

FRAC Code List ^{©*}2020: Fungal control agents sorted by cross resistance pattern and mode of action (including FRAC Code numbering)

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INTRODUCTION

The following table lists commercial fungicides, mainly for use in plant protection, according to their mode of action and resistance risk. The most important bactericides are also included. Grouping is considering the biochemical mode of action, but a main driver is to identify cross-resistance patterns between chemistries.

The Table headings are defined as:

MOA Code

Different letters (A to P, with added numbers) are used to distinguish fungicide groups according to their biochemical mode of action (MOA) in the biosynthetic pathways of plant pathogens. The grouping was made according to processes in the metabolism starting from nucleic acids synthesis (A) to secondary metabolism, e.g. melanin synthesis (I), followed by host plant defence inducers (P), recent molecules with an unknown mode of action and unknown resistance risk (U, transient status, until information about mode of action and mechanism of resistance becomes available), and chemical multi-site inhibitors (M). Fungicidal compositions of biological origin are grouped according to the main mode of action within the respective pathway categories. A more recently introduced category "Biologicals with multiple modes of action" (BM) is used for agents from biological origin showing multiple mechanisms of action.

Target Site and Code

If available, the biochemical mode of action is given. In several cases the precise target site may not be known, however, a grouping within a given pathway / functional cluster is still possible. Grouping can also be made due to cross resistance profiles within a group or in relation to other groups.

Group Name

The Group Names listed are based on chemical relatedness of structures which are accepted in literature (e.g. The Pesticide Manual). They are based on different sources (chemical structure, site of action, first important representative in group).

Chemical or Biological Group

Grouping is based on chemical considerations. Nomenclature is according to IUPAC and Chemical Abstract name. Taxonomic information may be used for agents of biological origin.

Common name

BSI/ISO accepted (or proposed) common name for an individual active ingredient expected to appear on the product label as definition of the product.

Comments on Resistance

Details are given for the (molecular) mechanism of resistance and the resistance risk. If field-resistance is known to one member of the Group, it is most likely but not exclusively valid that cross resistance to other group members will be present. There is increasing evidence that the degree of cross resistance can differ between group members and pathogen species or even within species. For the latest information on resistance and cross resistance status of a pathogen / fungicide combination, it is advised to contact local FRAC representatives, product manufacturer's representatives or crop protection advisors. The intrinsic risk for resistance evolution to a given fungicide group is estimated to be **low, medium or high** according to the principles described in FRAC Monographs 1, 2 and 3. Resistance management is driven by intrinsic risk of fungicide, pathogen risk and agronomic risk (see FRAC pathogen risk list).

Similar classification lists of fungicides have been published by T. Locke on behalf of FRAG – UK (Fungicide Resistance, August 2001), and by P. Leroux (Classification des fongicides agricoles et résistance, Phytoma, La Défense des Végétaux, No. 554, 43-51, November 2002).

FRAC Code

Numbers and letters are used to distinguish the fungicide groups according to their cross-resistance behaviour. This code should be used to define the GROUP Number on product labels. The numbers were assigned primarily according to the time of product introduction to the market. The letters refer to P = host plant defence inducers, M = chemical multi-site inhibitors, U = unknown mode of action and unknown resistance risk, and BM = biologicals with multiple modes of action. Reclassification of compounds based on new research may result in codes to expire. This is most likely in the U - section when the mode of actions gets clarified. These codes are not re-used for new groups; a note is added to indicate reclassification into a new code.

