A method of controlling phytopathogenic diseases on useful plants or on propagation material thereof, which comprises applying to the useful plants, the locus thereof or propagation material thereof a combination of components A) and B) in a synergistically effective amount, wherein component A) is a compound of formula (I), wherein R₁ is difluoromethyl or trifluoromethyl and R₂ is C₁-C₆ alkyl, C₁-C₄ alkoxy-C₁-C₆ alkyl or C₁-C₆ haloalkyl; or a tautomer of such a compound; and component B) is a compound selected from compounds known for their fungicidal and/or insecticidal activity, is particularly effective in controlling or preventing fungal diseases of useful plants.
(74) Agent: FETHERSTONHAUGH & CO.
Title: Fungicidal Compositions

Abstract: A method of controlling phytopathogenic diseases on useful plants or on propagation material thereof, which comprises applying to the useful plants, the locus thereof or propagation material thereof a combination of components A) and B) in a synergistically effective amount, wherein component A) is a compound of formula (I), wherein R₁ is difluoromethyl or trifluoromethyl and R₂ is C₃₋₅ alkyl, C₃₋₅ alkoxy-C₁₋₅ alkyl or C₃₋₅ haloalkyl; or a tautomer of such a compound; and component B) is a compound selected from compounds known for their fungicidal and/or insecticidal activity, is particularly effective in controlling or preventing fungal diseases of useful plants.
FUNGICIDAL COMPOSITIONS

The present invention relates to novel fungicidal compositions for the treatment of phytopathogenic diseases of useful plants, especially phytopathogenic fungi, and to a method of controlling phytopathogenic diseases on useful plants.

It is known from WO 04/058723 that certain diphenyl carboxamide derivatives have biological activity against phytopathogenic fungi. On the other hand various fungicidal compounds of different chemical classes are widely known as plant fungicides for application in various crops of cultivated plants. However, crop tolerance and activity against phytopathogenic plant fungi do not always satisfy the needs of agricultural practice in many incidents and aspects.

There is therefore proposed in accordance with the present invention a method of controlling phytopathogenic diseases on useful plants or on propagation material thereof, which comprises applying to the useful plants, the locus thereof or propagation material thereof a combination of components A) and B) in a synergistically effective amount, wherein component A) is a compound of formula I

![Chemical structure](image)

wherein

- $R_1$ is difluoromethyl or trifluoromethyl and
- $R_2$ is $C_1$-$C_6$alkyl, $C_1$-$C_4$alkoxy-$C_1$-$C_6$alkyl or $C_1$-$C_6$haloalkyl; or a tautomer of such a compound; and

component B) is a compound selected from the group consisting of
a strobilurin fungicide, such as Azoxystrobin (47), Dimoxystrobin (226), Fluoxastrobin (382), Kresoxim-methyl (485), Metominostrobin (551), Oryxastrobinn, Picoxystrobin (647), Pyraclostrobin (690); Trifloxystrobin (832) and a compound of formula B-6

\[
\begin{align*}
\text{B-6};
\end{align*}
\]

an azole fungicide, such as Azaconazole (40), Bromuconazole (96), Cyproconazole (207), Difenoconazole (247), Diniconazole (267), Diniconazole-M (267), Epoxiconazole (298), Fenbuconazole (329), Fluquinconazole (385), Flusilazole (393), Flutriafol (397), Hexaconazole (435), Imazalil (449), Imibenconazole (457), Ipconazole (468), Metconazole (525), Myclobutanil (564), Oxpoconazole (607), Pefurazolate (618), Penconazole (619), Prochloraz (659), Propiconazole (675), Prothioconazole (685), Simeconazole (731), Tebuconazole (761), Tetraconazole (778), Triadimefon (814), Triadimenol (815), Triflumizole (834) Trifluzoxide (842), Diclobutrazol (1068), Etaconazole (1129), Furconazole (1198), Furconazole-cis (1199) and Quinconazole (1378);

a phenyl pyrrole fungicide, such as Fenpiclonil (341) and Fludioxonil (368);
an anilino-pyrimidine fungicide, such as Cyprodinil (208), Mepanipyrim (508) and Pyrimethanil (705);
a morpholine fungicide, such as Aldimorph, Dodemorph (288), Fenpropimorph (344), Tridemorph (830), Fenpropidin (343), Spiroxamine (740); Piperalin (648) and a compound of formula B-7

\[
\begin{align*}
\text{B-7};
\end{align*}
\]
a compound of formula F-1

(F-2);

a racemic compound of formula F-3 (syn)

(F-3);

a racemic compound of formula F-4 (anti)
a compound of formula F-5

which represents an epimeric mixture of the racemic compounds of formula F-3 (syn) and F-4 (anti), wherein the ratio of racemic compounds of formula F-3 (syn) to racemic compounds of formula F-4 (anti) is from 1000 : 1 to 1 : 1000;

a compound of formula F-6

a racemic compound of formula F-7 (trans)
a racemic compound of formula F-8 (cis)

