorides, other reagents have been used, such as ketene, mixed carbox-

# O-Acyl Derivatives of Amidoximes R—C NO—R'

R	R'	M.p., °C.	Ref.	R	R'	M.p., °C.	Ref.
R = hydrocarbon				0-N			<del></del>
radical				C <sub>6</sub> H <sub>8</sub> —C C—	0.17	900	
<u>H</u>	— <u>н</u>	Viscous oil	103a	C <sub>6</sub> H <sub>5</sub> —C C—	$-C_6H_5$	206	3 <b>2</b>
H	−CH₃	77.5	36c				
H—	—C <sub>6</sub> H <sub>5</sub>	115–120	36c	N			
H	$-C_6H_4Br-p$	120	36c		31 1		
CH⊷	—H	29	36a	R = substituted hydrocarbon			
CH⊱	—CHt —C6H5	96 108	81a 36a	o-ClC <sub>6</sub> H <sub>4</sub> —	—C <sub>6</sub> H <sub>5</sub>	162	206
CH₂— (CH₂)₂CHCH₂CH₂—	—CfH <sub>1</sub>	87	55	p-BrC <sub>6</sub> H <sub>4</sub> —	—CH₃	145	22
(CH <sub>3</sub> ) <sub>2</sub> CHCH <sub>2</sub> CH <sub>2</sub> —	-CH <sub>2</sub> CH <sub>2</sub> CH(CH <sub>1</sub> ) <sub>2</sub>	115	55	p-BrC <sub>6</sub> H <sub>4</sub> —	—C₅H₅	161	22
(CH <sub>2</sub> ) <sub>2</sub> CHCH <sub>2</sub> CH <sub>2</sub> —	—C <sub>6</sub> H <sub>6</sub>	105106	55	CH <sub>5</sub> CHOH—	—C₅H₅	188–189	157
C <sub>6</sub> H <sub>5</sub> —	—U	103-100	36a	C.H.CHOH—	—CH₁	140	50, 175
C <sub>6</sub> H <sub>5</sub> —	—CH:	96	164	C6H6CHOH—	—C₅H₅	149	50, 175
C <sub>6</sub> H <sub>6</sub> —	—C <sub>2</sub> H <sub>5</sub>	93	164	o-HOC₀H.—	-CH <sub>3</sub>	117	168
C <sub>6</sub> H <sub>6</sub> —	—C2H7	94	164	o-HOC6H4 m-HOC6H4—	—C <sub>6</sub> H <sub>5</sub> —CH <sub>3</sub>	173 90	168
C <sub>6</sub> H <sub>6</sub> —	—C <sub>6</sub> H <sub>5</sub>	140-148	22, 193	m-HOC6H4—	—C <sub>6</sub> H <sub>6</sub>	166	23
C <sub>6</sub> H <sub>6</sub> —	-СОСООН	159ª	219	p-HOC <sub>6</sub> H <sub>4</sub> —	—CH2	122.5	62 62
C <sub>6</sub> H <sub>5</sub> —	—COCOOC₂H₅	118	219		—C111	122.5	04
o-CH₃C6H₄—	—C <sub>6</sub> H <sub>5</sub>	145	162	OH	CIT	1.0.1.0	
o-CH2C6H4-	—C6H4CH≥0	117-118	174	~~\	—CH₃	148–149	45
p-CH3C6H4-	—CH <sub>5</sub>	132	22	CH3 OH			
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	—CH₂Br		94	/ \ <u>\</u>	—C <sub>6</sub> H <sub>5</sub>	181-182	45
p-CH <sub>8</sub> C <sub>6</sub> H <sub>4</sub> —	—CH₂CN		94	CH3	—Cens	181-182	40
p-CH <sub>3</sub> C <sub>6</sub> H <sub>4</sub> —	—CH <sub>2</sub> —	188-189	96	0	~**		
$p\text{-}\mathrm{CH}_{5}\mathrm{C}_{6}\mathrm{H}_{4}$	$-C_6H_6$	173	161, 22	o-CH <sub>2</sub> OC <sub>6</sub> H <sub>4</sub> —	-CH:	130	22
$p\text{-}CH_8C_6H_4$ —	$-CH_2C_6H_5$	146-147	96	p-CH₃OC₀H←	—CH₃	106	100
$p\text{-}\mathrm{CH}_8\mathrm{C}_6\mathrm{H}_4$ —	$-CH_2C_6H_4NO_2-p$	160-161	96	p-CH₃OC₅H₄—	—C <sub>6</sub> H₅	148	100
$C_6H_5CH_2$ —	—CH₃	124	57	HOCOCH2	—C₅H₅	135	103
$C_6H_6CH_2$ —	$-C_6H_5$	144	57	C6H18CH(OCOC6H6)—	—C₀H₅	143	157
				$o\text{-}CH_2OCOC_6H_4$ — $C_6H_6CH(OCOCH_2)$ —	—CH₃	110	168
Сн₃∥ У—	-CH.	189	114	C <sub>6</sub> H <sub>6</sub> CH(OCOCH <sub>5</sub> )—	—CH3 —C4H4	113	50 50
CH <sub>3</sub>	<b>411</b>	200		o-CeHsOCOCeH4—	—C <sub>6</sub> H <sub>6</sub>	165	50