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Next update decisions: January 2021

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MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
metabolism	A1 RNA polymerase I	PA – fungicides (PhenylAmides)	acylalanines	benalaxyl benalaxyl-M (=kiralaxyl) furalaxyl metalaxyl metalaxyl-M (=mefenoxam)	Resistance and cross resistance well known in various Oomycetes but mechanism unknown. High risk.	4
tabo			oxazolidinones	oxadixyl	See FRAC Phenylamide Guidelines for resistance management	
met			butyrolactones	ofurace		
leic acids	A2 adenosin- deaminase	hydroxy- (2-amino-) pyrimidines	hydroxy- (2-amino-) pyrimidines	bupirimate dimethirimol ethirimol	Medium risk. Resistance and cross resistance known in powdery mildews. Resistance management required.	8
nucleic	A3 DNA/RNA synthesis	heterogromatics	isoxazoles	hymexazole		32
Ä	(proposed)	neteroaromatics	isothiazolones	octhilinone	Resistance not known.	JZ
	A4 DNA topoisomerase type II (gyrase)	carboxylic acids	carboxylic acids	oxolinic acid	Bactericide. Resistance known. Risk in fungi unknown. Resistance management required.	31

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
		B1 -tubulin assembly in mitosis Benzimidazole Carbamates)	benzimidazoles	benomyl carbendazim fuberidazole thiabendazole	Resistance common in many fungal species. Several target site mutations, mostly E198A/G/K, F200Y in β-tubulin gene.	
	ß-tubulin assembly		thiophanates	thiophanate thiophanate-methyl	Positive cross resistance between the group members. Negative cross resistance to N-phenyl carbamates. High risk. See FRAC Benzimidazole	1
					Guidelines for resistance management.	
r protein	B2 ß-tubulin assembly in mitosis	N-phenyl carbamates	N-phenyl carbamates	diethofencarb	Resistance known. Target site mutation E198K. Negative cross resistance to benzimidazoles. High risk. Resistance management required.	10
noto	B3 ß-tubulin assembly in mitosis	benzamides	toluamides	zoxamide	Low to medium risk.	00
and r		thiazole carboxamide	ethylamino-thiazole- carboxamide	ethaboxam	Resistance management required.	22
celeton	B4 cell division (unknown site)	phenylureas	phenylureas	pencycuron	Resistance not known.	20
B: Cytoskeleton and motor protein	B5 delocalisation of spectrin-like proteins	benzamides	pyridinylmethyl- benzamides	fluopicolide fluopimomide	Resistant isolates detected in grapevine downy mildew. Medium risk. Resistance management required	43
	B6 actin/myosin/fimbrin function	cyanoacrylates	aminocyanoacrylates	phenamacril	Resistance known in <i>Fusarium</i> graminearum. Target site mutations in the gene coding for myosin-5 found in lab studies. Medium to high risk. Resistance management required.	47
		tin/myosin/fimbrin	benzophenone	metrafenone	Less sensitive isolates detected in powdery mildews (<i>Blumeria</i> and <i>Sphaerotheca</i>)	
			ketones	pyriofenone	Medium risk. Resistance management required.	50
					Reclassified from U8 in 2018	

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
	C1 complex I NADH oxido-reductase	pyrimidinamines	pyrimidinamines	diflumetorim		
		pyrazole-MET1	pyrazole-5- carboxamides	tolfenpyrad	Resistance not known.	39
	UXIOU-TEORCIASE	Quinazoline	quinazoline	fenazaquin		
			phenyl-benzamides	benodanil flutolanil mepronil		
			phenyl-oxo-ethyl thiophene amide	isofetamid		
			pyridinyl-ethyl- benzamides	fluopyram		
			furan- carboxamides	fenfuram		
_		C2 complex II: ccinate-dehydro- genase SDHI (Succinate- dehydrogenase inhibitors)	oxathiin- carboxamides	carboxin oxycarboxin	Resistance known for several fungal species in field populations and lab mutants. Target site mutations in sdh gene, e.g. H/Y (or H/L) at 257, 267, 272 or P225L, dependent on fungal species. Resistance management required. Medium to high risk. See FRAC SDHI Guidelines	
atior	complex II: succinate-dehydro-		thiazole- carboxamides	thifluzamide		
C. respiration			pyrazole-4- carboxamides	benzovindiflupyr bixafen fluindapyr fluxapyroxad furametpyr inpyrfluxam isopyrazam penflufen penthiopyrad sedaxane		7
			N-cyclopropyl-N- benzyl-pyrazole- carboxamides	isoflucypram	for resistance management.	
			N-methoxy-(phenyl- ethyl)-pyrazole- carboxamides	pydiflumetofen	-	
			pyridine- carboxamides	boscalid		
			pyrazine- carboxamides	pyraziflumid		

