

Mode of Action Classification



The Insecticide Resistance Action Committee

Mode of Action Classification Brochure

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Foreword

Effective insecticide resistance management (IRM) in conjunction with integrated pest management (IPM) is vital to global crop protection, sustainable agriculture and improved public health, and it is an essential element of responsible product stewardship.

The Insecticide Resistance Action Committee (IRAC) was formed in 1984 and works as a specialist technical group of the industry association CropLife International, to provide a coordinated crop protection industry response to prevent or delay the development of resistance in insect and mite pests. There are now IRAC country group committees in many parts of the world, researching and responding to local resistance issues, as well as the parent IRAC International group, which provides a coordinating and supporting role at the global level (see also www.irac-online.org).

Developing new insecticides is becoming increasingly difficult and costly, so it is vital to protect those effective products in the marketplace from the development of resistance. Moreover, with fewer new insecticides being discovered and regulatory pressures reducing the number of older commercial chemistries available, the 'toolbox' of usable insecticides is being reduced, making effective IRM more important than ever. The Mode of Action Classification Scheme is a key part of IRAC's global IRM strategy.

Mode of Action Classification

IRAC promotes the use of a Mode of Action (MoA) Classification of insecticides and acaricides as the basis for effective and sustainable resistance management. Actives are allocated to specific groups based on their target site. Reviewed and re-issued periodically, the IRAC MoA Classification Scheme provides farmers, growers, advisors, extension staff, consultants and crop protection professionals with a guide to the selection of acaricides and insecticides in resistance management programs. Effective resistance management of this type preserves the utility and diversity of available insecticides and acaricides. A complete list of the different MoA groups is shown in the following pages, followed by a breakdown of MoAs available for Lepidoptera, aphids, whitefly, hoppers, mites and mosquitoes. For further information, please refer to the full IRAC MoA Classification Scheme on the IRAC website (www.irac-online.org).

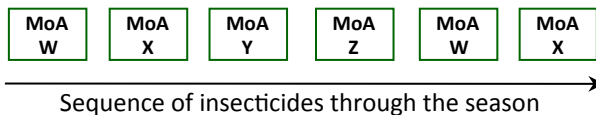
What is Resistance?

Resistance to insecticides may be defined as *‘a heritable change in the sensitivity of a pest population that is reflected in the repeated failure of a product to achieve the expected level of control when used according to the label recommendation for that pest species’* (IRAC). Resistance arises through the over-use or misuse of an insecticide or acaricide against a pest species, and results in the selection of resistant forms of the pest and the consequent evolution of populations that are resistant to that insecticide or acaricide.

Effective IRM Strategies: Sequences or Alternations of MoA

All effective insecticide resistance management (IRM) strategies seek to minimise the selection of resistance to any one type of insecticide. In practice, alternations, sequences or rotations of compounds from different MoA groups provide sustainable and effective IRM for insect and mite pests. This ensures that selection from compounds in the same MoA group is minimised, and resistance is less likely to evolve.

Example:



Applications are often arranged into MoA spray windows or blocks that are defined by the stage of crop development, together with the biology and phenology of the species of concern. Local expert advice should always be followed with regard to spray windows and timing. Several sprays may be possible within each spray window, but it is generally essential that successive generations of the pest are not treated with compounds from the same MoA group. IRAC also offers specific recommendations for some MoA groups. Metabolic resistance mechanisms may give cross-resistance between MoA groups; where this is known to occur, the above advice should be modified accordingly. For further information on the use of MoA groups and sub-groups, please see the notes at the end of the brochure and in the full MoA Classification Scheme.