which represents a mixture of racemic compounds of formula F-7 (trans) and F-8 (cis), wherein the ratio of racemic compounds of formula F-7 (trans) to racemic compounds of formula F-8 (cis) is from 2 : 1 to 100 : 1;
(796); Bupirimate (98); Dimethirimol (1082); Ethirimol (1133); Diethofencarb (245); Famoxadone (322); Fenamidone (325); Quinoxyfen (715); Biphenyl (81); Chloroneb (139); Dicloran (240); Etridiazole (321); Quintozene (PCNB) (716); Tecnazene (TCNB) (767); Tolclofos-methyl (808); Dimethomorph (263); Carproamid (122); Dioclocymet (237); Fenoxanil (338); Fthalide (643); Pyroquilon (710); Tricyclazole (828); Fenhexamid (334); Polyoxin (654); Pencycuron (620); Cyazofamid (185); Zoxamide (857); Blasticidin-S (85); Kasugamycin (483); Streptomyacin (744); Validamycin (846); Cymoxanil (200); Iodocarb (3-Iodo-2-propynyl butyl carbamate); Propamocarb (668); Prothiocarb (1361); Dinocap (270); Fluazinam (363); Fentin acetate (347); Fentin chloride; Fentin hydroxide (347); Oxolinic acid (606); Hymexazolene; Octhilinone (590); Fosetyl-Aluminium (407); Phosphoric acid; Tecloftalam; Triazoxide (821); Flusulfamide (394); Ferimzone (351); Diclomezine (239); Anilazine (878); Arsenates; Captan (113); Captan (114); Chlorothalonil (142); Copper (diverse salts); Copper Ammoniumcarbonate; Copper octanoate (170); Copper oleate; Copper sulphate (87; 172; 173); Copper hydroxide (169); Dichlofluanid (230); Dithianon (279); Dodine (289); Ferbam (350); Folpet (400); Guazatine (422); Iminoctadine (459); Mancozeb (496); Maneb (497); Mercury; Metiram (546); Propineb (676); Sulphur (754); Thiram (804); Tolyfluanid (810); Zineb (855); Ziram (856); Acibenzolar-S-methyl (6); Probenazole (658); Bentiavalcarb; Bentiavalcarb-isopropyl (68); Iprovalicarb (471); Diflumetorm (253); Ethaboxam (304); Flusulfamide (394); Methasulfoxcarb (528); Silthiofam (729); Bacillus pumilus GB34; Bacillus pumilus strain QST 2808; Bacillus subtilis (50); Bacillus subtilis + PCNB + Metalaxyl (50; 716; 516); Cadmium chloride; Carbon disulfide (945); Bordeaux mixture (87); Cedar leaf oil; Chlorine; Cinnamaldehyde; Cycloheximide (1022); Fenamnosul 1114; Fenamiphos (326); Dichloropropene (233); Dichlone (1052); Formaldehyde (404); Gliocladium virens GL-21 (417); Glyodin (1205); Hexachlorobenzene (434); Iprovalicarb (471); Manganese dimethylthiocarbamate; Mercuric chloride (511); Nabam (566); Neem oil (hydrophobic extract); Oxytetracycline (611); Chinomethionat (126); Paraformaldehyde; Pentachloronitrobenzene (716); Pentachlorphenol (623); paraffin oil (628); Polyoxyin D zinc salt (654); Sodium bicarbonate; Potassium bicarbonate; Sodium diacetate; Sodium propionate; TCMTB; Benalaxyl-M; Boscalid (88); Hexaconazole (435); Metrafenone; Oxine Copper (605); Penthiopeprad; Perfurazoate; Tolyfluanid; Trichoderma harzianum (825); Triphenyltin hydroxide (347); Xanthomonas campestris (852); Paclotrustrozol (612); 1,1-bis(4-chlorophenyl)-2-ethoxyethanol (IUPAC-Name) (910); 2,4-dichlorophenyl benzenesulfonate (IUPAC- / Chemical Abstracts-Name) (1059); 2-fluoro-N-methyl-N-1-naphthylacetamide (IUPAC-Name) (1295); 4-chlorophenyl phenyl sulfone
(IUPAC-Name) (981); abamectin (1); acequinocyl (3); acetoprole [CCN]; acrinathrin (9); aldicarb (16); aldoxycarb (863); alpha-cypermethrin (202); amidithion (870); amidoflumet [CCN]; amidothioate (872); amiton (875); amiton hydrogen oxalate (875); amitraz (24); aramite (881); arsenous oxide (882); AVI 382 (compound code); AZ 60541 (compound code); azinphos-ethyl (44); azinphos-methyl (45); azobenzene (IUPAC-Name) (888); azocyclotin (46); azothoate (889); benomyl (62); benoxafos (alternative name) [CCN]; benzoxyimate (71); benzyl benzoate (IUPAC-Name) [CCN]; bifenazate (74); bifenthrin (76); binapacryl (907); brofenvalerate (alternative name); bromocyclen (918); bromophos (920); bromophos-ethyl (921); bromopropylate (94); buprofezin (99); butocarboxim (103); butoxycarboxim (104); butylypyridaben (alternative name); calcium polysulfide (IUPAC-Name) (111); camphechlor (941); carbonolate (943); carbaryl (115); carbofuran (118); carbophenothion (947); CGA 50'439 (development code) (125); chinomethionat (126); chlorbenside (959); chlordimeform (964); chlordimeform hydrochloride (964); chlorfenapyr (130); chlorfenethol (968); chlorfenson (970); chlorfensulphide (971); chlorfenvinphos (131); chlorobenzilate (975); chloromebuform (977); chloromethion (978); chloropropylate (983); chlorpyrifos (145); chlorpyrifos-methyl (146); chlorthiophos (994); cinerin I (696); cinerin II (696); cinerins (696); clofentezine (156); closantel (alternative name) [CCN]; coumaphos (174); crotamiton (alternative name) [CCN]; crotaxyphos (1010); cufraneb (1013); cyanthoate (1020); cyhalothrin (196); cyhexatin (199); cypermethrin (201); DCPM (1032); DDT (219); demephion (1037); demephion-O (1037); demephion-S (1037); demeton (1038); demeton-methyl (224); demeton-O (1038); demeton-O-methyl (224); demeton-S (1038); demeton-S-methyl (224); demeton-S-methylsulphon (1039); diafenthiuron (226); dialifos (1042); diazinon (227); dichlofluanid (230); dichlorvos (236); dicliphos (alternative name); dicofol (242); dicropothos (243); dienochlor (1071); dimefox (1081); dimethoate (262); dinactin (alternative name) (653); dinex (1089); dinex-diclexine (1089); dinobuton (269); dinocap (270); dinocap-4 [CCN]; dinocap-6 [CCN]; dinocton (1090); dinopentox (1092); dinosulfon (1097); dinoterbon (1098); dioxathion (1102); diphenyl sulfone (IUPAC-Name) (1103); disulfiram (alternative name) [CCN]; disulfoton (278); DNOC (282); dofenapyn (1113); doramectin (alternative name) [CCN]; endosulfan (294); endothion (1121); EPN (297); eprinomectin (alternative name) [CCN]; ethion (309); ethoate-methyl (1134); etoxazole (320); etrimfos (1142); fenazaflor (1147); fenazaquin (328); fenbutatin oxide (330); fenothiocarb (337); fenprofprathrin (342); fenpyrad (alternative name); fenpyroximate (345); fenuron (1157); fentriflanil (1161); fenvalerate (349); fipronil (354); fluacrypyrim (360); fluazuron (1166); flubenzimine (1167); flucloxuron (366); flucytrilate (367); fluenetil
(1169); flufenoxuron (370); flumethrin (372); fluorbenside (1174); fluvalinate (1184); FMC 1137 (development code) (1185); formetanate (405); formetanate hydrochloride (405); formothion (1192); formparanate (1193); gamma-HCH (430); glyodin (1205); halifenprox (424); heptenophos (432); hexadecyl cyclopropanecarboxylate (IUPAC- / Chemical Abstracts-Name) (1216); hexythiazox (441); iodomethane (IUPAC-Name) (542); isocarbox- phos (alternative name) (473); isopropyl O-(methoxyaminothiophosphoryl)salicylate (IUPAC-Name) (473); ivermectin (alternative name) [CCN]; jasmolin I (696); jasmolin II (696); jodfenphos (1248); lindane (430); lufenuron (490); malathion (492); malonobufen (1254); mecarbam (502); mephosfolan (1261); mesulfen (alternative name) [CCN]; meth-carfipos (1266); methamidophos (527); methidathion (529); methiocarb (530); methomyl (531); methyl bromide (537); metolcarb (550); mevinphos (556); mexacarbate (1290); milbemectin (557); milbemycin oxime (alternative name) [CCN]; mipafox (1293); monocro- tophos (561); morphothion (1300); moxidectin (alternative name) [CCN]; naled (567); NC-184 (compound code); nifurturidide (1309); nikkomycins (alternative name) [CCN]; nitrilacarb (1313); nitrilacarb 1:1 zinc chloride complex (1313); NNI-0101 (compound code); NNI-0250 (compound code); omethoate (594); oxamyl (602); oxydeprofos (1324); oxydisulfoton (1325); pp'-DDT (219); parathion (615); permethrin (626); petroleum oils (alternative name) (628); phenkaptan (1330); phenothoate (631); phorate (636); phosphalon (637); phosfolan (1338); phosmet (638); phosphamidon (639); phoxim (642); pirimiphos-methyl (652); polychloro- terpenes (traditional name) (1347); polyactins (alternative name) (653); proclonal (1350); profenofos (662); promacyl (1354); propargite (671); propetamphos (673); propoxur (678); prothidathion (1360); prothoate (1362); pyrethrin I (696); pyrethrin II (696); pyrethrins (696); pyridaben (699); pyridaphenthion (701); pyrimidifen (706); pyrimicarb (1370); quinalphos (711); quintofos (1381); R-1492 (development code) (1382); RA-17 (development code) (1383); rotenone (722); schradan (1389); sebufos (alternative name); selamectin (alternative name) [CCN]; SI-0009 (compound code); sophamide (1402); spirodiclofen (738); spiromesifen (739); SSI-121 (development code) (1404); sulfiram (alternative name) [CCN]; sulfiramid (750); sulfotep (753); sulfur (754); SUI-121 (development code) (757); tau-fluvalinate (398); tebufenpyrad (763); TEPP (1417); terbarm (alternative name); tetrachlorvinphos (777); tetradifon (786); tetranactin (alternative name) (653); tetrasul (1425); thiafenox (alternative name); thiocarboxime (1431); thiocarvoxan (800); thiometon (801); thioquinox (1436); thuringiensin (alternative name) [CCN]; triamiphos (1441); triarathene (1443); triazophos (820); triazuron (alternative name); trichlorfon (824); triflumuron (1455); trinaclactin (alternative name) (653); vanidothion (847); vaniliprole [CCN];
YI-5302 (compound code); bethoxazin [CCN]; copper dioctanoate (IUPAC-Name) (170); copper sulfate (172); cybutryne [CCN]; dichlorone (1052); dichlorophen (232); endothal (295); fentin (347); hydrated lime [CCN]; nabam (566); quincloxamine (714); quinonamid (1379); simazine (730); triphenyltin acetate (IUPAC-Name) (347); triphenyltin hydroxide (IUPAC-Name) (347); abamectin (1); crufomate (1011); doramectin (alternative name) [CCN]; emamectin (291); emamectin benzoate (291); eprinomectin (alternative name) [CCN]; ivermectin (alternative name) [CCN]; milbemycin oxime (alternative name) [CCN]; moxidectin (alternative name) [CCN]; piperazine [CCN]; selamectin (alternative name) [CCN]; spinosad (737); thiophanate (1435); chloralose (127); endrin (1122); fenthion (346); pyridin-4-amine (IUPAC-Name) (23); strychnine (745); 1-hydroxy-1H-pyridine-2-thione (IUPAC-Name) (1222); 4-(quinoxalin-2-ylamino)benzenesulfonamide (IUPAC-Name) (748); 8-hydroxyquinoline sulfate (446); bronopol (97); copper dioctanoate (IUPAC-Name) (170); copper hydroxide (IUPAC-Name) (169); cresol [CCN]; dichlorophen (232); dipyrithione (1105); dodicin (1112); fenaminosulf (1144); formaldehyde (404); hydargaphen (alternative name) [CCN]; kasugamycin (483); kasugamycin hydrochloride hydrate (483); nickel bis(dimethylthiocarbamate) (IUPAC-Name) (1308); nitrapyrin (580); ochthilinone (590); oxolinic acid (606); oxytetracycline (611); potassium hydroxyquinoline sulfate (446); probenazole (658); streptomycin (744); streptomycin sesquisulfate (744); tecloftalam (766); thiomersal (alternative name) [CCN]; iodomethane (IUPAC-Name) (542); methyl bromide (537); apholate [CCN]; bisazir (alternative name) [CCN]; busulfan (alternative name) [CCN]; diflubensuron (250); dimatif (alternative name) [CCN]; hemel [CCN]; hempa [CCN]; metepa [CCN]; methiotepa [CCN]; methyl apholate [CCN]; morzid [CCN]; penfluron (alternative name) [CCN]; tepa [CCN]; thiohempa (alternative name) [CCN]; thiotepa (alternative name) [CCN]; tretamine (alternative name) [CCN]; uredepa (alternative name) [CCN]; (E)-dec-5-en-1-yl acetate with (E)-dec-5-en-1-ol (IUPAC-Name) (222); (E)-terdec-4-en-1-yl acetate (IUPAC-Name) (829); (E)-6-methylhept-2-en-4-ol (IUPAC-Name) (541); (E,Z)-tetradeca-4,10-dien-1-yl acetate (IUPAC-Name) (779); (Z)-dodec-7-en-1-yl acetate (IUPAC-Name) (285); (Z)-hexadec-11-enal (IUPAC-Name) (436); (Z)-hexadec-11-en-1-yl acetate (IUPAC-Name) (437); (Z)-hexadec-13-en-11-yn-1-yl acetate (IUPAC-Name) (438); (Z)-icos-13-en-10-one (IUPAC-Name) (448); (Z)-tetradec-7-en-1-al (IUPAC-Name) (782); (Z)-tetradec-9-en-1-ol (IUPAC-Name) (783); (Z)-tetradec-9-en-1-yl acetate (IUPAC-Name) (784); (7E,9Z)-dodeca-7,9-dien-1-yl acetate (IUPAC-Name) (283); (9Z,11E)-tetradeca-9,11-dien-1-yl acetate (IUPAC-Name) (780); (9Z,12E)-tetradeca-9,12-dien-1-yl acetate (IUPAC-Name) (781); 14-methylpentadec-1-ene (IUPAC-Name) (545); 4-
methylnonan-5-ol with 4-methylnonan-5-one (IUPAC-Name) (544); alpha-multistriatin
(alternative name) [CCN]; brevicomin (alternative name) [CCN]; codelure (alternative
name) [CCN]; codlemone (alternative name) (167); cuelure (alternative name) (179);
disparlure (277); dodec-8-en-1-yl acetate (IUPAC-Name) (286); dodec-9-en-1-yl acetate
(IUPAC-Name) (287); dodeca-8,10-dien-1-yl acetate (IUPAC-Name) (284); dominicalure
(alternative name) [CCN]; ethyl 4-methyloctanoate (IUPAC-Name) (317); eugenol
(alternative name) [CCN]; frontalin (alternative name) [CCN]; gossypure (alternative
name) (420); grandlure (421); grandlure I (alternative name) (421); grandlure II (alternative
name) (421); grandlure III (alternative name) (421); grandlure IV (alternative name) (421);
hexalure [CCN]; ipsdienol (alternative name) [CCN]; ipsenol (alternative name) [CCN];
japonilure (alternative name) (481); lineatin (alternative name) [CCN]; littlure (alternative
name) [CCN]; looplure (alternative name) [CCN]; medlure [CCN]; megatomoic acid
(alternative name) [CCN]; methyl eugenol (alternative name) (540); muscalure (563);
octadeca-2,13-dien-1-yl acetate (IUPAC-Name) (588); octadeca-3,13-dien-1-yl acetate
(IUPAC-Name) (589); orfralure (alternative name) [CCN]; oryctalure (alternative name)
(317); ostromone (alternative name) [CCN]; siglure [CCN]; sordidin (alternative name)
(736); sulcatol (alternative name) [CCN]; tetradec-11-en-1-yl acetate (IUPAC-Name) (785);
trimedlure (839); trimedlure A (alternative name) (839); trimedlure B1 (alternative name)
(839); trimedlure B2 (alternative name) (839); trimedlure C (alternative name) (839);
truncall (alternative name) [CCN]; 2-(octylthio)ethanol (IUPAC-Name) (591); butopyronoxyl
(933); butoxy(polypropylene glycol) (936); dibutyl adipate (IUPAC-Name) (1046); dibutyl
phthalate (1047); dibutyl succinate (IUPAC-Name) (1048); diethyglycolamide [CCN]; dimethyl
carbate [CCN]; dimethyl phthalate [CCN]; ethyl hexanediol (1137); hexamide [CCN];
methoquin-butyl (1276); methylneodecanamide [CCN]; oxamate [CCN]; picaridin [CCN]; 1,1-
dichloro-1-nitroethane (IUPAC- / Chemical Abstracts-Name) (1058); 1,1-dichloro-2,2-bis(4-
ethylphenyl)ethene (IUPAC-Name) (1056); 1,2-dichloropropane (IUPAC- / Chemical
Abstracts-Name) (1062); 1,2-dichloropropane with 1,3-dichloropropene (IUPAC-Name) (1063);
1-bromo-2-chloroethane (IUPAC- / Chemical Abstracts-Name) (916); 2,2,2-trichloro-1-(3,4-
dichlorophenyl)ethyl acetate (IUPAC-Name) (1451); 2,2-dichlorovinyl 2-ethylsulfinylethyl
methyl phosphate (IUPAC-Name) (1066); 2-(1,3-dithianol-2-yl)phenyl dimethylcarbamate
(IUPAC- / Chemical Abstracts-Name) (1109); 2-(2-butoxyethoxy)ethyl thiocyanate (IUPAC-/
Chemical Abstracts-Name) (935); 2-(4,5-dimethyl-1,3-dioxolan-2-yl)phenyl methylcarbamate
(IUPAC- / Chemical Abstracts-Name) (1084); 2-(4-chloro-3,5-xylloxy)ethanol (IUPAC-
Name) (986); 2-chlorovinyl diethyl phosphate (IUPAC-Name) (984); 2-imidazolidone (IUPAC-
Name) (1225); 2-isovalerylindan-1,3-dione (IUPAC-Name) (1246); 2-methyl(prop-2-ynyl)aminophenyl methylcarbamate (IUPAC-Name) (1284); 2-thiocyanatoethyl laureate (IUPAC-Name) (1433); 3-bromo-1-chloroprop-1-ene (IUPAC-Name) (917); 3-methyl-1-phenylpyrazol-5-yl dimethylcarbamate (IUPAC-Name) (1283); 4-methyl(prop-2-ynyl)amino-3,5-xylyl methylcarbamate (IUPAC-Name) (1285); 5,5-dimethyl-3-oxocyclohex-1-enyl dimethylcarbamate (IUPAC-Name) (1085); abamectin (1); acephate (2); acetamiprid (4); acethion (alternative name) [CCN]; acetoprole [CCN]; acrinathrin (9); acrylonitrile (IUPAC-Name) (861); alanycarb (15); aldicarb (16); aldoxycarb (863); aldrin (864); allethrin (17); allosamidin (alternative name) [CCN]; allyxycarb (866); alpha-cypermethrin (202); alpha-ecdyson (alternative name) [CCN]; aluminium phosphide (640); amidithion (870); amidthioate (872); aminocarb (873); amiton (875); amiton hydrogen oxalate (875); Amitraz (24); anabasine (877); athidathion (883); AVI 382 (compound code); AZ 60541 (compound code); azadirachtin (alternative name) (41); azamethiphos (42); azinphos-ethyl (44); azinphos-methyl (45); azothoate (889); Bacillus thuringiensis delta endotoxins (alternative name) (52); barium hexafluoroisilicate (alternative name) [CCN]; barium polysulfide (IUPAC- / Chemical Abstracts-Name) (892); barthrin [CCN]; BAS 320 I (compound code); Bayer 22/190 (development code) (893); Bayer 22408 (development code) (894); bendiocarb (58); benfuracarb (60); bensultap (66); beta-cyfluthrin (194); beta-cypermethrin (203); bifenthrin (76); bioallethrin (78); bioallethrin S-cyclopyrenyl isomer (alternative name) (79); bioethanolmetherin [CCN]; biopermethrin (908); bioremsmethrin (80); bis(2-chloroethyl) ether (IUPAC-Name) (909); bistrifluron (83); borax (86); brofenvalerate (alternative name); bromfenvinphos (914); bromocyclophos (918); bromo-DDT (alternative name) [CCN]; bromophos (920); bromophos-ethyl (921); buffencarb (924); buprofezin (99); butacarb (926); butathiofos (927); butocarboxim (103); butonate (932); butoxycarboxim (104); butylpyridaben (alternative name); cadusafos (109); calcium arsenate [CCN]; calcium cyanide (444); calcium polysulfide (IUPAC-Name) (111); camphenclocarb (941); carbonanolate (943); carbaryl (115); carbofuran (118); carbon disulfide (IUPAC- / Chemical Abstracts-Name) (945); carbon tetrachloride (IUPAC-Name) (946); carbophenothion (947); carbosulfan (119); cartap (123); cartap hydrochloride (123); cevadine (alternative name) (725); chlorbicyclen (960); chlordane (128); chlordecone (963); chlordezone (964); chlordezone hydrochloride (964); chloretroxyflos (129); chlorfenapyr (130); chlorfenphinphos (131); chlorfluazuron (132); chlorfosphos (136); chlorfoform [CCN]; chloropicrin (141); chlorphoxim (989); chlorprazophos (990); chlorpyrifos (145); chlorpyrifos-methyl (146); chlorthiophos (994); chromafenozone (150); cinerin I (696); cinerin II (696); cinerins (696); cis-resmethrin (alternative name);
cismethrin (80); clocythrin (alternative name); clothocarb (999); closantel (alternative name) [CCN]; clothianidin (165); copper acetoarsenite [CCN]; copper arsenate [CCN]; copper oleate [CCN]; coumaphos (174); coumihioate (1006); crotamiton (alternative name) [CCN]; crotuxyphos (1010); crufomate (1011); cryolite (alternative name) (177); CS 708 (development code) (1012); cyanofenphos (1019); cyanophos (184); cyanothioate (1020); cyclodrin [CCN]; cycloprothrin (188); cyfluthrin (193); cyhalothrin (196); cypermethrin (201); cyphenothrin (206); cyromazine (209); cythioate (alternative name) [CCN]; d-limonene (alternative name) [CCN]; d-tetramethrin (alternative name) (788); DAEP (1031); dazomet (216); DDT (219); decarbofuran (1034); deltamethrin (223); demephion (1037); demephion-O (1037); demephion-S (1037); demeton (1038); demeton-methyl (224); demeton-O (1038); demeton-O-methyl (224); demeton-S (1038); demeton-S-methyl (224); demeton-S-methylsulphon (1039); diafenthion (226); diafos (1042); dialidamol (1044); diazinon (227); dicapthon (1050); dichlofluanid (1051); dichlorvos (236); dicliphos (alternative name); dicresyl (alternative name) [CCN]; dicrotophos (243); dicyclanil (244); dieldrin (1070); diethyl 5-methylpyrazol-3-yl phosphate (IUPAC-Name) (1076); diflubenzuron (250); dilor (alternative name) [CCN]; dimefluthrin [CCN]; dimethoxy (1081); dimetan (1085); dimethoate (262); dimethrin (1083); dimethylvinphos (265); dimetilan (1086); dinex (1089); dinex-dicloxsline (1089); dinoprop (1093); dinosam (1094); dinoseb (1095); dinofuran (271); diofenolan (1099); dioxabenzofoos (1100); dioxacarb (1101); disulfoton (278); dithiocarb (1108); DNOC (282); doramectin (alternative name) [CCN]; DSP (1115); ecdysterone (alternative name) [CCN]; E1 642 (development code) (1118); emamectin (291); emamectin benzoate (291); EMPC (1120); empenthrin (292); endosulfan (294); endothion (1121); endrin (1122); EPBP (1123); EPN (297); epofenonanle (1124); epirimethrin (alternative name) [CCN]; esfenvalerate (302); etaphos (alternative name) [CCN]; ethofencarb (308); ethion (309); ethiprole (310); ethioate-methyl (1134); ethoprophos (312); ethyl formate (IUPAC-Name) [CCN]; ethyl-DDD (alternative name) (1056); ethylene dibromide (316); ethylene dichloride (chemical name) (1136); ethylene oxide [CCN]; etofenprox (319); etrilmfos (1142); EXD (1143); famphur (323); fenamiphos (326); fenazaflor (1147); fenchlorphos (1148); fenethacarb (1149); fenfurfurin (1150); fenitrothion (335); fenobucarb (336); fenoxacrin (1153); fenoxycarb (340); fenpirithrin (1155); fenpropadrin (342); fenpyrad (alternative name); fensulfothion (1158); fenthion (346); fentinphon-ethyl [CCN]; fenvalerate (349); fipronil (354); fipronil (358); fluocurcumon (1168); fluclycloxuron (366); flucypyr (367); fluenetil (1169); flufenoxuron [CCN]; flufenoxuron (370); flufenprox (1171); flumethrin (372); fluvulinate (1184); FMC 1137 (development code) (1185); fonofos (1191);
formetanate (405); formetanate hydrochloride (405); formothion (1192); formparanate (1193); fosmethylan (1194); fospirate (1195); fosthiazate (408); fosthietan (1196); furathiocarb (412); furethrin (1200); gamma-cyhalothrin (197); gamma-HCH (430); guazatine (422); guazatine acetates (422); GY-81 (development code) (423); halifenprox (424); halofenozide (425); HCH (430); HEOD (1070); heptachlor (1211); heptenophos (432); heterophos [CCN]; hexaflumuron (439); HHDN (864); hydramethylnon (443); hydrogen cyanide (444); hydroprene (445); hyquincarb (1223); imidacloprid (458); imiprothrin (460); indoxacarb (465); iodomethane (IUPAC-Name) (542); IPSP (1229); isazofos (1231); isobenzan (1232); isocarbophos (alternative name) (473); isodrin (1235); isofenphos (1236); isolane (1237); isopropcarb (472); isopropyl O-(methoxyaminothiophosphoryl)salicylate (IUPAC-Name) (473); isoprothiolane (474); isothioate (1244); isoxathion (480); ivermectin (alternative name) [CCN]; jasminol I (696); jasminol II (696); jodfenphos (1248); juvenile hormone I (alternative name) [CCN]; juvenile hormone II (alternative name) [CCN]; juvenile hormone III (alternative name) [CCN]; kelevan (1249); kinoprené (484); lambda-cyhalothrin (198); lead arsenate [CCN]; leptophos (1250); lindane (430); lidirimfos (1251); lufenuron (490); lythidathion (1253); m-cumyl methylcarbamate (IUPAC-Name) (1014); magnesium phosphide (IUPAC-Name) (640); malathion (492); malonoben (1254); mazidox (1255); mecarbam (502); mecarphon (1258); menazon (1260); mephosfolan (1261); mercurous chloride (513); mesulfenfos (1263); metam (519); metam-potassium (alternative name) (519); metam-sodium (519); methactrifos (1266); methamidophos (527); methanesulfonyl fluoride (IUPAC- / Chemical Abstracts-Name) (1268); methidathion (529); methiocarb (530); methocropophos (1273); methomyl (531); methoprene (532); methoquin-butyl (1276); methothrin (alternative name) (533); methoxychlor (534); methoxyfenozide (535); methyl bromide (537); methyl isothiocyanate (543); methylchloroform (alternative name) [CCN]; methylene chloride [CCN]; metofluthrin [CCN]; metolcarb (550); metoxadiazalone (1288); mevinphos (556); mexacarbate (1290); milbemectin (557); milbemycin oxime (alternative name) [CCN]; mipaflox (1293); mirex (1294); monocropophos (561); morphonothion (1300); moxidectin (alternative name) [CCN]; naftalofos (alternative name) [CCN]; naled (567); naphthalene (IUPAC- / Chemical Abstracts-Name) (1303); NC-170 (development code) (1306); NC-184 (compound code); nicotine (578); nicotine sulfate (578); nifluridide (1309); nitenpyram (579); nithiazine (1311); nitrilcarb (1313); nitrilcarb 1:1 zinc chloride complex (1313); NNI-0101 (compound code); NNI-0250 (compound code); normicotine (traditional name) (1319); novaluron (585); noviflumuron (586); O-2,5-dichloro-4-iodophenyl O-ethyl ethylphosphonothioate (IUPAC-Name) (1057); O,O-diethyl O-4-methyl-2-oxo-2H-chromen-7-
yl phosphorothioate (IUPAC-Name) (1074); O,O-diethyl O-6-methyl-2-propylpyrimidin-4-yl phosphorothioate (IUPAC-Name) (1075); O,O,O',O'-tetrapropyl dithiopyrophosphate (IUPAC-Name) (1424); oleic acid (IUPAC-Name) (593); omethoate (594); oxamyl (602); oxydemeton-methyl (609); oxydeprofos (1324); oxydisulfoton (1325); pp'-DDT (219); paradichlorobenzene [CCN]; parathion (615); parathion-methyl (616); penfluron (alternative name) [CCN]; pentachlorophenol (623); pentachlorophenyl laurate (IUPAC-Name) (623); permethrin (628); petroleum oils (alternative name) (628); PH 60-38 (development code) (1328); phenkapton (1330); phenothrin (630); phenthoate (631); phorate (636); phosalone (637); phosfolan (1338); phosphet (638); phosphichlorid (1339); phosphamidon (639); phosphate (IUPAC-Name) (640); phoxim (642); phoxim-methyl (1340); pirimphos (1344); pirimphos-ethyl (1345); pirimphos-methyl (652); polychlororodicyclopentadiene isomers (IUPAC-Name) (1346); polychloroterpenes (alternative name) (1347); potassium arsenite [CCN]; potassium thiocyanate [CCN]; prallethrin (655); precocene I (alternative name) [CCN]; precocene II (alternative name) [CCN]; precocene III (alternative name) [CCN]; primidophos (1349); profenofos (662); profenophos [CCN]; promacyl (1354); promecarb (1355); propaphos (1356); propetamphos (673); propoxur (678); prothidathion (1360); prothiofos (686); prothoate (1362); protrifonbutle [CCN]; pyrmetrozine (688); pyraclofos (689); pyrazofos (693); pyresmethrin (1367); pyrethrill (696); pyrethrins II (696); pyrethrins (696); pyridaben (699); pyridalyl (700); pyridaphenthion (701); pyrimidifen (706); pyrimidine (1370); pyriproxyfen (708); quassia (alternative name) [CCN]; quinalphos (711); quinalphos-methyl (1376); quinothion (1380); quintufo (1381); R-1492 (development code) (1382); rafoxanide (alternative name) [CCN]; resmethrin (719); rotenone (722); RU 15525 (development code) (723); RU 25475 (development code) (1386); ryania (alternative name) (1387); ryanodine (alternative name) (1387); sabadilla (alternative name) (725); schradan (1389); sebufos (alternative name); selamectin (alternative name) [CCN]; SI-0009 (compound code); silafflufen (728); SN 72129 (development code) (1397); sodium arsenite [CCN]; sodium cyanide (444); sodium fluoride (IUPAC- / Chemical Abstracts-Name) (1399); sodium hexafluorosilicate (1400); sodium pentachlorophenoxy (623); sodium selenite (IUPAC-Name) (1401); sodium thiocyanate [CCN]; sophamide (1402); spinosad (737); spiromesifen (739); sulfofon (746); sulfofon-sodium (746); sulfamid (750); sulfotep (753); sulfuryl fluoride (756); sulprofos (1408); tar oils (alternative name) (758); tau-fluvalinate (398); tacizincarb (1412); TDE (1414); tebufenoxid (762); tebufenpyrd (763); tebuiprimfos (764); tefubenzuron (768); tefluthrin (769); temephos (770); TEPP (1417); terallethrin (1418); terbam (alternative name); terbufos (773);
tetrachloroethane [CCN]; tetrachlorvinphos (777); tetramethrin (787); theta-cypermethrin (204); thiacloprid (791); thiafenox (alternative name); thiamethoxam (792); thicrofos (1428); thiocarboxime (1431); thicyclam (798); thiocyclam hydrogen oxalate (798); thiodicarb (799); thiophanox (800); thiometon (801); thionazin (1434); thiosultap (803); thiosultap-sodium (803); thuringiensin (alternative name) [CCN]; tolenpyrad (809); tralomethrin (812); transfluthrin (813); transpermethrin (1440); triamiphos (1441); triazamate (818); triazophos (820); triazuron (alternative name); trichlorfon (824); trichlorometaphos-3 (alternative name) [CCN]; trichloronat (1452); trifenofos (1455); triflumuron (835); trimethacarb (840); triprene (1459); vanidothion (847); vaniliprole [CCN]; veratridine (alternative name) (725); veratrine (alternative name) (725); XMC (853); xylilcarb (854); Y1-5302 (compound code); zeta-cypermethrin (205); zetamethrin (alternative name); zinc phosphide (640); zolaprofos (1469) und ZX1 8901 (development code) (858);