				m-C <sub>6</sub> H <sub>5</sub> OCOC <sub>6</sub> H <sub>4</sub> —	—C <sub>6</sub> H <sub>5</sub>	127 152.5	168 23
CH <sub>3</sub> ( )—	$-C_6H_{\bar{\mathfrak{g}}}$	158	114	p-C <sub>6</sub> H <sub>5</sub> OCOC <sub>8</sub> H <sub>4</sub> —	$-C_6H_5$	185	
CH <sub>3</sub>				p-cantococant—	-C6H6	180	62
$C_6H_5CH=CH-$	$-C_6H_5$	160	217	<b>/</b> \_	—C <sub>6</sub> H <sub>5</sub>	104	110
I I				CH3 OCOC <sub>6</sub> H <sub>5</sub>	—C6116	164	116
	-CH	129	150, 151	•			
	—CH	128	100, 101	CH <sub>3</sub>	~ ~		
	1			<u> </u>	$-C_6H_5$	143	45
				OCOC6H2			
		228	36	$p\text{-NO}_2\text{C}_6\text{H}_4\text{CH}_2$	—СН.	145	22
				p-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> CH <sub>2</sub> —	$-C_6H_5$	148	22
~ ^ .				C <sub>6</sub> H <sub>6</sub> CH <sub>2</sub> CH(NHCOCH <sub>2</sub> )—	-CH2	DL 160-162	118
	—CH₃	154	150, 151	C6H5CH2CH(NHCOC6H5)—	$-C_6H_5$	DL 206-207	118
			,	,		L 204-211	118
^^/				CNCH2—	$-CH_3$	142	158
	$-C_6H_5$	179	150, 151	CNCH <sub>2</sub> —	$-C_6H_6$	184182	158
				$p\text{-CNC}_6\text{H}_4\text{CH}_2$ —	$-C_6H_5$	172	154
				CH(:NOH)—	$-C_6H_5$	146-147	
R = heterocyclic radical				CH <sub>2</sub> C(:NOH)—	—C <sub>6</sub> H <sub>5</sub>	158	130
<b>◇</b> .				C6H5C(:NOH)—	$-C_6H_5$	β 168-169	
	—CH <sub>1</sub>	143-147	22 00	CH <sub>2</sub> C(:NOCH <sub>2</sub> )—	$-C_6H_5$	134-135	6
L North	—0m	140-147	22, 55	CH(: NOCOCH <sub>8</sub> )—	—CH₃	82	54, 144
IN				CH(:NOCOCH <sub>3</sub> )—	$-C_6H_5$	166-167	144
	—C <sub>6</sub> H <sub>5</sub>	194	22, 99	$CH^{3}C(:NOCOCH^{3})$ —	—CH₃	123	146
	-00110	194	44,00	CH <sub>2</sub> C(:NOCOCH <sub>2</sub> )—	$-C_6H_6$	143	130
N				$C_5H_5C(:NOCOCH_3)$ —	—CН <sub>5</sub>	β 133-138	132, 140
				$C_6H_6C(:NOCOCH_2)$ —	$-C_6H_5$	β 139-140	130
	-CH	115	13	CH(:NOCOC6H6)—	$-CH_3$	145–146	142
N.	<del>-</del>			CH(: NOCOC <sub>6</sub> H <sub>6</sub> )—	$-C_6H_5$	186-187	
				CH <sub>3</sub> C(:NOCOC <sub>6</sub> H <sub>6</sub> )—	$-C_6H_5$	206	130
NNC6H5				$C_6H_6C(:NOCOC_6H_6)$ —	$-C_6H_5$	<b>α</b> 190	140
11						β 185–186	
CHC C-	-СH	148	15				140
\ //				$C_6H_6COC(:NOCOC_6H_6)$ —	—C <sub>6</sub> H <sub>5</sub>	158-159	143
N				C <sub>6</sub> H <sub>6</sub> COONHCOCH <sub>2</sub> —	C <sub>6</sub> H <sub>5</sub>	165	103
VIII 0 0	~			$C_6H_6-N=N-$	—CH₃	168	86
NH:CC-	CH <sub>1</sub>	193-194	83	$C_6H_5-N=N-$	—C₅H₅	191–192	86
II II				$p\text{-}CH_3C_6H_4-N=N-$	$-CH_3$	193-194	86
NT NT				OTT O TT - **	A	100	
N N				p-CH <sub>3</sub> C <sub>6</sub> H <sub>4</sub> —N=N—	—C <sub>6</sub> H <sub>5</sub>	192–193	86
N N				$p-CH_3C_6H_4-N=N-$ $p-CH_8SC_6H_4-$ $p-NH_2SO_2C_6H_6-$	$egin{array}{l} \mathbf{C}_6 \mathbf{H}_5 \ \mathbf{C}_6 \mathbf{H}_5 \ \mathbf{C} \mathbf{H}_2 \mathbf{C} \mathbf{l} \end{array}$	192–193 210	86 20 12

<sup>&</sup>lt;sup>a</sup> Prepared by saponification of the ethyl ester.