IRAC Mode of Action Classification Scheme (Classification Version 8.1)

Targeted Physiology: Nerve & Muscle Growth & Development Respiration Midgut Unknown or Non-specific

Note: Rotations for resistance management should be based only on the numbered mode of action groups - see table footnotes for details

Main Group/Primary Site of Action	Chemical Subgroup or Exemplifying active	Active Ingredients
1 Acetylcholinesterase (AChE) inhibitors <i>See footnotes for further information on use of compounds between sub-groups.</i>	1A Carbamates	Alanycarb, Aldicarb, Bendiocarb, Benfuracarb, Butocarboxim, Butoxycarboxim, Carbaryl, Carbofuran, Carbosulfan, Ethiofencarb, Fenobucarb, Formetanate, Furathiocarb, Isoprocarb, Methiocarb, Methomyl, Metolcarb, Oxamyl, Pirimicarb, Propoxur, Thiodicarb, Thiofanox, Triazamate, Trimethacarb, XMC, Xyllycarb
	1B Organophosphates	Acephate, Azamethiphos, Azinphos-ethyl, Azinphos-methyl, Cadusafos, Chlorethoxyfos, Chlorfenvinphos, Chlormephos, Chlorpyrifos, Chlorpyrifos-methyl, Coumaphos, Cyanophos, Demeton-S-methyl, Diazinon, Dichlorvos/DDVP, Dicrotophos, Dimethoate, Dimethylvinphos, Disulfoton, EPN, Ethion, Ethoprophos, Famphur, Fenamiphos, Fenitrothion, Fenthion, Fosthiazate, Heptenophos, Imicyafos, Isufenphos, Isopropyl O-(methoxyaminothio-phosphoryl) salicylate, Isoxathion, Malathion, Mecarbam, Methamidophos, Methidathion, Mevinphos, Monocrotophos, Naled, Omethoate, Oxydemeton-methyl, Parathion, Parathion-methyl, Phenthoate, Phorate, Phosalone, Phosmet, Phosphamidon, Phoxim, Pirimiphos- methyl, Profenofos, Propetamphos, Prothiofos, Pyraclofos, Pyridaphenthion, Quinalphos, Sulfotep, Tebupirimfos, Temephos, Terbufos, Tetrachlorvinphos, Thiometon, Triazophos, Trichlorfon, Vamidothion
2 GABA-gated chloride channel blockers	2A Cyclodiene organochlorines	Chlordane, Endosulfan
	2B Phenylpyrazoles (Fiproles)	Ethiprole, Fipronil

3 Sodium channel modulators <i>See footnotes for further information on use of compounds between sub-groups.</i>	3A Pyrethroids Pyrethrins	Acrinathrin, Allethrin, d- <i>cis</i> -trans Allethrin, d- <i>trans</i> Allethrin, Bifenthrin, Bioallethrin, Bioallethrin S-cyclopentenyl, Bioresmethrin, Cycloprothrin, Cyfluthrin, <i>beta</i> -Cyfluthrin, Cyhalothrin, <i>lambda</i> -Cyhalothrin, <i>gamma</i> -Cyhalothrin, Cypermethrin, <i>alpha</i> -Cypermethrin, <i>beta</i> -Cypermethrin, <i>theta</i> -cypermethrin, <i>zeta</i> -Cypermethrin, Cyphenothrin [(1 <i>R</i>)- <i>trans</i> - isomers], Deltamethrin, Empenthrin [(<i>EZ</i>)- (1 <i>R</i>)- isomers], Esfenvalerate, Etofenprox, Fenpropathrin, Fenvalerate, Flucythrinate, Flumethrin, <i>tau</i> -Fluvalinate, Halfenprox, Imiprothrin, Kadethrin, Permethrin, Phenothrin [(1 <i>R</i>)- <i>trans</i> - isomer], Prallethrin, Pyrethrins (pyrethrum), Resmethrin, Silafluofen, Tefluthrin, Tetramethrin, Tetramethrin [(1 <i>R</i>)-isomers], Tralomethrin, Transfluthrin
	3B DDT Methoxychlor	DDT Methoxychlor
4 Nicotinic acetylcholine receptor (nAChR) competitive modulators <i>See footnotes for further information on use of compounds between sub-groups.</i>	4A Neonicotinoids	Acetamiprid, Clothianidin, Dinotefuran, Imidacloprid, Nitenpyram, Thiacloprid, Thiamethoxam
	4B Nicotine	Nicotine
	4C Sulfoximines	Sulfoxaflor
	4D Butenolides	Flupyradifurone
	4E Mesoionics	Triflumezopyrim
5 Nicotinic acetylcholine receptor (nAChR) allosteric modulators	Spinosyns	Spinetoram, Spinosad
6 Glutamate-gated chloride channel (GluCl) allosteric modulators	Avermectins, Milbemycins	Abamectin, Emamectin benzoate, Lepimectin, Milbemectin

