

Key to Targeted Physiology

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

Group 1: Acetylcholinesterase (AChE) inhibitors
(Only representative actives of the groups are shown)

1A Carbamates: Carbofuran, Carbosulfen, Methomyl

1B Organophosphates: Acaphate, Chlorpyrifos

Group 2: GABA-gated chloride channel antagonists

2A Cyclo diene Organochlorines: Chlordane, Endosulfan

2B Phenylpyrazoles (Fiproles): Ethiprole, Fipronil

Group 3: Sodium channel modulators (Only representative actives of group 3A are shown)

3A Pyrethroids Pyrethrins: DDT, Methoxychlor, 3B DDT, Methoxychlor

Others: Bifenthrin, Esfenvalerate, Permethrin, Deltamethrin, lambda-cyhalothrin, Etofenprox, Tefluthrin

Group 4: Nicotinic acetylcholine receptor (nAChR) competitive modulators

4A Neonicotinoids: Clothianidin, Imidacloprid, Thiacloprid, Dinotefuran

4B Nicotine

4C Sulfoximines: Sulfoxaflor

4D Butenolides: Flupyradifurone

4E Mesoionics: Triflumazopyrim, Flupyrifumin, Pyridylidene

Group 5: Nicotinic acetylcholine receptor (nAChR) allosteric modulators site I

5 Spinosyns: Spinetoram (R1 = ethyl, R2 = H), Spinosad (R1 = propargyl, R2 = H)

Group 6: Glutamate-gated chloride channel (GluCl) allosteric modulators

6 Avermectins & Milbemycins: Emamectin benzoate (R1 = ...), Milbexiclin

Group 7: Juvenile hormone mimics

7A Juvenile hormone analogues: Hydrogrenone (R1 = ethyl, R2 = H), Methoprene (R1 = isopropyl, R2 = OCH₃), Kiosgrenone (R1 = propargyl, R2 = H)

7B Fenoxycarb, 7C Pyriproxyfen, 7D Pyriproxyfen

Group 8: Miscellaneous non-specific (multi-site) inhibitors

8A Alkyl halides: Methyl bromide

8B Chlorpicrin

8C Fluorides: Sulfuryl fluoride

8D Borates: Borax

8E Tartar emetic: Tartrate

8F Methyl isothiocyanate generators: H₃C-Br, Diazomet

Use of Groups:

- Alternations, sequences or rotations of compounds between MoA groups reduce selection for **target site resistance**.
- Applications are arranged into MoA spray windows defined by crop growth stage and pest biology. Several sprays of a compound may be possible within each spray window, but successive generations of a pest should not be treated with compounds from the same MoA group. Local expert advice on spray windows and timings should always be followed.
- Groups in the classification whose members do not act at a common target site are grouped from the prescription against rotation within the group (Group 8, 13 and all UN groups: UN, UNB, UNE, UNF, UNM, UNP & UNV).

Use of Sub-Groups:

- Sub-groups represent distinct structural classes which are believed to have the same mode of action.
- Sub-groups provide differentiation between compounds that may bind at the same target site but are structurally different enough that risk of **metabolic cross-resistance** is lower than for close chemical analogs.
- Cross-resistance potential between sub-groups is higher than between groups, so rotation between sub-groups should be considered only when there are no alternatives, and only if cross-resistance does not exist, following consultation with local expert advice. These exceptions are not sustainable, and alternative options should be sought.

IRAC

Insecticide Resistance Action Committee Mode of Action Classification

Group 9: Chordotonal organ TRPV channel modulators

9B Pyridine azomethine derivatives: Pymetrozine, Pyflubiquazon

9D Pyropenes: Abdoxproten

Group 10: Mite growth inhibitors affecting CHS1

10A Clofentezine, Diflovidazin, Hexythiazox

10B Etoxazole

Group 11: Microbial disruptors of insect midgut membranes

Includes transgenic crops expressing *Bacillus thuringiensis* toxins (however, specific guidance for resistance management of transgenic crops is not based on rotation of modes of action)

Rotation between certain specific *B.t.* microbial products may provide resistance management benefits for some pests. Consult product-specific recommendations.

