

***Nomenclature of Organic Chemistry. IUPAC Recommendations
and Preferred Names 2013.***

Prepared for publication by Henri A. Favre and Warren H. Powell,
Royal Society of Chemistry, ISBN 978-0-85404-182-4

Appendix 2

DETACHABLE PREFIXES USED FOR SUBSTITUTIVE NOMENCLATURE

The symbol * designates the preferred prefix (for example: acetamido* = acetylamino; acetylamino = acetamido*) or the preselected prefix (for example: sulfanyl* = thio).

Prefixes that are not recommended are followed by the mention 'see' followed by the preferred or preselected prefix (for example: 'chloroxy: see chloryl*'). No formula is given for the entry 'chloroxy'. As a counterpart, the preferred or preselected prefix is followed by the mention 'not' and the prefix that is not recommended enclosed in appropriate enclosing marks [for example: chloryl* (not chloroxy)].

Name	Formula	Rule(s)
acetamido* = acetylamino	CH ₃ -CO-NH-	P-66.1.1.4.3
acetimidamido = ethanimidamido* = acetimidoylamino	CH ₃ -C(=NH)-NH-	P-66.4.1.3.5
acetimidoyl = ethanimidoyl* = 1-iminoethyl	CH ₃ -C(=NH)-	P-65.1.7.2.2
acetimidoylamino = ethanimidamido* = acetimidamido	CH ₃ -C(=NH)-NH-	P-66.4.1.3.5
acetohydrazido* = 2-acetylhydrazin-1-yl	CH ₃ -CO-NH-NH-	P-66.3.2.3
acetohydrazonoyl = ethanehydrazonoyl* = 1-hydrazinylideneethyl	CH ₃ -C(=N-NH ₂)-	P-65.1.7.2.2
acetohydroximoyl = <i>N</i> -hydroxyethanimidoyl* = <i>N</i> -hydroxyacetimidoyl	CH ₃ C(=N-OH)-	P-65.1.7.2.2
acetonyl = 2-oxopropyl*	$\begin{matrix} 3 & 2 & 1 \\ \text{CH}_3 & -\text{CO} & -\text{CH}_2- \end{matrix}$	P-64.5.1
acetonylidene: see 2-oxopropylidene*		
acetonylidyne: see 2-oxopropylidyne*		
acetoxy = acetyloxy*	CH ₃ -CO-O-	P-65.6.3.2.3
acetoxysulfonyl = (acetyloxy)sulfonyl*	CH ₃ -CO-O-SO ₂ -	P-65.3.2.3
acetyl* = ethanoyl = 1-oxoethyl	CH ₃ -CO-	P-65.1.7.2.1
<i>N</i> -acetylacetamido* = diacetylamino (not diacetylazanyl; not diacetamido)	(CH ₃ -CO) ₂ N-	P-66.1.2.1
acetylamino = acetamido*	CH ₃ -CO-NH-	P-66.1.1.4.3
acetylazanediyyl* (not acetylimino)	CH ₃ -CO-N<	P-66.1.1.4.4
acetylazanylidene* = acetylimino	CH ₃ -CO-N=	P-62.3.1.2

2-acetylhydrazin-1-yl = acetohydrazido*

acetylimino = acetylazanylidene* (not acetylazanediyl)

acetyloxy* = acetoxy

(acetyloxy)sulfonyl* = acetoxysulfonyl

acrylohydrazonoyl = prop-2-enehydrazonoyl* = 1-hydrazinylideneprop-2-en-1-yl

acryloyl = prop-2-enoyl* = 1-oxoprop-2-en-1-yl

adamantan-2-yl* = 2-adamantyl = tricyclo[3.3.1.1^{3,7}]decan-2-yl (also 1-isomer)

2-adamantyl = adamantan-2-yl* = tricyclo[3.3.1.1^{3,7}]decan-2-yl (also 1-isomer)

adipoyl = hexanedioyl* = 1,6-dioxohexane-1,6-diyl

allyl = prop-2-en-1-yl*

allylidene = prop-2-en-1-ylidene*

allylidyne = prop-2-enylidyne*

alumanyl*

alumanylidene*

amidino: see carbamimidoyl*

amidochlorophosphoryl = phosphoramidochloridoyl* (not chloroamidophosphoryl)

amidyl = azanidyl*

amidylidene = azanidylidene*

amino* = azanyl

(4'-amino[1,1'-biphenyl]-4-yl)amino* = benzidino

C-aminocarbonimidoyl = carbamimidoyl* = amino(imino)methyl (not amidino)

aminocarbonothioyl = carbamothioyl* = amino(sulfanylidene)methyl (not thiocarbamoyl)

aminocarbonyl = carbamoyl*

(aminocarbonyl)amino = carbamoylamino* (not ureido)

[(aminocarbonyl)amino]carbonyl = carbamoylcarbamoyl*

2-(aminocarbonyl)hydrazin-1-yl = 2-carbamoylhydrazin-1-yl* = semicarbazido

aminodichlorosilyl*

amino(hydrazinylidene)methyl = carbamohydrazonoyl*

[amino(hydroxy)methylidene]amino* (not 3-isoureido)

amino(imino)methyl = carbamimidoyl* = C-aminocarbonimidoyl (not amidino)

[amino(imino)methyl]amino = carbamimidoylamino* = carbamimidamido = guanidino

(aminomethylidene)amino*

CH₃-CO-NH-NH- P-66.3.2.3

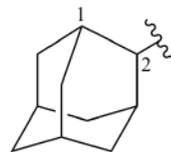
CH₃-CO-N= P-62.3.1.2

CH₃-CO-O- P-65.6.3.2.3

CH₃-CO-O-SO₂- P-65.3.2.3

CH₂=CH-C(=N-NH₂)- P-65.1.7.3.2

CH₂=CH-CO- P-65.1.7.3.1; P-65.1.7.4.1



-CO-[CH₂]₄-CO- P-65.1.7.3.1; P-65.1.7.4.1

CH₂=CH-CH₂- P-32.3

CH₂=CH-CH= P-32.3

CH₂=CH-C≡ P-32.3

H₂Al- P-29.3.1; P-68.1.2

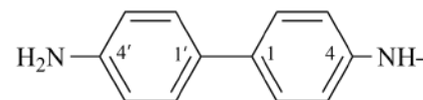
HAl= P-29.3.1; P-68.1.2

(H₂N)CIP(O)- P-67.1.4.1.1.4

-NH- P-72.6.3

-N= P-72.6.3

H₂N- P-62.2.3



H₂N-C(=NH)- P-65.2.1.5; P-66.4.1.3.1

H₂N-CS- P-65.2.1.5; P-66.1.4.4

H₂N-CO- P-65.2.1.5; P-66.1.1.4.1.1

H₂N-CO-NH- P-66.1.6.1.1.3

H₂N-CO-NH-CO- P-66.1.6.1.1.4

H₂N-CO-NH-NH- P-68.3.1.2.4

(H₂N)Cl₂Si- P-67.1.4.2

H₂N-C(=N-NH₂)- P-66.4.2.3.2

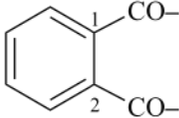
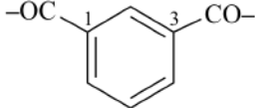
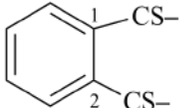
H₂N-C(OH)=N- P-66.1.6.1.2.2

H₂N-C(=NH)- P-65.2.1.5; P-66.4.1.3.1

H₂N-C(=NH)-NH- P-66.4.1.2.1.3

H₂N-CH=N- P-66.4.1.3.3

arsinetriyl: see arsanetriyl*		
arsinidine: see arsanylidene*		
arsino = arsanyl*	H ₂ As–	P-29.3.1; P-68.3.2.3.2.2
arsinoyl* = dihydroarsoryl (not arsinyll)	H ₂ As(O)–	P-67.1.4.1.1.2; P-67.1.4.1.2
arsinyll: see arsinoyl*		
arso: see dioxo-λ ⁵ -arsanyl*		
arsonato*	(–O) ₂ As(O)–	P-72.6.1
arsonio = arsaniumyl* = arsoniumyl	H ₃ As ⁺ –	P-73.6
arsoniumyl = arsaniumyl* = arsonio	H ₃ As ⁺ –	P-73.6
arsono* = dihydroxyarsoryl	(HO) ₂ As(O)–	P-67.1.4.1.1.1
arsonoyl* = hydroarsoryl	HAs(O)<	P-67.1.4.1.1.2; P-67.1.4.1.2
arsoranyl = λ ⁵ -arsanyl*	H ₄ As–	P-68.3.2.3.2.2
arsorimidoyl* = imidoarsoryl	As(=NH)<	P-67.1.4.1.1.4
arsoryl* (not arsenyl)	–As(O)<	P-67.1.4.1.1.2
azanediidyl*	N ^{2–} –	P-72.6.3
azanediyl* (not imino)	HN<	P-35.2.2; P-62.2.5.1
azanetriyl = nitrilo* (not azanylidyne; not azanylylidene)	–N<	P-35.2.1; P-62.2.5.1
azanidyl* = amidyl	–NH–	P-72.6.3
azanidylidene* = amidylidene	–N=	P-72.6.3
azaniumyl* = ammonio = ammoniumyl	H ₃ N ⁺ –	P-73.6
azanyl = amino*	H ₂ N–	P-62.2.3
azanylidene = imino*	HN=	P-35.2.1;;P-62.3.1.2
azanylidyne* (not nitrilo)	N≡	P-35.2.2
azanylylidene* (not nitrilo)	–N=	P-35.2.2; P-62.3.1.2
azido*	N ₃ –	P-61.7
azino: see hydrazinediylidene*		
azinoyl* = dihydronitroryl (not azinyll)	H ₂ N(O)–	P-67.1.4.1.1.2; P-67.1.4.1.2
azinyll: see azinoyl*		
azo = diazenediyl*	–N=N–	P-32.1.1; P-68.3.1.3.2.1; P-68.3.1.3.2.2
azonato*	(–O ₂ –N(O)–	P-72.6.1
azono* = dihydroxynitroryl	(HO) ₂ N(O)–	P-67.1.4.1.1.1; P-67.1.4.1.1.5
azonothiroyl* = thioazonoyl	HN(S)<	P-67.1.4.1.1.4
azonoyl* = hydronitroryl	HN(O)<	P-67.1.4.1.1.2; P-67.1.4.1.2
azoryl: see nitroryl*		
<i>NNO</i> -azoxy	–N=N(O)–	P-68.3.1.3.3.1
<i>NON</i> -azoxy	–N(O)=N– or –N=N(O)–	P-68.3.1.3.3.1

<i>ONN</i> -azoxy	$-\text{N}(\text{O})=\text{N}-$	P-68.3.1.3.3.1
benzal: see benzylidene*		
benzamido* = benzoylamino	$\text{C}_6\text{H}_5\text{-CO-NH}-$	P-66.1.1.4.3
benzenecarbohydroximoyl = <i>N</i> -hydroxybenzenecarboximidoyl* = <i>N</i> -hydroxybenzimidoyl = benzhydroximoyl	$\text{C}_6\text{H}_5\text{-C(=N-OH)-}$	P-65.1.7.2.2
benzenecarbonyl = benzoyl* = oxo(phenyl)methyl = phenylcarbonyl	$\text{C}_6\text{H}_5\text{-CO}-$	P-34.2.1.1; P-34.2.2; P-65.1.7.2.1
benzenecarbothioamido* = (benzenecarbothiyl)amino = thiobenzamido	$\text{C}_6\text{H}_5\text{-CS-NH}-$	P-66.1.4.4
benzenecarbothiyl* = thiobenzoyl = phenyl(sulfanylidene)methyl = phenyl(thioxo)methyl	$\text{C}_6\text{H}_5\text{-CS}-$	P-65.1.7.2.3
(benzenecarbothiyl)amino = benzenecarbothioamido* = thiobenzamido	$\text{C}_6\text{H}_5\text{-CS-NH}-$	P-66.1.4.4
benzenecarboximidohydrazido* = 2-(benzenecarboximidoyl)hydrazin-1-yl	$\text{C}_6\text{H}_5\text{-C(=NH)-NH-NH}-$	P-66.4.2.3.6
benzenecarboximidoyl* = benzimidoyl = imino(phenyl)methyl	$\text{C}_6\text{H}_5\text{-C(=NH)-}$	P-65.1.7.2.2
2-(benzenecarboximidoyl)hydrazin-1-yl = benzenecarboximidohydrazido*	$\text{C}_6\text{H}_5\text{-C(=NH)-NH-NH}-$	P-66.4.2.3.6
benzene-1,2-dicarbonyl* = phthaloyl = 1,2-phenylenedicarbonyl = 1,2-phenylenebis(oxomethylene)		P-65.1.7.3.1; P-65.1.7.4.2
benzene-1,3-dicarbonyl* = isophthaloyl = 1,3-phenylenedicarbonyl = 1,3-phenylenebis(oxomethylene)		P-65.1.7.3.1; P-65.1.7.4.2
benzene-1,4-dicarbonyl* = terephthaloyl = 1,4-phenylenedicarbonyl = 1,4-phenylenebis(oxomethylene)		P-65.1.7.3.1; P-65.1.7.4.2
benzene-1,2-dicarbothiyl* = 1,2-phenylenebis(sufanylidene)methylene) = 1,2-phenylenebis(thioxomethylene) (not dithiophthaloyl) (also 1,3- and 1,4-isomers)		P-65.1.7.3.1; P-65.1.7.4.3
benzene-1,4-dicarboximidoyl* = terephthalimidoyl = 1,4-phenylenebis(iminomethylene) = 1,4-phenylenedicarbonimidoyl (also phthalimidoyl = 1,2-isomer; and isophthalimidoyl = 1,3-isomer)		P-65.1.7.3.2
benzene-1,2-diyl: see 1,2-phenylene* (also 1,3- and 1,4-isomers)		
benzeneselenonyl* = phenylselenonyl	$\text{C}_6\text{H}_5\text{-SeO}_2-$	P-65.3.2.2.2
benzenesulfnamido* = (benzenesulfinyl)amino = (phenylsulfinyl)amino	$\text{C}_6\text{H}_5\text{-S(O)-NH}-$	P-66.1.1.4.3
benzenesulfinohydrazonamido* = (benzenesulfinohydrazonoyl)amino	$\text{C}_6\text{H}_5\text{-S(=N-NH}_2\text{)-NH}-$	P-66.4.2.3.5
(benzenesulfinohydrazonoyl)amino = benzenesulfinohydrazonamido*	$\text{C}_6\text{H}_5\text{-S(=N-NH}_2\text{)-NH}-$	P-66.4.2.3.5
benzenesulfinoselenoyl* = phenylsulfinoselenoyl	$\text{C}_6\text{H}_5\text{-S(Se)-}$	P-65.3.2.2.2
benzenesulfinyl* = phenylsulfinyl	$\text{C}_6\text{H}_5\text{-S(O)-}$	P-63.6; P-65.3.2.2.2
(benzenesulfinyl)amino = benzenesulfnamido* = (phenylsulfinyl)amino	$\text{C}_6\text{H}_5\text{-S(O)-NH}-$	P-66.1.1.4.3

benzenesulfonamido* = (benzenesulfonyl)amino = (phenylsulfonyl)amino
 benzenesulfonyl* = phenylsulfonyl
 (benzenesulfonyl)amino = benzenesulfonamido* = (phenylsulfonyl)amino
 benzhydryl: see diphenylmethyl*

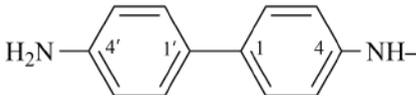
benzidino = (4'-amino[1,1'-biphenyl]-4-yl)amino*

benzimidoyl = benzenecarboximidoyl* = imino(phenyl)methyl
 benzohydrazido* = 2-benzoylhydrazin-1-yl
 benzohydroximoyl = *N*-hydroxybenzenecarboximidoyl* = *N*-hydroxybenzimidoyl
 = benzenecarbohydroximoyl
 benzoyl* = benzenecarbonyl = oxo(phenyl)methyl = phenylcarbonyl
 benzoylamino = benzamido*
 benzoylazediyyl*
 benzoylazanylidene = benzoylimino*
 2-benzoylhydrazin-1-yl = benzohydrazido*
 benzoylimino* = benzoylazanylidene
 benzoyloxy* = (phenylcarbonyl)oxy
 benzyl* = phenylmethyl
 benzylidene* = phenylmethylidene (not benzal)
 benzylidyne* = phenylmethylidyne
 benzyloxy* = phenylmethoxy

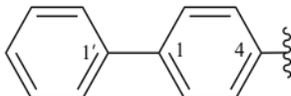
[1,1'-biphenyl]-4-yl* (not 4-phenylphenyl)

bis(acetyloxy)-λ³-iodanyl* (not diacetoxyiodo)
 bismuthaniumyl* = bismuthonio = bismuthoniumyl
 bismuthanyl* = bismuthino
 λ⁵-bismuthanylidene* = bismuthoranylidene
 bismuthino = bismuthanyl*
 bismuthonio = bismuthaniumyl* = bismuthoniumyl
 bismuthoniumyl = bismuthaniumyl* = bismuthonio
 bismuthoranylidene = λ⁵-bismuthanylidene*
 bis(selanyl)boranyl = diselenoborono*
 bis(silylamino)silyl* (not trisilazan-3-yl)
 1,4-bis(sulfanylidene)butane-1,4-diyl = butanebis(thioyl)* = 1,4-dithioxobutane-1,4-diyl

C₆H₅-SO₂-NH- P-66.1.1.4.3
 C₆H₅-SO₂- P-63.6; P-65.3.2.2.2
 C₆H₅-SO₂-NH- P-66.1.1.4.3

 P-62.2.4.1.1

C₆H₅-C(=NH)- P-65.1.7.2.2
 C₆H₅-CO-NH-NH- P-66.3.2.3
 C₆H₅-C(=N-OH)- P-65.1.7.2.2
 C₆H₅-CO- P-34.2.1.1; P-34.2.2; P-65.1.7.2.1
 C₆H₅-CO-NH- P-66.1.1.4.3
 C₆H₅-CO-N< P-66.1.1.4.4
 C₆H₅-CO-N= P-66.1.1.4.4
 C₆H₅-CO-NH-NH- P-66.3.2.3
 C₆H₅-CO-N= P-66.1.1.4.4
 C₆H₅-CO-O- P-65.6.3.2.3
 C₆H₅-CH₂- P-29.6.1; P-29.6.2.1
 C₆H₅-CH= P-29.6.1; P-29.6.2.1
 C₆H₅-C≡ P-29.6.1; P-29.6.2.1
 C₆H₅-CH₂-O- P-63.2.2.1.1

 P-29.3.5

(CH₃-CO-O)₂I- P-68.5.1
 H₃Bi⁺- P-73.6
 H₂Bi- P-29.3.1; P-68.3.3
 H₃Bi= P-68.3.3
 H₂Bi- P-29.3.1; P-68.3.3
 H₃Bi⁺- P-73.6
 H₃Bi⁺- P-73.6
 H₃Bi= P-68.3.3
 (HSe)₂B- P-68.1.4.2

 P-29.3.2.2
 -CS-CH₂-CH₂-CS- P-65.1.7.4.1; P-65.1.7.4.3

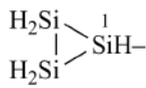
(not dithiosuccinyl)		
bis(sulfanylidene)ethanediyl = dithiooxalyl = ethanebis(thioyl)*	–CS-CS–	P-65.1.7.2.3
bis(sulfanyl)phosphoryl*	(HS) ₂ P(O)–	P-67.1.4.1.1.5
boranediyl* (not borylene; not borylidene; not boranylidene)	HB<	P-68.1.2
boranetriyl* (not borylidyne)	–B<	P-68.1.2
boranuidyl*	H ₃ B–	P-72.6.3
boranyl* (not boryl)	H ₂ B–	P-29.3.1; P-67.1.4.2; P-68.1.2
(boranylamino)boranyl* (not diborazan-1-yl)	H ₂ B-NH-BH–	P-68.1.2
boranylidene* (not borylidene)	HB=	P-29.3.1; P-67.1.4.2; P-68.1.2
boranylidyne* (not borylidyne)	B≡	P-29.3.1; P-67.1.4.2
borodiamidoyl: see diaminoboranyl*		
borono* = dihydroxyboranyl	(HO) ₂ B–	P-67.1.4.2; P-68.1.4.2
boryl: see boranyl*		
borylene: see boranediyl*		
borylidene: see boranylidene*		
borylidyne: see boranylidyne*		
bromo*	Br–	P-61.3.1
bromocarbonothioyl = carbonobromidothioyl*	Br-CS–	P-65.2.1.5
bromosyl*	BrO–	P-61.3.2.3
bromyl*	BrO ₂ –	P-61.3.2.3
butanamido* = butanoylamino = butyramido = butyrylamino	CH ₃ -[CH ₂] ₂ -CO-NH–	P-66.1.1.4.3
butanebis(thioyl)* = 1,4-bis(sulfanylidene)butane-1,4-diyl = 1,4-dithioxobutane-1,4-diyl (not dithiosuccinyl)	–CS-CH ₂ -CH ₂ -CS–	P-65.1.7.4.1; P-65.1.7.4.3
butanediimidoyl* = succinimidoyl = 1,4-diiminobutane-1,4-diyl	–C(=NH)-CH ₂ -CH ₂ -C(=NH)–	P-65.1.7.3.2
butanedioyl* = succinyl = 1,4-dioxobutane-1,4-diyl	–CO-CH ₂ -CH ₂ -CO–	P-65.1.7.3.1; P-65.1.7.4.1
butane-1,1-diyl*	CH ₃ -CH ₂ -CH ₂ -CH<	P-29.3.2.2
butane-1,4-diyl* (not tetramethylene)	–CH ₂ -CH ₂ -CH ₂ -CH ₂ –	P-29.3.2.2
butanethioyl* = thiobutyryl = 1-sulfanylidenebutyl = 1-thioxobutyl	CH ₃ -CH ₂ -CH ₂ -CS–	P-65.1.7.4.1
butanimidoyl* = butyrimidoyl = 1-iminobutyl	CH ₃ -CH ₂ -CH ₂ -C(=NH)–	P-65.1.7.3.2; P-65.1.7.4.1
butanoyl* = butyryl = 1-oxobutyl	CH ₃ -CH ₂ -CH ₂ -CO–	P-65.1.7.3.1; P-65.1.7.4.1
butanoylamino = butanamido* = butyramido = butyrylamino	CH ₃ -CH ₂ -CH ₂ -CO-NH–	P-66.1.1.4.3
butan-1-yl = butyl*	CH ₃ -CH ₂ -CH ₂ -CH ₂ –	P-29.3.2.1; P-29.3.2.2
butan-2-yl* = 1-methylpropyl (not <i>sec</i> -butyl; not but-2-yl)	CH ₃ -CH ₂ -CH(CH ₃)–	P-29.3.2.2; P-29.4.1; P-29.6.3
butan-1-ylidene = butylidene*	CH ₃ -CH ₂ -CH ₂ -CH=	P-29.3.2.1; P-29.3.2.2
butan-2-ylidene* = 1-methylpropylidene (not <i>sec</i> -butylidene)	CH ₃ -CH ₂ -C(CH ₃)=	P-29.3.2.2; P-29.4.1; P-29.6.3
butanylidyne = butylidyne*	CH ₃ -CH ₂ -CH ₂ -C≡	P-29.3.2.1; P-29.3.2.2