Main Group/Primary Site of Action	Chemical Subgroup or Exemplifying active	Active Ingredients
7 Juvenile hormone mimics	7A Juvenile hormone analogues	Hydroprene, Kinoprene, Methoprene
	7B Fenoxycarb	Fenoxycarb
	7C Pyriproxyfen	Pyriproxyfen
8 Miscellaneous non-* specific (multi-site) inhibitors	8A Alkyl halides	Methyl bromide and other alkyl halides
	8B Chloropicrin	Chloropicrin
	8C Fluorides	Cryolite (Sodium aluminum fluoride), Sulfuryl fluoride
	8D Borates	Borax, Boric acid, Disodium octaborate, Sodium borate, Sodium metaborate
	8E Tartar emetic	Tartar emetic
	8F Methyl isothiocyanate generators	Dazomet, Metam
9 Chordotonal organ TRPV channel modulators	9B Pyridine azomethine derivatives	Pymetrozine, Pyrifluquinazon
10 Mite growth inhibitors <i>See footnotes for further sub-grouping information for 10A</i>	10A Clofentezine Diflovidazin Hexythiazox	Clofentezine, Diflovidazin, Hexythiazox
	10B Etoxazole	Etoxazole

11 Microbial disruptors of insect midgut membranes	11A <i>Bacillus thuringiensis</i> and the insecticidal proteins they produce <i>See footnotes for further sub-grouping information</i>	<i>Bacillus thuringiensis</i> subsp. <i>israelensis</i> <i>Bacillus thuringiensis</i> subsp. <i>aizawai</i> <i>Bacillus thuringiensis</i> subsp. <i>kurstaki</i> <i>Bacillus thuringiensis</i> subsp. <i>tenebrionis</i> <i>Bt</i> crop proteins: (see footnote) Cry1Ab, Cry1Ac, Cry1Fa, Cry1A.105, Cry2Ab, Vip3A, mCry3A, Cry3Ab, Cry3Bb, Cry34Ab1/Cry35Ab1
	11B <i>Bacillus sphaericus</i>	<i>Bacillus sphaericus</i>
12 Inhibitors of mitochondrial ATP synthase	12A Diafenthiuron	Diafenthiuron
	12B Organotin miticides	Azocyclotin, Cyhexatin, Fenbutatin oxide
	12C Propargite	Propargite
	12D Tetradifon	Tetradifon
13 Uncouplers of * oxidative phosphorylation via disruption of the proton gradient	Pyrroles Dinitrophenols Sulfluramid	Chlorfenapyr, DNOC, Sulfluramid
14 Nicotinic acetylcholine receptor (nAChR) channel blockers	Nereistoxin analogues	Bensultap, Cartap hydrochloride, Thiocyclam, Thiosultap-sodium

Main Group/Primary Site of Action	Chemical Subgroup or Exemplifying active	Active Ingredients
15 Inhibitors of chitin biosynthesis, type 0	Benzoylureas	Bistrifluron, Chlorfluazuron, Diflubenzuron, Flucycloxuron, Flufenoxuron, Hexaflumuron, Lufenuron, Novaluron, Noviflumuron, Teflubenzuron, Triflumuron
16 Inhibitors of chitin biosynthesis, type 1	Buprofezin	Buprofezin
17 Moulting disruptors, Dipteran	Cyromazine	Cyromazine
18 Ecdysone receptor agonists	Diacylhydrazines	Chromafenozide, Halofenozide, Methoxyfenozide, Tebufenozide
19 Octopamine receptor agonists	Amitraz	Amitraz
20 Mitochondrial complex III electron transport inhibitors	20A Hydramethylnon	Hydramethylnon
	20B Acequinocyl	Acequinocyl
	20C Fluacrypyrim	Fluacrypyrim
	20D Bifenazate	Bifenazate
21 Mitochondrial complex I electron transport inhibitors	21A METI acaricides and insecticides	Fenazaquin, Fenpyroximate, Pyridaben, Pyrimidifen, Tebufenpyrad, Tolfenpyrad
	21B Rotenone	Rotenone (Derris)