TABLE 12
O-Diacyl Derivatives of Diamidoximes

R′	M.p., °C.	Ref.
—н	175–176	36a
—CH₃	184187 206212	40, 128, 221 54
$-COOC_2H_5$	168-169	36a
$-C_6H_5$	217-222	52, 221
—СH:	153-159	158
$-C_6H_5$	183-185	158
$-CH_8$	167-168	166
$-C_6H_b$		166
—CH3	115	14
—CH <sub>1</sub>	129	75
—CH₃	161.5–162	154
$-C_6H_6$	184	154
	HCHsCOOC2HsC6HsCHiC6HsCHsCHsCHsCHsCHsCHsCHs	R' °C.  —H 175–176  —CH <sub>4</sub> 184–187 206–212 —COOC <sub>2</sub> H <sub>5</sub> 168–169 —C <sub>6</sub> H <sub>5</sub> 217–222 —CH <sub>1</sub> 153–159 —C <sub>6</sub> H <sub>6</sub> 183–185 —CH <sub>8</sub> 167–168 —C <sub>6</sub> H <sub>6</sub> 115 —CH <sub>8</sub> 115 —CH <sub>8</sub> 129  —CH <sub>8</sub> 161.5–162

ylic-carbonic anhydrides, carboxylic acid azides, and thioacetic acid (36a, c).

Ketene reacts instantaneously with benzamidoxime in an inert solvent to give the O-acetylbenzamidoxime in quantitative yields

NOH NOCOCH: 
$$C_{0}H_{\delta}C = C + CH_{2} - C = C \rightarrow C_{0}H_{\delta}C$$
 NH<sub>2</sub>

Benzamidoxime and acetamidoxime give the corresponding O-acetyl derivative quantitatively when treated at room temperature with an ethereal solution of acetic-ethylcarbonic anhydride (LI).

NOH 
$$C_{\delta}H_{\delta}C + CH_{\delta}COOCOOC_{2}H_{\delta} \rightarrow \\ NH_{2} LI NOCOCH_{\delta} \\ C_{\delta}H_{\delta}C + CO_{2} + C_{2}H_{\delta}OH \\ NH_{2}$$

Also carboxylic acid azides are very efficient acylation agents which react readily at room temperature with amidoximes.

n amidoximes.

NOH

NOCOR'

$$+ N_3COR' \rightarrow RC + HN_3$$
 $NH_2$ 

NH<sub>2</sub>

In particular, formamidoxime, which cannot be benzoylated with benzoyl chloride, formed the expected derivative with benzoyl azide (36c).

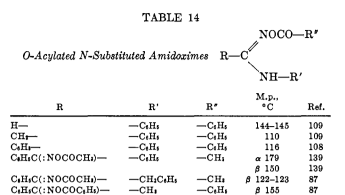
With thioacetic acid, benzamidoxime is acetylated at room temperature with evolution of hydrogen sulfide. The reaction can be conducted in water. However, sulfur appears during the reaction showing that simul-

TABLE 13
O-Acyl Derivatives of Imidoximes and Imidedioximes

Compound	M.p., °C.	Ref.
NOCOC <sub>6</sub> H <sub>5</sub>		<del></del>
CH=-C		
NH	194	40
	184	42
CH <sub>1</sub> —C		
°o		
NOCOC5H5		
CH <sub>2</sub> —C		
H <sub>2</sub> C NH	160	42
\ /		
CH <sub>2</sub> —C		
O		
NOCOCH		
ÇH₌—CŰ		
NH	170-171	166
CH-C		
NOCOCH:		
NOCOC <sub>6</sub> H <sub>6</sub>		
CHC		
NH	187–188	166
CH:-C		
NOCOC <sub>6</sub> H <sub>5</sub>		
NOCOCH.		
/		
CH <sub>2</sub> —C		
H₂C NH	127	14
CH.—C		
NOCOCH.		
NOCOC <sub>6</sub> H <sub>6</sub>		
CH <sub>2</sub> —C		
	170_190	14
H <sub>2</sub> C NH	179–180	14
CH <sub>2</sub> —C		
NOCOC <sub>6</sub> H <sub>5</sub>		
NOCOCH <sub>3</sub>		
NH CNH	192	22
NOCOCH <sub>3</sub>		
NOCOC <sub>6</sub> H <sub>5</sub>	248	22
NOCOC <sub>6</sub> H <sub>5</sub>	-10	

taneously a reduction occurs. A by-product of the formula  $C_9H_{12}O_2N_2$  can be isolated, which could be identified as a salt of benzamidine with acetic acid (36a).