(butan-2-yl)oxy* = 1-methylpropoxy (not <i>sec</i> -butoxy; not <i>sec</i> -butyloxy)	$\text{CH}_3\text{-CH}_2\text{-CH}(\text{CH}_3)\text{-O-}$	P-63.2.2.2
butan-2-yl-3-ylidene*	$\begin{array}{c} \parallel \quad \\ \text{CH}_3\text{-C-CH-CH}_3 \\ 4 \quad 3 \quad 2 \quad 1 \end{array}$	P-29.3.2.2
butan-3-yl-1-ylidene*	$\begin{array}{c} \\ \text{CH}_3\text{-CH-CH}_2\text{-CH=} \\ 4 \quad 3 \quad 2 \quad 1 \end{array}$	P-29.3.2.2
(2 <i>E</i>)-but-2-enedioyl* = fumaroyl = (2 <i>E</i>)-1,4-dioxobut-2-ene-1,4-diyl	$\begin{array}{c} 2 \quad 1 \\ \text{HC-CO-} \\ \parallel \\ \text{-OC-CH} \\ 4 \quad 3 \end{array}$	P-65.1.7.3.1; P-65.1.7.4.1
(2 <i>Z</i>)-but-2-enedioyl* = maleoyl = (2 <i>Z</i>)-1,4-dioxobut-2-ene-1,4-diyl	$\begin{array}{c} 2 \quad 1 \\ \text{HC-CO-} \\ \parallel \\ \text{HC-CO-} \\ 3 \quad 4 \end{array}$	P-65.1.7.3.1; P-65.1.7.4.1
but-2-ene-1,4-diyl*	$\begin{array}{c} 4 \quad 3 \quad 2 \quad 1 \\ \text{-CH}_2\text{-CH=CH-CH}_2\text{-} \end{array}$	P-32.1.1
but-2-enoyl* (not crotonyl)	$\text{CH}_3\text{-CH=CH-CO-}$	P-65.1.7.4
but-1-enyl: see but-1-en-1-yl*		
but-1-en-1-yl* (not but-1-enyl)	$\text{CH}_3\text{-CH}_2\text{-CH=CH-}$	P-32.1.1
but-2-enyl: see but-2-en-1-yl*		
but-2-en-1-yl* (not but-2-enyl)	$\text{CH}_3\text{-CH=CH-CH}_2\text{-}$	P-32.1.1
but-3-en-2-yl* = 1-methylprop-2-en-1-yl	$\begin{array}{c} \\ \text{CH}_2\text{=CH-CH-CH}_3 \\ 4 \quad 3 \quad 2 \quad 1 \end{array}$	P-32.1.1
butoxy* (not butyloxy)	$\begin{array}{c} 4 \quad 3 \quad 2 \quad 1 \\ \text{CH}_3\text{-CH}_2\text{-CH}_2\text{-CH}_2\text{-O-} \end{array}$	P-63.2.2.2
<i>sec</i> -butoxy: see (butan-2-yl)oxy*		
<i>tert</i> -butoxy* (unsubstituted) = (2-methylpropan-2-yl)oxy = 1,1-dimethylethoxy (not <i>tert</i> -butyloxy)	$(\text{CH}_3)_3\text{C-O-}$	P-63.2.2.2
butyl* = butan-1-yl	$\text{CH}_3\text{-CH}_2\text{-CH}_2\text{-CH}_2\text{-}$	P-29.3.2.1; P-29.3.2.2
but-2-yl: see butan-2-yl*		
<i>sec</i> -butyl: see butan-2-yl*		
<i>tert</i> -butyl* (unsubstituted) = 2-methylpropan-2-yl = 1,1-dimethylethyl	$(\text{CH}_3)_3\text{C-}$	P-29.4.1; P-29.6.1
butylidene* = butan-1-ylidene	$\text{CH}_3\text{-CH}_2\text{-CH}_2\text{-CH=}$	P-29.3.2.1; P-29.3.2.2
<i>sec</i> -butylidene: see butan-2-ylidene*		
butylidyne* = butanylidyne	$\text{CH}_3\text{-CH}_2\text{-CH}_2\text{-C}\equiv$	P-29.3.2.1; P-29.3.2.2
butyloxy: see butoxy*		
<i>sec</i> -butyloxy: see (butan-2-yl)oxy*		
<i>tert</i> -butyloxy: see <i>tert</i> -butoxy* (unsubstituted)		
butyramido = butanamido* = butyrylamino = butanoylamino	$\text{CH}_3\text{-[CH}_2\text{]}_2\text{-CO-NH-}$	P-66.1.1.4.3

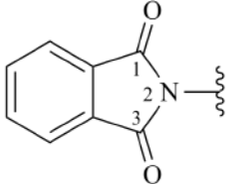
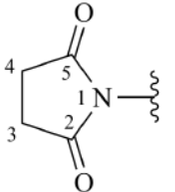
butyrimidoyl = butanimidoyl* = 1-iminobutyl	CH ₃ -CH ₂ -CH ₂ -C(=NH)-	P-65.1.7.3.2; P-65.1.7.4.1
butyryl = butanoyl* = 1-oxobutyl	CH ₃ -CH ₂ -CH ₂ -CO-	P-65.1.7.3.1; P-65.1.7.4.1
butyrylamino = butanamido* = butanoylamino = butyramido	CH ₃ -CH ₂ -CH ₂ -CO-NH-	P-66.1.1.4.3
carbamimidamido = carbamimidoylamino* = [amino(imino)methyl]amino = guanidino	H ₂ N-C(=NH)-NH-	P-66.4.1.2.1.3
carbamimidoyl* = C-aminocarbonimidoyl = amino(imino)methyl (not amidino)	H ₂ N-C(=NH)-	P-65.2.1.5; P-66.4.1.3.1
carbamimidoylamino* = carbamimidamido = [amino(imino)methyl]amino = guanidino	H ₂ N-C(=NH)-NH-	P-66.4.1.2.1.3
carbamohydrazonoyl* = amino(hydrazinylidene)methyl	H ₂ N-C(=N-NH ₂)-	P-66.4.2.3.2
carbamothioyl* = aminocarbonothioyl = amino(sulfanylidene)methyl (not thiocarbamoyl)	H ₂ N-CS-	P-65.2.1.5; P-66.1.4.4
carbamothioylamino* = [amino(sulfanylidene)methyl]amino	H ₂ N-CS-NH-	P-66.1.6.1.3.3
carbamoyl* = aminocarbonyl	H ₂ N-CO-	P-65.2.1.5; P-66.1.1.4.1.1
carbamoylamino* = (aminocarbonyl)amino (not ureido)	H ₂ N-CO-NH-	P-66.1.6.1.1.3
carbamoylcarbamoyl* = [(aminocarbonyl)amino]carbonyl	H ₂ N-CO-NH-CO-	P-66.1.6.1.1.4
carbamoylcarbonyl: see oxamoyl*		
carbamoylformamido: see oxamoylamino*		
carbamoylformyl: see oxamoyl*		
2-carbamoylhydrazin-1-yl* = 2-(aminocarbonyl)hydrazin-1-yl = semicarbazido	H ₂ N-CO-NH-NH-	P-68.3.1.2.4
carbamoylhydrazinylidene* = semicarbazono	H ₂ N-CO-NH-N=	P-68.3.1.2.5
carbazimidoyl: see hydrazinecarboximidoyl*		
carbazono: see diazenecarbohydrazido*		
carbazoyl: see hydrazinecarbonyl*		
carboethoxy: see ethoxycarbonyl*		
carbomethoxy: see methoxycarbonyl*		
carbonimidoyl*	-C(=NH)-	P-65.2.1.5
carbonobromidothioyl* = bromocarbonothioyl	Br-CS-	P-65.2.1.5
carbonochloridimidoyl* = C-chlorocarbonimidoyl	Cl-C(=NH)-	P-65.2.1.5
carbonochloridoyl* = chlorocarbonyl (not chloroformyl)	Cl-CO-	P-65.2.1.5
carbonocyanidoyl* = cyanocarbonyl = carbononitridoylcarbonyl	NC-CO-	P-65.2.1.5
carbonohydrazidimidoyl = hydrazinecarboximidoyl* = hydrazinyl(imino)methyl = C-hydrazinylcarbonimidoyl (not C-hydrazinocarbonimidoyl)	H ₂ N-NH-C(=NH)-	P-66.4.2.3.1
carbonohydrazidoyl = hydrazinecarbonyl* = hydrazinylcarbonyl (not carbazoyl; not hydrazinocarbonyl)	H ₂ N-NH-CO-	P-66.3.2.1
carbonohydrazonoyl*	-C(=N-NH ₂)-	P-65.2.1.5
carbononitridoyl = cyano*	NC-	P-65.2.2; P-66.5.1.1.4
carbononitridoylcarbonyl = carbonocyanidoyl* = cyanocarbonyl	NC-CO-	P-65.2.1.5
carbononitridoyl(disulfanyl) = cyanodisulfanyl* = carbononitridoyl(dithio (not thiocyanatosulfanyl)	NC-SS-	P-65.2.2
carbononitridoyl(dithio) = cyanodisulfanyl* = carbononitridoyl(disulfanyl)	NC-SS-	P-65.2.2

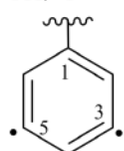
(not thiocyanatosulfanyl)		
carbononitridoxy = cyanato*	NC-O-	P-65.2.2
carbononitridoylperoxy = cyanoperoxy*	NC-OO-	P-65.2.2
carbononitridoylselanyl = selenocyanato*	NC-Se-	P-65.2.2
carbononitridoylsulfanyl = thiocyanato* = carbononitridoylthio	NC-S-	P-65.2.2
carbononitridoyltellanyl = tellurocyanato*	NC-Te-	P-65.2.2
carbononitridoylthio = thiocyanato* = carbononitridoylsulfanyl	NC-S-	P-65.2.2
carbonoperoxoxy* = (hydroperoxy)carbonyl (not peroxy-carboxy)	(HOO)-CO-	P-65.2.1.5
carbo(thio)peroxy* = (thiohydroperoxy)carbonyl	(HOS)-CO- or (HSO)-CO-	P-65.1.5.3; P-65.2.1.7
carbonothioyl* = thiocarbonyl	-CS-	P-65.2.1.5
carbonyl*	-CO-	P-65.2.1.5
carbonylbis(azanediyl)* (not ureylene)	-NH-CO-NH-	P-66.1.6.1.1.3
carboxy*	HOOC-	P-65.1.2.2.3; P-65.2.1.6
carboxyamino*	HOOC-NH-	P-65.2.1.6
carboxycarbonothioyl = carboxymethanethioyl*	HOOC-CS-	P-65.1.7.2.4; P-65.2.1.5
(carboxycarbonothioyl)sulfanyl = (carboxymethanethioyl)sulfanyl*	HOOC-CS-S-	P-65.1.7.2.4; P-65.2.1.5
carboxycarbonyl = oxalo* [not carboxyformyl; not hydroxyl(oxo)acetyl]	HOOC-CO-	P-65.1.2.2.3; P-65.1.7.2.1
(carboxycarbonyl)amino = oxaloamino*	HOOC-CO-NH-	P-65.1.7.2.4
(carboxycarbonyl)oxy = oxalooxy* [not (carboxyformyl)oxy]	HOOC-CO-O-	P-65.1.7.2.4
(carboxycarbonyl)sulfanyl = oxalosulfanyl* = (carboxycarbonyl)thio [not (carboxyformyl)sulfanyl; not (carboxyformyl)thio]	HOOC-CO-S-	P-65.1.7.2.4
(carboxycarbonyl)thio = oxalosulfanyl* = (carboxycarbonyl)sulfanyl [not (carboxyformyl)sulfanyl; not (carboxyformyl)thio]	HOOC-CO-S-	P-65.1.7.2.4
carboxyformyl: see oxalo*		
(carboxyformyl)oxy: see oxalooxy*		
(carboxyformyl)sulfanyl: see oxalosulfanyl*		
(carboxyformyl)thio: see oxalosulfanyl*		
carboxylato*	-O-CO-	P-72.6.1
carboxymethanethioyl* = carboxycarbonothioyl	HOOC-CS-	P-65.1.7.2.4; P-65.2.1.5
(carboxymethanethioyl)sulfanyl* = (carboxycarbonothioyl)sulfanyl	HOOC-CS-S-	P-65.1.7.2.4; P-65.2.1.5
3-carboxy-3-oxopropyl (not 2-oxaloethyl)	HOOC-CO-CH ₂ -CH ₂ -	P-65.1.2.2.3
carboxyoxy*	HOOC-O-	P-65.2.1.6
(carboxyoxy)carbonyl* [not (carboxyoxy)formyl]	HOOC-O-CO-	P-65.2.3.1.5
(carboxyoxy)formyl: see (carboxyoxy)carbonyl*		
carboxysulfanyl* = carboxythio	HOOC-S-	P-65.2.1.6
carboxythio = carboxysulfanyl*	HOOC-S-	P-65.2.1.6
chloro*	Cl-	P-61.3.1

chloroamidophosphoryl: see phosphoramidochloridoyl*		
chloroarsanyl*	ClAsH–	P-67.1.4.1.1.6
chloroboranyl* (not chloroboryl)	ClBH–	P-68.1.4.2
chloroboryl: see chloroboranyl*		
C-chlorocarbonimidoyl = carbonochloridimidoyl*	Cl-C(=NH)–	P-65.2.1.5
chlorocarbonyl = carbonochloridoyl* (not chloroformyl)	Cl-CO–	P-65.2.1.5
chloroformyl: see carbonochloridoyl*		
chlorooxalyl = chloro(oxo)acetyl*	ClCO-CO–	P-65.1.7.2.4
chloro(oxo)acetyl* = chlorooxalyl	ClCO-CO–	P-65.1.7.2.4
chloroso: see chlorosyl*		
chlorosulfinyl*	Cl-S(O)–	P-65.3.2.3; P-67.1.4.4.1
chlorosulfonyl* = sulfurochloridoyl	Cl-SO ₂ –	P-65.3.2.3; P-67.1.4.4.1
(chlorosulfonyl)oxy* = sulfurochloridoxyloxy	Cl-SO ₂ -O–	P-65.3.2.3; P-67.1.4.4.2
chlorosyl* (not chloroso)	OCl–	P-61.3.2.3
chloroxy: see chloryl*		
chloryl* (not chloroxy)	O ₂ Cl–	P-61.3.2.3
cinnamoyl = 3-phenylprop-2-enoyl*	C ₆ H ₅ -CH=CH-CO–	P-65.1.7.3.1; P-65.1.7.4.1
crotonyl: see but-2-enoyl*		
cyanato* = carbononitridoxyloxy	NC-O–	P-65.2.2
cyano* = carbononitridoyl	NC–	P-65.2.2; P-66.5.1.1.4
cyanocarbonyl = carbonocyanidoyl* = carbononitridoylcarbonyl	NC-CO–	P-65.2.1.5
cyanodisulfanyl* = carbononitridoyl(disulfanyl) = carbononitridoaldithio (not thiocyanatosulfanyl)	NC-SS–	P-65.2.2
cyano(isocyanato)phosphorothioyl = phosphorocyanidoisocyanatidothioyl* = cyano(isocyanato)(thiophosphoryl)	(OCN)(NC)P(S)–	P-67.1.4.1.1.4
cyano(isocyanato)(thiophosphoryl) = phosphorocyanidoisocyanatidothioyl* = cyano(isocyanato)phosphorothioyl	(OCN)(NC)P(S)–	P-67.1.4.1.1.4
cyanoperoxy* = carbononitridoylperoxy	NC-OO–	P-65.2.2
cyanosulfonyl* = sulfurocyanidoyl	NC-SO ₂ –	P-67.1.4.4.1
cyclohexanecarbonyl* = cyclohexylcarbonyl = cyclohexyl(oxo)methyl	C ₆ H ₁₁ -CO–	P-65.1.7.4.2
cyclohexanecarboximidoyl* = cyclohexylcarbonimidoyl = cyclohexyl(imino)methyl (not C-cyclohexylcarbonimidoyl)	C ₆ H ₁₁ -C(=NH)–	P-65.1.7.4.2
cyclohexane-1,1-diyl* (not cyclohexanylidene)	C ₆ H ₁₀ <	P-29.3.3
cyclohexane-1,4-diyl* (also 1,1-, 1,2-, and 1,3- isomers) (not 1,4-cyclohexylene)	–C ₆ H ₁₀ –	P-29.3.3
cyclohexanyl = cyclohexyl*	C ₆ H ₁₁ –	P-29.2; P-29.3.3
cyclohexanylidene = cyclohexylidene* (see also: cyclohexane-1,1-diyl)	C ₆ H ₁₀ =	P-29.3.3
cyclohexyl* = cyclohexanyl	C ₆ H ₁₁ –	P-29.2; P-29.3.3

cyclohexylcarbonimidoyl = cyclohexanecarboximidoyl* = cyclohexyl(imino)methyl (not <i>C</i> -cyclohexylcarbonimidoyl)	$C_6H_{11}-C(=NH)-$	P-65.1.7.4.2
<i>C</i> -cyclohexylcarbonimidoyl: see cyclohexanecarboximidoyl*		
cyclohexylcarbonyl = cyclohexanecarbonyl*	$C_6H_{11}-CO-$	P-65.1.7.4.2
1,4-cyclohexylene: see cyclohexane-1,4-diyl* (also 1,1-, 1,2- and 1,3-isomers)		
cyclohexylidene* = cyclohexanylidene (see also: cyclohexane-1,1-diyl)	$C_6H_{10}=$	P-29.3.3
cyclohexyl(imino)methyl = cyclohexanecarboximidoyl* = cyclohexylcarbonimidoyl (not <i>C</i> -cyclohexylcarbonimidoyl)	$C_6H_{11}-C(=NH)-$	P-65.1.7.4.2
cyclohexyl(oxo)methyl = cyclohexanecarbonyl* = cyclohexylcarbonyl	$C_6H_{11}-CO-$	P-65.1.7.4.2
cyclopentanecarbohydrazonoyl* = cyclopentyl(hydrazinylidene)methyl	$C_5H_9-C(=N-NH_2)-$	P-65.1.7.4.2
cyclopentanecarboximidoyl* = cyclopentyl(imino)methyl = cyclopentylcarbonimidoyl (not <i>C</i> -cyclopentylcarbonimidoyl)	$C_5H_9-C(=NH)-$	P-65.1.7.4.2
cyclopentylcarbonimidoyl = cyclopentanecarboximidoyl* = cyclopentyl(imino)methyl (not <i>C</i> -cyclopentylcarbonimidoyl)	$C_5H_9-C(=NH)-$	P-65.1.7.4.2
<i>C</i> -cyclopentylcarbonimidoyl: see cyclopentanecarboximidoyl*		
cyclopentyl(hydrazinylidene)methyl = cyclopentanecarbohydrazonoyl*	$C_5H_9-C(=N-NH_2)-$	P-65.1.7.4.2
cyclopentyl(imino)methyl = cyclopentanecarboximidoyl*	$C_5H_9-C(=NH)-$	P-65.1.7.4.2
cyclopropanyl = cyclopropyl*	C_3H_5-	P-29.3.3
cyclopropanylidene = cyclopropylidene*	$C_3H_4=$	P-29.3.3
cyclopropyl* = cyclopropanyl	C_3H_5-	P-29.3.3
cyclopropylidene* = cyclopropanylidene	$C_3H_4=$	P-29.3.3
		
cyclotrisilanyl = trisiliranyl*		P-68.2.2
decanedioyl* = 1,10-dioxodecane-1,10-diyl	$-CO-[CH_2]_8-CO-$	P-65.1.7.4.1
decanoyl* = 1-oxodecyl	$CH_3-[CH_2]_8-CO-$	P-65.1.7.4.1
decan-1-yl = decyl*	$CH_3-[CH_2]_8-CH_2-$	P-29.3.2.1; P-29.3.2.2
decan-1-ylidene = decylidene*	$CH_3-[CH_2]_8-CH=$	P-29.3.2.1; P-29.3.2.2
decanylidyne = decylidyne*	$CH_3-[CH_2]_8-C\equiv$	P-29.3.2.1; P-29.3.2.2
decyl* = decan-1-yl	$CH_3-[CH_2]_8-CH_2-$	P-29.3.2.1; P-29.3.2.2
decylidene* = decan-1-ylidene	$CH_3-[CH_2]_8-CH=$	P-29.3.2.1; P-29.3.2.2
decylidyne* = decanylidyne	$CH_3-[CH_2]_8-C\equiv$	P-29.3.2.1; P-29.3.2.2
diacetamido: see <i>N</i> -acetylacetamido*		
diacetoxyiodo: see bis(acetyloxy)- λ^3 -iodanyl*		
diacetylamino = <i>N</i> -acetylacetamido* (not diacetylazanyl; not diacetamido)	$(CH_3-CO)_2N-$	P-66.1.2.1
diacetylazanyl: see <i>N</i> -acetylacetamido*		
diaminoboranyl* (not borodiamidoyl)	$(H_2N)_2B-$	P-67.1.4.2
(diaminomethylidene)amino*	$(H_2N)_2C=N-$	P-66.4.1.2.1.3