22 Voltage-dependent sodium channel blockers <i>See footnotes for further information on sub-grouping</i>	22A Oxadiazines	Indoxacarb
	22B Semicarbazones	Metaflumizone
23 Inhibitors of acetyl CoA carboxylase	Tetronic and Tetramic acid derivatives	Spirodiclofen, Spiromesifen, Spirotetramat
24 Mitochondrial complex IV electron transport inhibitors	24A Phosphides	Aluminium phosphide, Calcium phosphide, Phosphine, Zinc phosphide
	24B Cyanides	Calcium cyanide, Potassium cyanide, Sodium cyanide
25 Mitochondrial complex II electron transport inhibitors <i>See footnotes for further information on sub-grouping</i>	25A <i>beta</i> -Ketonitrile derivatives	Cyenopyrafen, Cyflumetofen
	25B Carboxanilides	Pyflubumide
28 Ryanodine receptor modulators	Diamides	Chlorantraniliprole, Cyantraniliprole, Flubendiamide
29 Chordotonal organ modulators - undefined target site	Flonicamid	Flonicamid

Main Group/Primary Site of Action	Chemical Subgroup or Exemplifying active	Active Ingredients
UN * Compounds of unknown or uncertain mode of action	Azadirachtin	Azadirachtin
	Benzoximate	Benzoximate
	Bromopropylate	Bromopropylate
	Chinomethionat	Chinomethionat
	Dicofol	Dicofol
	GS-omega/kappa HXTX-Hv1a peptide	GS-omega/kappa HXTX-Hv1a peptide
	Lime sulfur	Lime sulfur
	Pyridalyl	Pyridalyl
	Sulfur	Sulfur

Targeted Physiology: Nerve & Muscle Growth & Development Respiration Midgut Unknown or Non-specific

The colour scheme in the table associates mode of action into broad categories based on the physiological functions affected, as an aid to understanding symptomology, speed of action and other properties of the insecticides, and not for any resistance management purpose. Rotations for resistance management should be based only on the numbered mode of action groups.

General Notes:

- Inclusion of a compound in the classification above does not necessarily signify regulatory approval.
- MoA assignments will usually involve identification of the target protein responsible for the biological effect, although groupings can be made where compounds share distinctive physiological effects and have related chemical structures.
- Groups 26 and 27 are unassigned at this time and have therefore been omitted from the table.
- A compound with an unknown or controversial MoA or an unknown mode of toxicity will be held in group 'UN' until evidence becomes available to enable that compound to be assigned to a more appropriate MoA class.
- Actives in groups marked with a * are thought not to share a common target site and therefore may be freely rotated with each other unless there is reason to expect cross-resistance. These groups are 8, 13, and UN.

Notes on Sub-Groups:

Sub-groups represent distinct chemical classes that are believed to have the same MoA but are different enough in chemical structure or mode of interaction with the target protein that the chance of selection for either metabolic or target-site cross-resistance is reduced compared to close analogs. Sub-groups may also distinguish compounds that are chemically similar but known to bind differently within the target or to have differential selectivity among multiple targets.

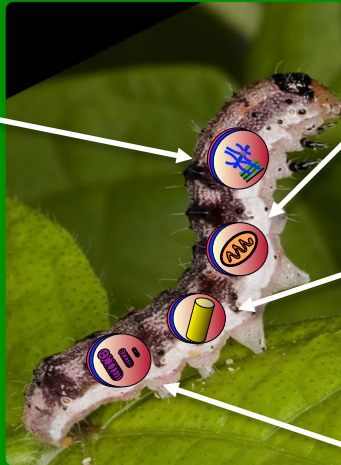
The cross-resistance potential between sub-groups is higher than that between different groups, so rotation between sub-groups should be avoided. In exceptional circumstances (i.e. where effective registered insecticides from other mode of action groups are unavailable) rotation may be considered following consultation with local expert advice and where cross-resistance does not exist. These exceptions should not be considered sustainable resistance management strategies, and alternative options should be sought to maintain pest susceptibility.