Acylation with a carboxylic acid is reported to occur



in a few cases. p-Nitrobenzamidoxime reportedly was acylated with boiling glacial acetic acid and p-sulf-aminobenzamidoxime reacted on heating with chloracetic acid to give the chloroacetyl derivative (12).

—С<sub>6</sub>Н<sub>5</sub>

 $-C_6H_6$ 

α 201

139

-C<sub>6</sub>H<sub>5</sub>

-CH<sub>2</sub>C<sub>6</sub>H<sub>5</sub>

CoHoC(: NOCOCoHo)-

CoHoC(: NOCOCoHo)-

Benzamidoxime reacts on heating with benzoic acid (193). The acylated amidoxime cannot be isolated but its dehydration product 3,5-diphenyl-1,2,4-oxadiazole (LII) is formed.

NOH 
$$C_6H_5C + C_6H_5COOH \rightarrow NH_2 + C_6H_5COOH NOO$$
 NH2 
$$C_6H_5C + 2H_2OO$$
 N LII

This compound is formed in all media where benzamidoxime and benzoic acid are present (193). However, no definite proof ever has been given that a normal acylation occurs. Indeed, when treated with acetic, propionic or butyric acid, benzamidoxime also is transformed into 3,5-diphenyl-1,2,4-oxadiazole and no trace of a 5-alkyl cyclic derivative can be isolated (162, 164)

$$C_6H_6C$$
 $NH_2$ 
 $RCOOH$ 
 $C_6H_6C$ 
 $N-O$ 
 $CC_6H_6$ 

This reaction has been interpreted as an autoxidation of benzamidoxime (64) (see p. 167).

O-Acylated amidoximes also have been prepared from acylated hydroximic acid chlorides (LIII) and ammonia. This reaction was applied to some derivatives of glyoxime (54, 144).

The formylation of amidoximes recently has been performed with the mixed anhydride of formic and acetic acids

The formyl esters of formamidoxime, acetamidoxime, and benzamidoxime have been synthesized but have not been isolated in a pure state: they were immediately dehydrated into the corresponding 3-monosubstituted oxadiazoles (LIV) (36b, 81a).

NOCOH 
$$\rightarrow$$
  $H_2O + RC$   $\rightarrow$   $N-O$   $\rightarrow$   $H_2O + RC$   $\rightarrow$   $N$ 

The diformyl ester of oxamidedioxime could be prepared in good yields using this mixed anhydride (81a).

HON NOH 
$$C-C + 2 \text{ HCOOCOCH}, \rightarrow \\ H_2N NH_2 \\ HCOON NOCOH \\ C-C + 2 CH_3COOH \\ H_2N NH_2$$

N-Monosubstituted amidoximes also give O-acylated derivatives (87, 108, 109, 139). These compounds are listed in Table 14.

Only one N-disubstituted amidoxime, N,N-dimethyl m-nitrobenzamidoxime, is reported to yield a benzoylated compound (18).

$$\begin{array}{ccc} & \text{NOH} & & \text{NOCOC}_6\text{H}_6\\ \\ \textit{m-NO}_2\text{C}_6\text{H}_4\text{C} & & \xrightarrow{\text{C}_6\text{H}_6\text{COCl}} & \textit{m-NO}_2\text{C}_6\text{H}_4\text{C} \\ & & \text{N(CH}_2)_2 & & \text{N(CH}_2)_2 \end{array}$$

The most important chemical properties of O-acylated amidoximes are the readiness with which they are hydrolyzed into the parent amidoximes and their ability to cyclize into the corresponding 1,2,4-oxadiazoles (see p. 177).

N-Acylated Amidoximes.—N-Acyl derivatives of amidoximes have been described by Ponzio and his collaborators who have published a great number of papers concerning the stereochemistry of  $\alpha$ -dioximes. Part of the glyoximes studied were amino glyoximes which can be considered as  $\alpha$ -oximino-amidoximes (LV).

Ponzio's conclusions must be included into this report

because they raise important questions about the tautomerism of amidoximes (see pp. 156 and 171).

Ponzio has claimed (132) that the two isomers  $\alpha$  and  $\beta$  of 1-phenyl-2-aminoglyoxime correspond to the structures LVI and LVII, respectively, and that on acylation, the amidoxime group of the former is N-acylated, while the latter undergoes O-acylation. In order to prove this point, cyclization with aqueous alkali is performed, which leads to a furazan LVIII in the first case and to an oxadiazole ring (LIX) in the second (129, 144).