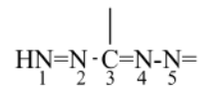
dihydrostiboryl = stibinoyl*	H ₂ Sb(O)–	P-67.1.4.1.1.2; P-67.1.4.1.2
dihydroxyarsoryl = arsono*	(HO) ₂ As(O)–	P-67.1.4.1.1.1
dihydroxyboranyl = borono*	(HO) ₂ B–	P-67.1.4.2; P-68.1.4.2
C,N-dihydroxycarbonimidoyl*	HO-C(=N-OH)–	P-65.1.3.3.2
dihydroxy-λ ³ -iodanyl* (not dihydroxyiodo)	(HO) ₂ I–	P-68.5.1
dihydroxyiodo: see dihydroxy-λ ³ -iodanyl*		
dihydroxynitroryl = azono*	(HO) ₂ N(O)–	P-67.1.4.1.1.1; P-67.1.4.1.1.5
dihydroxyphosphanyl* = dihydroxyphosphino	(HO) ₂ P–	P-67.1.4.1.1.6
dihydroxyphosphino = dihydroxyphosphanyl*	(HO) ₂ P–	P-67.1.4.1.1.6
dihydroxyphosphinothioyl: see dihydroxyphosphorothioyl*		
dihydroxyphosphorothioyl* (not dihydroxyphosphinothioyl)	(HO) ₂ P(S)–	P-67.1.4.1.1.5
dihydroxy(sulfanyl)silyl*	(HS)(HO) ₂ Si–	P-67.1.4.2
1,4-diiminobutane-1,4-diyl = butanediimidoyl* = succinimidoyl	–C(=NH)–CH ₂ –CH ₂ –C(=NH)–	P-65.1.7.3.2
diiminoethanediyl = ethanediimidoyl* = oxalimidoyl	–C(=NH)–C(=NH)–	P-65.1.7.2.2
1,3-diiminopropane-1,3-diyl = propanediimidoyl* = malonimidoyl	–C(=NH)–CH ₂ –C(=NH)–	P-65.1.7.4.1
dimethoxyphosphanyl*	(CH ₃ -O) ₂ P–	P-67.1.4.1.1.6
dimethoxyphosphoroselenoyl* = dimethoxy(selenophosphoryl)	(CH ₃ -O) ₂ P(Se)–	P-67.1.4.1.1.5
dimethoxyphosphoryl*	(CH ₃ -O) ₂ P(O)–	P-67.1.4.1.1.5
(dimethoxyphosphoryl)sulfanyl*	(CH ₃ -O) ₂ P(O)–S–	P-67.1.4.1.3
dimethoxy(selenophosphoryl) = dimethoxyphosphoroselenoyl*	(CH ₃ -O) ₂ P(Se)–	P-67.1.4.1.1.5
(dimethylamido)phosphoryl = N,N-dimethylphosphoramidoyl*	(CH ₃) ₂ N-P(O)<	P-67.1.4.1.1.4
dimethylammoniumylidene: see N-methylmethanaminiumylidene*		
2,3-dimethylanilino* = (2,3-dimethylphenyl)amino (not 2,3-xylidino) (also 2,4-, 2,5-, 2,6-, 3,4-, and 3,5-isomers)	2,3-(CH ₃) ₂ C ₆ H ₃ -NH–	P-62.2.1.1.2
dimethylazinoyl* (not dimethylnitroryl)	(CH ₃) ₂ N(O)–	P-67.1.6
(dimethylboranyl)oxy*	(CH ₃) ₂ B–O–	P-68.1.4.2
1,1-dimethylethoxy = (2-methylpropan-2-yl)oxy = <i>tert</i> -butoxy* (unsubstituted)	(CH ₃) ₃ C–O–	P-63.2.2.2
1,1-dimethylethyl = <i>tert</i> -butyl* (unsubstituted) = 2-methylpropan-2-yl	(CH ₃) ₃ C–	P-29.4.1; P-29.6.1
dimethylimmonio: see N-methylmethanaminiumylidene*		
dimethylnitroryl: see dimethylazinoyl*		
(2,3-dimethylphenyl)amino = 2,3-dimethylanilino* (not 2,3-xylidino) (also 2,4-, 2,5-, 2,6-, 3,4-, and 3,5-isomers)	2,3-(CH ₃) ₂ C ₆ H ₃ -NH–	P-62.2.1.1.2
dimethylphosphinoselenoyl* = dimethyl(selenophosphinoyl)	(CH ₃) ₂ P(Se)–	P-67.1.4.1.1.4
N,N-dimethylphosphoramidoyl* = (dimethylamido)phosphoryl	(CH ₃) ₂ N-P(O)<	P-67.1.4.1.1.4
1,1-dimethylpropyl = 2-methylbutan-2-yl* (not <i>tert</i> -pentyl)	CH ₃ –CH ₂ –C(CH ₃) ₂ –	P-29.6.3; P-57.1.4
2,2-dimethylpropyl* (not neopentyl)	CH ₃ –C(CH ₃) ₂ –CH ₂ –	P-57.1.4
dimethyl(selenophosphinoyl) = dimethylphosphinoselenoyl*	(CH ₃) ₂ P(Se)–	P-67.1.4.1.1.4

dioxo- λ^5 -arsanyl* (not arso)	O_2As-	P-61.6
1,4-dioxobutane-1,4-diyl = butanedioyl* = succinyl	$-CO-CH_2-CH_2-CO-$	P-65.1.7.3.1; P-65.1.7.4.1
(2E)-1,4-dioxobut-2-ene-1,4-diyl = (2E)-but-2-enedioyl* = fumaroyl	$\begin{array}{c} 2 \quad 1 \\ HC-CO- \\ \\ -OC-CH \\ 4 \quad 3 \end{array}$	P-65.1.7.3.1; P-65.1.7.4.1
(2Z)-1,4-dioxobut-2-ene-1,4-diyl = (2Z)-but-2-enedioyl* = maleoyl	$\begin{array}{c} 2 \quad 1 \\ HC-CO- \\ \\ HC-CO- \\ 3 \quad 4 \end{array}$	P-65.1.7.3.1; P-65.1.7.4.1
1,10-dioxodecane-1,10-diyl = decanedioyl*	$-CO-[CH_2]_8-CO-$	P-65.1.7.4.1
1,3-dioxo-1,3-dihydro-2H-isoindol-2-yl* = phthalimido		P-66.2.2
dioxoethanediyl = oxalyl* = ethanedioyl	$-CO-CO-$	P-65.1.7.2.1
1,6-dioxohexane-1,6-diyl = hexanedioyl* = adipoyl	$-CO-[CH_2]_4-CO-$	P-65.1.7.3.1; P-65.1.7.4.1
1,5-dioxopentane-1,5-diyl = pentanedioyl* = glutaryl	$-CO-CH_2-CH_2-CH_2-CO-$	P-65.1.7.3.1; P-65.1.7.4.1
dioxo- λ^5 -phosphanyl* (not phospho)	O_2P-	P-61.6; P-67.1.4.1.1.6
1,3-dioxopropane-1,3-diyl = propanedioyl* = malonyl	$-CO-CH_2-CO-$	P-65.1.7.3.1; P-65.1.7.4.1
1,2-dioxopropyl = 2-oxopropanoyl* (not pyruvoyl)	$CH_3-CO-CO-$	P-65.1.1.2.3; P-65.1.7.4.1
2,5-dioxopyrrolidin-1-yl* = succinimido		P-66.2.2
dioxy: see peroxy*		
diphenylmethyl* (not benzhydryl)	$(C_6H_5)_2CH-$	P-29.6.3
diphosphanyl* (not diphosphino)	H_2P-PH-	P-29.3.2.2; P-45.3.1; P-68.3.2.3.2.2
diphosphino: see diphosphanyl*		
diselanediyl* = diseleno	$-Se-Se-$	P-63.3.1
diselanyl* = diselenohydroperoxy	$HSeSe-$	P-63.4.2.2
diseleno = diselanediyl*	$-Se-Se-$	P-63.3.1
diselenoborono* = bis(selanyl)boranyl	$(HSe)_2B-$	P-68.1.4.2
diselenohydroperoxy = diselanyl*	$HSeSe-$	P-63.4.2.2
disilane-1,1-diyl*	$H_3Si-SiH<$	P-29.3.2.2; P-68.2.2
disilanyl* (disilyl)	$H_3Si-SiH_2-$	P-29.3.2.2; P-68.2.2

disilazan-1-yl: see (silylamino)silyl*		
disilazan-2-yl: see disilylamino*		
disiloxanyl*	$\text{H}_3\text{Si-O-SiH}_2\text{-}$	P-29.3.2.2; P-46.1.3
disilyl: see disilanyl*		
disilylamino* (not disilazan-2-yl)	$(\text{SiH}_3)_2\text{N-}$	P-29.3.2.2; P-68.2.2
disulfanediyl* = dithio	-S-S-	P-63.3.1
disulfanidyl*	-S-S-	P-72.6.3
disulfanyl* = dithiohydroperoxy (not thiosulfeno)	HS-S-	P-63.4.2.2
(disulfanylcarbonyl)oxy* = [(dithiohydroperoxy)carbonyl]oxy	HS-S-CO-O-	P-65.2.1.7
ditellanediyl* = ditelluro	-Te-Te-	P-63.3.1
ditellanyl* = ditellurohydroperoxy	HTe-Te-	P-63.4.2.2
ditelluro = ditellanediyl*	-Te-Te-	P-63.3.1
ditellurohydroperoxy = ditellanyl*	HTe-Te-	P-63.4.2.2
dithio = disulfanediyl*	-S-S-	P-63.3.1
dithiocarbonperoxy* (location of sulfur atoms unknown)	$\text{HOS}_2\text{C-}$	P-65.1.5.3
dithiocarboxy* = sulfanylcarbonothioyl	HS-CS-	P-65.2.1.6
[(dithiocarboxy)sulfanyl]carbonothioyl* = [sulfanyl(thiocarbonyl)sulfanyl](thiocarbonyl) = [(sulfanylcarbonothioyl)sulfanyl]carbonothioyl {not [(dithiocarboxy)sulfanyl]thioformyl}	HS-CS-S-CS-	P-65.2.3.1.5
[(dithiocarboxy)sulfanyl]thioformyl: see [(dithiocarboxy)sulfanyl]carbonothioyl*		
dithiohydroperoxy = disulfanyl* (not thiosulfeno)	HS-S-	P-63.4.2.2
[(dithiohydroperoxy)carbonyl]oxy = (disulfanylcarbonyl)oxy*	HS-S-CO-O-	P-65.2.1.7
1,2-dithiooxalo: see hydroxy(sulfanylidene)ethanethioyl*		
dithiooxalyl = ethanebis(thioyl)* = bis(sulfanylidene)ethanediyl	-CS-CS-	P-65.1.7.2.3
dithiophthaloyl: see benzene-1,2-dicarbothioyl*		
dithiosuccinyl: see butanebis(thioyl)*		
dithiosulfo* (unspecified)	$\text{HO-SS}_2\text{-}$ or HS-S(=S)(=O)-	P-65.3.2.1
1,4-dithioxobutane-1,4-diyl = butanebis(thioyl)* = 1,4-bis(sulfanylidene)butane-1,4-diyl (not dithiosuccinyl)	$\text{-CS-CH}_2\text{-CH}_2\text{-CS-}$	P-65.1.7.4.1; P-65.1.7.4.3
1,1-diyloethyl*	$\text{CH}_3\text{-C}^2\text{-}$	P-71.5
		P-71.5
3,5-diylophenyl*		
dodecanoyl* = 1-oxododecyl	$\text{CH}_3\text{-[CH}_2\text{]}_{10}\text{-CO-}$	P-65.1.7.4.1
dodecan-1-yl = dodecyl*	$\text{CH}_3\text{-[CH}_2\text{]}_{10}\text{-CH}_2\text{-}$	P-29.3.2.1; P-29.3.2.2
dodecyl* = dodecan-1-yl	$\text{CH}_3\text{-[CH}_2\text{]}_{10}\text{-CH}_2\text{-}$	P-29.3.2.1; P-29.3.2.2
episeleno = selano* (ring forming)	-Se-	P-25.4.2.1.4; P-63.5

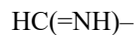
epitelluro = tellano* (ring forming)	-Te-	P-25.4.2.1.4; P-63.5
epithio = sulfano* (ring forming)	-S-	P-25.4.2.1.4; P-63.5
epoxidano: see epoxy* (ring forming)		
epoxy* (ring forming) (not epoxidano)	-O-	P-25.4.2.1.4; P-63.5
ethanebis(thioyl)* = dithiooxalyl = bis(sulfanylidene)ethanediyyl	-CS-CS-	P-65.1.7.2.3
ethanediimidoyl* = oxalimidoyl = diiminoethanediyyl	-C(=NH)-C(=NH)-	P-65.1.7.2.2
ethanedioyl = oxalyl* = dioxoethanediyyl	-CO-CO-	P-65.1.7.2.1
ethanedioylbis(azanediyyl) = oxalylbis(azanediyyl)*	-HN-CO-CO-NH-	P-66.1.1.4.5.2
ethanedioylbis(azanetriyl) = oxalyldinitrilo* = oxalylbis(azanetriyl) = ethanedioyldinitrilo	>N-CO-CO-N<	P-66.1.1.4.5.2
ethanedioylbis(azanylylidene) = oxalylbis(azanylylidene)*	=N-CO-CO-N=	P-66.1.1.4.5.2
ethanedioyldinitrilo = oxalyldinitrilo* = oxalylbis(azanetriyl) = ethanedioylbis(azanetriyl)	>N-CO-CO-N<	P-66.1.1.4.5.2
ethane-1,1-diyl*	CH ₃ -CH<	P-29.3.2.2
ethane-1,2-diyl* = ethylene	-CH ₂ -CH ₂ -	P-29.3.2.2; P-29.6.2.3
ethane-1,2-diylbis(oxy)* = ethylenebis(oxy) (not ethane-1,2-diylldioxy, not ethylenedioxy)	-O-CH ₂ -CH ₂ -O-	P-63.2.2.1.3
ethane-1,2-diylldioxy: see ethane-1,2-diylbis(oxy)*		
ethanehydrazonamido* = (ethanehydrazonoyl)amino	CH ₃ -C(=N-NH ₂)-NH-	P-66.4.2.3.5
ethanehydrazonoyl* = acetohydrazonoyl = 1-hydrazinylideneethyl	CH ₃ -C(=N-NH ₂)-	P-65.1.7.2.2
(ethanehydrazonoyl)amino = ethanehydrazonamido*	CH ₃ -C(=N-NH ₂)-NH-	P-66.4.2.3.5
ethaneselenoyl* = selenoacetyl = 1-selanylideneethyl	CH ₃ -CSe-	P-65.1.7.2.3
ethanesulfinimidoyl* = S-ethylsulfinimidoyl	CH ₃ -CH ₂ -S(=NH)-	P-65.3.2.2.2
ethanesulfinyl* = ethylsulfinyl	CH ₃ -CH ₂ -S(O)-	P-63.6; P-65.3.2.2.2
ethanesulfonimidoyl* = S-ethylsulfonimidoyl	CH ₃ -CH ₂ -S(O)(=NH)-	P-65.3.2.2.2
ethanesulfonodiimidamido* = ethanesulfonodiimidoylamino	CH ₃ -CH ₂ -S(=NH) ₂ -NH-	P-66.4.1.3.5
ethanesulfonodiimidoylamino = ethanesulfonodiimidamido*	CH ₃ -CH ₂ -S(=NH) ₂ -NH-	P-66.4.1.3.5
ethanesulfonothioyl* = ethylsulfonothioyl	CH ₃ -CH ₂ -S(O)(S)-	P-65.3.2.2.2
ethanesulfonyl* = ethylsulfonyl	CH ₃ -CH ₂ -SO ₂ -	P-63.6; P-65.3.2.2.2
ethanethioamido* = (ethanethioyl)amino = thioacetamido	CH ₃ -CS-NH-	P-66.1.4.4
ethanethioyl* = thioacetyl = 1-sulfanylideneethyl	CH ₃ -CS-	P-65.1.7.2.3
ethanethioamido* = (ethanethioyl)amino = thioacetamido	CH ₃ -CS-NH-	P-66.1.4.4
(ethanethioyl)amino = ethanethioamido* = thioacetamide	CH ₃ -CS-NH-	P-66.1.4.4
ethanimidamido* = acetimidamido = acetimidoylamino	CH ₃ -C(=NH)-NH-	P-66.4.1.3.5
ethanimidohydrazido* = 2-(ethanimidoyl)hydrazin-1-yl	CH ₃ -C(=NH)-NH-NH-	P-66.4.2.3.6
ethanimidoyl* = acetimidoyl = 1-iminoethyl	CH ₃ -C(=NH)-	P-65.1.7.2.2
2-(ethanimidoyl)hydrazin-1-yl = ethanimidohydrazido*	CH ₃ -C(=NH)-NH-NH-	P-66.4.2.3.6
ethanoyl = acetyl* = 1-oxoethyl	CH ₃ -CO-	P-65.1.7.2.1
ethanyl = ethyl*	CH ₃ -CH ₂ -	P-29.3.2.1; P-29.3.2.2
ethanylidene = ethylidene*	CH ₃ -CH=	P-29.3.2.1; P-29.3.2.2

formazan-3-yl-5-ylidene*



P-34.2.1.3; P-68.3.1.3.5.2

formimidoyl = methanimidoyl* = iminomethyl



P-65.1.7.2.2

formimidoylamino = methanimidamido* = (iminomethyl)amino



P-66.4.1.3.3

formohydrazido* = 2-formylhydrazin-1-yl



P-66.3.5.3

formohydrazonoyl = methanehydrazonoyl* = hydrazinylidenemethyl



P-65.1.7.2.2

formyl* = methanoyl = oxomethyl



P-65.1.7.2.1; P-66.6.1.3

formylamino = formamido*



P-66.1.1.4.3

formylazanedyl*



P-66.1.1.4.4

formylazanylidene = formylimino*



P-66.1.1.4.4

2-formylhydrazin-1-yl = formohydrazido*



P-66.3.5.3

formylimino* = formylazanylidene*



P-66.1.1.4.4

formyloxy*



P-65.1.8.3; P-65.6.3.2.3

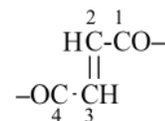
formylsulfanyl*



P-65.1.8.3

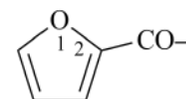
fulminato: see (λ^2 -methylideneamino)oxy*

fumaroyl = (2*E*)-but-2-enedioyl* = (2*E*)-1,4-dioxobut-2-ene-1,4-diyl



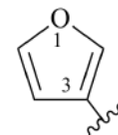
P-65.1.7.3.1; P-65.1.7.4.1

furan-2-carbonyl* = 2-furoyl = 2-furylcarbonyl (also 3-isomer)



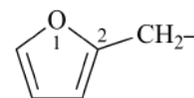
P-65.1.7.3.1; P-65.1.7.4.2

furan-3-yl* = 3-furyl (also 2-isomer)



P-29.6.2.3; P-57.1.5.3

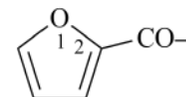
(furan-2-yl)methyl* (not furfuryl)



P-29.6.3

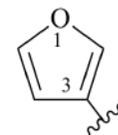
furfuryl (2-isomer only): see (furan-2-yl)methyl*

2-furoyl = furan-2-carbonyl* = 2-furylcarbonyl (also 3-isomer)



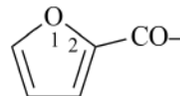
P-65.1.7.3.1; P-65.1.7.4.2

3-furyl = furan-3-yl* (also 2-isomer)



P-29.6.2.3; P-57.1.5.3

2-furylcarbonyl = furan-2-carbonyl* = 2-furoyl (also 3-isomer)



P-65.1.7.3.1; P-65.1.7.4.2

gallanyl*

H₂Ga-

P-29.3.1; P-68.1.2

germanediyl* (not germylene)