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
		Qol -fungicides	methoxy-acrylates methoxy-acetamide methoxy-carbamates	-acetamide mandestrobin pyraoxystrobin pyraoxystrobin -acetamide mandestrobin pyraclostrobin triclopyricarb	Resistance known in various fungal species. Target site mutations in cyt b gene (G143A, F129L) and additional mechanisms. Cross resistance shown	11
respiration	C3 complex III: cytochrome bc1	(Quinone outside Inhibitors)	oximino-acetates	trifloxystrobin dimoxystrobin fenaminstrobin	between all members of the Code 11 fungicides. High risk. See FRAC Qol Guidelines for resistance management.	
espi	(ubiquinol oxidase) at Qo site (cyt b		oxazolidine-diones	metominostrobin orysastrobin famoxadone		
	gene)		dihydro-dioxazines	fluoxastrobin		
0			imidazolinones	fenamidone		
			benzyl-carbamates	pyribencarb		
		Qol-fungicides (Quinone outside Inhibitors; Subgroup A)	tetrazolinones	metyltetraprole	Resistance not known. Not cross resistant with Code 11 fungicides on G143A mutants. High risk.	11A
					See FRAC Qol Guidelines for resistance management.	

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
	C4		cyano-imidazole	cyazofamid	Resistance risk unknown but assumed to be medium to high (mutations at target site known	
	complex III: cytochrome bc1 (ubiquinone		sulfamoyl-triazole	amisulbrom	in model organisms). Resistance management required.	21
	reductase) at Qi site		picolinamides	fenpicoxamid	No spectrum overlap with Oomycete fungicides cyazofamid and amisulbrom	
(pən	C5		dinitrophenyl- crotonates	binapacryl meptyldinocap dinocap	Resistance not known. Also acaricidal activity.	
contin	uncouplers of oxidative phos- phorylation		2,6-dinitro-anilines	fluazinam	Low risk. However, resistance claimed in <i>Botrytis</i> in Japan.	29
) uc			(pyrhydrazones)	(ferimzone)	Reclassified to U 14 in 2012.	
C: respiration (continued)	C6 inhibitors of oxidative phos- phorylation, ATP synthase	organo tin compounds	tri-phenyl tin compounds	fentin acetate fentin chloride fentin hydroxide	Some resistance cases known. Low to medium risk.	30
0	C7	thiophene-	thiophene-	silthiofam	Resistance reported. Risk low.	38
	ATP transport (proposed)	carboxamides	carboxamides	Sittioidin	Resistance reported. Risk low.	30
	C8 complex III: cytochrome bc1 (ubiquinone reductase) at Qo site, stigmatellin binding sub-site	QoSI fungicides (Quinone outside Inhibitor, stigmatellin binding type)	triazolo-pyrimidylamine	ametoctradin	Not cross resistant to Qol fungicides. Resistance risk assumed to be medium to high (single site inhibitor). Resistance management required.	45
protein synthesis	D1 methionine biosynthesis (proposed) (cgs gene)	AP - fungicides (Anilino- Pyrimidines)	anilino-pyrimidines	cyprodinil mepanipyrim pyrimethanil	Resistance known in <i>Botrytis</i> and <i>Venturia</i> , sporadically in <i>Oculimacula</i> . Medium risk. See FRAC Anilinopyrimidine Guidelines for resistance management.	9
protein s	D2 protein synthesis (ribosome, termination step)	enopyranuronic acid antibiotic	enopyranuronic acid antibiotic	blasticidin-S	Low to medium risk. Resistance management required.	23
amino acids and	D3 protein synthesis (ribosome, initiation step)	hexopyranosyl antibiotic	hexopyranosyl antibiotic	kasugamycin	Resistance known in fungal and bacterial (<i>P. glumae</i>) pathogens. Medium risk. Resistance management required.	24
D: amino	D4 protein synthesis (ribosome, initiation step)	glucopyranosyl antibiotic	glucopyranosyl antibiotic	streptomycin	Bactericide. Resistance known. High risk. Resistance management required.	25
	D5 protein synthesis (ribosome, elongation step)	tetracycline antibiotic	tetracycline antibiotic	oxytetracycline	Bactericide. Resistance known. High risk. Resistance management required.	41