Sub-group	Notes
3B	Because DDT is no longer used in agriculture, this is only applicable for the control of human disease vectors such as mosquitoes.
4A, 4B, 4C, 4D & 4E	Although these compounds are believed to have the same target site, current evidence indicates that the risk of metabolic cross-resistance between subgroups is low.
10A	Hexythiazox is grouped with clofentezine because they exhibit cross-resistance, even though they are structurally distinct, and the target site for these compounds is unknown. Diflovidazin has been added to this group because it is a close analogue of clofentezine and is expected to have the same mode of action.
11A	Different <i>Bacillus thuringiensis</i> products that target different insect orders may be used together without compromising their resistance management. Rotation between certain specific <i>Bacillus thuringiensis</i> microbial products may provide resistance management benefits for some pests. Consult product-specific recommendations. B.t. Crop Proteins: Where there are differences among the specific receptors within the midguts of target insects, transgenic crops containing certain combinations of the listed proteins provide resistance management benefits.
22A, 22B	Although these compounds are believed to have the same target site, current evidence indicates that the risk of metabolic cross-resistance between subgroups is low.
25A, 25B	Although these compounds are believed to have the same target site, current evidence indicates that the risk of metabolic cross-resistance between subgroups is low.

Nerve & Muscle Targets

1. Acetylcholinesterase (AChE) inhibitors
1A: *Carbamates*
1B: *Organophosphates*
2. GABA-gated chloride channel blockers
2A: *Cyclodiene Organochlorines*
2B: *Phenylpyrazoles*
3. Sodium channel modulators
3A: *Pyrethrins, Pyrethroids*
4. Nicotinic acetylcholine receptor (nAChR) competitive modulators
4A: *Neonicotinoids*
5. Nicotinic acetylcholine receptor (nAChR) allosteric modulators
5 *Spinosyns*
6. Glutamate-gated chloride channel (GluCl) allosteric modulators
6: *Avermectins, Milbemycins*
14. Nicotinic acetylcholine receptor (nAChR) channel blockers
14: *Nereistoxin analogues*
22. Voltage-dependent sodium channel blockers
22A: *Oxadiazines*
22B: *Semicarbazones*
28. Ryanodine receptor modulators
28: *Diamides*

Lepidoptera - Mode of Action Classification by Target Site



Respiration Targets

13. Uncouplers of oxidative phosphorylation via disruption of the proton gradient
13: *Chlorfenapyr*
21. Mitochondrial complex I electron transport inhibitors
21A: *METI acaracides and insecticides (Tolfenpyrad)*

Midgut Targets

11. Microbial disruptors of insect midgut membranes
11A: *Bacillus thuringiensis*,
11B: *Bacillus sphaericus*

Growth & Development Targets

7. Juvenile hormone mimics
7A: *Juvenile hormone analogues (Hydroprene)*
7B: *Fenoxycarb*
15. Inhibitors of chitin biosynthesis, type 0
15: *Benzoylureas*
18. Ecdysone receptor agonists
18: *Diacylhydrazines*

Unknown or uncertain MoA

Azadirachtin, Pyridalyl

Nerve & Muscle Targets

1. Acetylcholinesterase (AChE) inhibitors
1A: *Carbamates*
1B: *Organophosphates*
2. GABA-gated chloride channel blockers
2A: *Cyclodiene Organochlorines*
2B: *Phenylpyrazoles*
3. Sodium channel modulators
3A: *Pyrethrins, Pyrethroids*
4. Nicotinic acetylcholine receptor (nAChR) competitive modulators
4A: *Neonicotinoids*
4C: *Sulfoximines*
4D: *Butenolides*
4E: *Mesoionics*
9. Chordotonal organ TRPV channel modulators
9B: *Pyridine azomethine derivatives*
22. Voltage-dependent sodium channel blockers
22A: *Oxadiazines*
28. Ryanodine receptor modulators
28: *Diamides (Cyantraniliprole)*
29. Chorodotonal organ modulators – undefined target site
29: *Fonicamid*