H₂Ge<

P-68.2.2

germanediylidene*

=Ge=

P-68.2.2

germanetetrayl*

>Ge<

P-68.2.2

germanetriyl*

-GeH<

P-68.2.2

germanyyl = germyl*

H₃Ge-

P-29.3.1; P-68.2.2

germanyliidene = germylidene*

H₂Ge=

P-29.3.1; P-68.2.2

germanyliidyne = germylidyne*

HGe≡

P-29.3.1; P-68.2.2

germanylylidene*

-GeH=

P-68.2.2

germyl* = germanyyl

H₃Ge-

P-29.3.1; P-68.2.2

germylene: see germanediyl

germyliidene* = germanyliidene

H₂Ge=

P-29.3.1; P-68.2.2

germyliidyne* = germanyliidyne

HGe≡

P-29.3.1; P-68.2.2

glutaryl = pentanedioyl* = 1,5-dioxopentane-1,5-diyl

-CO-CH₂-CH₂-CH₂-CO-

P-65.1.1.2.2; P-65.1.7.3.1

guanidino = carbamimidoylamino* = carbamimidamido = [amino(imino)methyl]amino

H₂N-C(=NH)-NH-

P-66.4.1.2.1.3

heptanoyl* = 1-oxoheptyl

CH₃-[CH₂]₅-CO-

P-65.1.7.4.1

heptan-1-yl = heptyl*

CH₃-[CH₂]₅-CH₂-

P-29.3.2.1; P-29.3.2.2

heptan-1-ylidene = heptylidene*

CH₃-[CH₂]₅-CH=

P-29.3.2.1; P-29.3.2.2

heptanyliidyne = heptyliidyne*

CH₃-[CH₂]₅-C≡

P-29.3.2.1; P-29.3.2.2

heptyl* = heptan-1-yl

CH₃-[CH₂]₅-CH₂-

P-29.3.2.1; P-29.3.2.2

heptylidene* = heptan-1-ylidene

CH₃-[CH₂]₅-CH=

P-29.3.2.1; P-29.3.2.2

heptyliidyne* = heptanyliidyne

CH₃-[CH₂]₅-C≡

P-29.3.2.1; P-29.3.2.2

hexadecanoyl* = palmitoyl = 1-oxohexadecyl

CH₃-[CH₂]₁₄-CO-

P-65.1.7.3.1

hexadecan-1-yl = hexadecyl*

CH₃-[CH₂]₁₄-CH₂-

P-29.3.2.1; P-29.3.2.2

hexadecyl* = hexadecan-1-yl

CH₃-[CH₂]₁₄-CH₂-

P-29.3.2.1; P-29.3.2.2

hexamethylene: see hexane-1,6-diyl*

hexanedioyl* = adipoyl = 1,6-dioxohexane-1,6-diyl

-CO-[CH₂]₄-CO-

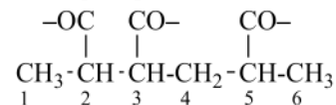
P-65.1.7.3.1; P-65.1.7.4.1

hexane-1,6-diyl* (not hexamethylene)

-CH₂-[CH₂]₄-CH₂-

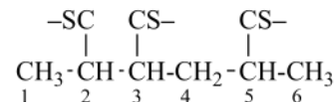
P-29.3.2.2

hexane-2,3,5-tricarbonyl* = hexane-2,3,5-triyltris(oxomethylene) = hexane-2,3,5-triyltricarbonyl



P-65.1.7.4.2

hexane-2,3,5-tricarbothioyl* = hexane-2,3,5-triyltris(sulfanyliidene)methylene) = hexane-2,3,5-triyltris(thioxomethylene)



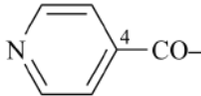
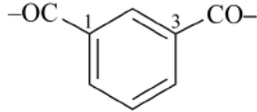
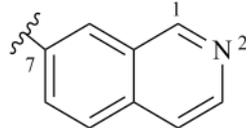
P-65.1.7.4.2

hexane-2,3,5-triyltris(oxomethylene) = hexane-2,3,5-tricarbonyl* = hexane-2,3,5-triyltricarbonyl	$\begin{array}{cccccc} & \text{-OC} & \text{CO-} & & \text{CO-} & \\ & & & & & \\ \text{CH}_3 & \text{-CH} & \text{-CH} & \text{-CH}_2 & \text{-CH} & \text{-CH}_3 \\ & 1 & 2 & 3 & 4 & 5 & 6 \end{array}$	P-65.1.7.4.2
hexane-2,3,5-triyltricarbonyl = hexane-2,3,5-tricarbonyl* = hexane-2,3,5-triyltris(oxomethylene)		
hexane-2,3,5-triyltris(thioxomethylene) = hexane-2,3,5-tricarbothioyl* = hexane-2,3,5-triyltris(sulfanylidene)methylene)	$\begin{array}{cccccc} & \text{-SC} & \text{CS-} & & \text{CS-} & \\ & & & & & \\ \text{CH}_3 & \text{-CH} & \text{-CH} & \text{-CH}_2 & \text{-CH} & \text{-CH}_3 \\ & 1 & 2 & 3 & 4 & 5 & 6 \end{array}$	P-65.1.7.4.2
hexane-2,3,5-triyltris(sulfanylidene)methylene) = hexane-2,3,5-tricarbothioyl* = hexane-2,3,5-triyltris(thioxomethylene)		
hexanoyl* = 1-oxohexyl	$\text{CH}_3\text{-[CH}_2\text{]}_4\text{-CO-}$	P-65.1.7.4.1
hexan-1-yl = hexyl*	$\text{CH}_3\text{-[CH}_2\text{]}_5\text{-}$	P-29.3.2.1; P-29.3.2.2
hexan-1-ylidene = hexylidene*	$\text{CH}_3\text{-[CH}_2\text{]}_4\text{-CH=}$	P-29.3.2.1; P-29.3.2.2
hexanylidyne = hexylidyne*	$\text{CH}_3\text{-[CH}_2\text{]}_4\text{-C}\equiv$	P-29.3.2.1; P-29.3.2.2
hexyl* = hexan-1-yl	$\text{CH}_3\text{-[CH}_2\text{]}_5\text{-}$	P-29.3.2.1; P-29.3.2.2
hexylidene* = hexan-1-ylidene	$\text{CH}_3\text{-[CH}_2\text{]}_4\text{-CH=}$	P-29.3.2.1; P-29.3.2.2
hexylidyne* = hexanylidyne	$\text{CH}_3\text{-[CH}_2\text{]}_4\text{-C}\equiv$	P-29.3.2.2; P-29.3.2.1
hydrazi: (not to be used to form heterocycles)		
hydrazidimidophosphoryl = phosphorohydrazidimidoyl*	$(\text{H}_2\text{N-NH})\text{-P(=NH)}<$	P-67.1.4.1.1.4
hydrazinecarbohydrazido* = 2-(hydrazinecarbonyl)hydrazin-1-yl = 2-(hydrazinylcarbonyl)hydrazin-1-yl	$\text{H}_2\text{N-NH-CO-NH-NH-}$	P-66.3.5.3; P-68.3.1.2.6
hydrazinecarbohydrazonoyl* = C-hydrazinylcarbohydrazonoyl = hydrazinyl(hydrazinylidene)methyl	$\text{H}_2\text{N-NH-C(=N-NH}_2\text{)-}$	P-66.4.3.4.1
hydrazinecarbonyl* = hydrazinylcarbonyl = carbonohydrazidoyl (not carbazoyl; not hydrazinocarbonyl)	$\text{H}_2\text{N-NH-CO-}$	P-66.3.2.1
(hydrazinecarbonyl)diazenyl* = (hydrazinylcarbonyl)diazenyl	$\text{H}_2\text{N-NH-CO-N=N-}$	P-68.3.1.3.4
2-(hydrazinecarbonyl)hydrazin-1-yl* = hydrazinecarbohydrazido = 2-(hydrazinylcarbonyl)hydrazin-1-yl	$\text{H}_2\text{N-NH-CO-NH-NH-}$	P-66.3.5.3; P-68.3.1.2.6
(hydrazinecarbonyl)hydrazinylidene* = (hydrazinylcarbonyl)hydrazinylidene	$\text{H}_2\text{N-NH-CO-NH-N=}$	P-68.3.1.2.6
hydrazinecarboximidoyl* = hydrazinyl(imino)methyl = C-hydrazinylcarbonimidoyl = carbonohydrazidimidoyl (not carbazimidoyl; not C-hydrazinocarbonimidoyl)	$\text{H}_2\text{N-NH-CO(=NH)-}$	P-66.4.2.3.1
hydrazine-1,2-diyl* = diazane-1,2-diyl (not hydrazo)	-NH-NH-	P-29.3.2.2; P-68.3.1.2.1
hydrazinediylidene* = diazannediylidene (not azino)	$=\text{N-N=}$	P-29.3.2.2; P-68.3.1.2.1
hydrazinesulfinyl* = hydrazinylsulfinyl (not hydrazinosulfinyl)	$\text{H}_2\text{N-NH-S(O)-}$	P-66.3.2.1
hydrazinesulfonyl* = hydrazinylsulfonyl (not hydrazinosulfonyl)	$\text{H}_2\text{N-NH-SO}_2\text{-}$	P-66.3.2.1
hydrazino: see hydrazinyl*		
C-hydrazinocarbonimidoyl: see hydrazinecarboximidoyl*		
hydrazinocarbonyl: see hydrazinecarbonyl*		
hydrazinosulfinyl: see hydrazinesulfinyl*		
hydrazinosulfonyl: see hydrazinesulfonyl*		
hydrazinyl* = diazanyl (not hydrazino)	$\text{H}_2\text{N-NH-}$	P-29.3.2.2; P-68.3.1.2.1

C-hydrazinylcarbonimidoyl = hydrazinecarboximidoyl* = hydrazinyl(imino)methyl = carbonohydrazidimidoyl (not carbazimidoyl; not C-hydrazinocarbonimidoyl)	$\text{H}_2\text{N-NH-CO(=NH)-}$	P-66.4.2.3.1
C-hydrazinylcarbonohydrazonoyl = hydrazinecarbohydrazonoyl* = hydrazinyl(hydrazinylidene)methyl	$\text{H}_2\text{N-NH-C(=N-NH}_2\text{)-}$	P-66.4.3.4.1
hydrazinylcarbonyl = hydrazinecarbonyl* = carbonohydrazidoyl (not carbazoyl; not hydrazinocarbonyl)	$\text{H}_2\text{N-NH-CO-}$	P-66.3.2.1
(hydrazinylcarbonyl)diazenyl = (hydrazinecarbonyl)diazenyl*	$\text{H}_2\text{N-NH-CO-N=N-}$	P-68.3.1.3.4
2-(hydrazinylcarbonyl)hydrazin-1-yl = hydrazinecarbohydrazido* = 2-(hydrazinecarbonyl)hydrazin-1-yl	$\text{H}_2\text{N-NH-CO-NH-NH-}$	P-66.3.5.3; P-68.3.1.2.6
(hydrazinylcarbonyl)hydrazinylidene = (hydrazinecarbonyl)hydrazinylidene*	$\text{H}_2\text{N-NH-CO-NH-N=}$	P-68.3.1.2.6
hydrazinyl(hydrazinylidene)methyl = hydrazinecarbohydrazonoyl* = C-hydrazinylcarbonohydrazonoyl	$\text{H}_2\text{N-NH-C(=N-NH}_2\text{)-}$	P-66.4.3.4.1
hydrazinylidene* = diazanylidene (not hydrazono)	$\text{H}_2\text{N-N=}$	P-29.3.2.2; P-68.3.1.2.1
1-hydrazinylideneethyl = ethanehydrazonoyl* = acetohydrazonoyl	$\text{CH}_3\text{-C(=N-NH}_2\text{)-}$	P-65.1.7.2.2
hydrazinylidene(hydroxy)methyl = C-hydroxycarbonohydrazonoyl* [not hydrazono(hydroxy)methyl]	$\text{HO-C(=N-NH}_2\text{)-}$	P-65.1.3.2.2
hydrazinylidenemethyl = methanehydrazonoyl* = formohydrazonoyl	$\text{HC(=N-NH}_2\text{)-}$	P-65.1.7.2.2
(hydrazinylidenemethyl)amino = methanehydrazonamido* = methanehydrazonoylamino	$\text{HC(=N-NH}_2\text{)-NH-}$	P-66.4.2.3.3
(hydrazinylidenemethyl)diazenyl = formazan-1-yl*	$\text{HC(=N-NH}_2\text{)-N=N-}$	P-34.2.1.3; P-68.3.1.3.5.2
2-(hydrazinylidenemethyl)hydrazin-1-yl = methanehydrazonohydrazido* = 2-(methanehydrazonoyl)hydrazin-1-yl	$\text{HC(=N-NH}_2\text{)-NH-NH-}$	P-66.4.3.4.2
hydrazinylidenemethylidene* = diazanylidene(methylidene) (not hydrazonomethylidene)	$\text{H}_2\text{N-N=C=}$	P-65.2.1.8
1-hydrazinylideneprop-2-en-1-yl = prop-2-enehydrazonoyl* = acrylohydrazonoyl	$\text{CH}_2\text{=CH-C(=N-NH}_2\text{)-}$	P-65.1.7.3.2
hydrazinyl(imino)methyl = hydrazinecarboximidoyl* = carbonohydrazidimidoyl = C-hydrazinylcarbonimidoyl	$\text{H}_2\text{N-NH-C(=NH)-}$	P-66.4.2.3.1
hydrazinylsulfinyl = hydrazinesulfinyl* (not hydrazinosulfinyl)	$\text{H}_2\text{N-NH-S(O)-}$	P-66.3.2.1
hydrazinylsulfonyl = hydrazinesulfonyl* (not hydrazinosulfonyl)	$\text{H}_2\text{N-NH-SO}_2\text{-}$	P-66.3.2.1
hydrazo: see hydrazine-1,2-diyl*		
hydrazono: see hydrazinylidene*		
hydrazono(hydroxy)methyl: see C-hydroxycarbonohydrazonoyl		
hydrazonomethylidene:: see hydrazinylidenemethylidene*		
hydrazonostiboryl = stiborohydrazonoyl*	$\text{-Sb(=N-NH}_2\text{)<}$	P-67.1.4.1.1.4
hydroarsoryl = arsonoyl*	HAs(O)<	P-67.1.4.1.1.2; P-67.1.4.1.2
hydromethoxyboryl: see methoxyboranyl*		
hydronitrolyl = azonoyl*	HN(O)<	P-67.1.4.1.1.2; P-67.1.4.1.2
hydroperoxy*	HOO-	P-63.4.2.2
(hydroperoxy)carbonyl = carbonoperoxoyl* (not peroxy-carboxy)	(HOO)-CO-	P-65.2.1.5
(hydroperoxy)phosphoryl = phosphoroperoxoyl* = peroxyphosphoryl	(HOO)-P(O)<	P-67.1.4.1.1.4

hydrophosphoryl = phosphonyl*	HP(O)<	P-67.1.4.1.1.2; P-67.1.4.1.2
hydroseleninyl*	HSe(O)–	P-65.3.2.3
hydroseleno: see selanyl*		
hydrostiboryl = stibonoyl*	HSb(O)<	P-67.1.4.1.1.2; P-67.1.4.1.2
hydrosulfinyl*	HS(O)–	P-65.3.2.3
hydrosulfonyl*	HSO ₂ –	P-65.3.2.3
hydrotelluro: see tellanyl*		
hydro(thiophosphoryl) = phosphonothioyl*	HP(S)<	P-67.1.4.1.2
hydrotrioxy = trioxidanyl*	HO-O-O–	P-68.4.1.3
hydrotriseleno = triselanyl*	HSe-Se-Se–	P-68.4.1.3
hydrotritelluro = tritellanyl*	HTe-Te-Te–	P-68.4.1.3
hydrotrithio = trisulfanyl*	HS-S-S–	P-68.4.1.3
hydroxy* (not oxidanyl)	HO–	P-63.1.4
<i>N</i> -hydroxyacetimidoyl = <i>N</i> -hydroxyethanimidoyl* = acetohydroximoyl	CH ₃ C(=N-OH)–	P-65.1.7.2.2
hydroxyamino* (not hydroxylamino)	HO-NH–	P-68.3.1.1.1.5
hydroxyarsanyl*	(HO)AsH–	P-67.1.4.1.1.6
hydroxyarsoryl*	(HO)As(O)<	P-67.1.4.1.1.5
hydroxyazanediyyl*	HO-N<	P-68.3.1.1.1.5
hydroxyazonoyl*	(HO)HN(O)–	P-67.1.4.1.1.5
<i>N</i> -hydroxybenzenecarboximidoyl* = <i>N</i> -hydroxybenzimidoyl = benzenecarbohydroximoyl = benzhydroximoyl	C ₆ H ₅ -C(=N-OH)–	P-65.1.7.2.2
<i>N</i> -hydroxybenzimidoyl = <i>N</i> -hydroxybenzenecarboximidoyl* = benzenecarbohydroximoyl = benzhydroximoyl	C ₆ H ₅ -C(=N-OH)–	P-65.1.7.2.2
hydroxybis(sulfanylidene)ethyl = hydroxy(sulfanylidene)ethanethioyl* (not 1,2-dithiooxalo)	HO-CS-CS–	P-65.1.7.2.4
hydroxyboranyl*	(HO)-HB–	P-67.1.4.2
<i>C</i> -hydroxycarbonimidoyl* = hydroxy(imino)methyl	HO-C(=NH)–	P-35.3.2; P-65.1.3.1.2
(<i>C</i> -hydroxycarbonimidoyl)amino* = [hydroxy(imino)methyl]amino (not 1-isoureido)	HO-C(=NH)-NH–	P-66.1.6.1.2.2
<i>C</i> -hydroxycarbonohydrazonoyl* = hydrazinylidene(hydroxy)methyl [not hydrazono(hydroxy)methyl]	HO-C(=N-NH ₂)–	P-65.1.3.2.2
hydroxycarbonothioyl*	HO-CS–	P-65.2.1.6
(hydroxycarbonothioyl)carbonyl = hydroxy(thiocarbonyl)carbonyl = hydroxy(sulfanylidene)acetyl* (not 2-thiooxalo; not 2-hydroxy-2-thiooxaly)	HO-CS-CO–	P-65.1.7.2.4
<i>N</i> -hydroxyethanimidoyl* = <i>N</i> -hydroxyacetimidoyl = acetohydroximoyl	CH ₃ -C(=N-OH)–	P-65.1.7.2.2
hydroxyimino*	HO-N=	P-68.3.1.1.2
hydroxy(imino)methyl = <i>C</i> -hydroxycarbonimidoyl*	HO-C(=NH)–	P-35.3.2; P-65.1.3.1.2
[hydroxy(imino)methyl]amino = (<i>C</i> -hydroxycarbonimidoyl)amino* (not 1-isoureido)	HO-C(=NH)-NH–	P-66.1.6.1.2.2
hydroxylamino: see hydroxyamino*		

hydroxy(mercapto)phosphoryl: see hydroxy(sulfanyl)phosphoryl*		
hydroxy(methyl)boranyl*	CH ₃ (HO)B–	P-68.1.4.2
hydroxy(methylphosphonoyl)* = hydroxy(methyl)phosphoryl	CH ₃ -P(O)(OH)–	P-67.1.4.1.1.5
hydroxy(methyl)phosphoryl = hydroxy(methylphosphonoyl)*	CH ₃ -P(O)(OH)–	P-67.1.4.1.1.5
hydroxy(oxo)acetyl: see oxalo*		
hydroxy(oxo)-λ ⁵ -arsanylidene*	HO-As(O)=	P-67.1.4.1.1.6
hydroxy(oxo)-λ ⁵ -azanylidene* = <i>aci</i> -nitro	HO-N(O)=	P-61.5.3; P-67.1.4.1.1.6; P-67.1.6
hydroxy(oxo)-λ ⁵ -phosphanylidene*	HO-P(O)=	P-67.1.4.1.1.6
hydroxy(oxo)-λ ⁵ -stibanediyyl = hydroxystiboryl*	HO-Sb(O)<	P-67.1.4.1.1.5; P-67.1.4.1.1.6
hydroxy(oxo)-λ ⁵ -stibanylidene*	HO-Sb(O)=	P-67.1.4.1.1.6
hydroxyphosphanylidene*	HO-P=	P-67.1.4.1.1.6
hydroxyphosphoryl*	HO-P(O)<	P-67.1.4.1.1.5
hydroxyselanyl* = <i>OSe</i> -selenohydroperoxy (not seleneno)	HO-Se–	P-63.4.2.2
(hydroxyselanyl)methyl* = (<i>OSe</i> -selenohydroperoxy)methyl	(HO-Se)-CH ₂ –	P-63.4.2.2
hydroxystibanediyyl*	HO-Sb<	P-67.1.4.1.1.6
hydroxystiboryl* = hydroxy(oxo)-λ ⁵ -stibanediyyl	HO-Sb(O)<	P-67.1.4.1.1.5; P-67.1.4.1.1.6
hydroxysulfanyl* = <i>OS</i> -thiohydroperoxy (not sulfeno; not hydroxythio)	HO-S–	P-63.4.2.2
hydroxy(sulfanyl)boranyl = thioborono*	(HO)(HS)B–	P-68.1.4.2
(hydroxysulfanyl)carbonoselenoyl* = (<i>OS</i> -thiohydroperoxy)carbonoselenoyl	(HOS)-C(Se)–	P-65.2.1.7
(hydroxysulfanyl)carbonyl* = (<i>OS</i> -thiohydroperoxy)carbonyl	(HOS)-CO–	P-65.1.5.3; P-65.2.1.7
hydroxy(sulfanylidene)acetyl* = (hydroxycarbonothioyl)carbonyl = hydroxy(thiocarbonyl)carbonyl (not 2-thiooxalo; not 2-hydroxy-2-thiooxalyl)	HO-CS-CO–	P-65.1.7.2.4
hydroxy(sulfanylidene)ethanethioyl* = hydroxybis(sulfanylidene)ethyl (not 1,2-dithiooxalo)	HO-CS-CS–	P-65.1.7.2.4
(hydroxysulfanyl)phosphorothioyl* = (<i>OS</i> -thiohydroperoxy)phosphorothioyl	(HOS)-P(S)<	P-67.1.4.1.1.5
hydroxy(sulfanyl)phosphoryl* [not hydroxy(mercapto)phosphoryl]	(HO)(HS)P(O)–	P-67.1.4.1.1.5
hydroxysulfonothioyl*	HO-S(O)(S)–	P-65.3.2.3
hydroxytellanyl* = <i>OTe</i> -tellurohydroperoxy (not tellureno)	HO-Te–	P-63.4.2.2
hydroxythio: see hydroxysulfanyl*		
hydroxy(thiocarbonyl)carbonyl = hydroxy(sulfanylidene)acetyl* = (hydroxycarbonothioyl)carbonyl (not 2-thiooxalo; not 2-hydroxy-2-thiooxalyl)	HO-CS-CO–	P-65.1.7.2.4
2-hydroxy-2-thiooxalyl: see hydroxy(sulfanylidene)acetyl		
imidoarsoryl = arsorimidoyl*	As(=NH)<	P-67.1.4.1.1.4
imidophosphinoyl = phosphinimidoyl* = dihydrophosphorimidoyl	H ₂ P(=NH)–	P-67.1.4.1.1.4; P-67.1.4.1.2
imidostibinoyl = stibinimidoyl* = dihydrostiborimidoyl	H ₂ Sb(=NH)–	P-67.1.4.1.1.2; P-67.1.4.1.2
imino* = azanylidene (see also azanediyyl)	HN=	P-35.2.1; P-62.3.1.2
1-iminobutyl = butanimidoyl* = butyrimidoyl	CH ₃ -CH ₂ -CH ₂ -C(=NH)–	P-65.1.7.3.2; P-65.1.7.4.1