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
	E1	aza- naphthalenes	aryloxyquinoline	quinoxyfen	Resistance to quinoxyfen known. Medium risk.	
ч	signal transduction (mechanism unknown)		quinazolinone	proquinazid	Resistance management required. Cross resistance found in <i>Erysiphe (Uncinula)</i> <i>necator</i> but not in <i>Blumeria</i> <i>graminis</i> .	13
signal transduction	E2 MAP/Histidine- Kinase in osmotic signal transduction (os-2, HOG1)	PP-fungicides (PhenylPyrroles)	phenylpyrroles	fenpiclonil fludioxonil	Resistance found sporadically, mechanism speculative. Low to medium risk. Resistance management required.	12
E: signal tr	E3 MAP/Histidine- Kinase in osmotic signal transduction (os-1, Daf1)	dicarboximides	dicarboximides	chlozolinate dimethachlone iprodione procymidone vinclozolin	Resistance common in <i>Botrytis</i> and some other pathogens. Several mutations in OS-1, mostly I365S. Cross resistance common between the group members. Medium to high risk. See FRAC Dicarboximide Guidelines for resistance management.	2

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE		
	F1		forme	rly dicarboximides				
	F2 phospholipid	phosphoro- thiolates	phosphoro-thiolates	edifenphos iprobenfos (IBP) pyrazophos	Resistance known in specific fungi. Low to medium risk.	6		
_	biosynthesis, methyltransferase	Dithiolanes	dithiolanes	isoprothiolane	Resistance management required if used for risky pathogens.			
synthesis or transport / membrane integrity or function	F3 cell peroxidation (proposed)	AH-fungicides (Aromatic Hydrocarbons) (chlorophenyls, nitroanilines)	aromatic hydrocarbons	biphenyl chloroneb dicloran quintozene (PCNB) tecnazene (TCNB) tolclofos-methyl	Resistance known in some fungi. Low to medium risk. Cross resistance patterns complex due to different	14		
ntegrit		heteroaromatics	1,2,4-thiadiazoles	etridiazole	activity spectra.			
eir	F4			is de servis	Leve to meeting with			
embran	cell membrane permeability, fatty acids (proposed)	Carbamates	carbamates	iodocarb propamocarb prothiocarb	Low to medium risk. Resistance management required.	28		
_ m	F5		formerly CAA-fungicides					
r transport	F6 microbial disrupters of pathogen cell membranes	f	formerly <i>Bacillus amyloliquefaciens</i> strains (FRAC Code 44); reclassified to BM02 in 2020					
id synthesis o	F7 cell membrane disruption	plant extract	terpene hydrocarbons, terpene alcohols and terpene phenols	extract from Melaleuca alternifolia (tea tree) plant oils (mixtures): eugenol, geraniol, thymol	Resistance not known.	46		
F: lipid	F8 ergosterol binding	Polyene	amphoteric macrolide antifungal antibiotic from <i>Streptomyces</i> <i>natalensis</i> or <i>S. chattanoogensis</i>	natamycin (pimaricin)	Resistance not known. Agricultural, food and topical medical uses.	48		
	F9 lipid homeostasis and transfer/storage	OSBPI oxysterol binding protein homologue inhibition	¥	oxathiapiprolin fluoxapiprolin	Resistance risk assumed to be medium to high (single site inhibitor). Resistance management required. (Previously U15).	49		