This dissimilar behavior in cyclization is Ponzio's main argument to prove the existence of forms LVI and LVII.

On the other hand he established (140) the possibility of transformation of the oxadiazole derivative into the corresponding furazan by treatment with alkali

so that the above proof of the structure appears to be less convincing.

Another N-acyl amidoxime, N-acetylcyanoform-amidoxime (LX), was claimed to have been isolated by Ponzio when he tried to prepare a monosubstituted furazan (LXI), which he did not obtain; for this reason he considered the latter structure too unstable to exist in alkaline medium (135).

Steinkopf (169, 171), who studied the halogenated acetamidoximes, claimed that their acylation occurs on the amino nitrogen, but no proof was given.

p-Nitrobenzamidoxime has been reported (12) to give an N-acetyl derivative (LXII) when treated with boiling acetic acid

NOH 
$$p\text{-NO}_2\text{C}_6\text{H}_4\text{C} + \text{CH}_4\text{COOH} \rightarrow \\ \text{NH}_2 \\ p\text{-NO}_2\text{C}_6\text{H}_4\text{C} + \text{H}_3\text{O} \\ \\ \text{NHCOCH}_3 \\ \text{LXII} \\ \\ \end{array}$$

The proposed structure has been based on the interpretation of the infrared spectrum, but the arguments are not wholly convincing.

Recently, both N- and O-acetylated formamidoximes have been synthesized (36c). When the acylation of formamidoxime is carried out with mixed acetic-ethylcarbonic anhydride (LXIII), the two isomers are present in almost equal quantities.

The infrared spectra of both isomers are quite distinct: the NH<sub>2</sub> doublet at 3295 and 3420 cm.<sup>-1</sup> and the —OC=O band at 1735 cm.<sup>-1</sup> characterize the O-acetyl formamidoxime molecule, whereas the N-acetylated compound shows only a simple band at 3335 cm.<sup>-1</sup> as well as a broad absorption region between 3300 and 2900 cm.<sup>-1</sup>, while the carbonyl absorption band of the amide function is visible at 1730 cm.<sup>-1</sup>.

Solubility and chemical behavior of the two isomers also are dissimilar: for example, the O-derivative can be dehydrated into the 5-methyl-1,2,4-oxadiazole, whereas the N-derivative cannot.

$$NOCOCH_3$$
  $\rightarrow$   $HC$   $CCH_3$   $NH_2$ 

If the acylation is carried out with acetic anhydride, the same isomers are obtained but in addition a small amount of acetyl urea is formed. A Lossen transformation may explain the presence of this product (see p. 180).

The formylation of formamidoxime also yields both O and N-formylated isomers. Only the latter has been isolated and identified by its infrared spectrum, while the former has been dehydrated immediately into unsubstituted 1,2,4-oxadiazole (36b, 103a).

Derivatives of amidoximes reported in the literature as having the N-acyl structure are listed in Table 15.

TABLE 15 NOH N-Acyl Derivatives of Amidoximes RC NH—CO—R' M.p., R' Ref. —н 146-147 103a н--CH 175 36c CHCl2-—CH₃ª 169, 171 114-115 −CH<sub>2</sub>a CH.I-103-105 169, 171 -CH₃ª 186 144 p-NO2C6H4-—CH₃ª 226 CH(:NOCOCH:)-CH 144, 169, 171 154 C6H6C(:NOCOCH3)-—CH₃ª α 150-151 132, 140 p-CH<sub>2</sub>C<sub>5</sub>H<sub>4</sub>C(:NOCOCH<sub>3</sub>)— —CH₃ª 168 132 C6H6C(: NOCOC6H6)-—CH₃ª 190-191 129 C6H6C(: NOCOC6H6)-—С<sub>6</sub>Н₅<sup>а</sup>  $\alpha$  189 132

-CaHa

199-200

132

p-CH<sub>2</sub>C<sub>6</sub>H<sub>4</sub>C(: NOCOCOC<sub>6</sub>H<sub>6</sub>)-

H-

CN-

Diacyl Derivatives of Amidoximes.—The benzoylation of o-methoxybenzamidoxime gives the N,Odibenzoyl derivative (LXIV), even at room temperature, when treated with an equivalent amount of benzoyl chloride (22).

When treated with benzoyl chloride at 100°, phenylaminoglyoxime is reported to yield a tribenzoyl derivative (LXV) (132).