1-iminoethyl = ethanimidoyl* = acetimidoyl	$\text{CH}_3\text{-C(=NH)-}$	P-65.1.7.2.2
iminomethyl = methanimidoyl* = formimidoyl	HC(=NH)-	P-65.1.7.2.2
(iminomethyl)amino = methanimidamido* = formimidoylamino	HN=CH-NH-	P-66.4.1.3.3
iminomethylidene*	HN=C=	P-65.2.1.8
imino(phenyl)methyl = benzenecarboximidoyl* = benzimidoyl	$\text{C}_6\text{H}_5\text{-C(=NH)-}$	P-65.1.7.2.2
1-iminopropyl = propanimidoyl* = propionimidoyl	$\text{CH}_3\text{-CH}_2\text{-C(=NH)-}$	P-65.1.7.3.2; P-65.1.7.4.1
1-imino-2-selanylideneethane-1,2-diyl*	-C(=NH)-C(Se)-	P-65.1.7.5
[imino(sulfanyl)methyl]amino = (C-sulfanylcarbonimidoyl)amino*	HS-C(=NH)-NH-	P-66.1.6.1.3.3
indiganyl*	$\text{H}_2\text{In-}$	P-29.3.1; P-68.1.2
iodoso: see iodosyl*		
iodosyl* (not iodoso)	OI-	P-61.3.2.3
iodyl*	$\text{O}_2\text{I-}$	P-61.3.2.3
isobutoxy: see 2-methylpropoxy*		
isobutyl: see 2-methylpropyl*		
isocyanato*	OCN-	P-61.8
isocyano*	CN-	P-61.9
isofulminato: see (oxo- λ^5 -azanylidyne)methyl*		
isonicotinoyl = pyridine-4-carbonyl* = 4-pyridylcarbonyl = oxo(pyridine-4-yl)methyl		P-65.1.7.3.1; P-65.1.7.4.2
isopentyl: see 3-methylbutyl*		
isophthaloyl = benzene-1,3-dicarbonyl* = 1,3-phenylenedicarbonyl = 1,3-phenylenebis(oxomethylene)		P-65.1.7.3.1; P-65.1.7.4.2
isopropenyl = prop-1-en-2-yl* = 1-methylethen-1-yl	$\text{CH}_2=\text{C(CH}_3\text{)-}$	P-32.1.1; P-32.3
isopropoxy = (propan-2-yl)oxy* = 1-methylethoxy	$\text{(CH}_3\text{)}_2\text{CH-O-}$	P-63.2.2.2
isopropyl = propan-2-yl* = 1-methylethyl	$\text{(CH}_3\text{)}_2\text{CH-}$	P-29.3.2.2; P-29.4.1; P-29.6.2.2
isopropylidene = propan-2-ylidene* = 1-methylethylidene	$\text{(CH}_3\text{)}_2\text{C=}$	P-29.3.2.2; P-29.4.1; P-29.6.2.2
isoquinolin-7-yl* = 7-isoquinolyl (also 1-, 3-, 4-, 5-, 6- and 8-isomers)		P-29.3.4.1; P-57.1.5.3
7-isoquinolyl = isoquinolin-7-yl* (also 1-, 3-, 4-, 5-, 6- and 8-isomers)		
isoselenocyanato*	SeCN-	P-61.8
isotellurocyanato*	TeCN-	P-61.8
isothiocyanato*	SCN-	P-61.8
isothiocyanatosulfonothioyl* = sulfur(isothiocyanatido)thioyl	(SCN)-S(O)(S)-	P-67.1.4.4.1
isothiocyanatosulfonyl* = sulfur(isothiocyanatidoyl)	$\text{(SCN)-SO}_2\text{-}$	P-67.1.4.4.1

1-isoureido: see (*C*-hydroxycarbonimidoyl)amino*
 3-isoureido: see [amino(hydroxy)methylidene]amino*
 keto (not to be used): see oxo*

maleoyl = (2*Z*)-but-2-enediyl* = (2*Z*)-1,4-dioxobut-2-ene-1,4-diyl

malonimidoyl = propanediimidoyl* = 1,3-diiminopropane-1,3-diyl

malonyl = propanediyl* = 1,3-dioxopropane-1,3-diyl

mercapto: see sulfanyl*

mercaptocarbonyl: see sulfanylcarbonyl*

mercaptooxy: see sulfanyloxy*

methacryloyl = 2-methylprop-2-enoyl* = 2-methyl-1-oxoprop-2-en-1-yl

methanediyl: see methylene*

methanehydrazonamido* = methanehydrazonoylamino = (hydrazinylidenemethyl)amino

methanehydrazonohydrazido* = 2-(methanehydrazonoyl)hydrazin-1-yl
 = 2-(hydrazinylidenemethyl)hydrazin-1-yl

methanehydrazonoyl* = formohydrazonoyl = hydrazinylidenemethyl

methanehydrazonoylamino = methanehydrazonamido* = (hydrazinylidenemethyl)amino

2-(methanehydrazonoyl)hydrazin-1-yl = methanehydrazonohydrazido*
 = 2-(hydrazinylidenemethyl)hydrazin-1-yl

methaneseleninyl* = methylseleninyl

methaneselenonyl* = methylselenonyl

methaneselenoyl* = selenoformyl = selanylidenemethyl

methanesulfinamido* = (methanesulfinyl)amino

methanesulfinimidoyl* = *S*-methylsulfinimidoyl

methanesulfinyl* = methylsulfinyl

(methanesulfinyl)amino = methanesulfinamido*

methanesulfonamido* = (methanesulfonyl)amino

methanesulfonimidoyl* = *S*-methylsulfonimidoyl

methanesulfonyl* = methylsulfonyl

(methanesulfonyl)amino = methanesulfonamido*

(methanesulfonyl)azanylidene = (methanesulfonyl)imino* = (methylsulfonyl)imino

(methanesulfonyl)imino* = (methylsulfonyl)imino = (methanesulfonyl)azanylidene

methanetelluroyl* = telluroformyl = tellanylidenemethyl

methanetetrayl*

methanethioamido* = (methanethiyl)amino = thioformamido

methanethiyl* = thioformyl = sulfanylidenemethyl



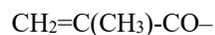
P-65.1.7.3.1; P-65.1.7.4.1



P-65.1.7.4.1



P-65.1.7.3.1; P-65.1.7.4.1



P-65.1.7.3.1; P-65.1.7.4.1



P-66.4.2.3.3



P-66.4.3.4.2



P-65.1.7.2.2



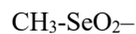
P-66.4.2.3.3



P-66.4.3.4.2



P-65.3.2.2.2



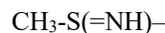
P-65.3.2.2.2



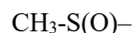
P-65.1.7.2.3; P-66.6.3



P-66.1.1.4.3



P-65.3.2.2.2



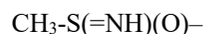
P-65.3.2.2.2



P-66.1.1.4.3



P-66.1.1.4.3



P-65.3.2.2.2



P-65.3.2.2.2



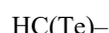
P-66.1.1.4.3



P-66.1.1.4.4



P-66.1.1.4.4



P-65.1.7.2.3; P-66.6.3



P-29.3.1



P-66.1.4.4



P-65.1.7.2.3; P-66.6.3

(methanethiyl)amino = methanethioamido* = thioformamido	HCS-NH-	P-66.1.4.4
methanetriyl*	-CH<	P-29.3.1
methanidyl*	H ₂ C-	P-72.6.3
methanimidamido* = (iminomethyl)amino = formimidoylamino	HC(=NH)-NH-	P-66.4.1.3.3
methanimidoyl* = formimidoyl = iminomethyl	HC(=NH)-	P-65.1.7.2.2
methanoyl = formyl* = oxomethyl	HCO-	P-65.1.7.2.1; P-66.6.1.3
methanyl = methyl*	CH ₃ -	P-29.3.1
methanylidene = methylidene*	CH ₂ =	P-29.3.1
methanylidyne = methylidyne*	CH≡	P-29.3.1
methanylylidene*	-CH=	P-29.3.1
methoxy* (not methyloxy)	CH ₃ -O-	P-63.2.2.2
2-methoxyanilino* = (2-methoxyphenyl)amino (also 3- and 4-methoxy isomers) (not 2-anisidino; not <i>o</i> -anisidino)	2-(CH ₃ -O)-C ₆ H ₄ -NH-	P-62.2.1.1.2
methoxyboranyl* (not hydromethoxyboryl)	CH ₃ -O-BH-	P-67.1.4.2
methoxyboranylidene*	CH ₃ -O-B=	P-67.1.4.1.1.6
C-methoxycarbonimidoyl*	CH ₃ -O-C(=NH)-	P-65.2.1.5
methoxycarbonothioyl* = methoxythiocarbonyl	CH ₃ -O-CS-	P-65.2.1.5
methoxycarbonyl* (not carbomethoxy)	CH ₃ -O-CO-	P-65.6.3.2.3
methoxy(isocyanato)phosphoryl*	(CH ₃ -O)(OCN)P(O)-	P-67.1.4.1.1.5
methoxy(oxo)-λ ⁵ -arsanylidene*	CH ₃ -O-As(O)=	P-67.1.4.1.1.6
(2-methoxyphenyl)amino = 2-methoxyanilino* (also <i>m</i> = 3- and <i>p</i> = 4-isomers)	2-(CH ₃ -O)-C ₆ H ₄ -NH-	P-62.2.1.1.2
methoxysulfanyl* (not methoxythio)	CH ₃ -O-S-	P-63.3.2
<i>S</i> -methoxysulfinimidoyl*	CH ₃ -O-S(=NH)-	P-65.3.2.3
(methoxysulfinyl)oxy*	CH ₃ -O-S(O)-O-	P-67.1.4.4.2
methoxysulfonyl* = methoxysulfuryl	CH ₃ -O-SO ₂ -	P-65.3.2.3; P-67.1.4.4.1
(methoxysulfonyl)amino*	CH ₃ -O-S(O) ₂ -NH-	P-67.1.4.4.2
methoxysulfuryl = methoxysulfonyl*	CH ₃ -O-SO ₂ -	P-65.3.2.3; P-67.1.4.4.1
methoxythio: see methoxysulfanyl*		
methoxythiocarbonyl = methoxycarbonothioyl*	CH ₃ -O-CS-	P-65.2.1.5
methyl* = methanyl	CH ₃ -	P-29.3.1
(methylamino)sulfinyl*	CH ₃ -NH-S(O)-	P-66.1.1.4.2
2-methylanilino* = (2-methylphenyl)amino (not <i>o</i> -toluidino; not 2-toluidino) (also 3- and 4-isomers)	2-CH ₃ -C ₆ H ₄ -NH-	P-62.2.1.1.2
(methylboranyl)amino*	CH ₃ -BH-NH-	P-68.1.4.2
2-methylbutan-2-yl* = 1,1-dimethylpropyl (not <i>tert</i> -pentyl)	CH ₃ -CH ₂ -C(CH ₃) ₂ -	P-29.4.1; P-29.6.3; P-57.1.4
1-methylbutyl = pentan-2-yl*	CH ₃ -CH ₂ -CH ₂ -CH(CH ₃)-	P-29.3.2.2; P-29.4.1
2-methylbutyl*	CH ₃ -CH ₂ -CH(CH ₃)-CH ₂ -	P-29.4.1

3-methylbutyl* (not isopentyl)	$(\text{CH}_3)_2\text{CH}-\text{CH}_2-\text{CH}_2-$	P-29.4.1; P-29.6.3
methyldioxy: see methylperoxy*		
methyldiselanyl* = methyldiseleno	$\text{CH}_3-\text{Se}-\text{Se}-$	P-63.3.1
methyldiseleno = methyldiselanyl*	$\text{CH}_3-\text{Se}-\text{Se}-$	P-63.3.1
methyldisulfanyl* = methyldithio	$\text{CH}_3-\text{S}-\text{S}-$	P-63.3.1
methylditellanyl* = methylditelluro	$\text{CH}_3-\text{Te}-\text{Te}-$	P-63.3.1
methylditelluro = methylditellanyl*	$\text{CH}_3-\text{Te}-\text{Te}-$	P-63.3.1
methyldithio = methyldisulfanyl*	$\text{CH}_3-\text{S}-\text{S}-$	P-63.3.1
methylene* (not methanediy)	$-\text{CH}_2-$	P-29.6.1
methylenebis(oxy)* (not methylenedioxy)	$-\text{O}-\text{CH}_2-\text{O}-$	P-63.2.2.1.3
methylenebis(sulfaneyl)* = methylenebis(thio)	$-\text{S}-\text{CH}_2-\text{S}-$	P-63.2.2.1.3
methylenebis(thio) = methylenebis(sulfaneyl)*	$-\text{S}-\text{CH}_2-\text{S}-$	P-63.2.2.1.3
methylenedioxy: see methylenebis(oxy)*		
1-methylethane-1,2-diyl = propane-1,2-diyl* (not propylene)	$-\text{CH}_2-\text{CH}(\text{CH}_3)-$	P-29.3.2.2
1-methylethen-1-yl = prop-1-en-2-yl* = isopropenyl	$\text{CH}_2=\text{C}(\text{CH}_3)-$	P-32.1.1; P-32.3
1-methylethoxy = (propan-2-yl)oxy* = isopropoxy	$(\text{CH}_3)_2\text{CH}-\text{O}-$	P-63.2.2.2
1-methylethyl = propan-2-yl* = isopropyl	$(\text{CH}_3)_2\text{CH}-$	P-29.3.2.2; P-29.4.1; P-29.6.2.2
1-methylethylidene = propan-2-ylidene* = isopropylidene	$(\text{CH}_3)_2\text{C}=\text{C}-$	P-29.3.2.2; P-29.4.1; P-29.6.2.2
methylidene* = methanylidene	$\text{CH}_2=\text{C}-$	P-29.3.1
$(\lambda^2\text{-methylideneamino})\text{oxy}^*$ (not fulminato)	$\text{C}=\text{N}-\text{O}-$	P-61.10
methylidyne* = methanylidyne	$\text{CH}\equiv\text{C}-$	P-29.3.1
<i>N</i> -methylmethanaminiumylidene* (not dimethylammoniumylidene; not dimethylimmonio)	$(\text{CH}_3)_2\text{N}^+=\text{C}-$	P-73.6
2-methyl-1-oxoprop-2-en-1-yl = 2-methylprop-2-enoyl* = methacryloyl	$\text{CH}_2=\text{C}(\text{CH}_3)-\text{CO}-$	P-65.1.7.3.1; P-65.1.7.4.1
methyloxy = methoxy*		
methylperoxy* (not methyldioxy)	$\text{CH}_3-\text{OO}-$	P-63.3.1
2-methylphenyl* = <i>o</i> -tolyl (also <i>m</i> - = 3- and <i>p</i> - = 4-isomers)	$2-\text{CH}_3-\text{C}_6\text{H}_4-$	P-29.6.2.3; P-57.1.5.3
(2-methylphenyl)amino = 2-methylanilino* (not <i>o</i> -toluidino; not 2-toluidino) (also 3- and 4-isomers)	$2-\text{CH}_3-\text{C}_6\text{H}_4-\text{NH}-$	P-62.2.1.1.2
methyl(phenyl)arsinoyl*	$(\text{C}_6\text{H}_5)(\text{CH}_3)\text{As}(\text{O})-$	P-67.1.4.1.1.3
methylphosphonoyl*	$\text{CH}_3-\text{P}(\text{O})<$	P-67.1.4.1.1.3
2-methylpropan-2-yl = <i>tert</i> -butyl* (unsubstituted) = 1,1-dimethylethyl	$(\text{CH}_3)_3\text{C}-$	P-29.4.1; P-29.6.1
2-methylpropan-2-ylum-1-yl*	$\text{CH}_3-\text{C}^+(\text{CH}_3)-\text{CH}_2-$	P-73.6
(2-methylpropan-2-yl)oxy = <i>tert</i> -butoxy* (unsubstituted) = 1,1-dimethylethoxy	$(\text{CH}_3)_3\text{C}-\text{O}-$	P-63.2.2.2
2-methylprop-2-enoyl* = methacryloyl = 2-methyl-1-oxoprop-2-en-1-yl	$\text{CH}_2=\text{C}(\text{CH}_3)-\text{CO}-$	P-65.1.7.3.1; P-65.1.7.4.1
1-methylprop-2-en-1-yl = but-3-en-2-yl*	$\text{CH}_2=\text{CH}-\text{CH}(\text{CH}_3)-$	P-32.1.1
1-methylpropoxy = (butan-2-yl)oxy* (not <i>sec</i> -butoxy; not <i>sec</i> -butyloxy)	$\text{CH}_3-\text{CH}_2-\text{CH}(\text{CH}_3)-\text{O}-$	P-63.2.2.2
2-methylpropoxy* (not isobutoxy)	$\text{CH}_3-\text{CH}(\text{CH}_3)-\text{CH}_2-\text{O}-$	P-63.2.2.2

1-methylpropyl = butan-2-yl* (not *sec*-butyl; not but-2-yl)
 2-methylpropyl* (not isobutyl)
 1-methylpropylidene = butan-2-ylidene* (not *sec*-butylidene)

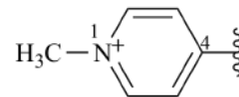
1-methylpyridin-1-ium-4-yl*

methylselanyl* = methylseleno
 methylseleninyl = methaneseleninyl*
 methylseleno = methylselanyl*
 methylselenonyl = methaneselenonyl*
 methylsulfaniumdiyl* = methylsulfoniumdiyl
 methylsulfanyl* = methylthio
 (methylsulfanyl)oxy* [not (methylthio)oxy]
 (methylsulfanyl)sulfonyl* = (methylthio)sulfonyl
S-methylsulfinimidoyl = methanesulfinimidoyl*
 methylsulfinyl = methanesulfinyl*
S-methylsulfonimidoyl = methanesulfonimidoyl*
 methylsulfoniumdiyl = methylsulfaniumdiyl*
 methylsulfonyl = methanesulfonyl*
 (methylsulfonyl)imino = (methanesulfonyl)imino* [not (methanesulfonyl)azanylidene]
 methyltellanyl* = methyltelluro
 methyltelluro = methyltellanyl*
 1-methyltetrasilan-1-yl*
 methylthio = methylsulfanyl*
 (methylthio)oxy: see (methylsulfanyl)oxy*
 (methylthio)sulfonyl = (methylsulfanyl)sulfonyl*
 methyltrisulfanyl* = methyltrithio
 methyltrithio = methyltrisulfanyl*
 morpholino: see morpholin-4-yl*

morpholin-4-yl* (not morpholino)

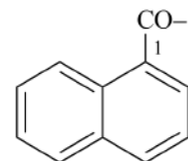
naphthalene-1-carbonyl* = 1-naphthoyl = 1-naphthylcarbonyl = naphthalen-1-yl(oxo)methyl
 (also 2-isomer)

CH₃-CH₂-CH(CH₃)–
 CH₃-CH(CH₃)-CH₂–
 CH₃-CH₂-C(CH₃)=



CH₃-Se–
 CH₃-Se(O)–
 CH₃-Se–
 CH₃-SeO₂–
 CH₃-S⁺<
 CH₃-S–
 CH₃-S-O–
 CH₃-S-SO₂–
 CH₃-S(=NH)–
 CH₃-S(O)–
 CH₃-S(=NH)(O)–
 CH₃-S⁺<
 CH₃-SO₂–
 CH₃-SO₂-N=

CH₃-S-SO₂–
 CH₃-S-S-S–
 CH₃-S-S-S–



P-29.3.2.2; P-29.4.1; P-29.6.3
 P-29.6.3; P-57.1.4
 P-29.3.2.2; P-29.4.1; P-29.6.3

P-73.6

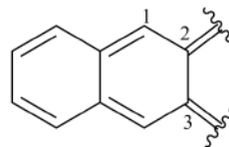
P-63.2.2.1.2; P-63.2.5
 P-65.3.2.2.2
 P-63.2.2.1.2; P-63.2.5
 P-65.3.2.2.2
 P-73.6
 P-63.2.2.1.2; P-63.2.5
 P-63.3.2
 P-65.3.2.3; P-65.6.3.2.3
 P-65.3.2.2.2
 P-65.3.2.2.2
 P-65.3.2.2.2
 P-73.6
 P-65.3.2.2.2
 P-66.1.1.4.4
 P-63.2.5
 P-63.2.5
 P-29.4.1
 P-63.2.2.1.2; P-63.2.5

P-65.3.2.3; P-65.6.3.2.3
 P-68.4.1.3
 P-68.4.1.3

P-29.3.3; P-29.6.2.3; P-64.7.1

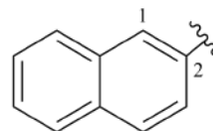
P-65.1.7.3.1; P-65.1.7.4.2

naphthalene-2,3-diylidene*



P-29.3.4.1

naphthalen-2-yl* = 2-naphthyl (also 1-isomer)