11A *Bacillus thuringiensis* (includes *B.t. israelensis*, *B.t. altissalis*, *B.t. kurstaki*, *B.t. terrestris*)

11B *Bacillus sphaericus*

Group 12: Inhibitors of mitochondrial ATP synthase

12A Diafenthurion

12B Organotin miticides: Cyhexatin, Fenbutatin oxide

12C Propargite

12D Tetradifon

Group 13: Uncouplers of oxidative phosphorylation via disruption of proton gradient

13 Pyroles, Dinitrophenols, Sulfuramid: Chlorfentrapryl, DNOC, Sulfuramid

Group 14: Nicotinic acetylcholine receptor (nAChR) channel blockers

14 Nereistoxin analogues: Bensusalt, Thiocyclam

12B Carptap hydrochloride, 12C Thiostapsodium

Group 15: Inhibitors of chitin biosynthesis affecting CHS1
(Only representative actives of group are shown)

15 Benzoylureas: Diflubenzuron, Flufenoxuron, Lufenuron, Novaluron, Teflubenzuron

Group 16: Inhibitors of chitin biosynthesis, type 1

16 Buprofezin

Group 17: Moulting disruptors, Dipteran

17 Cyromazine

Group 18: Ecdysone receptor agonists

18 Diacylhydrazines: Chromanfenozide, Halofenozide, Methoxyfenozide, Tebufenozide

Group 19: Octopamine receptor agonists

19 Amitraz

Group 20: Mitochondrial complex III electron transport inhibitors – Qo site

20A Hydranmethylinon

20B Acequinonyl

20C Fluacrypyrim

20D Bifenazate

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Group 21: Mitochondrial complex I electron transport inhibitors

Fenazaquin, Pyridaben, Pyrimidifen, Tolfenpyrad, Fenpyroximate

21A METI acaricides and insecticides: Rotenone, 21B Rotenone

Group 22: Voltage-dependent sodium channel blockers

Indoxacarb, 22A Oxadiazines

22B Semicarbazones, 22C Metaflumizone

Group 23: Inhibitors of acetyl CoA carboxylase

23 Tetroneic and Tetramic acid derivatives: Spiromesifen, Spirotropin

Group 24: Mitochondrial complex IV electron transport inhibitors

AIP: Aluminum phosphide

Ca₃P₂: Calcium phosphide

PH₃: Phosphine

24A Phosphides: Zinc phosphide

CN⁻: Cyanide salts

24B Cyanides: 24A Zinc phosphide

Group 25: Mitochondrial complex II electron transport inhibitors

25A beta-Ketonitrile derivatives: Cyanoacrylate, 25B Carboxanilides: Cyflumetofen

Group 28: Ryanodine receptor modulators

Chlorantraniliprole R-CI, Cyantraniliprole R-Cy

28 Diamides: Flubendamide

Tetraniliprole

Group 29: Chordotonal organ modulators – undefined target site

29 Flonicamid

Group 30: GABA-gated chloride channel allosteric modulators

30 Meta-diamides & Isoxazolines: Broflanilide, Fluzametanilide, Isoxycloseram

Group 31: Baculoviruses

31 Granuloviruses & Nucleopolyhedroviruses: *Cydia pomonella* GV, *Thaumetobia leucocreta* GV, *Anticarsia gemmatilis* MNPV, *Helicoverpa armigera* NPV

Group 32: Nicotinic acetylcholine receptor (nAChR) allosteric modulators site II

32 GS-omega-kappa HXTX-Hv1a peptide

Group 33: Calcium-activated potassium channel (KCa2) modulators

33 Acynonapyr

Group 34: Mitochondrial complex III electron transport inhibitors – Q1 site

34 Flometoquin

UN: Unknown or uncertain mode of action

UNF: Fungal agents: Beauveria bassiana strains, Metarhizium brunneum strain FS2, Paecilomyces fumosoroseus, Koppica strain 97

UNB: Bacterial agents (non-BI): Burkholderia spp., Wobaschia pipentis (Zap)

UNM: Non-specific mechanical and physical disruptors: Diatomaceous earth, Mineral oil

UNE: Botanical essence including synthetic, extracts and unrefined oils: Chenopodium ambrosioides, neem ambrosioides extract, Fatty acid monomers with glycerol or propyl acetate, Neem oil

Other UN agents: Azadirachtin, Bromopropylate, Benzoximate, Oxazoxyflor, Chinomethionol, Dicofol, Mancozeb, Sulfur, Benzoic acid, Pyridalil

Poster Notes:

- Sub-group 3B: DDT is no longer used in agriculture and therefore this is only applicable for the control of insect vectors of human disease, such as mosquitoes, because of a lack of alternatives.
- Sub-group 10A: Hexythiazox is grouped with Clofentezine because they exhibit cross-resistance even though they are structurally distinct. 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.
- Group 20: While there is strong evidence that Bifenazate acts on the Qo site of Mitochondrial Complex III and some Bifenazate resistance mutations confer cross-resistance to Acequinonyl, the sites of action of Fluacrypyrim and Hydranmethylinon have not been determined.
- Groups 26 and 27 are unassigned.
- In some cases, only representative actives are shown.
- Please visit www.irc-online.org for the complete IRAC classification.