a compound of formula A-1

![Image of compound A-1](attachment:image1)

(A-1);

a compound of formula A-2

![Image of compound A-2](attachment:image2)

(A-2);

a compound of formula A-3

![Image of compound A-3](attachment:image3)
a compound of formula A-4

a compound of formula A-5

a compound of formula A-6
a compound of formula A-7

\[ \text{Chemical Structure of A-7} \]

a compound of formula A-8

\[ \text{Chemical Structure of A-8} \]

a compound of formula A-9

\[ \text{Chemical Structure of A-9} \]

a compound of formula A-10
a compound of formula A-10;

a compound of formula A-11;

a compound of formula A-12;

a compound of formula A-13;
a compound of formula A-14

\[ \text{(A-14)} \]

a compound of formula A-15

\[ \text{(A-15)} \]

a compound of formula A-15A

\[ \text{(A-15A)} \]

a compound of formula (A-16)
a compound of formula (A-16);

a compound of formula (A-17);

a compound of formula (A-18);

a compound of formula (A-19)
a compound of formula (A-19);

(A-19);

a compound of formula (A-20)

(A-20);

a compound of formula (A-21)

(A-21);

a compound of formula (A-22)
a compound of formula (A-22)

a compound of formula (A-23)

a compound of formula (A-24)

a compound of formula (A-25)
bis(tributyltin) oxide (IUPAC-Name) (913); bromoacetamide [CCN]; calcium arsenate [CCN]; cloethocarb (999); copper acetatoarsenite [CCN]; copper sulfate (172); fentin (347); ferric phosphate (IUPAC-Name) (352); metaldehyde (518); methiocarb (530); niclosamide (576); niclosamide-olamine (576); pentachlorophenol (623); sodium pentachlorophenoxide (623); tazimcarb (1412); thiodicarb (799); tributyltin oxide (913); trifenmorph (1454); trimethacarb (840); triphenyltin acetate (IUPAC-Name) (347); triphenyltin hydroxide (IUPAC-Name) (347); 1,2-dibromo-3-chloropropane (IUPAC- / Chemical Abstracts-Name) (1045); 1,2-dichloropropane (IUPAC- / Chemical Abstracts-Name) (1062); 1,2-dichloropropane with 1,3-dichloropropene (IUPAC-Name) (1063); 1,3-dichloropropene (233); 3,4-dichlorotetrahydrothiophene 1,1-dioxide (IUPAC- / Chemical Abstracts-Name) (1065); 3-(4-chlorophenyl)-5-methylrhodanine (IUPAC-Name) (980); 5-methyl-6-thioxo-1,3,5-thiadiazinan-3-ylacetic acid (IUPAC-Name) (1286); 6-isopentenylaminopurine (alternative name) (210); abamectin (1); acetoprole [CCN]; alancarb (15); aldicarb (16); aldioxycarb (863); AZ 60541 (compound code); benclotiaz [CCN]; benomyl (62); butlypyridaben (alternative name); cadusafos (109); carbofuran (118); carbon disulfide (945); carbosulfan (119); chloropicrin (141);
chlorpyrifos (145); cloethocarb (999); cytokinins (alternative name) (210); dazomet (216); DBCP (1045); DCIP (218); diamidofos (1044); dichlofenthion (1051); dicliphos (alternative name); dimethoate (262); doramectin (alternative name) [CCN]; emamectin (291); emamectin benzoate (291); eprinomectin (alternative name) [CCN]; ethoprophos (312); ethylene dibromide (316); fenamiphos (326); fenpyrad (alternative name); fensulfothion (1158); fosthiazate (408); fothialtan (1196); furfural (alternative name) [CCN]; GY-81 (development code) (423); heterophos [CCN]; iodomethane (IUPAC-Name) (542); isamidofos (1230); isazofos (1231); ivermectin (alternative name) [CCN]; kinetin (alternative name) (210); mecarphon (1258); metam (519); metam-potassium (alternative name) (519); metam-sodium (519); methyl bromide (537); methyl isothiocyanate (543); milbemycin oxime (alternative name) [CCN]; moxidectin (alternative name) [CCN]; Myrothecium verrucaria composition (alternative name) (565); NC-184 (compound code); oxamyl (602); phorate (636); phosphamidon (639); phosphocarb [CCN]; sebufos (alternative name); selamectin (alternative name) [CCN]; spinosad (737); terbuthram (alternative name); terbufos (773); tetrachlorothiophene (IUPAC- / Chemical Abstracts-Name) (1422); thiafenox (alternative name); thionazin (1434); triazophos (820); triazuron (alternative name); xylenols [CCN]; YI-5302 (compound code); zeatin (alternative name) (210); potassium ethylenethiuram [CCN]; nitrapyrin (580); acibenzolar (6); acibenzolar-S-methyl (6); probenazole (658); Reynoutria sachalinensis extract (alternative name) (720); 2-isovalerylindan-1,3-dione (IUPAC-Name) (1246); 4-(quinoxalin-2-ylamino)benzenesulfonamide (IUPAC-Name) (748); alphachlorohydrin [CCN]; aluminium phosphide (640); antu (880); arsenous oxide (882); barium carbonate (891); bisthiosemic (912); brodifacoum (89); bromadiolone (91); bromethalin (92); calcium cyanide (444); chloralose (127); chlorophacinone (140); chlorecalciferol (alternative name) (850); coumarchlor (1004); coumafuryl (1005); coumatetralyl (175); crimeidine (1009); difenacoum (246); difethialone (249); diphacinone (273); ergocalciferol (301); flocoumafen (357); fluoroacetamide (379); flupropadine (1183); flupropadine hydrochloride (1183); gamma-HCH (430); HCH (430); hydrogen cyanide (444); iodomethane (IUPAC-Name) (542); lindane (430); magnesium phosphide (IUPAC-Name) (640); methyl bromide (537); norbormide (1318); phosacetim (1336); phosphine (IUPAC-Name) (640); phosphorus [CCN]; pindone (1341); potassium arsenite [CCN]; pyrinuron (1371); scilioside (1390); sodium arsenite [CCN]; sodium cyanide (444); sodium fluoroacetate (735); strychnine (745); thallium sulfate [CCN]; warfarin (851); zinc phosphide (640); 2-(2-butoxyethoxy)ethyl piperonylate (IUPAC-Name) (934); 5-(1,3-benzodioxol-5-yl)-3-hexylcyclohex-2-enone (IUPAC-Name) (903); farnesol with nerolidol (alternative name) (324); MB-599 (development code) (498);
MGK 264 (development code) (296); piperonyl butoxide (649); piprotal (1343); propyl isome (1358); S421 (development code) (724); sesamex (1393); sesasmolin (1394); sulfoxide (1406); anthraquinone (32); chloralose (127); copper napthenate [CCN]; copper oxychloride (171); diazinon (227); dicyclopentadiene (chemical name) (1069); guazatine (422); guazatine acetates (422); methiocarb (530); pyridin-4-amine (IUPAC-Name) (23); thiram (804); trimethacarb (840); zinc napthenate [CCN]; ziram (856); imanin (alternative name) [CCN]; ribavirin (alternative name) [CCN]; mercuric oxide (512); octhilinone (590); thiophanate-methyl (802);