Aphids, Whiteflies & Hoppers - Mode of Action Classification by Target Site



MoA Group	Aphids	Whiteflies	Hoppers
1A	X	X	X
1B	X	X	X
2A	X	X	X
2B			X
3A	X	X	X
4A	X	X	X
4C	X	X	X
4D	X	X	X
4E			X
7A	X	X	
7C		X	
9B	X	X	X
12A	X	X	
15		X	
16		X	X
21A		X	
22A			X
23	X	X	
28	X	X	X
29	X	X	X

Respiration Targets

12. Inhibitors of mitochondrial ATP synthesis
12A: *Difenthiuron*
21. Mitochondrial complex I electron transport inhibitors
21A: *METI acaricides and insecticides (Pyridaben, Tolfenpyrad)*

Growth & Development Targets

7. Juvenile hormone mimics
7A: *Kinoprene*
7C: *Pyriproxyfen*
15. Inhibitors of chitin biosynthesis, type 0
15: *Benzoylureas*
16. Inhibitors of chitin biosynthesis, type 1
16: *Buprofezin*
23. Inhibitors of acetyl CoA carboxylase
23: *Tetronic & Tetramic acid derivatives*

The table lists the main mode of action groups for the control of aphids, whiteflies and hoppers. However, the availability may differ regionally due to registration status.

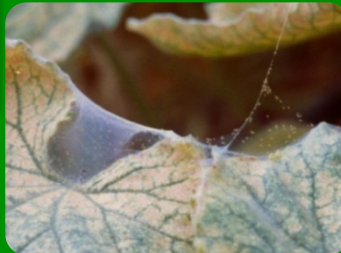
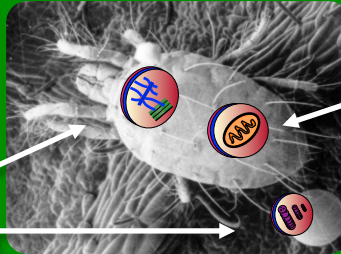
Nerve & Muscle Targets

1. Acetylcholinesterase (AChE) inhibitors
1A: *Carbamates*
1B: *Organophosphates*
2. GABA-gated chloride channel blockers
2A: *Cyclodiene Organochlorines*
3. Sodium channel modulators
3A: *Pyrethrins, Pyrethroids*
5. Nicotinic acetylcholine receptor (nAChR) allosteric modulators
5: *Spinosyns*
6. Glutamate-gated chloride channel (GluCl) allosteric modulators
6: *Avermectins, Milbemycins*
19. Octopamine receptor agonists
19: *Amitraz*

Growth & Development Targets

10. Mite growth inhibitors
10A: *Clofentezine, Diflovidazin*
Hexythiazox
10B: *Etoxazole*
15. Inhibitors of chitin biosynthesis, type O
15: *Benzoylureas*
23. Inhibitors of acetyl CoA carboxylase
23: *Tetronic & Tetramic acid derivatives*

Mites - Mode of Action Classification by Target Site



Respiration Targets

12. Inhibitors of mitochondrial ATP synthesis
12A: *Difenthiuron*
12B: *Organotin miticides*
12C: *Propargite*
13. Uncouplers of oxidative phosphorylation via disruption of the proton gradient
13: *Chlorfenapyr*
20. Mitochondrial complex III electron transport inhibitors
20B: *Acequinocyl*
20C: *Fluacrypyrim*
20D: *Bifenazate*
21. Mitochondrial complex I electron transport inhibitors
21A: *METI acaricides*
25. Mitochondrial complex II electron transport inhibitors
25A: *Cyenoptyrafen, Cyflumetofen*
25B: *Pyflubumide*

Unknown or uncertain MoA

Benzoximate, Chinomethionat, Dicofof

Mosquitoes - Mode of Action Classification by Target Site

Nerve & Muscle Targets (Larvae)

1. Acetylcholinesterase (AChE) inhibitors
1B: *Organophosphates*
3. Sodium channel modulators
3A: *Pyrethrins, Pyrethroids*
5. Nicotinic acetylcholine receptor (nAChR) allosteric modulators
5: *Spinosyns*