A hexaacetyl compound (LXVII) is formed when the diaminotetraoxime LXVI is treated with hot acetic anhydride (83).

$$\begin{array}{c} \operatorname{NH_2(C-)_{NH_2}} & \xrightarrow{\operatorname{CH_3CONH}(C-)_{NOCOCH_3}} \\ & & \xrightarrow{\operatorname{LXVI}} & \xrightarrow{\operatorname{LXVII}} \end{array}$$

O,O'-dimethyloxamidedioxime is reported to form a tetraacetyl derivative (LXVIII) (6).

$$\begin{array}{c} \text{CH}_{2}\text{ON} & \text{NOCH}_{3} \\ \text{NH}_{2} & \xrightarrow{\text{CCH}_{2}\text{CO})_{2}\text{O}} \\ \text{NH}_{2} & \text{CH}_{2}\text{OH} & \text{NOCH}_{3} \\ \\ & \text{CC-C} & \\ \text{(CH}_{3}\text{CO})_{2}\text{N} & \text{N(COCH}_{3})_{2} \end{array}$$

Finally, although the structure of the compounds is not well elucidated, it is worth mentioning that Longo (87) reported the formation of diacetyl amidoximes LXIX and LXX in which the two acetyl radicals are linked to the hydroxamino group NHOH.

$$C_{\delta}H_{\delta}C(:NOCOCH_{2})C OCOCH_{\delta}$$

$$LXIX C_{\delta}H_{\delta}C(:NOCOCH_{\delta})C OCOCH_{\delta}$$

$$LXIX NCH_{2}C_{\delta}H_{\delta}$$

$$LXX$$

The  $\beta$ -form of LXX has been formulated by Longo as

Formation of 1,2,4-Oxadiazoles.—Acyl derivatives of the amidoximes are in most cases dehydrated easily into the corresponding 3,5-disubstituted 1,2,4-oxadiazole

$$RC$$
 $N-O$ 
 $CR' + H_2C$ 
 $NH_2$ 

The dehydration of the acyl amidoximes generally is accomplished by heating these compounds either in the dry state or in solution in glacial acetic acid, acetic anhydride, water, dilute NaOH, or H<sub>2</sub>SO<sub>4</sub>. If the acylation of amidoximes is carried out at 100° or above, spontaneous cyclization occurs.

#### (1) Carbonic Acid Derivatives of Amidoximes

Ethyl chloroformate reacts with the isonitroso group of the amidoximes to give carbonic acid derivatives (38, 177).

On heating ethanol is eliminated and a 5-hydroxy-1,2,4-oxadiazole LXXI is formed (38, 39, 178).

a No proof of the structure is given.

Similarly oxamidedioxime forms a disubstituted derivative (221).

HON NOH 
$$C-C + 2CICOOC_2H_5 \rightarrow NH_2 NH_2 C_2H_5OCOON NOCOOC_2H_5 + 2HCINH_2 NH_2$$

This compound could not by cyclized.

N-Substituted amidoximes also react in the same way (18). For example, N-dimethyl *m*-nitrobenzamidoxime treated with ethyl chloroformate yields a carbonic ester.

NOH 
$$m\text{-NO}_2\text{C}_6\text{H}_4\text{C} + \text{ClCOOC}_2\text{H}_5 \rightarrow \\ \text{N(CH}_3)_2 \\ m\text{-NO}_2\text{C}_6\text{H}_4\text{C} \\ \text{N(CH}_3)_2$$

Phosgene reacts with amidoximes to give derivatives of carbonic acid; on cyclization one molecule of amidoxime is regenerated (194).

Thiophosgene and amidoximes yield derivatives of thiocarbonic acid, which on cyclization give 5-mercapto-1,2,4-oxadiazoles (64).

Table 16 contains the list of the carbonic acid derivatives of amidoximes.

# (m) Reaction with $\beta$ -Ketocarboxylic Esters

Although amidoximes are indifferent toward non-activated esters, they react on heating with an excess of ethyl acetoacetate. Water and ethanol are eliminated and 5-aryl-3-acetonyl-1,2,4-oxadiazoles are formed (151, 162, 182, 201).

TABLE 16
Carbonic Acid Derivatives of Amidoximes
NOCOOC:H:

M.p.

R	°C.	Ref.		
C <sub>6</sub> H <sub>5</sub> —	127	1, 38, 39		
p-CH <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	130	162		
	142	114		
C₀H₅CH=CH→	101	218		
	111	151		
	121	151		
N	136	99		
	97	13		
o-HOCoH4-	96	100		
C <sub>6</sub> H <sub>6</sub> CHOH—	106-107	51		
p-CH <sub>4</sub> OC <sub>6</sub> H <sub>4</sub> —	119-120	100		
m-NO <sub>2</sub> C <sub>6</sub> H <sub>6</sub> —	152-153	159, 160		
p-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	169	201		
CO NCH2CH2-	142	36a		

NO—CO—ON R—C C—R					
	NH: NH:	· · · · · · · · · · · · · · · · · · ·			
(CH <sub>3</sub> ) <sub>2</sub> CHCH <sub>2</sub> CH <sub>2</sub>	114	55			
C <sub>6</sub> H <sub>5</sub> —	128-129	38, 39			
C <sub>6</sub> H <sub>6</sub> CHOH—	131	51			
p-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	232	201			
	NO-CS-ON				

	NO-CS-ON R-C C-	D.	
	NH <sub>2</sub> NH <sub>3</sub>	ĸ	
C6H6—	96	64	
p-CH <sub>4</sub> C <sub>6</sub> H <sub>4</sub> —	115	64	

Highest yields are obtained when the reaction is carried out in boiling toluene until all water and ethanol are eliminated by distillation (95).