a compound of formula B-1A

\[
 \text{R'} \quad \text{N} \quad \text{F} \quad \text{F} \\
\text{Cl} \quad \text{N} \quad \text{Fluorescent Blue} \quad \text{Cl} \\
(\text{B-1A}) 
\]

wherein \( \text{R'} \) is hydrogen, \( \text{C}_{14}\text{alkyl} \) or \( \text{C}_{14}\text{haloalkyl} \);

a compound of formula B-2

\[
\text{Cl} \quad \text{Cl} \quad \text{Cl} \quad \text{CF}_3 \quad \text{O} \\
\text{Cl} \quad \text{N} \quad \text{N} \quad \text{H} \quad \text{O} \\
\text{Cl} \quad \text{Cl} \\
(\text{B-2}) 
\]

a compound of formula B-3

\[
\begin{align*}
\text{CF}_3 & \quad \text{N} \quad \text{O} \\
\text{F} & \quad \text{N} \quad \text{O} \\
\text{CF}_3 & \\
\text{F} & \\
\text{F} & \\
\text{B-3})
\end{align*}
\]
a compound of formula B-4

![Chemical structure](image)

(B-4);

a compound of formula B-5

![Chemical structure](image)

(B-5);

a compound of formula B-8

![Chemical structure](image)

(B-8);

and a compound of formula B-9
It has now been found, surprisingly, that the active ingredient mixture according to the invention not only brings about the additive enhancement of the spectrum of action with respect to the phytopathogen to be controlled that was in principle to be expected but achieves a synergistic effect which extends the range of action of the component (A) and of the component (B) in two ways. Firstly, the rates of application of the component (A) and of the component (B) are lowered whilst the action remains equally good. Secondly, the active ingredient mixture still achieves a high degree of phytopathogen control even where the two individual components have become totally ineffective in such a low application rate range. This allows, on the one hand, a substantial broadening of the spectrum of phytopathogens that can be controlled and, on the other hand, increased safety in use.

However, besides the actual synergistic action with respect to fungicidal activity, the pesticidal compositions according to the invention also have further surprising advantageous properties which can also be described, in a wider sense, as synergistic activity. Examples of such advantageous properties that may be mentioned are: a broadening of the spectrum of fungicidal activity to other phytopathogens, for example to resistant strains; a reduction in the rate of application of the active ingredients; synergistic activity against animal pests, such as insects or representatives of the order Acarina; a broadening of the spectrum of pesticidal activity to other animal pests, for example to resistant animal pests; adequate pest control with the aid of the compositions according to the invention, even at a rate of application at which the individual compounds are totally ineffective; advantageous behaviour during formulation and/or upon application, for example upon grinding, sieving, emulsifying, dissolving or dispensing; increased storage stability; improved stability to light; more advantageous degradability; improved toxicological and/or ecotoxicological behaviour; improved characteristics of the useful plants including: emergence, crop yields, more
developed root system, tillering increase, increase in plant height, bigger leaf blade, less
dead basal leaves, stronger tillers, greener leaf colour, less fertilizers needed, less seeds
needed, more productive tillers, earlier flowering, early grain maturity, less plant verse
(lodging), increased shoot growth, improved plant vigor, and early germination; or any other
advantages familiar to a person skilled in the art.

The alkyl groups appearing in the substituent definitions may be straight-chain or branched
and are, for example, methyl, ethyl, n-propyl, isopropyl, n-butyl, sec-butyl, isobutyl, tert-butyl,
pentyl, hexyl and the branched isomers thereof, preferred alkyl groups are methyl, isopropyl
and tert-butyl. Alkoxy radicals are derived from the mentioned alkyl radicals, for example
methoxy.

Halogen is generally fluorine, chlorine, bromine or iodine, preferably fluorine. The same is
true of halogen in haloalkyl.

Haloalkyl groups have a chain length of from 1 to 6 carbon atoms. Haloalkyl is, for example,
fluoromethyl, difluoromethyl, trifluoromethyl, chloromethyl, dichloromethyl, trichloromethyl,
1,1-difluoroethyl, 2,2,2-trifluoroethyl, 2-fluoroethyl, 2-chloroethyl, pentafluoroethyl, 1,1-
difluoro-2,2,2-trichloroethyl, 2,2,3,3-tetrafluoroethyl or 2,2,2-trichloroethyl, preferably 1,1-
difluoroethyl.

Alkoxy groups preferably have a chain length of from 1 to 4 carbon atoms. Alkoxy is, for
eexample, methoxy, ethoxy, propoxy, isoproxy, n-butoxy, isobutoxy, sec-butoxy and tert-
butoxy; preferably methoxy.

Since compounds of formula I may also contain asymmetric carbon atoms in the definition of
the substituent R₂, all the stereoisomers and all chiral <R> and <S> forms are also included.

The components B) are described in „The Pesticide Manual“ [The Pesticide Manual - A
World Compendium; Thirteenth Edition; Editor: C. D. S. Tomlin; The British Crop Protection
Council]. The numbers mentioned after the common names are the entry numbers of said
reference.

The following components B) are registered under a CAS-Reg. No.: Orysastrobin (CAS
248593-16-0); Aldimorph (CAS 91315-15-0); Iodocarb (3-iodo-2-propynyl butyl carbamate)
(CAS 55406-53-6); Fentin chloride (CAS 668-34-8); Hymexazole (CAS 10004-44-1); Phosphoric acid (CAS 7664-38-2); Tedoflam (CAS 76280-91-6); Arsenates (CAS 1327-53-3); Copper Ammoniumcarbonate (CAS 33113-08-5); Copper olate (CAS 1120-44-1); Mercury (CAS 7487-94-7; 21908-53-2; 7546-30-7); Benthiavalcarb (CAS 413615-35-7); Cadmium chloride (CAS 10108-64-2); Cedar leaf oil (CAS 8007-20-3); Chlorine (CAS 7782-50-5); Cinnamaldehyde (CAS: 104-55-2); Manganous dimethylthiocarbamate (CAS 15339-36-3); Neem oil (hydrophobic extract) (CAS 8002-65-1); Paraformaldehyde (CAS 30525-89-4); Sodium bicarbonate (CAS 144-55-8); Potassium bicarbonate (CAS 298-14-6); Sodium diacetate (CAS 127-09-3); Sodium propionate (CAS 137-40-6); TCMTB (CAS 21564-17-0); Benalaxyl -M (CAS 98243-83-5); Metrafoline (CAS 220899-03-6); Penthionpyrad (CAS 183675-82-3) and Tolyfluaniid (CAS 731-27-1).

The compound of formula F-1 is described in WO-01/87822. The compound of formula F-2 is described in WO-04/058723. The compounds of formula F-3, F-4, F-5 and F-6 are described in WO-04/035589. The compounds of formula F-7, F-8 and F-9 are described in WO-03/074491.


Bacillus pumilus GB34 and Bacillus pumilus strain QST are described at the U.S. Environmental Protection Agency, U.S. EPA PC Code 006493 and U.S. EPA PC Code 006485, respectively (see: http://www.epa.gov).

Compounds of formula B-1A and the compound of formula B-1 are described in WO 98/46607. The compound of formula B-2 is described in WO 99/042447. The compound of formula B-3 is described in WO 96/19442. The compound of formula B-4 is described in WO 99/14187. The compound of formula B-5 is described in US-5,945,423 and WO 94/26722. The compound of formula B-6 is described in EP-0-936-213. The compound of formula B-7 is described in US-6,020,332, CN-1-167-568, CN-1-155-977 and EP-0-860-438. The compound of formula B-8 is registered under CAS-Reg. No.: 325156-49-8 and is also known as Pyribencarb. The compound of formula B-9 is registered under CAS-Reg. No.: 348635-87-0 and is also known as Ambromdole or Amisulbrom.

According to the instant invention, a "racemic compound" means a mixture of two enantiomers in a ratio of substantially 50:50 of the two enantiomers.
Throughout this document the expression "combination" stands for the various combinations of components A) and B), for example in a single "ready-mix" form, in a combined spray mixture composed from separate formulations of the single active ingredient components, such as a "tank-mix", and in a combined use of the single active ingredients when applied in a sequential manner, i.e. one after the other with a reasonably short period, such as a few hours or days. The order of applying the components A) and B) is not essential for working the present invention.

The combinations according to the invention may also comprise more than one of the active components B), if, for example, a broadening of the spectrum of phytopathogenic disease control is desired. For instance, it may be advantageous in the agricultural practice to combine two or three components B) with any of the compounds of formula I, or with any preferred member of the group of compounds of formula I.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is trifluoromethyl and one component B) as described above.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and one component B) as described above.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₁-C₆alkyl, and one component B) as described above.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₁-C₆haloalkyl, and one component B) as described above.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₁-C₆alkoxy-C₁-C₆alkyl, and one component B) as described above.
A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ia

![Chemical structure of Ia](image)

and one component B) as described above.

The preferred compound of formula Ia and its manufacturing process starting from known and commercially available compounds is described in WO 04/058723. It is described in WO 04/058723 that the compound of formula Ia can be prepared by reacting an acid chloride of formula II

![Chemical structure of II](image)

with an amine of formula III
Acids of formula IV

are used for the production of the acid chlorides of formula II, via reaction steps as described in WO 04/058723. When producing the acids of the formula IV using said methodology impurities of formula IVA, IVB and/or IVC may be formed:

When applying the described manufacturing processes for compounds of formula I some/all of those impurities may be carried through different steps of said manufacturing processes. This then can lead to the formation of the corresponding acid chlorides (IIIA, IIB and/or IIC)
and to the formation of the corresponding amides (VA, VB and/or VC)

as further impurities of compounds of formula Ia. The presence/amount of said impurities in preparations of said compounds of formula Ia varies dependent on purification steps used.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ib
and one component B) as described above.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ic

and one component B) as described above.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Id
and one component B) as described above.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula (le)

and one component B) as described above.

Preferred components B) are selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrobin, Dimoxystrobin, Fluoastrobio, Kresoxim-methyl, Metominostrobin, Oryxstrobin, Picoxystrobin, Pyraclostrobin; Trifoxystrobin; and a compound of formula B-6
an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triticonazole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;
a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;
an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil;
a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperaline and a compound of formula B-7

a compound of formula F-1
- 37 -

a compound of formula B-1

Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxyl; Benalaxyl-M; Benomyl; Biteranol; Boscalid; Carboxin; Carpropanid; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamide; Fenoxycarb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazole; Iprodione; Lufenuron; Mancozeb; Metalaxyl; Mefenoxyam; Metrafenone; Nuarimol; Paclorotrazol; Pencycuron; Pentiopyrad; Procymidone; Pyroquilon; Quinoxyfen; Silthiofam; Sulfur; Thiacabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A more preferred component B) is Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

(F-1);
a compound of formula B-1

\[ \text{CH}_3 \]

\[
\text{N} \quad \text{F} \quad \text{F} \\
\text{N} \quad \text{Cl} \quad \text{F} \\
\text{N} \quad \text{F} \\
\text{N}
\]

(B-1);

Chlorothalonil, Epoxiconazole or Prothioconazole.

A further more preferred component B) is Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

\[ \text{CH} \]

\[
\text{O} \quad \text{N} \quad \text{CH}_3 \\
\text{O} \quad \text{CH} \quad \text{CH}
\]

(F-1);

or Chlorothalonil.