Growth & Development Targets (Larvae)

7. Juvenile hormone mimics
7A: *Juvenile hormone analogues*
7C: *Pyriproxyfen*
15. Inhibitors of chitin biosynthesis, type 0
15: *Benzoylureas*



Nerve & Muscle Targets (Adults)

1. Acetylcholinesterase (AChE) inhibitors
1A: *Carbamates*
1B: *Organophosphates*
3. Sodium channel modulators
3A: *Pyrethrins, Pyrethroids*
3B: *DDT*

Midgut Targets (Larvae)

11. Microbial disruptors of insect midgut membranes
11A: *Bacillus thuringiensis*,
11B: *Bacillus sphaericus*



Active Ingredients (Alphabetical Order) with MOA Classification

Abamectin	6
Acephate	1B
Acequinocyl	20B
Acetamiprid	4A
Acrinathrin	3A
Alanycarb	1A
Aldicarb	1A
Allethrin	3A
<i>alpha</i> -Cypermethrin	3A
Aluminium phosphide	24A
Amitraz	19
Azadirachtin	UN
Azamethiphos	1B
Azinphos-ethyl	1B
Azinphos-methyl	1B
Azocyclotin	12B
<i>Bacillus thuringiensis</i>	11A
<i>Bacillus sphaericus</i>	11B
Bendiocarb	1A
Benfuracarb	1A
Bensultap	14
Benzoximate	UN
<i>beta</i> -Cyfluthrin	3A
<i>beta</i> -Cypermethrin	3A
Bifenazate	20D
Bifenthrin	3A
Bioallethrin	3A
Bioallethrin S-cyclopentenyl isomer	3A
Bioresmethrin	3A

Bistrifluron	15
Borax	8D
Boric acid	8D
Bromopropylate	UN
Buprofezin	16
Butocarboxim	1A
Cadusafos	1B
Calcium cyanide	24B
Calcium phosphide	24A
Carbaryl	1A
Carbofuran	1A
Carbosulfan	1A
Cartap hydrochloride	14
Chinomethionat	UN
Chlorantraniliprole	28
Chlordane	2A
Chlorethoxyfos	1B
Chlorfenapyr	13
Chlorfenvinphos	1B
Chlorfluazuron	15
Chlormephos	1B
Chloropicrin	8B
Chlorpyrifos	1B
Chlorpyrifos-methyl	1B
Chromafenozide	18
Clofentezine	10A
Clothianidin	4A
Coumaphos	1B
Cryolite	8C

Cyanide	24B
Cyanophos	1B
Cyantraniliprole	28
Cycloprothrin	3A
Cyenoxyrafen	25A
Cyflumetofen	25A
Cyfluthrin	3A
Cyhalothrin	3A
Cyhexatin	12B
Cypermethrin	3A
Cyphenothrin (1 <i>R</i>)- <i>trans</i> -isomers]	3A
Cyromazine	17
<i>d-cis-trans</i> Allethrin	3A
Dazomet	8F
DDT	3B
Deltamethrin	3A
Demeton-S-methyl	1B
Diafenthiuron	12A
Diazinon	1B
Dichlorvos/ DDVP	1B
Dicofol	UN
Dicrotophos	1B
Diflovidazin	10A
Diflubenzuron	15
Dimethoate	1B
Dimethylvinphos	1B
Dinotefuran	4A
Disodium octaborate	8D
Disulfoton	1B

DNOC	13
<i>d-trans</i> Allethrin	3A
Emamectin benzoate	6
Empenthrin [(<i>EZ</i>)-(1 <i>R</i>)-isomers]	3A
Endosulfan	2A
EPN	1B
Esfenvalerate	3A
Ethiofencarb	1A
Ethion	1B
Ethiprole	2B
Ethoprophos	1B
Etofenprox	3A
Etoazole	10B
Famphur	1B
Fenamiphos	1B
Fenazaquin	21A
Fenbutatin oxide	12B
Fenitrothion	1B
Fenobucarb	1A
Fenoxycarb	7B
Fenpropathrin	3A
Fenpyroximate	21A
Fenthion	1B
Fenvalerate	3A
Fipronil	2B
Fonicamid	29
Fluacrypyrim	20C
Flubendimide	28