Other  $\beta$ -ketocarboxylic esters such as ethyl benzoylacetate, o-methoxybenzoylacetate, and acetonedicarboxylate react similarly with aromatic amidoximes; e.g. (95).

$$C_6H_5C$$
 $N-O$ 
 $CCH_2COCH_2C$ 
 $N-O$ 
 $CC_6H$ 

#### (n) Reaction with Aldehydes

The aliphatic aldehydes react with amidoximes yielding 3,5-disubstituted 4,5-dihydro-1,2,4-oxadiazoles (182, 220).

The reaction occurs when a solution of the reagents in water or aqueous ethanol is left for a few hours at room temperature or on a water-bath. Aromatic aldehydes do not react with amidoximes with the exception of o-hydroxybenzaldehyde which yields the expected product with benzamidoxime after several weeks of standing (220).

The dihydro-oxadiazoles are crystalline products with basic properties, forming salts with mineral acids. They are hydrolyzed with diluted acids and bases into the corresponding aldehydes and amidoximes and are easily oxidized by potassium permanganate into the corresponding oxadiazoles (182, 220). Analogous dicyclic compounds are obtained from oxamidedioxime (199). The 3,5-disubstituted 4,5-dihydro-1,2,4-oxadiazoles reported in literature are listed in Table 17.

TABLE 17 N—O 4,5-Dihydro-1,2,4-oxadiazoles R—C NH CH—R'

R	R'	M.p., °C.	Ref.
C <sub>5</sub> H <sub>5</sub> —	-CH2	82	182
C6H6—	C2H5	64	220
C <sub>6</sub> H <sub>6</sub> —	-CH(CH <sub>2</sub> ) <sub>2</sub>	96	220
C <sub>6</sub> H <sub>5</sub> —	-CH2CH(CH2)2	83	220
C <sub>5</sub> H <sub>5</sub> —	$-CH_2C_6H_5$	136	220
C <sub>5</sub> H <sub>5</sub> —	-C6H4OH-0	155	220
p-CH₃C₀H₄—	—CH <sub>1</sub>	127.5	162
	—CH₃	121	151
p-CH <sub>2</sub> OC <sub>6</sub> H <sub>4</sub> —	—CH2	127.5	100
p-NO <sub>2</sub> C <sub>6</sub> H <sub>4</sub> —	-CH1	153	201
p-NO <sub>2</sub> C <sub>5</sub> H <sub>4</sub> —	$-CH_2Cl$	176	201

The exceptional complex forming reaction of chloral with amidoximes has been mentioned already (page 166).

Acetone and acetone dimethyl acetal also are reported to give 5-dimethyl-4,5-dihydroöxadiazoles (20b, 36a).

# (o) Action of Cyanogen

Cyanogen and benzamidoxime or  $\beta$ -naphthamidoxime yields addition products (111) formulated as LXXII and LXXIII. They are rather unstable and hydrolyze readily into the original amidoximes. The same compounds treated with acetic anhydride give the corresponding O-acetylamidoximes:

# (p) Action of Sulfur Dioxide

Sulfur dioxide gives with amidoximes derived from long chain fatty acids unstable addition compounds (LXXIV) (35).

The structure of these substances is not well established.

# (q) Action of Carbon Disulfide

The reaction of carbon disulfide with amidoximes in alkaline alcoholic solution produces cyclic compounds known as 5-mercapto-1,2,4-thiadiazoles (25) (LXXV).

In boiling ethanol, benzamidoxime gives with an excess of CS<sub>2</sub> the thiobenzamidoxime salt of the thioloxime of benzoyldithiocarbamic acid (LXXVI). Dilute hydrochloric acid hydrolyzes this compound into benzamidine hydrochloride.

$$\begin{array}{c|c} & N-S \\ & C_6H_5C \\ \hline \\ C_6H_5C \\ \hline \\ NSH \\ \hline \\ C_6H_5C \\ \hline \\ NSH \\ \\ NSH \\ \hline \\ NSH \\ NSH \\ \hline \\ NSH \\ NSH$$

Methylbenzamidoxime shows a similar behavior (25).

# (r) Action of Phenylhydrazine

Phenyl formazyl (LXXVII) is formed when benzamidoxime is refluxed with phenylhydrazine in dilute acetic acid (8).