Most preferred component B) is Azoxystrobin; Cyproconazole; Propiconazole; Cyprodinil; Chlorothalonil; Fludioxonil or Fenpropidin.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is trifluoromethyl and one component B) selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrobin, Dimoxystrobin, Flouxastrobin, Kresoxim-methyl, Metominostrobin, Orysastrobin, Picoxystrobin, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6
an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triticonzole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;

a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;

an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil;

a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7

a compound of formula F-1
a compound of formula B-1

Chlorothalonil; Famoxadone; Fenamidone; Aclbensolar; Benalaxyl; Benalaxyl-M; Benomyl; Bitertanol; Boscalid; Carboxin; Carpropanid ; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamide; Fenoxy carb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazole; Iprodione; Lufenuron; Mancozeb; Metalaxyl; Mefeno xam; Metrafenone; Nuarimol; Paclobutrazol; Pencycuron; Penthitoryd; Procymidone; Pyroqilone; Quinoxyfen; Silthiofam; Sulfur; Thiabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R_1 is difluoromethyl and one component B) selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrobin, Dimoxystrobin, Flu oxystrobin, Kresoxim-methyl, Metominostrobin, Ory sastrobin, Picoxystrobin, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6
an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetaconazole, Triadimefon, Triadimenol, Triflumizole, Triticonazole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;
a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;
an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil;
a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7

a compound of formula F-1
a compound of formula B-1

Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxyl; Benalaxyl-M; Benomyl; Bitertanol; Boscalid; Carboxin; Carprolamid ; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamide; Fenoxycarb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazole; Ipodione; Lufenuron; Mancozeb; Metalaxyl; Mefenoxyd; Metrafenone; Nuarimol; Paclotrazol; Pancrycuron; Penthiozyraid; Procymidon; Pyroquilon; Quinoxyfen; Silthiofan; Sulfur; Thiabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₁-C₆alkyl, and one component B) selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrrobín, Dimoxystrobin, Fluoxastrobin, Kresoxim-methyl, Metominostrobin, Orysastrobin, Picoxystrrobín, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6
an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diconazole, Diconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalin, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefluazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triticonazole, Diclobutanil, Etaconazole, Furconazole, Furconazole-cis and Quinconazole; a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil; an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil; a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7

a compound of formula F-1
Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxyl; Benalaxyl-M; Benomyl; Bitertanol; Boscalid; Carboxin; Carpropanid; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamide; Fenoxycarb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazole; Iprodione; Lufenuron; Mancozeb; Metalaxyl; Mefenoxam; Metrafenone; Nuarimol; Paclobutrazol; Pencycuron; Penthiopyrad; Procymidine; Pyroquilon; Quinoxyfen; Silthiofam; Sulfur; Thiabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₃-C₆ haloalkyl, and one component B) selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrobín, Dimoxystrobin, Fluoxastrobin, Kresoxim-methyl, Metominostrobin, Oryastrobin, Picoxystrobin, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6.
an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenoconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triticonazole, Dichlobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;

a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;

an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil;

a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7

a compound of formula F-1
a compound of formula B-1

Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxyl; Benalaxyl-M; Benomyl; Bitertanol; Boscalid; Carboxin; Carpropanid; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamide; Fenoxycarb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazol; Iprodione; Lufenuron; Mancozeb; Metalaxyl; Mefenoxam; Metrafenone; Nuarimol; Paclobutrazol; Pencycuron; Penthionpyrad; Procymidine; Pyroquilon; Quinoxyfen; Silthiofam; Sulfur; Thiacendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₁-C₆alkoxy-C₁-C₆alkyl, and one component B) selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrobin, Dimoxystrobin, Fluoastrobine, Kresoxim-methyl, Metominostrobin, Oryastrobine, Picoxystrobin, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6
an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxpconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triticonazole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;

a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;

an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil;

a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperaliln and a compound of formula B-7

a compound of formula F-1
A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ia
and one component B) selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrobin, Dimoxystrobin, Fluoastrobins, Kresoxim-methyl, Metominostrobin, Orysatrobin, Picoxystrobin, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6

a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;
an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil,
Mepanipyrim and Pyrimethanil;
a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7
- 50 -

![Chemical structure](image)

(B-7);

a compound of formula F-1

![Chemical structure](image)

(F-1);

a compound of formula B-1

![Chemical structure](image)

(B-1);

Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxyl; Benalaxyl-M; Benomyl; Bitertanol; Boscalid; Carboxin; Carpropanid ; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamide; Fenoxy carb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazol e; Iprodione; Lufenuron; Mancozeb; Metalaxyl; Mefenoxam; Metrafenone; Nuarimol; Paclobutrazol; Pencycuron; Penthiopyrad; Procymidone; Pyroquilon; Quinoxyfen; Silthiofam; Sulfur; Thiabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.
A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula (lb)

![Chemical Structure](image)

and one component B) selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrobin, Dimoxystrobin, Fluoxastrobin, Kresoxim-methyl, Metominostrobin, Orysastrobin, Picoxystrobin, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6

![Chemical Structure](image)

an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triflumazole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;
a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;
an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil;
a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7

![Chemical structure of B-7]

a compound of formula F-1

![Chemical structure of F-1]

a compound of formula B-1

![Chemical structure of B-1]

Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxyl; Benalaxyl-M; Benomyl; Bitertanol; Boscalid; Carboxin; Carprocamid; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamide; Fenoxy carb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazol e; Iprodione; Lufenuron; Mancozeb; Metalaxyl; Mefeno xam; Metrafenone; Nuar imol; Paclobutrazol; Pencycuron; Penth iopyrad; Procymidone; Pyroquinol;
Quinoxyfen; Silthiofam; Sulfur; Thiabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula 1c

![Chemical Structure](image)

(B-6);

an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fruquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Tiritconazole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;
a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fluoxonil;  
an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil,  
Mepanipyrim and Pyrimethanil;  
a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph,  
Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of  
formula B-7

![Chemical Structure B-7](image)

(F-7);

a compound of formula F-1

![Chemical Structure F-1](image)

(F-1);

a compound of formula B-1

![Chemical Structure B-1](image)

(B-1);

Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxy; Benalaxy-M; Benomyl; 
Bitertanol; Boscalid; Carboxin; Carpropamid; Copper; Cyazofamid; Cymoxanil;
Diethofencarb; Dithianon; Fenhexamide; Fenoxycarb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazole; Iprodione; Lufenuron; Mancozeb; Metalaxyl; Mefenoxam; Metrafenone; Nuarimol; Paclobutrazol; Pancurion; Penthiopyrad; Procyclidone; Pyroquilon; Quinoxyfen; Silthiofam; Sulfur; Thiabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Id

\[
\text{Id),}
\]

and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoystrobin, Dimoxystrobin, Fluoxastrobin, Kresoxim-methyl, Metominostrobin, Oryasastrobin, Picoxystrobin, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6

\[
\text{(B-6);}
\]

an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flutilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole,
Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triticonazole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole; a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil; an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil; a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7

![Chemical structure](image)

(B-7);

a compound of formula F-1

![Chemical structure](image)

(F-1);

a compound of formula B-1

![Chemical structure](image)

(B-1);
Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxyl; Benalaxyl-M; Benomyl; Biteranol; Boscald; Carboxin; Carpropanid ; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamide; Fenoxycarb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazol; Ipodione; Lufenuron; Mancozeb; Metalaxyl; Mefenoxyam; Metrafenone; Nuarimol; Paclorbutrazol; Pencycuron; Penthioctrad; Procymidone; Pyroquilon; Quinoxyfen; Silthiofam; Sulfur; Thiabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ie

![Chemical structure of formula Ie](image1)

and one component B) selected from the group consisting of strobilurin fungicide, selected from the group consisting of Azoxystrobin, Dimoxystrobin, Fluoxastrobin, Kresoxim-methyl, Metominostrobin, Orysastrobin, Picoxystrobin, Pyraclostrobin; Trifloxystrobin; and a compound of formula B-6

![Chemical structure of formula B-6](image2)

an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil,
Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoate, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triticonazole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;
a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;
an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil;
a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7

![Chemical Structure B-7](image)

(B-7);

a compound of formula F-1

![Chemical Structure F-1](image)

(F-1);

a compound of formula B-1
Chlorothalonil; Famoxadone; Fenamidone; Acibenzolar; Benalaxyl; Benalaxyl-M; Benomyl; Biteranol; Boscalid; Carboxin; Carproamid; Copper; Cyazofamid; Cymoxanil; Diethofencarb; Dithianon; Fenhexamid; Fenoxycarb; Fluazinam; Flutolanil; Folpet; Guazatine; Hymexazol; Iprodione; Lufenuron; Mancozeb; Metalaxyl; Mefenoxam; Metrafenone; Nuarimol; Pactobutrazol; Pencycuron; Penthiopyrad; Procymidone; Pyroquilon; Quinoxyfen; Silthiofam; Sulfur; Thiabendazole; Thiram; Triazoxide; Tricyclazole; Abamectin; Emamectin benzoate; Tefluthrin and Thiamethoxam.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is trifluoromethyl and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

\[
\text{(F-1);}
\]

a compound of formula B-1
Chlorothalonil, Epoxiconazole or Prothioconazole.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is difluoromethyl and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

![Chemical structure](image)

(F-1);

a compound of formula B-1

![Chemical structure](image)

Chlorothalonil, Epoxiconazole or Prothioconazole.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is difluoromethyl
and R₂ is C₁-C₆alkyl, and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

\[
\begin{align*}
\text{(F-1)};
\end{align*}
\]

a compound of formula B-1

\[
\begin{align*}
\text{(B-1)};
\end{align*}
\]

Chlorothalonil, Epoxiconazole or Prothioconazole.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₁-C₆haloalkyl, and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

\[
\begin{align*}
\text{(F-1)};
\end{align*}
\]

a compound of formula B-1
Chlorothalonil, Epoxiconazole or Prothioconazole.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein $R_1$ is difluoromethyl and $R_2$ is $C_1-C_4$-alkoxy-$C_1-C_6$-alkyl, and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

$$\text{(F-1);}$$

a compound of formula B-1

$$\text{(B-1);}$$

Chlorothalonil, Epoxiconazole or Prothioconazole.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ia
and one component B) selected from the group consisting of Azoxytrobin; Picoxytrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

(F-1);

a compound of formula B-1

(B-1);

Chlorothalonil, Epoxiconazole or Prothioconazole.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula 1b
and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ic

Chlorothalonil, Epoxiconazole or Prothioconazole.
and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

\[
\begin{align*}
\text{(F-1);} \\
\end{align*}
\]

a compound of formula B-1

\[
\begin{align*}
\text{(B-1);} \\
\end{align*}
\]

Chlorothalonil, Epoxiconazole or Prothioconazole.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Id
and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

\[
\begin{align*}
\text{CH} & \text{CH} \\
\text{O} & \text{N} \\
\text{O} & \text{O-CH}_3
\end{align*}
\]

a compound of formula B-1

\[
\begin{align*}
\text{CH}_3 & \\
\text{F} & \text{Cl} \\
\text{N} & \text{N}
\end{align*}
\]

Chlorothalonil, Epoxiconazole or Prothioconazole.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ie
and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is trifluoromethyl
and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

\[ \text{(F-1)}; \]

or Chlorothalonil.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is difluoromethyl and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

\[ \text{(F-1)}; \]

or Chlorothalonil.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is difluoromethyl and \( R_2 \) is \( C_1-C_6 \) alkyl, and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1
or Chlorothalonil.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₁-C₆ haloalkyl, and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

or Chlorothalonil.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein R₁ is difluoromethyl and R₂ is C₁-C₆ alkoxy-C₁-C₆ alkyl, and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1
A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ia

and one component B) selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

or Chlorothalonil.
A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ib

![Chemical Structure](image)

and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

![Chemical Structure](image)

or Chlorothalonil.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ic
and one component B) selected from the group consisting of Azoxytrobin; Picoxystrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

or Chlorothalonil.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Id
and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin; Picoxytrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1

![Chemical Structure](image)

or Chlorothalonil.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula Ie

![Chemical Structure](image)

and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin; Picoxytrobin; Cyproconazole; Difenoconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1
or Chlorothalonil.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is trifluoromethyl and one component B) selected from the group consisting of Azoxyostrobin; Cyproconazole; Propiconazole; Cypinidinil; Chlorothalonil; Fludioxonil or Fenpropidin.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is difluoromethyl and one component B) selected from the group consisting of Azoxyostrobin; Cyproconazole; Propiconazole; Cypinidinil; Chlorothalonil; Fludioxonil or Fenpropidin.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is difluoromethyl and \( R_2 \) is \( C_1-C_6 \)alkyl, and one component B) selected from the group consisting of Azoxyostrobin; Cyproconazole; Propiconazole; Cypinidinil; Chlorothalonil; Fludioxonil or Fenpropidin.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula I, wherein \( R_1 \) is difluoromethyl and \( R_2 \) is \( C_1-C_6 \)haloalkyl, and one component B) selected from the group consisting of Azoxyostrobin; Cyproconazole; Propiconazole; Cypinidinil; Chlorothalonil; Fludioxonil or Fenpropidin.
of Azoxystrobin; Cyproconazole; Propiconazole; Cyprodinil; Chlorothalonil; Fludioxonil or Fenpropidin.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula la

![Chemical structure of la](image)

and one component B) selected from the group consisting of Azoxystrobin; Cyproconazole; Propiconazole; Cyprodinil; Chlorothalonil; Fludioxonil or Fenpropidin.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula lb

![Chemical structure of lb](image)

and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin; Cyproconazole; Propiconazole; Cyprodinil; Chlorothalonil; Fludioxonil or Fenpropidin.
A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula \( \text{Ic} \)

\[
\begin{align*}
\text{CF}_2\text{H} & \quad \text{O} \\
\text{N} & \quad \text{H} \\
\text{N} & \quad \text{CH}_3 \\
\text{CH}_3 & \quad \text{O} \\
\text{CH}_3 & \quad \text{CH}_3 \\
\text{(Ic)} &
\end{align*}
\]

and one component B) selected from the group consisting of Azoxystrobin; Cyproconazole; Propiconazole; Cyprodinil; Chlorothalonil; Fludioxonil or Fenpropidin.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula \( \text{Id} \)

\[
\begin{align*}
\text{CF}_2\text{H} & \quad \text{O} \\
\text{N} & \quad \text{H} \\
\text{N} & \quad \text{CH}_3 \\
\text{CH}_3 & \quad \text{CH}_3 \\
\text{(Id)} &
\end{align*}
\]

and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin; Cyproconazole; Propiconazole; Cyprodinil; Chlorothalonil; Fludioxonil or Fenpropidin.

A preferred embodiment of the present invention is represented by those combinations which comprise as component A) a compound of the formula \( \text{Ie} \)
and one component B) selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobilin; Cyproconazole; Propiconazole; Cyproconil; Chlorothalonil; Fludioxonil or Fenpropidin.

The active ingredient combinations are effective against harmful microorganisms, such as microorganisms, that cause phytopathogenic diseases, in particular against phytopathogenic fungi and bacteria.

The active ingredient combinations are effective especially against phytopathogenic fungi belonging to the following classes: Ascomycetes (e.g. Venturia, Podosphaera, Erysiphe, Monilinia, Mycosphaerella, Uncinula); Basidiomycetes (e.g. the genus Hemileia, Rhizoctonia, Phakopsora, Puccinia, Ustilago, Tilletia); Fungi imperfecti (also known as Deuteromycetes; e.g. Botrytis, Helminthosporium, Rhynchosporium, Fusarium, Septoria, Cercospora, Alternaria, Pyricularia and Pseudocercosporella); Oomycetes (e.g. Phytophthora, Peronospora, Pseudoperonospora, Albugo, Bremia, Pythium, Pseudosclerospora, Plasmopara).

According to the invention “useful plants” typically comprise the following species of plants: grape vines; cereals, such as wheat, barley, rye or oats; beet, such as sugar beet or fodder beet; fruits, such as pomes, stone fruits or soft fruits, for example apples, pears, plums, peaches, almonds, cherries, strawberries, raspberries or blackberries; leguminous plants, such as beans, lentils, peas or soybeans; oil plants, such as rape, mustard, poppy, olives, sunflowers, coconut, castor oil plants, cocoa beans or groundnuts; cucumber plants, such as marrows, cucumbers or melons; fibre plants, such as cotton, flax, hemp or jute; citrus fruit,
such as oranges, lemons, grapefruit or mandarins; vegetables, such as spinach, lettuce, asparagus, cabbages, carrots, onions, tomatoes, potatoes, cucurbits or paprika; lauraceae, such as avocados, cinnamon or camphor; maize; tobacco; nuts; coffee; sugar cane; tea; vines; hops; durian; bananas; natural rubber plants; turf or ornamentals, such as flowers, shrubs, broad-leaved trees or evergreens, for example conifers. This list does not represent any limitation.