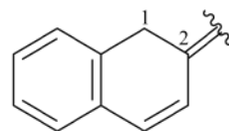
P-29.3.4.1; P-29.6.2.3; P-57.1.5.3

naphthalen-1-yl(oxo)methyl = naphthalene-1-carbonyl* = 1-naphthoyl = 1-naphthylcarbonyl (also 2-isomer)

1-C₁₀H₇-CO-

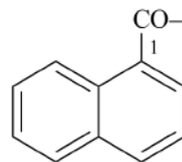
P-65.1.7.3.1; P-65.1.7.4.2

naphthalen-2(1*H*)-ylidene* [also 1(2*H*)-isomer]



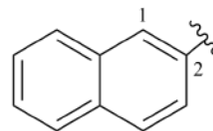
P-29.3.4.1

1-naphthoyl = naphthalene-1-carbonyl* = 1-naphthylcarbonyl = naphthalen-1-yl(oxo)methyl (also 2-isomer)



P-65.1.7.3.1; P-65.1.7.4.2

2-naphthyl = naphthalen-2-yl* (also 1-isomer)



P-29.3.4.1; P-29.6.2.3

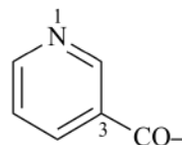
1-naphthylcarbonyl = naphthalene-1-carbonyl* = 1-naphthoyl = naphthalen-1-yl(oxo)methyl (also 2-isomer)

1-C₁₀H₇-CO-

P-65.1.7.3.1; P-65.1.7.4.2; P-57.1.5.3

neopentyl: see 2,2-dimethylpropyl*

nicotinoyl = pyridine-3-carbonyl* = 3-pyridylcarbonyl = oxo(pyridin-3-yl)methyl



P-65.1.7.3.1; P-65.6.3.2.3

nitramido* = nitroamino

O₂N-NH-

P-67.1.4.3.2

nitridophosphoryl = phosphoronitridoyl*

N≡P<

P-67.1.4.1.1.4

nitridostiboryl = stiboronitridoyl*

N≡Sb<

P-67.1.4.1.1.4

nitriolo* = azanetriyl (not azanylidene; not azanylylidene)

-N<

P-35.2.1; P-62.2.5.1

nitro*

O₂N-

P-61.5.1

aci-nitro = hydroxy(oxo)-λ⁵-azanylidene*

HO-N(O)=

P-61.5.3; P-67.1.4.1.1.6; P-67.1.6

nitroamino = nitramido*

O₂N-NH-

P-67.1.4.3.2

nitroazanediyyl*

O₂N-N<

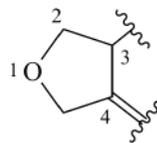
P-67.1.4.3.2

1-nitrohydrazin-1-yl*

H₂N-N(NO₂)-

P-67.1.4.3.3

2-nitrohydrazin-1-yl*	O ₂ N-NH-NH-	P-67.1.4.3.3
nitroimino*	O ₂ N-N=	P-67.1.4.3.2
nitrooxy*	O ₂ N-O-	P-67.1.4.3.1
nitroaryl* (not azoryl)	-N(O)<	P-67.1.4.1.1.2
nitroso*	O=N-	P-61.5.1
nitrosoamino*	ON-NH-	P-67.1.4.3.2
nitrosohydrazinylidene*	ON-NH-N=	P-67.1.4.3.3
nitrosooxy*	ON-O-	P-67.1.4.3.1
nitrososelanyl*	ON-Se-	P-67.1.4.3.1
nitrosulfanyl*	O ₂ N-S-	P-67.1.4.3.1
nonanoyl* = 1-oxononyl	CH ₃ -[CH ₂] ₇ -CO-	P-65.1.7.4.1
nonan-1-yl = nonyl*	CH ₃ -[CH ₂] ₇ -CH ₂ -	P-29.3.2.1; P-29.3.2.2
nonan-1-ylidene = nonylidene*	CH ₃ -[CH ₂] ₇ -CH=	P-29.3.2.1; P-29.3.2.2;
nonanylidyne = nonylidyne*	CH ₃ -[CH ₂] ₇ -C≡	P-29.3.2.1; P-29.3.2.2
nonyl* = nonan-1-yl	CH ₃ -[CH ₂] ₇ -CH ₂ -	P-29.3.2.1; P-29.3.2.2
nonylidene* = nonan-1-ylidene	CH ₃ -[CH ₂] ₇ -CH=	P-29.3.2.1; P-29.3.2.2
nonylidyne* = nonanylidyne	CH ₃ -[CH ₂] ₇ -C≡	P-29.3.2.1; P-29.3.2.2
octadecanoyl* = stearoyl = 1-oxooctadecyl	CH ₃ -[CH ₂] ₁₆ -CO-	P-65.1.7.3.1; P-65.1.7.4.1
octadecan-1-yl = octadecyl*	CH ₃ -[CH ₂] ₁₇ -	P-29.3.2.1; P-29.3.2.2
	$\begin{array}{c} 10 \quad 11-17 \quad 18 \\ \text{HC}-[\text{CH}_2]_7-\text{CH}_3 \\ \\ \text{HC}-[\text{CH}_2]_7-\text{CO}- \\ 9 \quad 8-2 \quad 1 \end{array}$	P-65.1.7.3.1; P-65.1.7.4.1
(9Z)-octadec-9-enoyl* = oleoyl = (9Z)-1-oxooctadec-9-en-1-yl		
octadecyl* = octadecan-1-yl	CH ₃ -[CH ₂] ₁₇ -	P-29.3.2.1; P-29.3.2.2
octanoyl* = 1-oxooctyl	CH ₃ -[CH ₂] ₆ -CO-	P-65.1.7.4.1
octan-1-yl = octyl*	CH ₃ -[CH ₂] ₆ -CH ₂ -	P-29.3.2.1; P-29.3.2.2
octan-1-ylidene = octylidene*	CH ₃ -[CH ₂] ₆ -CH=	P-29.3.2.1; P-29.3.2.2
octanylidyne = octylidyne*	CH ₃ -[CH ₂] ₆ -C≡	P-29.3.2.1; P-29.3.2.2
octyl* = octan-1-yl	CH ₃ -[CH ₂] ₆ -CH ₂ -	P-29.3.2.1; P-29.3.2.2
octylidene* = octan-1-ylidene	CH ₃ -[CH ₂] ₆ -CH=	P-29.3.2.1; P-29.3.2.2
octylidyne* = octanylidyne	CH ₃ -[CH ₂] ₆ -C≡	P-29.3.2.1; P-29.3.2.2
	$\begin{array}{c} 10 \quad 11-17 \quad 18 \\ \text{HC}-[\text{CH}_2]_7-\text{CH}_3 \\ \\ \text{HC}-[\text{CH}_2]_7-\text{CO}- \\ 9 \quad 8-2 \quad 1 \end{array}$	P-65.1.7.3.1; P-65.1.7.4.1
oleoyl = (9Z)-octadec-9-enoyl* = (9Z)-1-oxooctadec-9-en-1-yl		
oxalimidoyl = ethanediimidoyl* = diiminoethanediyl	-C(=NH)-C(=NH)-	P-65.1.7.2.2
oxalo* = carboxycarbonyl (not carboxyformyl; not hydroxy(oxo)acetyl)	HO-CO-CO-	P-65.1.2.2.3; P-65.1.7.2.1

oxaloamino* = (carboxycarbonyl)amino	HOOC-CO-NH-	P-65.1.7.2.4
2-oxaloethyl: see 3-carboxy-3-oxopropyl*		
oxaloxy* = (carboxycarbonyl)oxy [not (carboxyformyl)oxy]	HO-CO-CO-O-	P-65.1.7.2.4
oxalosulfanyl* = (carboxycarbonyl)sulfanyl = (carboxycarbonyl)thio [not (carboxyformyl)sulfanyl; not (carboxyformyl)thio]	HOOC-CO-S-	P-65.1.7.2.4
oxalyl* = ethanedioyl = dioxoethanediy	-CO-CO-	P-65.1.7.2.1
oxalylbis(azanediyl)* = ethanedioylbis(azanediyl)	-HN-CO-CO-NH-	P-66.1.1.4.5.2
oxalylbis(azanetriyl) = oxalylidinitrilo* = ethanedioyldinitrilo = ethanedioylbis(azanetriyl)	>N-CO-CO-N<	P-66.1.1.4.5.2
oxalylbis(azanylylidene)* = ethanedioylbis(azanylylidene)	=N-CO-CO-N=	P-66.1.1.4.5.2
oxalylidinitrilo* = oxalylbis(azanetriyl) = ethanedioyldinitrilo = ethanedioylbis(azanetriyl)	>N-CO-CO-N<	P-66.1.1.4.5.2
oxamoyl* = aminooxalyl = amino(oxo)acetyl (not carbamoylformyl; not carbamoylcarbonyl)	H ₂ N-CO-CO-	P-66.1.1.4.1.2
oxamoylamino* = amino(oxo)acetamido (not carbamoylformamido)	H ₂ N-CO-CO-NH-	P-66.1.1.4.5.1
oxamoylazanediyl*	H ₂ N-CO-CO-N<	P-66.1.1.4.5.2
oxamoylimino* = [amino(oxo)acetyl]imino	H ₂ N-CO-CO-N=	P-66.1.1.4.5.1
oxidanyl: see hydroxy*		
oxido*	-O-	P-72.6.2
oxo* (not keto)	O=	P-64.5.1
oxoacetyl*	OCH-CO-	P-65.1.6.3; P-65.1.7.2.4
oxoarsanyl* (not arsenoso)	O=As-	P-61.6
oxo-λ ⁵ -azanyl*	H ₂ N(O)-	P-62.5
(oxo-λ ⁵ -azanylidyne)methyl* (not isofulminato)	ON≡C-	P-61.10; P-66.5.4.2
1-oxobutyl = butanoyl* = butyryl	CH ₃ -CH ₂ -CH ₂ -CO-	P-65.1.7.3.1; P-65.1.7.4.1
1-oxodecyl = decanoyl*	CH ₃ -[CH ₂] ₈ -CO-	P-65.1.7.4.1
1-oxododecyl = dodecanoyl*	CH ₃ -[CH ₂] ₁₀ -CO-	P-65.1.7.4.1
1-oxoethyl = acetyl* = ethanoyl	CH ₃ -CO-	P-65.1.7.2.1
1-oxoheptyl = heptanoyl*	CH ₃ -[CH ₂] ₅ -CO-	P-65.1.7.4.1
1-oxohexadecyl = hexadecanoyl* = palmitoyl	CH ₃ -[CH ₂] ₁₄ -CO-	P-65.1.7.3.1; P-65.1.7.4.1
1-oxohexyl = hexanoyl*	CH ₃ -[CH ₂] ₄ -CO-	P-65.1.7.4.1
		P-29.3.3
oxolan-3-yl-4-ylidene*		
oxomethyl = formyl* = methanoyl	HCO-	P-65.1.7.2.1; P-66.6.1.3
oxomethylidene*	O=C=	P-65.2.1.8
1-oxononyl = nonanoyl*	CH ₃ -[CH ₂] ₇ -CO-	P-65.1.7.4.1

(9Z)-1-oxooctadec-9-en-1-yl = (9Z)-octadec-9-enoyl* = oleoyl

1-oxooctadecyl = octadecanoyl* = stearoyl

1-oxooctyl = octanoyl*

1-oxopentyl = pentanoyl*

oxo(phenyl)methyl = benzoyl* = benzenecarbonyl = phenylcarbonyl

oxophosphanyl* (not phosphoroso)

oxo- λ^5 -phosphanylidene*

oxo- λ^5 -phosphanylidyne*

2-oxopropanoyl* = 1,2-dioxopropyl (not pyruvoyl)

1-oxoprop-2-en-1-yl = prop-2-enoyl* = acryloyl

1-oxopropyl = propanoyl* = propionyl

2-oxopropyl* = acetonyl

2-oxopropylidene* (not acetonylidene)

2-oxopropylidyne* (not acetonylidyne)

oxo(pyridin-3-yl)methyl = nicotinyl = pyridine-3-carbonyl* = 3-pyridylcarbonyl

oxo(pyridin-4-yl)methyl = isonicotinyl = pyridine-4-carbonyl* = 4-pyridylcarbonyl

oxostibanyl*

1-oxo-4-sulfanylidenebutane-1,4-diyl*

1-oxotetradecyl = tetradecanoyl*

oxy*

oxylcarbonyl* = (ylooxidanyl)formyl

palmitoyl = hexadecanoyl* = 1-oxohexadecyl

pentanedioyl* = glutaryl = 1,5-dioxopentane-1,5-diyl

pentanoyl* = 1-oxopentyl

pentan-1-yl = pentyl*

pentan-2-yl* = 1-methylbutyl

pentan-1-ylidene = pentylidene*

pentan-3-ylidene* = 1-ethylpropylidene

pentanylidyne = pentylidyne*

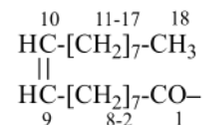
pent-2-enoyl*

pentyl* = pentan-1-yl

tert-pentyl: see 2-methylbutan-2-yl*

pentylidene* = pentan-1-ylidene

pentylidyne* = pentanylidyne



P-65.1.7.3.1; P-65.1.7.4.1



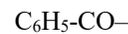
P-65.1.7.3.1; P-65.1.7.4.1



P-65.1.7.4.1



P-65.1.7.4.1



P-34.2.1.1; P-34.2.2; P-65.1.7.2.1



P-61.6; P-67.1.4.1.1.6



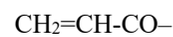
P-67.1.4.1.1.6



P-67.1.4.1.1.6



P-65.1.1.2.3; P-65.1.7.4.1



P-65.1.7.3.1; P-65.1.7.4.1



P-65.1.7.3.1; P-65.1.7.4.1



P-64.5.1



P-64.5



P-64.5



P-65.1.7.3.1; P-65.6.3.2.3



P-65.1.7.3.1; P-65.1.7.4.2



P-67.1.4.1.1.6



P-65.1.7.5



P-65.1.7.4.1



P-15.3.1.2.1.1; P-63.2.2.1.1



P-71.5



P-65.1.7.3.1; P-65.1.7.4.1



P-65.1.7.3.1; P-65.1.7.4.1



P-65.1.7.4.1



P-29.3.2.1; P-29.3.2.2



P-29.3.2.2; P-29.4.1



P-29.3.2.1; P-29.3.2.2



P-29.3.2.2; P-29.4



P-29.3.2.1; P-29.3.2.2



P-65.1.7.4.1



P-29.3.2.1; P-29.3.2.2



P-29.3.2.1; P-29.3.2.2



P-29.3.2.1; P-29.3.2.2

pentyloxy*
 perbromyl*
 perchloryl*
 perfluoryl*
 periodyl*
 peroxy* (not dioxy)
 peroxycarboxy: see carbonoperoxoyl*
 peroxyphosphoryl = phosphoroperoxoyl* = (hydroperoxy)phosphoryl

phenanthren-9-yl* = 9-phenanthryl (also 1-, 2-, 3-, and 4-isomers)

9-phenanthryl = phenanthren-9-yl* (also 1-, 2-, 3-, and 4-isomers)

phenethyl: see 2-phenylethyl

o-phenetidino: see 2-ethoxyanilino* (also *m* = 3 and *p* = 4 isomers)

phenoxy* (not phenyloxy)

phenyl*

phenylamino = anilino*

(phenylamino)sulfonyl = phenylsulfamoyl* = anilinosulfonyl

phenylazo = phenyldiazenyl*

phenylcarbonyl = benzoyl* = benzenecarbonyl = oxo(phenyl)methyl

(phenylcarbonyl)oxy = benzoyloxy*

phenyl(chlorophosphonyl) = phenylphosphonochloridoyl*

phenyldiazenyl* = phenylazo

1,2-phenylene* (not benzene-1,2-diyl) (also 1,3- and 1,4-isomers)

1,4-phenylenebis(iminomethylene) = benzene-1,4-dicarboximidoyl* = terephthalimidoyl
= 1,4-phenylenedicarboximidoyl

1,2-phenylenebis(oxomethylene) = benzene-1,2-dicarbonyl* = phthaloyl
= 1,2-phenylenedicarbonyl

1,3-phenylenebis(oxomethylene) = benzene-1,3-dicarbonyl* = isophthaloyl
= 1,3-phenylenedicarbonyl

1,4-phenylenebis(oxomethylene) = benzene-1,4-dicarbonyl* = terephthaloyl

CH₃-[CH₂]₃-CH₂-O-

O₃Br-

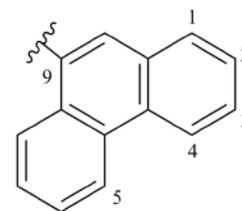
O₃Cl-

O₃F-

O₃I-

-OO-

(HOO)-P(O)<



9-C₁₄H₉-

C₆H₅-O-

C₆H₅-

C₆H₅-NH-

C₆H₅-NH-SO₂-

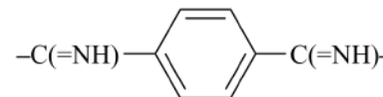
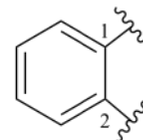
C₆H₅-N=N-

C₆H₅-CO-

C₆H₅-CO-O-

(C₆H₅)CIP(O)-

C₆H₅-N=N-



1,2-C₆H₄(CO-)₂

1,3-C₆H₄(CO-)₂

1,4-C₆H₄(CO-)₂

P-63.2.2.1.1

P-61.3.2.3

P-61.3.2.3

P-61.3.2.3

P-61.3.2.3

P-63.3.1

P-67.1.4.1.1.4

P-29.3.4.1; P-29.6.2.3; P-57.1.5.3

P-29.3.4.1; P-29.6.2.3; P-57.1.5.3

P-63.2.2.2

P-29.6.1

P-62.2.1.1.1

P-66.1.1.4.2

P-68.3.1.3.2.2

P-34.2.1.1; P-34.2.2; P-65.1.7.2.1

P-65.6.3.2.3

P-67.1.4.1.1.4

P-68.3.1.3.2.2

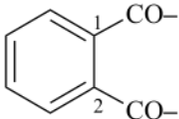
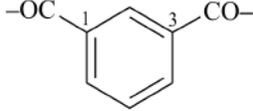
P-29.6.1

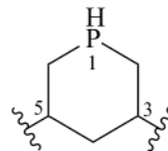
P-65.1.7.3.2

P-65.1.7.3.1; P-65.1.7.4.2

P-65.1.7.3.1; P-65.1.7.4.2

P-65.1.7.3.1; P-65.1.7.4.2

= 1,4-phenylenedicarbonyl		
1,2-phenylenebis(sulfanylidene)methylene) = benzene-1,2-dicarbothioyl*	$1,2-C_6H_4(CS-)_2$	P-65.1.7.3.1; P-65.1.7.4.3
= 1,2-phenylenebis(thioxomethylene) (not dithiophthaloyl)		
1,2-phenylenebis(thioxomethylene) = benzene-1,2-dicarbothioyl*	$1,2-C_6H_4(CS-)_2$	P-65.1.7.3.1; P-65.1.7.4.3
= 1,2-phenylenebis(sulfanylidene)methylene) (not dithiophthaloyl)		
1,4-phenylenedicarbonimidoyl = benzene-1,4-dicarboximidoyl* = terephthalimidoyl	$1,4-C_6H_4(C=NH)_2-$	P-65.1.7.3.2
= 1,4-phenylenebis(iminomethylene)		
1,2-phenylenedicarbonyl = benzene-1,2-dicarbonyl* = phthaloyl		P-65.1.7.3.1; P-65.1.7.4.2
= 1,2-phenylenebis(oxomethylene)		
1,3-phenylenedicarbonyl = benzene-1,3-dicarbonyl* = isophthaloyl		P-65.1.7.3.1; P-65.1.7.4.2
= 1,3-phenylenebis(oxomethylene)		
1,4-phenylenedicarbonyl = benzene-1,4-dicarbonyl* = terephthaloyl		P-65.1.7.3.1; P-65.1.7.4.2
= 1,4-phenylenebis(oxomethylene)		
2-phenylethenyl* = 2-phenylvinyl = styryl	$C_6H_5-CH=CH-$	P-32.3
2-phenylethyl* (not phenethyl)	$C_6H_5-CH_2-CH_2-$	P-29.6.3
phenylmethoxy = benzyloxy*	$C_6H_5-CH_2-O-$	P-63.2.2.1.1
phenylmethyl = benzyl*	$C_6H_5-CH_2-$	P-29.6.1; P-29.6.2.1
phenylmethylidene = benzylidene* (not benzal)	$C_6H_5-CH=$	P-29.6.1; P-29.6.2.1
phenylmethylidyne = benzylidyne*	$C_6H_5-C\equiv$	P-29.6.1; P-29.6.2.1
phenyloxy: see phenoxy*		
4-phenylphenyl: see [1,1'-biphenyl]-4-yl		
phenylphosphonochloridoyl* = phenyl(chlorophosphonoyl)	$(C_6H_5)CIP(O)-$	P-67.1.4.1.1.4
3-phenylprop-2-enoyl* = cinnamoyl	$C_6H_5-CH=CH-CO-$	P-65.1.7.3.1
phenylselanyl* = phenylseleno	C_6H_5-Se-	P-63.2.2.1.2; P-63.2.5
(phenylselanyl)oxy*	$C_6H_5-Se-O-$	P-63.3.2
phenylseleno = phenylselanyl*	C_6H_5-Se-	P-63.2.2.1.2; P-63.2.5
phenylselenonyl = benzeneselenonyl*	$C_6H_5-SeO_2-$	P-65.3.2.2.2
phenylsulfamoyl* = (phenylamino)sulfonyl = anilinosulfonyl	$C_6H_5-NH-SO_2-$	P-66.1.1.4.2
phenylsulfanyl* = phenylthio	C_6H_5-S-	P-63.2.2.1.2; P-63.2.5
phenyl(sulfanylidene)methyl = benzenecarbothioyl* = thiobenzoyl = phenyl(thio)oxomethyl	C_6H_5-CS-	P-65.1.7.2.3
(phenylsulfanyl)oxy*	C_6H_5-S-O-	P-63.3.2
phenylsulfinoselenoyl = benzenesulfinoselenoyl*	$C_6H_5-S(Se)-$	P-65.3.2.2.2
phenylsulfinyl = benzenesulfinyl*	$C_6H_5-S(O)-$	P-63.6; P-65.3.2.2.2
(phenylsulfinyl)amino = benzenesulfinamido* = (benzenesulfinyl)amino	$C_6H_5-S(O)-NH-$	P-66.1.1.4.3