NOH
$$C_{6}H_{5}C + 3C_{6}H_{5}NH-NH_{2} \rightarrow NH_{2}$$

$$N-NH-C_{6}H_{5}$$

$$C_{6}H_{5}C + NH_{2}OH + 2NH_{3} + C_{6}H_{5}NH_{2}$$

$$N=N-C_{6}H_{5}$$

$$LXXVII$$

Other reactions of an amidoxime with a hydrazine derivative have not been reported.

#### (s) Beckmann Transformation

The presence of urea among the decomposition products of formamidoxime (91) has been explained tentatively by a Lossen rearrangement, related to the Beckmann transformation (41).

The formation of acetylurea when formamidoxime is acetylated with acetic anhydride also could be interpreted by a Lossen-Beckmann transformation (36c) (see page 176).

Amidoximes have been converted into unsymmetrical ureas when treated first with benzenesulfonyl chloride and then with water. The mechanism postulated by Tiemann (191) involves the conversion of the amidoxime into its O-benzene-sulfonyl derivative which then undergoes a Beckmann transformation into an O-benzenesulfonylisourea. Subsequent hydrolysis yields the asymmetrical urea.

In support of this reaction mechanism it was found (120, 121) that certain amidoximes yielded isolable O-benzenesulfonyl derivatives which on heating with water gave the expected ureas (or their decomposition products) and benzenesulfonic acid.

Nevertheless, Partridge and Turner (115) claimed in 1953 that the isourea derivative (LXXVIII) decomposes spontaneously into benzenesulfonic acid and a substituted cyanamide (LXXVIX). Hydrolysis of this cyanamide would then yield a urea, whereas its reaction with an amine yielded a guanidine derivative.

The urea is therefore to be considered as a secondary reaction product and the isolable primary products should be benzenesulfonic acid and a substituted cyanamide.

The formation of the O-benzenesulfonyl derivative of the amidoxime is demonstrated readily in the case of phenylacetamidoxime, since the derivative is stable and can be purified without decomposition, whereas O-benzenesulfonylbenzamidoxime is too unstable to be isolated. On heating in an inert solvent, O-benzenesulfonylphenylacetamidoxime readily undergoes a Beckmann transformation into benzyl cyanamide and benzenesulfonic acid. When the transformation is carried out in pyridine with the sulfonyl derivative of the amidoxime formed in situ, the yield of benzylcyanamide is 61%. The benzenesulfonyl ester of benzamidoxime prepared in pyridine similarly affords phenylcyanamide and its trimer.

When the products of the transformation are heated with an amine, a N,N'-disubstituted guanidine is obtained in good yield.

The mechanism may be represented as an intramolecular process in which the hydrogen bond controls the subsequent electronic displacements.

$$C_6H_5NHC$$
 $H \rightarrow C_6H_5NHCN + C_6H_5SO_2OH$ 
 $SO_2C_6H_5$ 

When the amino group is substituted by two methyl radicals, no transformation occurs and the O-benzene-sulfonyl derivative can be isolated easily (18).

Table 18 summarizes the results obtained in this field.

# 3. Chemotherapeutic Properties of Amidoximes

Several papers have been published concerning the anti-bacterial activity of amidoximes, compared with amidines. *p*-Sulfamidobenzamidoxime for example has a slightly superior activity over the corresponding

Yield, %

			Beckmani	n Transforn	nation of A	lmidoximes		
	No	Tiema: SO <sub>2</sub> C <sub>6</sub> H <sub>5</sub>	nn reaction				-Partridge reaction-	···
R	RC NH. M.p., °C.	Yield,	—RNH M.p., °C.	CONH2— Yield, %	M.p., °C.	Yield,	RN R'	NH    HCNHR' M.p., °C.
CH	130	68	147	43-44	40.5	15 60 (trimer)	C <sub>6</sub> H <sub>6</sub> —  p-BrC <sub>6</sub> H <sub>4</sub> —  p-CH <sub>1</sub> C <sub>1</sub> H <sub>4</sub> —	141-142 167-168 123-124
≻CH₃C6H4— C6H6CH2—	128	73	180 148	43–44 8	42.5	5061	C <sub>6</sub> H <sub>5</sub> — p-CH <sub>5</sub> OC <sub>6</sub> H <sub>4</sub> —	122-123 122-123 172

TABLE 18
Beckmann Transformation of Amidoximes

amidine on experimental typhus infections in mice (3, 200). However, in most cases, the introduction of the oxygen atom decreases the anti-bacterial power of the amidines, together with their toxicity (20).

p-CH2(CH2)5OC6H4-

Lamb and White (75) have studied the trypanocide activity of several diamidoximes.

2-Methoxy-9-aminoacridine-6-amidoxime and 9-anil-acridine-3-amidoxime are patented as products having pharmacological properties and useful in therapeutics (47).

Some halogenated phenols carrying an amidoxime group are active against *Mycobacterium tuberculosis*, in vitro (20a).

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