The term "useful plants" is to be understood as including also useful plants that have been rendered tolerant to herbicides like bromoxynil or classes of herbicides (such as, for example, HPPD inhibitors, ALS inhibitors, for example primisulfuron, prosulfuron and trifloxysulfuron, EPSPS (5-enol-pyrovl-shikimate-3-phosphate-synthase) inhibitors, GS (glutamine synthetase) inhibitors or PPO (protoporphyrinogen-oxidase) inhibitors) as a result of conventional methods of breeding or genetic engineering. An example of a crop that has been rendered tolerant to imidazolinones, e.g. imazamox, by conventional methods of breeding (mutagenesis) is Clearfield® summer rape (Canola). Examples of crops that have been rendered tolerant to herbicides or classes of herbicides by genetic engineering methods include glyphosate- and glufosinate-resistant maize varieties commercially available under the trade names RoundupReady®, Herculex I® and LibertyLink®.

The term "useful plants" is to be understood as including also useful plants which have been so transformed by the use of recombinant DNA techniques that they are capable of synthesising one or more selectively acting toxins, such as are known, for example, from toxin-producing bacteria, especially those of the genus Bacillus.

Toxins that can be expressed by such transgenic plants include, for example, insecticidal proteins, for example insecticidal proteins from Bacillus cereus or Bacillus popilliae; or insecticidal proteins from Bacillus thuringiensis, such as δ-endotoxins, e.g. CryIA(b), CryIA(c), CryIF, CryIF(a2), CryIIB, CryIII A, CryIIB(b1) or Cry9c, or vegetative insecticidal proteins (VIP), e.g. VIP1, VIP2, VIP3 or VIP3A; or insecticidal proteins of bacteria colonising nematodes, for example Photorhabdus spp. or Xenorhabdus spp., such as Photorhabdus luminescens, Xenorhabdus nematophilus; toxins produced by animals, such as scorpion toxins, arachnid toxins, wasp toxins and other insect-specific neurotoxins; toxins produced by fungi, such as Streptomyces toxins, plant lectins, such as pea lectins, barley lectins or snowdrop lectins; agglutinins; proteinase inhibitors, such as trypsin inhibitors, serine
protease inhibitors, patatin, cystatin, papain inhibitors; ribosome-inactivating proteins (RIP), such as ricin, maize-RIP, abrin, luffin, saporin or bryodin; steroid metabolism enzymes, such as 3-hydroxysteroidoxidase, ecodysteroid-UDP-glycosyl-transferase, cholesterol oxidases, ecdysone inhibitors, HMG-COA-reductase, ion channel blockers, such as blockers of sodium or calcium channels, juvenile hormone esterase, diuretic hormone receptors, stilbene synthase, bibenzyl synthase, chitinases and glucanases.

In the context of the present invention there are to be understood by δ-endotoxins, for example CryIA(b), CryIA(c), CryIF, CryLF(a2), CryIIIA(b), CryIIIA, CryIIIB(b1) or Cry9c, or vegetative insecticidal proteins (VIP), for example VIP1, VIP2, VIP3 or VIP3A, expressly also hybrid toxins, truncated toxins and modified toxins. Hybrid toxins are produced recombinantly by a new combination of different domains of those proteins (see, for example, WO 02/15701). An example for a truncated toxin is a truncated CryIA(b), which is expressed in the Bt11 maize from Syngenta Seed SAS, as described below. In the case of modified toxins, one or more amino acids of the naturally occurring toxin are replaced. In such amino acid replacements, preferably non-naturally present protease recognition sequences are inserted into the toxin, such as, for example, in the case of CryIIIA055, a cathepsin-D-recognition sequence is inserted into a CryIIIA toxin (see WO 03/018810)


The processes for the preparation of such transgenic plants are generally known to the person skilled in the art and are described, for example, in the publications mentioned above. Cryl-type deoxyribonucleic acids and their preparation are known, for example, from WO 95/34656, EP-A-0 367 474, EP-A-0 401 979 and WO 90/13651.

The toxin contained in the transgenic plants imparts to the plants tolerance to harmful insects. Such insects can occur in any taxonomic group of insects, but are especially commonly found in the beetles (Coleoptera), two-winged insects (Diptera) and butterflies (Lepidoptera).
Transgenic plants containing one or more genes that code for an insecticidal resistance and express one or more toxins are known and some of them are commercially available. Examples of such plants are: YieldGard® (maize variety that expresses a CryIA(b) toxin); YieldGard Rootworm® (maize variety that expresses a CryIIC(b1) toxin); YieldGard Plus® (maize variety that expresses a CryIA(b) and a CryIIC(b1) toxin); Starlink® (maize variety that expresses a Cry9(c) toxin); Herculex I® (maize variety that expresses a CryIF(a2) toxin and the enzyme phosphinothricine N-acetyltransferase (PAT) to achieve tolerance to the herbicide glufosinate ammonium); NuCOTN 33B® (cotton variety that expresses a CryIA(c) toxin); Bollgard I® (cotton variety that expresses a CryIA(c) toxin); Bollgard II® (cotton variety that expresses a CryIA(c) and a CryIIA(b) toxin); VIPCOT® (cotton variety that expresses a VIP toxin); NewLeaf® (potato variety that expresses a CryIIIa toxin); Nature-Gard® and Protecta®.

Further examples of such transgenic crops are:
1. **Bt11 Maize** from Syngenta Seeds SAS, Chemin de l'Hobit 27, F-31 790 St. Sauveur, France, registration number C/FR/96/05/10. Genetically modified *Zea mays* which has been rendered resistant to attack by the European corn borer (*Ostrinia nubilalis* and *Sesamia nonagrioides*) by transgenic expression of a truncated CryIA(b) toxin. Bt11 maize also transgenically expresses the enzyme PAT to achieve tolerance to the herbicide glufosinate ammonium.

2. **Bt176 Maize** from Syngenta Seeds SAS, Chemin de l'Hobit 27, F-31 790 St. Sauveur, France, registration number C/FR/96/05/10. Genetically modified *Zea mays* which has been rendered resistant to attack by the European corn borer (*Ostrinia nubilalis* and *Sesamia nonagrioides*) by transgenic expression of a CryIA(b) toxin. Bt176 maize also transgenically expresses the enzyme PAT to achieve tolerance to the herbicide glufosinate ammonium.

3. **MIR604 Maize** from Syngenta Seeds SAS, Chemin de l'Hobit 27, F-31 790 St. Sauveur, France, registration number C/FR/96/05/10. Maize which has been rendered insect-resistant by transgenic expression of a modified CryIIIa toxin. This toxin is Cry3A055 modified by insertion of a cathepsin-D-protease recognition sequence. The preparation of such transgenic maize plants is described in WO 03/018810.

4. **MON 863 Maize** from Monsanto Europe S.A. 270-272 Avenue de Tervuren, B-1150 Brussels, Belgium, registration number C/DE/02/9. MON 863 expresses a CryIIC(b1) toxin and has resistance to certain Coleoptera insects.
5. **IPC 531 Cotton** from Monsanto Europe S.A. 270-272 Avenue de Tervuren, B-1150 Brussels, Belgium, registration number C/ES/96/02.

6. **1507 Maize** from Pioneer Overseas Corporation, Avenue Tedesco, 7 B-1160 Brussels, Belgium, registration number C/NL/00/10. Genetically modified maize for the expression of the protein Cry1F for achieving resistance to certain Lepidoptera insects and of the PAT protein for achieving tolerance to the herbicide glufosinate ammonium.

7. **NK603 x MON 810 Maize** from Monsanto Europe S.A. 270-272 Avenue de Tervuren, B-1150 Brussels, Belgium, registration number C/GB/02/M3/03. Consists of conventionally bred hybrid maize varieties by crossing the genetically modified varieties NK603 and MON 810. NK603 x MON 810 Maize transgenically expresses the protein CP4 EPSPS, obtained from *Agrobacterium sp.* strain CP4, which imparts tolerance to the herbicide Roundup® (contains glyphosate), and also a Cry1A(b) toxin obtained from *Bacillus thuringiensis subsp. kurstaki* which brings about tolerance to certain Lepidoptera, include the European corn borer.

Transgenic crops of insect-resistant plants are also described in BATS (Zentrum für Biosicherheit und Nachhaltigkeit, Zentrum BATS, Clarastrasse 13, 4058 Basel, Switzerland) Report 2003, (http://bats.ch).

The term "useful plants" is to be understood as including also useful plants which have been so transformed by the use of recombinant DNA techniques that they are capable of synthesising antipathogenic substances having a selective action, such as, for example, the so-called "pathogenesis-related proteins" (PRPs, see e.g. EP-A-0 392 225). Examples of such antipathogenic substances and transgenic plants capable of synthesising such antipathogenic substances are known, for example, from EP-A-0 392 225, WO 95/33818, and EP-A-0 353 191. The methods of producing such transgenic plants are generally known to the person skilled in the art and are described, for example, in the publications mentioned above.

Antipathogenic substances which can be expressed by such transgenic plants include, for example, ion channel blockers, such as blockers for sodium and calcium channels, for example the viral KP1, KP4 or KP6 toxins; stilbene synthases; bibenzyl synthases; chitinases; glucanases; the so-called "pathogenesis-related proteins" (PRPs; see e.g. EP-A-0 392 225); antipathogenic substances produced by microorganisms, for example peptide antibiotics or heterocyclic antibiotics (see e.g. WO 95/33818) or protein or polypeptide
factors involved in plant pathogen defence (so-called "plant disease resistance genes", as described in WO 03/000906).

Useful plants of elevated interest in connection with present invention are cereals; soybean; rice; oil seed rape; pome fruits; stone fruits; peanuts; coffee; tea; strawberries; turf; vines and vegetables, such as tomatoes, potatoes, cucurbits and lettuce.

The term "locus" of a useful plant as used herein is intended to embrace the place on which the useful plants are growing, where the plant propagation materials of the useful plants are sown or where the plant propagation materials of the useful plants will be placed into the soil. An example for such a locus is a field, on which crop plants are growing.

The term "plant propagation material" is understood to denote generative parts of a plant, such as seeds, which can be used for the multiplication of the latter, and vegetative material, such as cuttings or tubers, for example potatoes. There may be mentioned for example seeds (in the strict sense), roots, fruits, tubers, bulbs, rhizomes and parts of plants. Germinated plants and young plants which are to be transplanted after germination or after emergence from the soil, may also be mentioned. These young plants may be protected before transplantation by a total or partial treatment by immersion. Preferably "plant propagation material" is understood to denote seeds.

A further aspect of the instant invention is a method of protecting natural substances of plant and/or animal origin, which have been taken from the natural life cycle, and/or their processed forms against attack of fungi, which comprises applying to said natural substances of plant and/or animal origin or their processed forms a combination of components A) and B) in a synergistically effective amount.

According to the instant invention, the term "natural substances of plant origin, which have been taken from the natural life cycle" denotes plants or parts thereof which have been harvested from the natural life cycle and which are in the freshly harvested form. Examples of such natural substances of plant origin are stalks, leaves, tubers, seeds, fruits or grains. According to the instant invention, the term "processed form of a natural substance of plant origin" is understood to denote a form of a natural substance of plant origin that is the result of a modification process. Such modification processes can be used to transform the natural
substance of plant origin in a more storable form of such a substance (a storage good). Examples of such modification processes are pre-drying, moistening, crushing, comminuting, grounding, compressing or roasting. Also falling under the definition of a processed form of a natural substance of plant origin is timber, whether in the form of crude timber, such as construction timber, electricity pylons and barriers, or in the form of finished articles, such as furniture or objects made from wood.

According to the instant invention, the term "natural substances of animal origin, which have been taken from the natural life cycle and/or their processed forms" is understood to denote material of animal origin such as skin, hides, leather, furs, hairs and the like.

The combinations according the present invention can prevent disadvantageous effects such as decay, discoloration or mold.

A preferred embodiment is a method of protecting natural substances of plant origin, which have been taken from the natural life cycle, and/or their processed forms against attack of fungi, which comprises applying to said natural substances of plant and/or animal origin or their processed forms a combination of components A) and B) in a synergistically effective amount.

A further preferred embodiment is a method of protecting fruits, preferably pomes, stone fruits, soft fruits and citrus fruits, which have been taken from the natural life cycle, and/or their processed forms, which comprises applying to said fruits and/or their processed forms a combination of components A) and B) in a synergistically effective amount.

The combinations of the present invention may also be used in the field of protecting industrial material against attack of fungi. According to the instant invention, the term "industrial material" denotes non-live material which have been prepared for use in industry. For example, industrial materials which are intended to be protected against attack of fungi can be glues, sizes, paper, board, textiles, carpets, leather, wood, constructions, paints, plastic articles, cooling lubricants, aqueous hydraulic fluids and other materials which can be infested with, or decomposed by, microorganisms. Cooling and heating systems, ventilation and air conditioning systems and parts of production plants, for example cooling-water circuits, which may be impaired by multiplication of microorganisms may also be
mentioned from amongst the materials to be protected. The combinations according the
present invention can prevent disadvantageous effects such as decay, discoloration or mold.

The combinations of the present invention may also be used in the field of protecting
technical material against attack of fungi. According to the instant invention, the term
"technical material" includes paper; carpets; constructions; cooling and heating systems;
ventilation and air conditioning systems and the like. The combinations according the
present invention can prevent disadvantageous effects such as decay, discoloration or mold.

The combinations according to the present invention are particularly effective against
powdery mildews; rusts; leafspot species; early blights and molds; especially against
Septoria, Puccinia, Erysiphe, Pyrenophora and Tapesia in cereals; Phakopsora in soybeans;
Hemileia in coffee; Phragmidium in roses; Alternaria in potatoes, tomatoes and cucurbits;
Sclerotinia in turf, vegetables, sunflower and oil seed rape; black rot, red fire, powdery
mildew, grey mold and dead arm disease in vine; Botrytis cinerea in fruits; Monilinia spp. in
fruits and Penicillium spp. in fruits.

The combinations according to the present invention are furthermore particularly effective
against seedborne and soilborne diseases, such as Alternaria spp., Ascochyta spp., Botrytis
cinerea, Cercospora spp., Claviceps purpurea, Cochliobolus sativus, Colletotrichum spp.,
Epicoccum spp., Fusarium graminearum, Fusarium moniliforme, Fusarium oxysporum,
Fusarium proliferatum, Fusarium solani, Fusarium subglutinans, Gäumannomyces graminis,
Helminthosporium spp., Microdochium nivale, Phoma spp., Pyrenophora graminea,
Pyricularia oryzae, Rhizoctonia solani, Rhizoctonia cerealis, Sclerotinia spp., Septoria spp.,
Sphacelotheca reiliana, Tilletia spp., Typhula incarnata, Urocystis occulta, Ustilago spp. or
Verticillium spp.; in particular against pathogens of cereals, such as wheat, barley, rye or
oats; maize; rice; cotton; soybean; turf; sugarbeet; oil seed rape; potatoes; pulse crops, such
as peas, lentils or chickpea; and sunflower.