phenylsulfonyl = benzenesulfonyl*	$C_6H_5-SO_2-$	P-63.6; P-65.3.2.2.2
(phenylsulfonyl)amino = benzenesulfonamido* = (benzenesulfonyl)amino	$C_6H_5-SO_2-NH-$	P-66.1.1.4.3
phenyltellanyl* = phenyltelluro	C_6H_5-Te-	P-63.2.2.1.2; P-63.2.5
(phenyltellanyl)oxy*	$C_6H_5-Te-O-$	P-63.3.2
phenyltelluro = phenyltellanyl*	C_6H_5-Te-	P-63.2.2.1.2; P-63.2.5
phenylthio = phenylsulfanyl*	C_6H_5-S-	P-63.2.2.1.2; P-63.2.5
phenyl(thio)oxymethyl = benzenecarbothioyl* = thiobenzoyl = phenyl(sulfanylidene)methyl	C_6H_5-CS-	P-65.1.7.2.3
2-phenylvinyl = 2-phenylethenyl* = styryl	$C_6H_5-CH=CH-$	P-32.3
phosphanediyl* (not phosphinediyl)	HP<	P-68.3.2.3.2.2
phosphanetriyl* (not phosphinetriyl)	-P<	P-68.3.2.3.2.2
phosphaniumyl* = phosphonio = phosphoniumyl	H_3P^+-	P-73.6; P-74.1.3
phosphanyl* = phosphino	H_2P-	P-29.3.1; P-68.3.2.3.2.2
λ^5 -phosphanyl* = phosphoranyl	H_4P-	P-68.3.2.3.2.2
phosphanylidene*	HP=	P-29.3.1; P-68.3.2.3.2.2
phosphanylylidene*	-P=	P-68.3.2.3.2.2
phosphinane-3,5-diyl*		P-29.3.3
phosphinediyl: see phosphanediyl*		
phosphinetriyl: see phosphanetriyl*		
phosphinimidoyl* = imidophosphinoyl = dihydrophosphorimidoyl	$H_2P(=NH)-$	P-67.1.4.1.1.4; P-67.1.4.1.2
phosphino = phosphanyl*	H_2P-	P-29.3.1; P-68.3.2.3.2.2
phosphinothioyl* = thiophosphinoyl = dihydrophosphorothioyl	$H_2P(S)-$	P-67.1.4.1.1.4; P-67.1.4.1.2
phosphinoyl* = dihydrophosphoryl (not phosphinyl)	$H_2P(O)-$	P-67.1.4.1.1.2; P-67.1.4.1.2
phosphinyl: see phosphinoyl*		
phospho: see dioxo- λ^5 -phosphanyl*		
phosphonato*	$(O^-)_2P(O)-$	P-72.6.1
phosphonio = phosphaniumyl* = phosphoniumyl	H_3P^+-	P-73.6; P-74.1.3
phosphoniumyl = phosphaniumyl* = phosphonio	H_3P^+-	P-73.6; P-74.1.3
phosphono*	$(HO)_2P(O)-$	P-67.1.4.1.1.1
phosphonooxy*	$(HO)_2P(O)-O-$	P-67.1.4.1.3
phosphonothioyl* = hydro(thiophosphoryl)	HP(S)<	P-67.1.4.1.2
phosphonoyl* = hydrophosphoryl	HP(O)<	P-67.1.4.1.1.2; P-67.1.4.1.2
phosphoramidochloridoyl* = amidochlorophosphoryl (not chloroamidophosphoryl)	$(H_2N)CIP(O)-$	P-67.1.4.1.1.4; P-67.1.5.2
phosphoranyl = λ^5 -phosphanyl*	H_4P-	P-68.3.2.3.2.2
phosphorocyanidoisocyanatidothioyl* = cyano(isocyanato)phosphorothioyl	$(OCN)(NC)P(S)-$	P-67.1.4.1.1.4

plumbylidyne* = plumbanylidyne	HPb≡	P-29.3.1; P-68.2.2
propanamido* = propanoylamino = propionamido = propionylamino	CH ₃ -CH ₂ -CO-NH-	P-66.1.1.4.3
propanediimidoyl* = malonimidoyl = 1,3-diiminopropane-1,3-diyl	-C(=HN)-CH ₂ -C(=NH)-	P-65.1.7.4.1
propanedioyl* = malonyl = 1,3-dioxopropane-1,3-diyl	-CO-CH ₂ -CO-	P-65.1.7.3.1; P-65.1.7.4.1
propane-1,3-diyl* (not trimethylene)	-CH ₂ -CH ₂ -CH ₂ -	P-29.3.2.2
propane-1,2-diyl* = 1-methylethane-1,2-diyl (not propylene)	-CH ₂ -CH(CH ₃)-	P-29.3.2.2
propane-1,1,1-triyl*	$\begin{array}{c} \\ \text{CH}_3\text{-CH}_2\text{-C-} \\ \end{array}$	P-29.3.2.2
propanethioyl* = thiopropionyl = 1-sulfanylidenepropyl = 1-thioxopropyl	CH ₃ -CH ₂ -CS-	P-65.1.7.4.1
propanimidoyl* = propionimidoyl = 1-iminopropyl	CH ₃ -CH ₂ -C(=NH)-	P-65.1.7.3.2; P-65.1.7.4.1
propanoyl* = propionyl = 1-oxopropyl	CH ₃ -CH ₂ -CO-	P-65.1.7.3.1; P-65.1.7.4.1
propanoylamino = propanamido* = propionamido = propionylamino	CH ₃ -CH ₂ -CO-NH-	P-66.1.1.4.3
propanoyloxy* = propionyloxy	CH ₃ -CH ₂ -CO-O-	P-65.6.3.2.3
propan-1-yl = propyl*	CH ₃ -CH ₂ -CH ₂ -	P-29.3.2.1; P-29.3.2.2
propan-2-yl* = isopropyl = 1-methylethyl	(CH ₃) ₂ CH-	P-29.3.2.2; P-29.4.1; P-29.6.2.2
propan-1-ylidene = propylidene*	CH ₃ -CH ₂ -CH=	P-29.3.2.1; P-29.3.2.2
propan-2-ylidene* = 1-methylethylidene = isopropylidene	(CH ₃) ₂ C=	P-29.3.2.2; P-29.4.1; P-29.6.2.2
propanylidyne = propylidyne*	CH ₃ -CH ₂ -C≡	P-29.3.2.1; P-29.3.2.2
(propan-2-yl)oxy* = isopropoxy = 1-methylethoxy	(CH ₃) ₂ CH-O-	P-63.2.2.2
propan-1-yl-1-ylidene*	$\begin{array}{c} \\ \text{CH}_3\text{-CH}_2\text{-C=} \end{array}$	P-29.3.2.2
prop-2-enehydrazonoyl* = acrylohydrazonoyl = 1-hydrazinylideneprop-2-en-1-yl	CH ₂ =CH-C(=NNH ₂)-	P-65.1.7.3.2
prop-2-eneselenoyl* = selenoacryloyl = 1-selanylideneprop-2-en-1-yl	CH ₂ =CH-C(Se)-	P-65.1.7.3.3
prop-2-enoyl* = acryloyl = 1-oxoprop-2-en-1-yl	CH ₂ =CH-CO-	P-65.1.7.3.1; P-65.1.7.4.1
prop-1-en-1-yl*	CH ₃ -CH=CH-	P-32.1.1
prop-1-en-2-yl* = 1-methylethen-1-yl = isopropenyl	CH ₂ =C(CH ₃)-	P-32.1.1; P-32.3
prop-2-en-1-yl* = allyl	CH ₂ =CH-CH ₂ -	P-32.1.1; P-32.3
prop-2-en-1-ylidene* = allylidene	CH ₂ =CH-CH=	P-32.1.1; P-32.3
prop-2-en-1-ylidyne* = allylidyne	CH ₂ =CH-C≡	P-32.1.1; P-32.3
propionamido = propanamido* = propionylamino = propanoylamino	CH ₃ -CH ₂ -CO-NH-	P-66.1.1.4.3
propionimidoyl = propanimidoyl* = 1-iminopropyl	CH ₃ -CH ₂ -C(=NH)-	P-65.1.7.3.2; P-65.1.7.4.1
propionyl = propanoyl* = 1-oxopropyl	CH ₃ -CH ₂ -CO-	P-65.1.7.3.1; P-65.1.7.4.1
propionylamino = propanamido* = propanoylamino = propionamido	CH ₃ -CH ₂ -CO-NH-	P-66.1.1.4.3
propionyloxy = propanoyloxy*	CH ₃ -CH ₂ -CO-O-	P-65.6.3.2.3
propoxy* (not propyloxy)	CH ₃ -CH ₂ -CH ₂ -O-	P-63.2.2.2

propyl* = propan-1-yl

propylene: see propane-1,2-diyl*

propylidene* = propan-1-ylidene

propylidyne* = propanylidyne

propyloxy: see propoxy*

pyridine-3-carbonyl* = nicotinoyl = 3-pyridylcarbonyl = oxo(pyridin-3-yl)methyl

pyridine-4-carbonyl* = 4-pyridylcarbonyl = isonicotinoyl = oxo(pyridin-4-yl)methyl

pyridinio = pyridin-1-ium-1-yl*

pyridin-1-ium-1-yl* = pyridinio

pyridin-1(4*H*)-yl* (also 1(2*H*)-isomer)

pyridin-2-yl* = 2-pyridyl (also 3- and 4- isomers)

2-pyridyl = pyridin-2-yl* (also 3- and 4- isomers)

3-pyridylcarbonyl = pyridine-3-carbonyl* = nicotinoyl = oxo(pyridin-3-yl)methyl

4-pyridylcarbonyl = pyridine-4-carbonyl * = isonicotinoyl = oxo(pyridin-4-yl)methyl

pyruvoyl: see 2-oxopropanoyl*

quinolin-2-yl* = 2-quinolyl (and 3-, 4-, 5-, 6-, 7-, and 8-isomers)

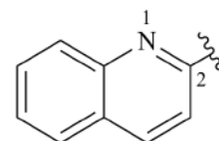
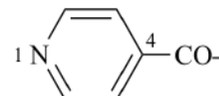
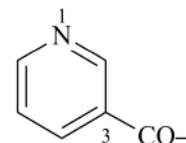
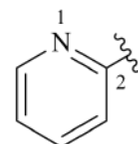
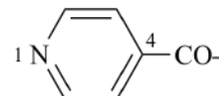
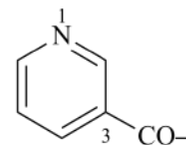
2-quinolyl = quinolin-2-yl*

selenediyl* (not seleno)

seleniumyl* = selenonio = selenoniumyl

selano* = episeleno (ring forming)

selanyl* (not hydroseleno)



P-29.3.2.1; P-29.3.2.2

P-29.3.2.1; P-29.3.2.2

P-29.3.2.1; P-29.3.2.2

P-65.1.7.3.1; P-65.6.3.2.3

P-65.1.7.3.1; P-65.1.7.4.2

P-73.6

P-29.3.4.1; P-29.6.2.3

P-29.3.4.1; P-29.6.2.3; P-57.1.5.3

P-65.1.7.3.1; P-65.6.3.2.3

P-65.1.7.3.1; P-65.1.7.4.2

P-29.6.2.3; P-57.1.5.3

P-63.2.5

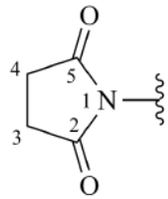
P-73.6

P-25.4.2.1.4; P-63.5

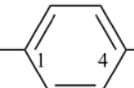
P-63.1.5

selanylidene* = selenoxo	Se=	P-29.3.1; P-64.6.1
1-selanylideneethyl = ethaneselenoyl* = selenoacetyl	CH ₃ -C(Se)-	P-65.1.7.2.3
selanylidenemethyl = methaneselenoyl* = selenoformyl	HC(Se)-	P-65.1.7.2.3; P-66.6.3
1-selanylideneprop-2-en-1-yl = prop-2-eneselenoyl* = selenoacryloyl	CH ₂ =CH-C(Se)-	P-65.1.7.3.3
selanylphosphonoyl*	HP(O)(SeH)-	P-67.1.4.1.1.5
seleneno: see hydroxyselanyl*		
selenino* (unmodified)	HO-Se(O)-	P-65.3.0; P-65.3.2.1
seleninyl*	O=Se<	P-65.3.2.3
seleno: see selanediyl*		
selenoacetyl = ethaneselenoyl* = 1-selanylideneethyl	CH ₃ -C(Se)-	P-65.1.7.2.3
selenoacryloyl = prop-2-eneselenoyl* = 1-selanylideneprop-2-en-1-yl	CH ₂ =CH-C(Se)-	P-65.1.7.3.3
selenocyanato* = carbononitridoyl selanyl	NC-Se-	P-65.2.2
selenoformyl = methaneselenoyl* = selanylidenemethyl	HC(Se)-	P-65.1.7.2.3; P-66.6.3
<i>OSe</i> -selenohydroperoxy = hydroxyselanyl* (not seleneno)	HO-Se-	P-63.4.2.2
(<i>OSe</i> -selenohydroperoxy)methyl = (hydroxyselanyl)methyl*	(HO-Se)-CH ₂ -	P-63.4.2.2
selenonimidothioyl*	Se(=NH)(=S)<	P-65.3.2.3
selenonio = selaniumyl* = selenoniumyl	H ₂ Se ⁺ -	P-73.6
selenoniumyl = selaniumyl* = selenonio	H ₂ Se ⁺ -	P-73.6
selenono* (unmodified)	HO-SeO ₂ -	P-65.3.0; P-65.3.2.1
selenonohydrazonoyl*	Se(O)(=N-NH ₂)<	P-65.3.2.3
selenonyl*	-SeO ₂ -	P-65.3.2.3
selenoxo = selanylidene*	Se=	P-29.3.1; P-64.6.1
semicarbazido = 2-carbamoylhydrazin-1-yl* = 2-(aminocarbonyl)hydrazine-1-yl	H ₂ N-CO-NH-NH-	P-68.3.1.2.4
semicarbazono = carbamoylhydrazinylidene*	H ₂ N-CO-NH-N=	P-68.3.1.2.5
silanediyl* (not silylene)	H ₂ Si<	P-29.3.1; P-68.2.2
silanediyl di(ethane-2,1-diyl)* = silanediyl diethylene	-CH ₂ -CH ₂ -SiH ₂ -CH ₂ -CH ₂ -	P-29.4.2
silanediyl diethylene = silanediyl di(ethane-2,1-diyl)*	-CH ₂ -CH ₂ -SiH ₂ -CH ₂ -CH ₂ -	P-29.4.2
silanediylidene*	=Si=	P-68.2.2
silanetetrayl*	>Si<	P-68.2.2
silanetriyl*	-SiH<	P-68.2.2
silanyl = silyl*	H ₃ Si-	P-29.3.1; P-68.2.2
silanylidene = silylidene*	H ₂ Si=	P-29.3.1; P-68.2.2
silanylidyne = silylidyne*	HSi≡	P-29.3.1; P-68.2.2
silanylylidene*	-SiH=	P-68.2.2
siloxy: see silyloxy		
silyl* = silanyl	H ₃ Si-	P-29.3.1; P-68.2.2

(silylamino)silyl* (not disilazan-1-yl)	HSi-NH-SiH ₂ -	P-29.3.2.2
silylene: see silanediyl*		
silylidene* = silanylidene	H ₂ Si=	P-29.3.1; P-68.2.2
silylidyne* = silanylidyne	HSi≡	P-29.3.1; P-68.2.2
silyloxy* (not siloxy)	H ₃ Si-O-	P-63.2.2.1.1
3-silyltetrasilan-1-yl*	⁴ SiH ₃ - ³ SiH(SiH ₃)- ² SiH ₂ - ¹ SiH ₂ -	P-29.4.1
stannanediyl* (not stannylene)	H ₂ Sn<	P-68.2.2
stannanediylidene*	=Sn=	P-68.2.2
stannanetetrayl*	>Sn<	P-68.2.2
stannanetriyl*	-SnH<	P-68.2.2
stannanyl = stannyl*	H ₃ Sn-	P-29.3.1; P-68.2.2
stannanylidene = stannylidene*	H ₂ Sn=	P-29.3.1; P-68.2.2
stannanylidyne = stannylidyne*	HSn≡	P-29.3.1; P-68.2.2
stannanylylidene*	-SnH=	P-68.2.2
stannyl* = stannanyl	H ₃ Sn-	P-29.3.1; P-68.2.2
stannylene: see stannanediyl		
stannylidene* = stannanylidene	H ₂ Sn=	P-29.3.1; P-68.2.2
stannylidyne* = stannanylidyne	HSn≡	P-29.3.1; P-68.2.2
stearoyl = octadecanoyl* = 1-oxooctadecyl	CH ₃ -[CH ₂] ₁₆ -CO-	P-65.1.7.3.1; P-65.1.7.4.1
stibanediyl* (not stibinediyl)	HSb<	P-56.4; P-68.3.2.3.2.2
stibanetriyl* (not stibinetriyl)	-Sb<	P-68.3.2.3.2.2
stibaniumyl* = stibonio = stiboniumyl	H ₃ Sb ⁺ -	P-73.6
stibanyl* = stibino	H ₂ Sb-	P-29.3.1; P-68.3.2.3.2.2
λ ⁵ -stibanyl* = stiboranyl	H ₄ Sb-	P-68.3.2.3.2.2
stibanylidene* (not stibinylidene)	HSb=	P-29.3.1; P-56.4; P-68.3.2.3.2.2
stibanylylidene*	-Sb=	P-68.3.2.3.2.2
stibinediyl: see stibanediyl*		
stibinetriyl: see stibanetriyl*		
stibinimidoyl* = imidostibinoyl = dihydrostiborimidoyl	H ₂ Sb(=NH)-	P-67.1.4.1.1.2; P-67.1.4.1.2
stibino = stibanyl*	H ₂ Sb-	P-29.3.1; P-68.3.2.3.2.2
stibinothioyl* = dihydrostiborothioyl	H ₂ Sb(S)-	P-67.1.4.1.1.2; P-67.1.4.1.2
stibinoyl* = dihydrostiboryl	H ₂ Sb(O)-	P-67.1.4.1.1.2; P-67.1.4.1.2
stibinylidene: see stibanylidene*		
stibonato*	(⁻ O) ₂ Sb(O)-	P-72.6.1
stibonio = stibaniumyl* = stiboniumyl	H ₃ Sb ⁺ -	P-73.6
stiboniumyl = stibaniumyl* = stibonio	H ₃ Sb ⁺ -	P-73.6

stibono*	$(\text{HO})_2\text{Sb}(\text{O})-$	P-67.1.4.1.1.1
stibonoyl* = hydrostiboryl	$\text{HSb}(\text{O})<$	P-67.1.4.1.1.2; P-67.1.4.1.2
stiboranyl = λ^5 -stibanyl*	$\text{H}_4\text{Sb}-$	P-68.3.2.3.2.2
stiborodiamidothioyl*	$(\text{H}_2\text{N})_2\text{Sb}(\text{S})-$	P-67.1.4.1.1.4
stiborohydrazonoyl* = hydrazonostiboryl	$\text{Sb}(=\text{NNH}_2)<$	P-67.1.4.1.1.4
stiboronitridoyl* = nitridostiboryl	$\text{N}\equiv\text{Sb}<$	P-67.1.4.1.1.4
stiboryl* (not antimonyl)	$-\text{Sb}(\text{O})<$	P-67.1.4.1.1.2
styryl = 2-phenylethenyl* = 2-phenylvinyl	$\text{C}_6\text{H}_5-\text{CH}=\text{CH}-$	P-32.3
		