Flucycloxuron	15
Flucythrinate	3A
Flufenoxuron	15
Flumethrin	3A
Flupyradifurone	4D
Formetanate	1A
Fosthiazate	1B
Furathiocarb	1A
<i>gamma</i> -Cyhalothrin	3A
GS-omega/kappa HXTX-Hv1a	UN
Halfenprox	3A
Halofenozide	18
Heptenophos	1B
Hexaflumuron	15
Hexythiazox	10A
Hydramethylnon	20A
Hydroprene	7A
Imicyafos	1B
Imidacloprid	4A
Imiprothrin	3A
Indoxacarb	22A
Isofenphos	1B
Isoprocarb	1A
Isopropyl O- (methoxy-aminothio-phosphoryl) salicylate	1B
Isoxathion	1B
Kadethrin	3A
Kinoprene	7A
<i>lambda</i> -Cyhalothrin	3A
Lepimectin	6

Lime sulfur	UN
Lufenuron	15
Malathion	1B
Mecarbam	1B
Metaflumizone	22B
Metam	8F
Methamidophos	1B
Methidathion	1B
Methiocarb	1A
Methomyl	1A
Methoprene	7A
Methoxychlor	3B
Methoxyfenozide	18
Methyl bromide	8A
Metolcarb	1A
Mevinphos	1B
Milbemectin	6
Monocrotophos	1B
Naled	1B
Nicotine	4B
Nitenpyram	4A
Novaluron	15
Noviflumuron	15
Omethoate	1B
Oxamyl	1A
Oxydemeton-methyl	1B
Parathion	1B
Parathion-methyl	1B
Permethrin	3A
Phenothrin [(1 <i>R</i>)- <i>trans</i> - isomer]	3A
Penthotoate	1B

Phorate	1B
Phosalone	1B
Phosmet	1B
Phosphamidon	1B
Phosphine	24A
Phoxim	1B
Pirimicarb	1A
Pirimiphos- methyl	1B
Potassium cyanide	24B
Prallethrin	3A
Profenofos	1B
Propargite	12C
Propetamphos	1B
Propoxur	1A
Prothiofos	1B
Pyflubumide	25B
Pymetrozine	9B
Pyraclofos	1B
Pyrethrins (pyrethrum)	3A
Pyridaben	21A
Pyridalyl	UN
Pyridaphenthion	1B
Pyrifluquinazon	9B
Pyrimidifen	21A
Pyriproxyfen	7C
Quinalphos	1B
Resmethrin	3A
Rotenone (Derris)	21B
Silafluofen	3A
Sodium borate	8D
Sodium cyanide	24B
Sodium metaborate	8D

Spinetoram	5
Spinosad	5
Spirodiclofen	23
Spiromesifen	23
Spirotetramat	23
Sulfotep	1B
Sulfoxaflor	4C
Sulfur	UN
Sulfuramid	13
Sulfuryl fluoride	8C
Tartar emetic	8E
<i>tau</i> -Fluvalinate	3A
Tebufenozide	18
Tebufenpyrad	21A
Tebupirimfos	1B
Teflubenzuron	15
Tefluthrin	3A
Temephos	1B
Terbufos	1B
Tetrachlorvinphos	1B
Tetradifon	12D
Tetramethrin	3A
Tetramethrin [(1 <i>R</i>)-isomers]	3A
<i>theta</i> -cypermethrin	3A
Thiacloprid	4A
Thiamethoxam	4A
Thiocyclam	14
Thiofanox	1A
Thiometon	1B
Thiosultap-sodium	14

Active Ingredients (Alphabetical Order) with MOA Classification

Tolfenpyrad	21A
Tralomethrin	3A
Transfluthrin	3A
Triazamate	1A

Triazophos	1B
Trichlorfon	1B
Triflumuron	15
Triflumezopyrim	4E

Trimethacarb	1A
Vamidothion	1B
XMC	1A
Xylycarb	1A

<i>zeta</i> -Cypermethrin	3A
Zinc phosphide	24A

Photograph
Acknowledgements:

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