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
			piperazines	triforine pyrifenox		
			pyridines	pyrisoxazole		
				fenarimol		
			pyrimidines	nuarimol	-	
				imazalil	There are big differences in	
				oxpoconazole	the activity spectra of DMI fungicides.	
			imidazoles	pefurazoate prochloraz	iungiciues.	
				triflumizole	Resistance is known in various	
				azaconazole	fungal species. Several	
				bitertanol	resistance mechanisms are	
				bromuconazole	known incl. target site mutations in cyp51 (erg 11)	
				cyproconazole	gene, e.g. V136A, Y137F,	
				difenoconazole	A379G, I381V; cyp51	
	G1	DMI-fungicides		diniconazole epoxiconazole	promotor; ABC transporters	
		(DeM ethylation		etaconazole	and others.	
	C14- demethylase	Inhibitors)		fenbuconazole		3
	in sterol	,		fluquinconazole	Generally wise to accept that cross resistance is present	
S	biosynthesis (erg11/cyp51)	(SBI: Class I)		flusilazole	between DMI fungicides active	
biosynthesis in membranes	(eiginoypor)			flutriafol	against the same fungus.	
raı			triazoles	hexaconazole imibenconazole		
qu				ipconazole	DMI fungicides are Sterol	
ler				mefentrifluconazole	Biosynthesis Inhibitors (SBIs),	
E				metconazole	but show no cross resistance to other SBI classes.	
Ľ.				myclobutanil	to other SDI classes.	
sis				penconazole	Medium risk.	
Je:				propiconazole simeconazole		
ntl				tebuconazole	See FRAC SBI Guidelines	
sy				tetraconazole	for resistance management.	
io				triadimefon		
				triadimenol		
erc			triazolinthiones	triticonazole		
G: sterol			utazoiinutiiones	prothioconazole aldimorph	Decreased consitivity for	
G	G2			dodemorph	Decreased sensitivity for powdery mildews. Cross resistance within the	
	. 14		morpholines	fenpropimorph		
	Δ^{14} -reductase	amines		tridemorph	group generally found but not	
	and $\Delta^8 \rightarrow \Delta^{7-}$	("morpholines")		fenpropidin	to other	5
	isomerase	(SBI: Class II)	piperidines	piperalin	SBI classes.	v
	in sterol	(SDI. Class II)			Low to medium risk.	
	biosynthesis		spiroketal-amines	spiroxamine	See FRAC SBI Guidelines	
	(erg24, erg2)		,		for resistance management.	
	G3	KRI fungicides (KetoReductase	hydroxyanilides	fenhexamid	Low to medium risk.	
	3-keto reductase,	Inhibitors)			Resistance management	17
	C4- de-methylation		amino-pyrazolinone	fennyrazamino	required.	17
	(erg27)	(SBI: Class III)	аншю-ругадошноне	fenpyrazamine		
	G4	,	thiocarbamates	pyributicarb	Resistance not known, fungicidal and herbicidal	
	auglono onovidore		1110001001110100	Pyributiodia	activity.	
	squalene-epoxidase in sterol	(SBI class IV)			· · ·	18
	biosynthesis		allylamines	naftifine	Medical fungicides only.	
	(erg1)		anylaninoo	terbinafine	moaroar langiolado only.	