succinimido = 2,5-dioxopyrrolidin-1-yl*		P-66.2.2
succinimidoyl = butanediimidoyl* = 1,4-diiminobutane-1,4-diyl	$-\text{C}(=\text{NH})-\text{CH}_2-\text{CH}_2-\text{C}(=\text{NH})-$	P-65.1.7.3.2
succinyl = butanedioyl* = 1,4-dioxobutane-1,4-diyl	$-\text{CO}-\text{CH}_2-\text{CH}_2-\text{CO}-$	P-65.1.7.3.1; P-65.1.7.4.1
sulfamoyl* = aminosulfonyl = sulfuramidoyl	$\text{H}_2\text{N}-\text{SO}_2-$	P-65.3.2.3; P-66.1.1.4.2
sulfamoyloxy* = sulfuramidoyloxy	$\text{H}_2\text{N}-\text{SO}_2-\text{O}-$	P-67.1.4.4.2
sulfanediyl* (not thio; not sulfenyl)	$-\text{S}-$	P-63.2.5
sulfanediylbis(methylene)* (not sulfanediyl dimethylene; not thiodimethylene)	$-\text{CH}_2-\text{S}-\text{CH}_2-$	P-63.2.2.1.3
sulfanediyl dimethylene: see sulfanediylbis(methylene)*		
sulfaniumyl* = sulfoniumyl = sulfonio	H_2S^+-	P-73.6
sulfano* = epithio (ring forming)	$-\text{S}-$	P-25.4.2.1.4; P-63.5
sulfanyl* (not mercapto)	$\text{HS}-$	P-29.3.1; P-63.1.5
sulfanylbis(sulfanylidene)ethyl = sulfanyl(sulfanylidene)ethanethioyl* = trithiooxalo	$\text{HS}-\text{CS}-\text{CS}-$	P-65.1.7.2.4; P-65.1.7.3.3
sulfanylboranyl*	$\text{HS}-\text{BH}-$	P-67.1.4.2
(C-sulfanylcarbonimidoyl)amino* = [imino(sulfanyl)methyl]amino	$\text{HS}-\text{C}(=\text{NH})-\text{NH}-$	P-66.1.6.1.3.3
sulfanylcarbonothioyl = dithiocarboxy*	$\text{HS}-\text{CS}-$	P-65.2.1.6
[(sulfanylcarbonothioyl)sulfanyl]carbonothioyl = [(dithiocarboxy)sulfanyl]carbonothioyl* = [sulfanyl(thiocarbonyl)sulfanyl](thiocarbonyl) {not [(dithiocarboxy)sulfanyl]thioformyl}	$\text{HS}-\text{CS}-\text{S}-\text{CS}-$	P-65.2.3.1.5
sulfanylcarbonyl* (not mercaptocarbonyl)	$\text{HS}-\text{CO}-$	P-65.2.1.6
(sulfanylcarbonyl)oxy*	$\text{HS}-\text{CO}-\text{O}-$	P-65.2.1.6
sulfanylidene* = thioxo	$\text{S}=\text{S}=\text{S}$	P-29.3.1; P-64.6.1
sulfanylideneamino* = thionitroso = thioxoamino	$\text{S}=\text{N}-$	P-67.1.4.3.2
(sulfanylideneamino)sulfanyl* = (thionitroso)sulfanyl = (thioxoamino)sulfanyl	$\text{S}=\text{N}-\text{S}-$	P-67.1.4.3.2
1-sulfanylidenebutyl = butanethioyl* = thiobutyryl = 1-thioxobutyl	$\text{CH}_3-\text{CH}_2-\text{CH}_2-\text{CS}-$	P-65.1.7.4.1
1-sulfanylideneethyl = ethanethioyl* = thioacetyl	$\text{CH}_3-\text{CS}-$	P-65.1.7.2.3

sulfanylidene-methyl = methanethiyl* = thioformyl	HCS–	P-65.1.7.2.3; P-66.6.3
sulfanylidene-methylidene* = thioxomethylidene	S=C=	P-65.2.1.8
1-sulfanylidene-propyl = propanethiyl* = thiopropionyl = 1-thioxopropyl	CH ₃ -CH ₂ -CS–	P-65.1.7.4.1
sulfanyloxy* = SO-thiohydroperoxy (not mercaptooxy)	HS-O–	P-63.4.2.2
[(sulfanyloxy)carbonyl]oxy* = [(SO-thiohydroperoxy)carbonyl]oxy	(HSO)-CO-O–	P-65.2.1.7
(sulfanyloxy)phosphoryl* = (SO-thiohydroperoxy)phosphoryl	(HSO)-P(O)<	P-67.1.4.1.1.5
sulfanylphosphonothioyl* = sulfanyl(thiophosphonoyl)	HS-HP(S)–	P-67.1.4.1.1.5
sulfanyl(sulfanylidene)ethanethiyl* = sulfanylbis(sulfanylidene)ethyl = trithiooxalo	HS-CS-CS–	P-65.1.7.2.4; P-65-1.7.3.3
(sulfanylsulfanyl)oxy*	HS-S(O)-O–	P-65.3.2.3
sulfanylsulfonodithiyl = trithiosulfo*	HS-S(S) ₂ –	P-65.3.2.1
[sulfanyl(thiocarbonyl)sulfanyl](thiocarbonyl) = [(dithiocarboxy)sulfanyl]carbonothioyl*	HS-CS-S-CS–	P-65.2.3.1.5
= [(sulfanylcarbonothioyl)sulfanyl]carbonothioyl {not [(dithiocarboxy)sulfanyl]thioformyl}	HS-HP(S)–	P-67.1.4.1.1.5
sulfanyl(thiophosphonoyl) = sulfanylphosphonothioyl*		
sulfeno: see hydroxysulfanyl*		
sulfenyl: see sulfanediyl*		
sulfido*	–S–	P-72.6.2
sulfenamoyl: see aminosulfanyl*		
sulfenamoyloxy: see (aminosulfanyl)oxy*		
sulfenimidoyl*	–S(=NH)–	P-65.3.2.3
sulfino* (when unmodified)	HO-S(O)–	P-65.3.0; P-65.3.2.1
sulfinothiyl*	–S(S)–	P-65.3.2.3
sulfanyl* = thionyl	–S(O)–	P-65.3.2.3
sulfanylbis(oxy)* (not sulfanyldioxy)	–O-S(O)-O–	P-65.3.2.3
sulfanyldioxy: see sulfanylbis(oxy)*		
sulfo* (when unmodified)	HO-SO ₂ –	P-65.3.0; P-65.3.2.1
sulfonato*	–O-SO ₂ –	P-72.6.1
sulfonimidoyl* = sulfurimidoyl	–S(O)(=NH)–	P-65.3.2.3; P-67.1.4.4.1
sulfonio = sulfaniumyl* = sulfoniumyl	H ₂ S ⁺ –	P-73.6
sulfoniumyl = sulfaniumyl* = sulfonio	H ₂ S ⁺ –	P-73.6
sulfonodihydrazonoyl* = sulfurodihydrazonoyl	–S(=N-NH ₂) ₂ –	P-65.3.2.3; P-67.1.4.4.1
sulfonodiimidoyl* = sulfurodiimidoyl	–S(=NH) ₂ –	P-65.3.2.3; P-67.1.4.4.1
sulfonodithiyl* = sulfurodithiyl	–S(=S) ₂ –	P-65.3.2.3; P-67.1.4.4.1
sulfonohydrazonoyl* = sulfurohydrazonoyl	–S(O)(=NNH ₂)–	P-65.3.2.3; P-67.1.4.4.1
sulfonothioyl* = sulfurothiyl	–S(O)(S)–	P-65.3.2.3; P-67.1.4.4.1
sulfonyl* = sulfuryl	–SO ₂ –	P-65.3.2.3; P-67.1.4.4.1
sulfonylbis(methylene)* (not sulfonyldimethylene)	–CH ₂ -SO ₂ -CH ₂ –	P-65.3.2.3
sulfonylbis(oxy)* (not sulfonyldioxy)	–O-SO ₂ -O–	P-65.3.2.3

sulfonylbis(sulfanediy)* (not sulfonyldisulfanediy)	–S-SO ₂ -S–	P-65.3.2.3
sulfonyldimethylene: see sulfonylbis(methylene)*		
sulfonyldioxy: see sulfonylbis(oxy)*		
sulfonyldisulfanediy: see sulfonylbis(sulfanediy)*		
sulfooxy*	HO-SO ₂ -O–	P-65.3.2.3; P-67.1.4.4.2
sulfuramidoyl = sulfamoyl* = aminosulfonyl	H ₂ N-SO ₂ –	P-65.3.2.3; P-66.1.1.4.2
sulfuramidoyloxy = sulfamoyloxy*	H ₂ N-SO ₂ -O–	P-67.1.4.4.2
sulfurimidoyl = sulfonimidoyl*	–S(O)(=NH)–	P-65.3.2.3; P-67.1.4.4.1
sulfur(isothiocyanatido)thioyl = isothiocyanatosulfonothioyl*	(SCN)-S(O)(S)–	P-67.1.4.4.1
sulfur(isothiocyanatidoyl) = isothiocyanatosulfonyl*	(SCN)-SO ₂ –	P-67.1.4.4.1
sulfurochloridoyl = chlorosulfonyl*	Cl-SO ₂ –	P-65.3.2.3; P-67.1.4.4.1
sulfurochloridoyloxy = (chlorosulfonyl)oxy*	Cl-SO ₂ -O–	P-65.3.2.3; P-67.1.4.4.2
sulfurocyanidoyl = cyanosulfonyl*	NC-SO ₂ –	P-67.1.4.4.1
sulfurodihydrazonoyl = sulfonodihydrazonoyl*	–S(=NNH ₂) ₂ –	P-65.3.2.3; P-67.1.4.4.1
sulfurodiimidoyl = sulfonodiimidoyl*	–S(=NH) ₂ –	P-67.1.4.4.1
sulfurodithioyl = sulfonodithioyl*	–S(S) ₂ –	P-65.3.2.3; P-67.1.4.4.1
sulfurohydrazonoyl = sulfonohydrazonoyl*	–S(O)(=NNH ₂)–	P-65.3.2.3; P-67.1.4.4.1
sulfurothioyl = sulfonothioyl*	–S(O)(S)–	P-65.3.2.3; P-67.1.4.4.1
sulfuryl = sulfonyl*	–SO ₂ –	P-65.3.2.3; P-67.1.4.4.1
tellanediy* (not telluro)	–Te–	P-63.2.2.1.2; P-63.2.5
tellano* = epitelluro (ring forming)	–Te–	P-25.4.2.1.4; P-63.5
tellanyl* (not hydrotelluro)	HTe–	P-63.1.5
tellanylidene* = telluroxo	Te=	P-29.3.1; P-64.6.1; P-66.6.3
tellanylidenemethyl = methanetelluroyl* = telluroformyl	HC(Te)–	P-65.1.7.2.3; P-66.6.3
tellureno: see hydroxytellanyl*		
tellurino* (unmodified)	HO-Te(O)–	P-65.3.0; P-65.3.2.1
tellurinyl*	–Te(O)–	P-65.3.2.3
telluro: see tellanediy*		
tellurocyanato* = carbononitridoyltellanyl	NC-Te–	P-65.2.2
telluroformyl = methanetelluroyl* = tellanylidenemethyl	HC(Te)–	P-65.1.7.2.3; P-66.6.3
<i>OTe</i> -tellurohydroperoxy = hydroxytellanyl* (not tellureno)	HO-Te–	P-63.4.2.2
tellurono* (unmodified)	HO-TeO ₂ –	P-63.3.0; P-65.3.2.1
telluronyl*	–TeO ₂ –	P-65.3.2.3
telluroxo = tellanylidene*	Te=	P-29.3.1, P-64.6.1; P-66.6.3
terephthalimidoyl = benzene-1,4-dicarboximidoyl* = 1,4-phenylenebis(iminomethylene) = 1,4-phenylenedicarbonimidoyl	–(HN=C)–  –C(=NH)–	P-65.1.7.3.2

terephthaloyl = benzene-1,4-dicarbonyl* = 1,4-phenylenedicarbonyl
= 1,4-phenylenebis(oxomethylene)

tetraazan-1-yl*

tetradecanoyl* = 1-oxotetradecyl

tetramethylene: see butane-1,4-diyl*

tetrasulfanediyyl* = tetrathio

tetrathio = tetrasulfanediyyl*

thallanyl*

thenyl (2-isomer only): see (thiophen-2-yl)methyl*

2-thienyl = thiophen-2-yl* (also 3-isomer)

thio: see sulfanediyyl*

thioacetamido = ethanethioamido* = (ethanethioy)amino

thioacetyl = ethanethioyl* = 1-sulfanylideneethyl

thioazonoyl = azonothioyl*

thiobenzamido = benzenecarbothioamido* = (benzenecarbothioyl)amino

thiobenzoyl = benzenecarbothioyl* = phenyl(sulfanylidene)methyl = phenyl(thioxo)methyl

thioborono* = hydroxy(sulfanyl)boranyl

thiobutyryl = butanethioyl* = 1-sulfanylidenebutyl = 1-thioxobutyl

thiocarbamoyl: see carbamothioyl*

thiocarbonyl = carbonothioyl*

thiocarboxy* (unspecified)

(thiocarboxy)carbonyl*

thiochlorosyl*

thiocyanato* = carbononitridoysulfanyl = carbononitridoylthio

thiocyanatosulfanyl: see cyanodisulfanyl*

thiodimethylene: see sulfanediyylbis(methylene)*

thioformamido = methanethioamido* = (methanethioyl)amino

thioformyl = methanethioyl* = sulfanylideneethyl

OS-thiohydroperoxy = hydroxysulfanyl* (not hydroxythio, not sulfeno)

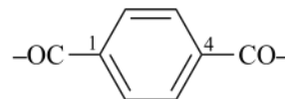
SO-thiohydroperoxy = sulfanyloxy* (not mercaptooxy)

(*OS*-thiohydroperoxy)carbonoselenoyl = (hydroxysulfanyl)carbonoselenoyl*

(thiohydroperoxy)carbonyl = carbono(thioperoxy)*

(*OS*-thiohydroperoxy)carbonyl = (hydroxysulfanyl)carbonyl*

[(*SO*-thiohydroperoxy)carbonyl]oxy = [(sulfanyloxy)carbonyl]oxy*



P-65.1.7.3.1; P-65.1.7.4.2



P-68.3.1.4.1



P-65.1.7.4.1



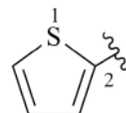
P-68.4.1.2



P-68.4.1.2



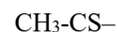
P-29.3.1; P-68.1.2



P-29.6.2.3; P-57.1.5.3



P-66.1.4.4



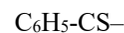
P-65.1.7.2.3



P-67.1.4.1.1.4



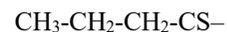
P-66.1.4.4



P-65.1.7.2.3



P-68.1.4.2



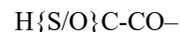
P-65.1.7.4.1



P-65.2.1.5



P-65.2.1.6



P-65.1.7.2.1; P-65.1.7.2.4



P-67.1.4.5



P-65.2.2



P-66.1.4.4



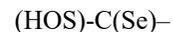
P-65.1.7.2.3; P-66.6.3



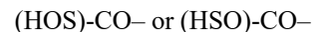
P-63.4.2.2



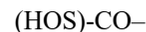
P-63.4.2.2



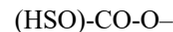
P-65.2.1.7



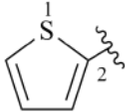
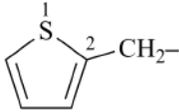
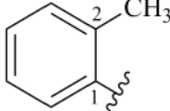
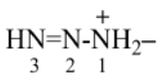
P-65.1.5.3; P-65.2.1.7



P-65.1.5.3; P-65.2.1.7



P-65.2.1.7

(<i>OS</i> -thiohydroperoxy)phosphorothioyl = (hydroxysulfanyl)phosphorothioyl*	(HOS)-P(S)<	P-67.1.4.1.1.4; P-67.1.4.1.1.5
(thiohydroperoxy)phosphoryl = phosphoro(thioperoxoyl)* = (thioperoxy)phosphoryl	(HSO)-P(O)< or (HOS)-P(O)<	P-67.1.4.1.1.4
(<i>SO</i> -thiohydroperoxy)phosphoryl = (sulfanyloxy)phosphoryl*	(HSO)-P(O)<	P-67.1.4.1.1.5
thionitroso = sulfanylideneamino* = thioxoamino	S=N-	P-67.1.4.3.2
(thionitroso)sulfanyl = (sulfanylideneamino)sulfanyl* = (thioxoamino)sulfanyl	S=N-S-	P-67.1.4.3.2
thionyl = sulfanyl*	-S(O)-	P-65.3.2.3
2-thiooxalo: see hydroxy(sulfanylidene)acetyl*		
(thioperoxy)phosphoryl = phosphoro(thioperoxoyl)* = (thiohydroperoxy)phosphoryl	(HSO)-P(O)< or (HOS)-P(O)<	P-67.1.4.1.1.4
thiophen-2-yl* = 2-thienyl (also 3- isomer)		P-29.6.2.3; P-57.1.5.3
(thiophen-2-yl)methyl* (not thenyl)		P-29.6.3
thiophosphinoyl = phosphinothioyl* = dihydrophosphorothioyl	H ₂ P(S)-	P-67.1.4.1.1.4; P-67.1.4.1.2
thiophosphono* (unspecified)	(HO)(HS)P(O)- or (HO) ₂ P(S)-	P-67.1.4.1.1.1
thiophosphoryl = phosphorothioyl*	-P(S)<	P-67.1.4.1.1.4
thiopropionyl = propanethioyl* = 1-sulfanylidenepropyl = 1-thioxopropyl	CH ₃ -CH ₂ -CS-	P-65.1.7.4.1
thiosulfeno: see disulfanyl*		
thiosulfino* (unspecified)	H{S/O}S-	P-65.3.2.1
thiosulfo* (unspecified)	HO ₂ S ₂ -	P-65.3.2.1
thioxo = sulfanylidene*	S=	P-29.3.1; P-64.6.1
thioxoamino = thionitroso = sulfanylideneamino*	S=N-	P-67.1.4.3.2
(thioxoamino)sulfanyl = (sulfanylideneamino)sulfanyl* = (thionitroso)sulfanyl	S=N-S-	P-67.1.4.3.2
1-thioxobutyl = butanethioyl* = 1-sulfanylidenebutyl = thiobutyl	CH ₃ -CH ₂ -CH ₂ -CS-	P-65.1.7.4.1
thioxomethylidene = sulfanylidenemethylidene*	S=C=	P-65.2.1.8
1-thioxopropyl = propanethioyl* = thiopropionyl = 1-sulfanylidenepropyl	CH ₃ -CH ₂ -CS-	P-65.1.7.4.1
<i>o</i> -toluidino: see 2-methylanilino* (also <i>m</i> - and <i>p</i> -isomers)		
<i>o</i> -tolyl = 2-methylphenyl* (also <i>m</i> - = 3- and <i>p</i> - = 4-isomers)		P-29.6.2.3; P-57.1.5.3
triazano: see triazan-1-yl*		
triazan-1-yl* (not triazano)	H ₂ N-NH-NH-	P-29.3.2.2; P-68.3.1.4.1
triaz-1-ene-1,3-diyl* (not diazoamino)	-N=N-NH-	P-68.3.1.4.2
triaz-2-en-1-ium-1-yl* = triaz-2-en-1-io		P-73.6
triaz-2-en-1-io = triaz-2-en-1-ium-1-yl*		

triaz-2-eno: see triaz-2-en-1-yl*		
triaz-2-en-1-yl* (not triaz-2-eno)	HN=N-NH-	P-32.1.1; P-68.3.1.4.1
triboran(5)-1-yl*	H ₂ B-BH-BH-	P-68.1.2
tricyclo[3.3.1.1 ^{3,7}]decan-2-yl = adamantan-2-yl* = 2-adamantyl (also 1-isomer)	C ₁₀ H ₁₅ -	P-29.6.2.3
trihydroxysilyl*	(HO) ₃ Si-	P-67.1.4.2
trimethoxysilyl*	(CH ₃ O) ₃ Si-	P-67.1.4.2
trimethylene: see propane-1,3-diyl*		
trioxidenediyl* = trioxy	-O-O-O-	P-68.4.1.2
trioxidanyl* = hydrotrioxy	HO-O-O-	P-68.4.1.3
trioxy = trioxidenediyl*	-O-O-O-	P-68.4.1.2
triphenylmethyl* = trityl (unsubstituted)	(C ₆ H ₅) ₃ C-	P-29.6.2.2
triselanediyl* = triseleno	-Se-Se-Se-	P-68.4.1.2
triselanyl* = hydrotriseleno	HSeSeSe-	P-68.4.1.3
triseleno = triselanediyl	-Se-Se-Se-	P-68.4.1.2
trisilan-2-yl*	(SiH ₃) ₂ SiH-	P-29.3.2.2
trisilazan-3-yl: see bis(silylamino)silyl*		
trisiliranyl* = cyclotrisilanyl	$\begin{array}{c} \text{H}_2\text{Si} \\ \\ \text{H}_2\text{Si} \end{array} \text{SiH}-$	P-68.2.2
trisulfanediyl* = trithio	-S-S-S-	P-68.4.1.2
trisulfanyl* = hydrotrithio	HS-S-S-	P-68.4.1.3
tritellanediyl* = tritelluro	-Te-Te-Te-	P-68.4.1.2
tritellanyl* = hydrotritelluro	HTe-Te-Te-	P-68.4.1.3
tritelluro = tritellanediyl*	-Te-Te-Te-	P-68.4.1.2
trithio = trisulfanediyl*	-S-S-S-	P-68.4.1.2
trithiooxalo = sulfanyl(sulfanylidene)ethanethiyl* = sulfanylbis(sulfanylidene)ethyl	HS-CS-CS-	P-65.1.7.2.4; P-65.1.7.3.3
trithiophosphono*	(HS) ₂ P(S)-	P-67.1.4.1.1.1
trithiosulfo* = sulfanylsulfonodithiyl	HS-S(S) ₂ -	P-65.3.2.1
trityl (unsubstituted) = triphenylmethyl*	(C ₆ H ₅) ₃ C-	P-29.6.2.2
undecan-1-yl = undecyl*	CH ₃ -[CH ₂] ₉ -CH ₂ -	P-29.3.2.1; P-29.3.2.2
undecyl* = undecan-1-yl	CH ₃ -[CH ₂] ₉ -CH ₂ -	P-29.3.2.1; P-29.3.2.2
ureido: see carbamoylamino*		
ureylene: see carbonylbis(azanediyl)*		
vinyl = ethenyl*	CH ₂ =CH-	P-32.3
vinylidene: see ethenylidene*		
vinylidene = ethenylidene*	CH ₂ =C=	P-32.3
2,3-xylydino: see 2,3-dimethylanilino*		

yloamino* = yloazanyl
yloazanyl = yloamino*
yloformyl*
ylohydroxy: see ylooxidanyl*
ylomethyl*
ylooxidanyl* = ylooxy (not ylohydroxy)
(ylooxidanyl)formyl = oxylcarbonyl*
ylooxy = ylooxidanyl* (not ylohydroxy)

HN [•] -	P-71.5
HN [•] -	P-71.5
O=C [•] -	P-71.5
H ₂ C [•] -	P-71.5
[•] O-	P-71.5
[•] O-CO-	P-71.5
[•] O-	P-71.5