The combinations according to the present invention are furthermore particularly effective
against post harvest diseases such as Botrytis cinerea, Colletotrichum musae, Curvularia
lunata, Fusarium semitectum, Geotrichum candidum, Monilinia fructicola, Monilinia
fructigena, Monilinia laxa, Mucor piriformis, Penicillium italicum, Penicillium solitum,
Penicillium digitatum or Penicillium expansum in particular against pathogens of fruits, such
as pomefruits, for example apples and pears, stone fruits, for example peaches and plums,
citrus, melons, papaya, kiwi, mango, berries, for example strawberries, avocados, pomegranates and bananas, and nuts.

The amount of a combination of the invention to be applied, will depend on various factors, such as the compounds employed; the subject of the treatment, such as, for example plants, soil or seeds; the type of treatment, such as, for example spraying, dusting or seed dressing; the purpose of the treatment, such as, for example prophylactic or therapeutic; the type of fungi to be controlled or the application time.

It has been found that the use of components B) in combination with the compound of formula I surprisingly and substantially enhance the effectiveness of the latter against fungi, and vice versa. Additionally, the method of the invention is effective against a wider spectrum of such fungi that can be combated with the active ingredients of this method, when used solely.

The weight ratio of A):B) is so selected as to give a synergistic activity. In general the weight ratio of A) : B) is between 2000 : 1 and 1 : 1000, preferably between 100 : 1 and 1: 100, more preferably between 20 : 1 and 1: 50.

The synergistic activity of the combination is apparent from the fact that the fungicidal activity of the composition of A) + B) is greater than the sum of the fungicidal activities of A) and B).

The method of the invention comprises applying to the useful plants, the locus thereof or propagation material thereof in admixture or separately, a synergistically effective aggregate amount of a compound of formula I and a compound of component B).

Some of said combinations according to the invention have a systemic action and can be used as foliar, soil and seed treatment fungicides.

With the combinations according to the invention it is possible to inhibit or destroy the phytopathogenic microorganisms which occur in plants or in parts of plants (fruit, blossoms, leaves, stems, tubers, roots) in different useful plants, while at the same time the parts of plants which grow later are also protected from attack by phytopathogenic microorganisms.
The combinations of the present invention are of particular interest for controlling a large number of fungi in various useful plants or their seeds, especially in field crops such as potatoes, tobacco and sugarbeets, and wheat, rye, barley, oats, rice, maize, lawns, cotton, soybeans, oil seed rape, pulse crops, sunflower, coffee, sugarcane, fruit and ornamentals in horticulture and viticulture, in vegetables such as cucumbers, beans and cucurbits.

The combinations according to the invention are applied by treating the fungi, the useful plants, the locus thereof, the propagation material thereof, the natural substances of plant and/or animal origin, which have been taken from the natural life cycle, and/or their processed forms, or the industrial materials threatened by fungus attack with combination of of components A) and B) in a synergistically effective amount.

The combinations according to the invention may be applied before or after infection of the useful plants, the propagation material thereof, the natural substances of plant and/or animal origin, which have been taken from the natural life cycle, and/or their processed forms, or the industrial materials by the fungi.

The combinations according to the invention are particularly useful for controlling the following plant diseases:
- Alternaria species in fruit and vegetables,
- Ascochyta species in pulse crops,
- Botrytis cinerea in strawberries, tomatoes, sunflower, pulse crops, vegetables and grapes,
- Cercospora arachidica in peanuts,
- Cochliobolus sativus in cereals,
- Colletotrichum species in pulse crops,
- Erysiphe species in cereals,
- Erysiphe cichoracearum and Sphaerotheca fuliginea in cucurbits,
- Fusarium species in cereals and maize,
- Gãumannomyces graminis in cereals and lawns,
- Helminthosporium species in maize, rice and potatoes,
- Hemileia vastatrix on coffee,
- Microdochium species in wheat and rye,
- Phakopsora species in soybean,
- Puccinia species in cereals, broadleaf crops and perrenial plants,
Pseudocercosporella species in cereals,
Phragmidium mucronatum in roses,
Podosphaera species in fruits,
Pyrenophora species in barley,
Pyricularia oryzae in rice,
Ramularia collo-cygni in barley,
Rhizoctonia species in cotton, soybean, cereals, maize, potatoes, rice and lawns,
Rhynchosporium secalis in barley and rye,
Sclerotinia species in lawns, lettuce, vegetables and oil seed rape,
Septoria species in cereals, soybean and vegetables,
Sphacelotheca reiliana in maize,
Tilletia species in cereals,
Uncinula necator, Guignardia bidwellii and Phomopsis viticola in vines,
Urocystis occulta in rye,
Usitilago species in cereals and maize,
Venturia species in fruits,
Monilinia species on fruits,
Penicillium species on citrus and apples.

The combinations according to the invention are preventively and/or curatively valuable active ingredients in the field of pest control, even at low rates of application, which have a very favorable biocidal spectrum and are well tolerated by warm-blooded species, fish and plants. The active ingredients according to the invention which are partially known for their insecticidal action act against all or individual developmental stages of normally sensitive, but also resistant, animal pests, such as insects or representatives of the order Acarina. The insecticidal or acaricidal activity of the combinations according to the invention can manifest itself directly, i.e. in destruction of the pests, which takes place either immediately or only after some time has elapsed, for example during ecdysis, or indirectly, for example in a reduced oviposition and/or hatching rate, a good activity corresponding to a destruction rate (mortality) of at least 50 to 60%.

Examples of the abovementioned animal pests are:
from the order Acarina, for example,
from the order Anoplura, for example,
Haematopinus spp., Linognathus spp., Pediculus spp., Pemphigus spp. and Phylloxera spp.;
from the order Coleoptera, for example,
from the order Diptera, for example,
from the order Heteroptera, for example,
from the order Homoptera, for example,
from the order Hymenoptera, for example,
Acromyrmex, Atta spp., Cephus spp., Diprion spp., Diprionidae, Gilpinia polytoma, Hoplo-
campa spp., Lasius spp., Monomorium pharaonis, Neodiprion spp., Solenopsis spp. and
Vespa spp.;
from the order Isoptera, for example,
Reticulitermes spp.;
from the order Lepidoptera, for example,
Acleris spp., Adoxophyes spp., Aeigeria spp., Agrotis spp., Alabama argillacea, Amylois
spp., Anticarsia gemmatalis, Archips spp., Argyrotaenia spp., Autographa spp., Busseola
fusca, Cadra cautella, Carposina nipponensis, Chilo spp., Choristoneura spp., Clysia ambi-
guella, Cnaphalocrocis spp., Cnephasia spp., Cochylis spp., Coleophora spp., Crocidolomia
binotalis, Cryptophlebia leucotreta, Cydia spp., Diatraea spp., Dicopomorpha castanea, Earias
spp., Ephesia spp., Eucoelia spp., Euboea ambigua, Eupoecilia ambiguella, Eucoilis spp., Eu-
xoia spp., Grapholita spp., Hedya nubiferana, Heliothis spp., Hellula undalis, Hyphantria cunea, Keiferia
lycopersicella, Leuctopera scitella, Lithocolletis spp., Lobesia botrana, Lymantria spp., Ly-
onetia spp., Malacosoma spp., Mamestra brassicae, Manduca sexta, Operophtera spp.,
Ostrinia nubilalis, Pammene spp., Pandemis spp., Panolis flammea, Pectinophora gossypi-
ela, Phthorimaea operculella, Pieris rapae, Pieris spp., Plutella xylostella, Prays spp., Scir-
pophaga spp., Sesamia spp., Sparganothis spp., Spodoptera spp., Synanthedon spp.,
Thaumetopoea spp., Tortrix spp., Trichoplusia ni and Yponomeuta spp.;
from the order Mallophaga, for example,
Damalinae spp. and Trichodectes spp.;
from the order Orthoptera, for example,
Blatta spp., Blattella spp., Gryllotalpa spp., Leucoptera maderae, Locusta spp., Periplaneta
spp. and Schistocerca spp.;
from the order Psocoptera, for example,
Liposcelis spp.;
from the order Siphonaptera, for example,
Ceratophyllus spp., Ctenocephalides spp. and Xenopsylla cheopis;
from the order Thysanoptera, for example,
Frankliniella spp., Hercinothrips spp., Scirtothrips aurantii, Taeniothrips spp., Thrips palmi
and Thrips tabaci;
from the order Thysanura, for example,
Lepisma saccharina;
nematodes, for example root knot nematodes, stem eelworms and foliar nematodes; especially Heterodera spp., for example Heterodera schachtii, Heterodora avenae and Heterodora trifolii; Globodera spp., for example Globodera rostochiensis; Meloidogyne spp., for example Meloidogyne incognita and Meloidogyne javanica; Radopholus spp., for example Radopholus similis; Pratylenchus, for example Pratylenchus neglectans and Pratylenchus penetrans; Tylenchulus, for example Tylenchulus semipenetrans; Longidorus, Trichodorus, Xiphinema, Ditylenchus, Aphelenchoides and Anguina; crucifer flea beetles (Phyllotreta spp.); root maggots (Delia spp.) and cabbage seedpod weevil (Ceutorhynchus spp.).

The combinations according to the invention can be used for controlling, i.e. containing or destroying, animal pests of the abovementioned type which occur on useful plants in agriculture, in horticulture and in forests, or on organs of useful plants, such as fruits, flowers, foliage, stalks, tubers or roots, and in some cases even on organs of useful plants which are formed at a later point in time remain protected against these animal pests.

When applied to the useful plants the compound of formula I is applied at a rate of 5 to 2000 g a.i./ha, particularly 10 to 1000 g a.i./ha, e.g. 50, 75, 100 or 200 g a.i./ha, in association with 1 to 5000 g a.i./ha, particularly 2 to 2000 g a.i./ha, e.g. 100, 250, 500, 800, 1000, 1500 g a.i./ha of a compound of component B), depending on the class of chemical employed as component B).

In agricultural practice the application rates of the combination according to the invention depend on the type of effect desired, and typically range from 20 to 4000 g of total combination per hectare.

When the combinations of the present invention are used for treating seed, rates of 0.001 to 50 g of a compound of formula I per kg of seed, preferably from 0.01 to 10g per kg of seed, and 0.001 to 50 g of a compound of component B), per kg of seed, preferably from 0.01 to 10g per kg of seed, are generally sufficient.

The invention also provides fungicidal compositions comprising a compound of formula I and a compound of component B) in a synergistically effective amount.
The composition of the invention may be employed in any conventional form, for example in the form of a twin pack, a powder for dry seed treatment (DS), an emulsion for seed treatment (ES), a flowable concentrate for seed treatment (FS), a solution for seed treatment (LS), a water dispersible powder for seed treatment (WS), a capsule suspension for seed treatment (CF), a gel for seed treatment (GF), an emulsion concentrate (EC), a suspension concentrate (SC), a suspo-emulsion (SE), a capsule suspension (CS), a water dispersible granule (WG), an emulsifiable granule (EG), an emulsion, water in oil (EO), an emulsion, oil in water (EW), a micro-emulsion (ME), an oil dispersion (OD), an oil miscible flowable (OF), an oil miscible liquid (OL), a soluble concentrate (SL), an ultra-low volume suspension (SU), an ultra-low volume liquid (UL), a technical concentrate (TK), a dispersible concentrate (DC), a wettable powder (WP) or any technically feasible formulation in combination with agriculturally acceptable adjuvants.

Such compositions may be produced in conventional manner, e.g. by mixing the active ingredients with appropriate formulation inerts (diluents, solvents, fillers and optionally other formulating ingredients such as surfactants, biocides, anti-freeze, stickers, thickeners and compounds that provide adjuvancy effects). Also conventional slow release formulations may be employed where long lasting efficacy is intended. Particularly formulations to be applied in spraying forms, such as water dispersible concentrates (e.g. EC, SC, DC, OD, SE, EW, EO and the like), wettable powders and granules, may contain surfactants such as wetting and dispersing agents and other compounds that provide adjuvancy effects, e.g. the condensation product of formaldehyde with naphthalene sulphonate, an alkylaryl/sulphonate, a lignin sulphonate, a fatty alkyl sulphate, and ethoxylated alkylphenol and an ethoxylated fatty alcohol.

A seed dressing formulation is applied in a manner known per se to the seeds employing the combination of the invention and a diluent in suitable seed dressing formulation form, e.g. as an aqueous suspension or in a dry powder form having good adherence to the seeds. Such seed dressing formulations are known in the art. Seed dressing formulations may contain the single active ingredients or the combination of active ingredients in encapsulated form, e.g. as slow release capsules or microcapsules.
In general, the formulations include from 0.01 to 90% by weight of active agent, from 0 to 20% agriculturally acceptable surfactant and 10 to 99.99% solid or liquid formulation inerts and adjuvant(s), the active agent consisting of at least the compound of formula I together with a compound of component B), and optionally other active agents, particularly microbiocides or conservatives or the like. Concentrated forms of compositions generally contain in between about 2 and 80%, preferably between about 5 and 70% by weight of active agent. Application forms of formulation may for example contain from 0.01 to 20% by weight, preferably from 0.01 to 5% by weight of active agent. Whereas commercial products will preferably be formulated as concentrates, the end user will normally employ diluted formulations.

The Examples which follow serve to illustrate the invention, "active ingredient" denoting a mixture of compound I and a compound of component B) in a specific mixing ratio.

### Formulation Examples

<table>
<thead>
<tr>
<th>Wettable powders</th>
<th>a)</th>
<th>b)</th>
<th>c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>active ingredient ([I : \text{comp B}) = 1:3(a), 1:2(b), 1:1(c)])</td>
<td>25%</td>
<td>50%</td>
<td>75%</td>
</tr>
<tr>
<td>sodium lignosulfonate</td>
<td>5%</td>
<td>5%</td>
<td>-</td>
</tr>
<tr>
<td>sodium lauryl sulfate</td>
<td>3%</td>
<td>-</td>
<td>5%</td>
</tr>
<tr>
<td>sodium diisobutylnaphthalenesulfonate</td>
<td>-</td>
<td>6%</td>
<td>10%</td>
</tr>
<tr>
<td>phenol polyethylene glycol ether</td>
<td>-</td>
<td>2%</td>
<td>-</td>
</tr>
<tr>
<td>(7-8 mol of ethylene oxide)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>highly dispersed silicic acid</td>
<td>5%</td>
<td>10%</td>
<td>10%</td>
</tr>
<tr>
<td>Kaolin</td>
<td>62%</td>
<td>27%</td>
<td>-</td>
</tr>
</tbody>
</table>

The active ingredient is thoroughly mixed with the adjuvants and the mixture is thoroughly ground in a suitable mill, affording wettable powders that can be diluted with water to give suspensions of the desired concentration.