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
s	H3		Formerly glucopyranos antibiotic (validamycin		reclassified to U18	26
H: cell wall biosynthesis	H4 chitin synthase	polyoxins	peptidyl pyrimidine nucleoside	polyoxin	Resistance known. Medium risk. Resistance management required.	19
wall bid	Н5	CAA-fungicides	cinnamic acid amides	dimethomorph flumorph pyrimorph	Resistance known in Plasmopara viticola but not in Phytophthora infestans.	
H: cell	(C arb	(Carboxylic Acid Amides)	valinamide carbamates	benthiavalicarb iprovalicarb valifenalate	Cross resistance between all members of the CAA group. Low to medium risk. See FRAC CAA Guidelines for resistance management.	40
			mandelic acid amides	mandipropamid		
_	11	MBI-R	isobenzo-furanone	fthalide	Resistance not known.	
wal	reductase in	(Melanin Biosynthesis	pyrrolo-quinolinone	pyroquilon		16.1
cell	melanin biosynthesis	Inhibitors – R eductase)	triazolobenzo- thiazole	tricyclazole		
is in	12	MBI-D	cyclopropane- carboxamide	carpropamid	Resistance known.	
thes	dehydratase in	(Melanin Biosynthesis Inhibitors –	carboxamide	diclocymet	Medium risk. Resistance management	16.2
syn	melanin biosynthesis	Dehydratase)	propionamide	fenoxanil	required.	
I: melanin synthesis in cell wall	I3 polyketide synthase in melanin biosynthesis	MBI-P (Melanin Biosynthesis Inhibitors – Polyketide synthase)	trifluoroethyl- carbamate	tolprocarb	Resistance not known. Additional activity against bacteria and fungi through induction of host plant defence	16.3

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
	P 1 salicylate-related	benzo- thiadiazole (BTH)	benzo-thiadiazole (BTH)	acibenzolar-S-methyl	Resistance not known.	P 01
- -	P 2 salicylate-related	benzisothiazole	benzisothiazole	probenazole (also antibacterial and antifungal activity)	Resistance not known.	P 02
duction	P 3 salicylate-related	thiadiazole- carboxamide	thiadiazole- carboxamide	tiadinil isotianil	Resistance not known.	P 03
ence in	P 4 polysaccharide elicitors	natural compound	polysaccharides	laminarin	Resistance not known.	P 04
host plant defence induction	P 5 anthraquinone elicitors	plant extract	complex mixture, ethanol extract (anthraquinones, resveratrol)	extract from <i>Reynoutria</i> <i>sachalinensis</i> (giant knotweed)	Resistance not known.	P 05
ost			bacterial Bacillus spp.	Bacillus mycoides isolate J		
Р: Ч	P 6 microbial elicitors	P 6 microbial	fungal Saccharomyces spp.	cell walls of Saccharomyces cerevisiae strain LAS117	Resistance not known.	P 06
	Р7		ethyl phosphonates	fosetyl-Al	Few resistance cases reported in few pathogens.	P 07
	phosphonates	phosphonates		phosphorous acid and salts	Low risk. Reclassified from U33 in 2018	(33)

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
	unknown	cyanoacetamide- oxime	cyanoacetamide- oxime	cymoxanil	Resistance claims described. Low to medium risk. Resistance management required.	27
		formerly phosp	honates (FRAC code 33	3), reclassified to P (07 in 2018	
	unknown	phthalamic acids	phthalamic acids	teclofthalam (Bactericide)	Resistance not known.	34
ides)	unknown	benzotriazines	benzotriazines	triazoxide	Resistance not known.	35
d fungic	unknown	benzene- sulfonamides	benzene- sulphonamides	flusulfamide	Resistance not known.	36
ssified	unknown	pyridazinones	pyridazinones	diclomezine	Resistance not known.	37
i on eclas		formerly methas	sulfocarb (FRAC code 4	2), reclassified to M	12 in 2018	
e of act ive from r	unknown	phenyl- acetamide	phenyl-acetamide	cyflufenamid	Resistance in <i>Sphaerotheca</i> . Resistance management required	U 06
U: Unknown mode of action appearing in the list derive from reclassified fungicides)	cell membrane disruption (proposed)	guanidines	guanidines	dodine	Resistance known in Venturia inaequalis. Low to medium risk. Resistance management recommended.	U 12
J: Unk l pearing	unknown	thiazolidine	cyano-methylene- thiazolidines	flutianil	Resistance in <i>Sphaerotheca</i> . Resistance management required	U 13
l not ap	unknown	pyrimidinone- hydrazones	pyrimidinone- hydrazones	ferimzone	Resistance not known (previously C5).	U 14
(U numbers not	complex III: cytochrome bc1, unknown binding site (proposed)	4-quinolyl- acetate	4-quinolyl-acetates	tebufloquin	Not cross resistant to Qol. Resistance risk unknown but assumed to be medium. Resistance management required.	U 16
	Unknown	tetrazolyloxime	tetrazolyloximes	picarbutrazox	Resistance not known. Not cross resistant to PA, QoI, CAA.	U 17
	Unknown (Inhibition of trehalase)	glucopyranosyl antibiotic	glucopyranosyl antibiotics	validamycin	Resistance not known. Induction of host plant defense by trehalose proposed (previously H3).	U 18

MOA	TARGET SITE AND CODE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
NC: not clas- si- fied	unknown	diverse	diverse	mineral oils, organic oils, inorganic salts, material of biological origin	Resistance not known.	NC
		inorganic (electrophiles)	inorganic	copper (different salts)	Also applies to organic copper complexes	M 01
		inorganic (electrophiles)	inorganic	sulphur		M 02
		dithiocarbamates and relatives (electrophiles)	dithio-carbamates and relatives	ferbam mancozeb maneb metiram propineb thiram zinc thiazole zineb ziram		M 03
activity		phthalimides (electrophiles)	phthalimides	captan captafol folpet		M 04
nulti-site	and the star	chloronitriles (phthalonitriles) (unspecified mechanism)	chloronitriles (phthalonitriles)	chlorothalonil	generally considered as a low risk group without any signs of resistance developing to the	M 05
rith m	multi-site contact activity	sulfamides (electrophiles)	sulfamides	dichlofluanid tolylfluanid	fungicides.	M 06
Chemicals with multi-site activity		bis-guanidines (membrane disruptors, detergents)	bis-guanidines	guazatine iminoctadine		M 07
M: Che		triazines (unspecified mechanism)	triazines	anilazine		M 08
		quinones (anthraquinones) (electrophiles)	quinones (anthraquinones)	dithianon		M 09
		quinoxalines (electrophiles)	quinoxalines	chinomethionat / quinomethionate		M 10
		maleimide (electrophiles)	maleimide	fluoroimide		M 11
		thiocarbamate (electrophiles)	thiocarbamate	methasulfocarb	reclassified from U42 in 2018	M 12

MOA	TARGET SITE	GROUP NAME	CHEMICAL OR BIOLOGICAL GROUP	COMMON NAME	COMMENTS	FRAC CODE
BM: Biologicals with multiple modes of action	multiple effects on cell wall, ion membrane transporters; chelating effects	plant extract	polypeptide (lectin)	extract from the cotyledons of lupine plantlets ("BLAD")	Resistance not known (previously M12).	BM 01
	affects fungal spores and germ tubes, induced plant defence	plant extract	Phenols, Sesquiterpenes, Triterpenoids, Coumarins	extract from Swinglea glutinosa	Resistance not known	
	multiple effects described (examples, not all apply to all biological groups): competition, mycoparasitism, antibiosis, membrane disruption by fungicidal lipopeptides, lytic enzymes, induced plant defence	microbial (living microbes or extract, metabolites)	fungal <i>Trichoderma</i> spp.	Trichoderma atroviride strain I-1237 Trichoderma atroviride strain LU132 Trichoderma atroviride strain SC1 Trichoderma asperellum strain T34 Gliocladium	Resistance not known	
			fungal <i>Clonostachys</i> spp.	catenulatum strain J1446 Clonostachys rosea strain CR-7		
			bacterial <i>Bacillus</i> spp.	Bacillus amyloliquefaciens strain QST713 strain FZB24 strain MBI600 strain D747 strain F727 Bacillus subtilis strain AFS032321	synonyms for Bacillus amyloliquefaciens are Bacillus subtilis and B. subtilis var. amyloliquefaciens (previous taxonomic classification). Bacillus amyloliquefaciens reclassified from F6, Code 44 in 2020	BM 02
			bacterial Pseudomonas spp.	Pseudomonas chlororaphis strain AFS009		
			bacterial Streptomyces spp.	Streptomyces griseovirides strain K61 Streptomyces lydicus strain WYEC108		