### Powders for dry seed treatment

<table>
<thead>
<tr>
<th>Powders for dry seed treatment</th>
<th>a)</th>
<th>b)</th>
<th>c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>active ingredient ([I : \text{comp B}) = 1:3(a), 1:2(b), 1:1(c)])</td>
<td>25%</td>
<td>50%</td>
<td>75%</td>
</tr>
<tr>
<td>light mineral oil</td>
<td>5%</td>
<td>5%</td>
<td>5%</td>
</tr>
<tr>
<td>highly dispersed silicic acid</td>
<td>5%</td>
<td>5%</td>
<td>-</td>
</tr>
<tr>
<td>Kaolin</td>
<td>65%</td>
<td>40%</td>
<td>-</td>
</tr>
</tbody>
</table>
Talcum

The active ingredient is thoroughly mixed with the adjuvants and the mixture is thoroughly ground in a suitable mill, affording powders that can be used directly for seed treatment.

**Emulsifiable concentrate**

- active ingredient \((I : \text{comp B}) = 1:6\) 10 %
- octylphenol polyethylene glycol ether 3 %
- (4-5 mol of ethylene oxide) calcium dodecylbenzenesulfonate 3 %
- castor oil polyglycol ether (35 mol of ethylene oxide) 4 %
- Cyclohexanone 30 %
- xylene mixture 50 %

Emulsions of any required dilution, which can be used in plant protection, can be obtained from this concentrate by dilution with water.

**Dusts**

<table>
<thead>
<tr>
<th></th>
<th>a)</th>
<th>b)</th>
<th>c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active ingredient ((I : \text{comp B}) = 1:6(a), 1:2(b), 1:10(c))</td>
<td>5 %</td>
<td>6 %</td>
<td>4 %</td>
</tr>
<tr>
<td>Talcum</td>
<td>95 %</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Kaolin</td>
<td>-</td>
<td>94 %</td>
<td>-</td>
</tr>
<tr>
<td>Mineral filler</td>
<td>-</td>
<td>-</td>
<td>96 %</td>
</tr>
</tbody>
</table>

Ready-for-use dusts are obtained by mixing the active ingredient with the carrier and grinding the mixture in a suitable mill. Such powders can also be used for dry dressings for seed.

**Extruder granules**

- Active ingredient \((I : \text{comp B}) = 2:1\) 15 %
- Sodium lignosulfonate 2 %
- Carboxymethylcellulose 1 %
- Kaolin 82 %

The active ingredient is mixed and ground with the adjuvants, and the mixture is moistened with water. The mixture is extruded and then dried in a stream of air.

**Coated granules**

- Active ingredient \((I : \text{comp B}) = 1:10\) 8 %
polyethylene glycol (mol. wt. 200) 3 %
Kaolin 89 %

The finely ground active ingredient is uniformly applied, in a mixer, to the kaolin moistened with polyethylene glycol. Non-dusty coated granules are obtained in this manner.

**Suspension concentrate**
active ingredient (I : comp B) = 1:8) 40 %
propylene glycol 10 %
nonylphenol polyethylene glycol ether (15 mol of ethylene oxide) 6 %
Sodium lignosulfonate 10 %
carboxymethylcellulose 1 %
silicone oil (in the form of a 75 % emulsion in water) 1 %
Water 32 %

The finely ground active ingredient is intimately mixed with the adjuvants, giving a suspension concentrate from which suspensions of any desired dilution can be obtained by dilution with water. Using such dilutions, living plants as well as plant propagation material can be treated and protected against infestation by microorganisms, by spraying, pouring or immersion.

**Flowable concentrate for seed treatment**
active ingredient (I : comp B) = 1:8) 40 %
propylene glycol 5 %
copolymer butanol PO/EO 2 %
tristyrenephenole with 10-20 moles EO 2 %
1,2-benzisothiazolin-3-one (in the form of a 20% solution in water) 0.5 %
monoazo-pigment calcium salt 5 %
Silicone oil (in the form of a 75 % emulsion in water) 0.2 %
Water 45.3 %

The finely ground active ingredient is intimately mixed with the adjuvants, giving a suspension concentrate from which suspensions of any desired dilution can be obtained by dilution with water. Using such dilutions, living plants as well as plant propagation material can be treated and protected against infestation by microorganisms, by spraying, pouring or immersion.
Slow Release Capsule Suspension

28 parts of a combination of the compound of formula I and a compound of component B), or of each of these compounds separately, are mixed with 2 parts of an aromatic solvent and 7 parts of toluene diisocyanate/polymethylene-polyphenylisocyanate-mixture (8:1). This mixture is emulsified in a mixture of 1.2 parts of polyvinylalcohol, 0.05 parts of a defoamer and 51.6 parts of water until the desired particle size is achieved. To this emulsion a mixture of 2.8 parts 1,6-diaminohexane in 5.3 parts of water is added. The mixture is agitated until the polymerization reaction is completed.

The obtained capsule suspension is stabilized by adding 0.25 parts of a thickener and 3 parts of a dispersing agent. The capsule suspension formulation contains 28% of the active ingredients. The medium capsule diameter is 8-15 microns.

The resulting formulation is applied to seeds as an aqueous suspension in an apparatus suitable for that purpose.

Biological Examples

A synergistic effect exists whenever the action of an active ingredient combination is greater than the sum of the actions of the individual components.

The action to be expected E for a given active ingredient combination obeys the so-called COLBY formula and can be calculated as follows (COLBY, S.R. "Calculating synergistic and antagonistic responses of herbicide combination", Weeds, Vol. 15, pages 20-22; 1967):

ppm = milligrams of active ingredient (= a.i.) per liter of spray mixture

\[ X = \% \text{ action by active ingredient A) using } p \text{ ppm of active ingredient} \]

\[ Y = \% \text{ action by active ingredient B) using } q \text{ ppm of active ingredient.} \]

According to COLBY, the expected (additive) action of active ingredients A)+B) using

\[ p+q \text{ ppm of active ingredient is } E = X + Y - \frac{X \cdot Y}{100} \]

If the action actually observed \(O\) is greater than the expected action \(E\), then the action of the combination is super-additive, i.e. there is a synergistic effect. In mathematical terms the synergism factor \(SF\) corresponds to \(O/E\). In the agricultural practice an \(SF\) of \(\geq 1.2\) indicates significant improvement over the purely complementary addition of activities (expected
activity), while an SF of \( \leq 0.9 \) in the practical application routine signals a loss of activity compared to the expected activity.

Example B-1: Action against Botrytis cinerea on grapes

a) Fungal growth assay
Conidia of the fungus from cryogenic storage were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24\(^\circ\)C and the inhibition of growth was determined photometrically after 48-72hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

b) Protective Treatment
5 week old grape seedlings cv. Gutedel are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. Two days after application, the grape plants are inoculated by spraying a spore suspension (1\( \times \)10\(^6\) conidia/ml) on the test plants. After an incubation period of 4 days at 21\(^\circ\)C and 95% relative humidity in a greenhouse the disease incidence is assessed. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-2: Action against Septoria tritici on wheat

a) Fungal growth assay
Conidia of the fungus from cryogenic storage were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24\(^\circ\)C and the inhibition of growth was determined photometrically after 72 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

b) Protective Treatment
2 week old wheat plants cv. Riband are treated with the formulated test compound (0.2% active ingredient) in a spray chamber. One day after application, wheat plants are inoculated by spraying a spore suspension (10\( \times \)10\(^5\) conidia/ml) on the test plants. After an incubation period of 1 day at 23\(^\circ\)C and 95% relative humidity, the plants are kept for 16 days at 23\(^\circ\)C and 60% relative humidity in a greenhouse. The disease incidence is assessed 18
days after inoculation. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-3: Action against Pyricularia oryzae on rice

a) Fungal growth assay

Conidia of the fungus from cryogenic storage were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24°C and the inhibition of growth was determined photometrically after 72 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

b) Protective Treatment

Rice leaf segments are placed on agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus. After appropriate incubation the activity of a compound is assessed 96 hrs after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-4: Action against Alternaria solani (early blight)

a) Fungal growth assay

Conidia harvested from a freshly grown colony- of the fungus were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24°C and the inhibition of growth was determined photometrically after 48 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

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<th>Observed control</th>
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### Control of Alternaria solani

*Dosage in mg active ingredient / liter final medium*

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### Control of Alternaria solani

*Dosage in mg active ingredient / liter final medium*

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### Control of *Alternaria solani*

**Dosage in mg active ingredient / liter final medium**

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### Control of *Alternaria solani*

**Dosage in mg active ingredient / liter final medium**

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### Control of *Alternaria solani*

**Dosage in mg active ingredient / liter final medium**

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### b) Protective Treatment

4 week old tomato plants cv. Roter Gnom are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. Two days after application, the tomato plants are inoculated by spraying a spore suspension (2x10^5 conidia/ml) on the test plants. After an incubation period of 3 days at 20°C and 95% relative humidity in a growth chamber the disease incidence is assessed. The fungicide interactions in the combinations are calculated according to COLBY method.

**Example B-5: Action against Pyrenophora teres (Net blotch)**

### a) Fungal growth assay
Conidia of the fungus from cryogenic storage were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24°C and the inhibition of growth was determined photometrically after 48 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

b) Protective Treatment

Barley leaf segments are placed on agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus. After appropriate incubation the activity of a compound is assessed 96 hrs after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-6: Action against Venturia inaequalis on apple

a) Fungal growth assay

Conidia of the fungus from cryogenic storage were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24°C and the inhibition of growth was determined photometrically after 144 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

b) Protective Treatment

4 week old apple seedlings cv. McIntosh are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. One day after application, the apple plants are inoculated by spraying a spore suspension (4x10^5 conidia/ml) on the test plants. After an incubation period of 4 days at 21°C and 95% relative humidity the plants are placed for 4 days at 21°C and 60% relative humidity in a greenhouse. After another 4 day incubation period at 21°C and 95% relative humidity the disease incidence is assessed. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-7: Action against Pythium ultimum (Damping off) - fungal growth assay

Mycelial fragments of the fungus, prepared from a fresh liquid culture, were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24°C and the inhibition of growth was
determined photometrically after 48 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-8: Action against Leptosphaeria nodorum (glume blotch) - fungal growth assay
Conidia of the fungus from cryogenic storage were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24°C and the inhibition of growth was determined photometrically after 48 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

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Control of *Leptosphaeria nodorum*

Dosage in mg active ingredient / liter final medium

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Control of *Leptosphaeria nodorum*

Dosage in mg active ingredient / liter final medium

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Control of *Leptosphaeria nodorum*

Dosage in mg active ingredient / liter final medium

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Example B-9: Action against Pseudocercospora herpotrichoides var. acuformis (eyespot/cereals) - fungal growth assay
Conidia of the fungus from cryogenic storage were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24°C and the inhibition of growth was determined photometrically after 72 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-10: Action against Ustilago maydis (corn smut) - fungal growth assay
Conidia of the fungus from cryogenic storage were directly mixed into nutrient broth (PDB potato dextrose broth). After placing a (DMSO) solution of the test compounds into a microtiter plate (96-well format) the nutrient broth containing the fungal spores was added. The test plates were incubated at 24°C and the inhibition of growth was determined photometrically after 48 hrs. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-11: Action against Phytophthora infestans (late blight) on tomato – protective treatment
Tomato leaf disks are placed on water agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus. After appropriate incubation the activity of a compound is assessed 96 hrs after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-12: Action against Plasmopara viticola (downy mildew) on grape vines – protective treatment
Grape vine leaf disks are placed on agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus. After appropriate incubation the activity of a compound is assessed 7 days after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-13: Action against Botrytis cinerea (Grey mould) on beans – protective treatment
Bean leaf disks are placed on agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus.
After appropriate incubation the activity of a compound is assessed 96 hrs after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

**Example B-14: Action against Erysiphe graminis f.sp. hordei (Barley powdery mildew) on barley – protective treatment**

Barley leaf segments are placed on agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus. After appropriate incubation the activity of a compound is assessed 96 hrs after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

**Example B-15: Action against Erysiphe graminis f.sp. tritici (Wheat powdery mildew) on barley – protective treatment**

Barley leaf segments are placed on agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus. After appropriate incubation the activity of a compound is assessed 96 hrs after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

**Example B-16: Action against Puccinia recondita (Brown rust) on wheat**

a) **Protective Treatment of leaf segments**

Wheat leaf segments are placed on agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus. After appropriate incubation the activity of a compound is assessed 9 days after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

b) **Protective Treatment of plants**

1 week old wheat plants cv. Arina are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. One day after application, the wheat plants are inoculated by spraying a spore suspension (1x10⁵ uredospores/ml) on the test plants. After an incubation period of 2 days at 20°C and 95% relative humidity the plants are kept in a greenhouse for 8 days at 20°C and 60% relative humidity. The disease incidence is assessed 10 days after inoculation. The fungicide interactions in the combinations are calculated according to COLBY method.
Example B-17: Action against Septoria nodorum on wheat

a) Protective Treatment of leaf segments
Wheat leaf segments are placed on agar in multiwell plates (24-well format) and sprayed with test solutions. After drying, the leaf disks are inoculated with a spore suspension of the fungus. After appropriate incubation the activity of a compound is assessed 96 hrs after inoculation as preventive fungicidal activity. The fungicide interactions in the combinations are calculated according to COLBY method.

b) Protective Treatment of plants
1 week old wheat plants cv. Arina are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. One day after application, the wheat plants are inoculated by spraying a spore suspension (5x10^6 conidia/ml) on the test plants. After an incubation period of 1 day at 20°C and 95% relative humidity the plants are kept for 10 days at 20°C and 60% relative humidity in a greenhouse. The disease incidence is assessed 11 days after inoculation. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-18: Action against Podosphaera leucotricha (Powdery mildew) on apple – protective treatment
5 week old apple seedlings cv. McIntosh are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. One day after application, the application apple plants are inoculated by shaking plants infected with apple powdery mildew above the test plants. After an incubation period of 12 days at 22°C and 60% relative humidity under a light regime of 14/10 hours (light/dark) the disease incidence is assessed. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-19: Action against Erysiphe graminis (Powdery mildew) on barley – protective treatment
1 week old barley plants cv. Regina are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. One day after application, the barley plants are inoculated by shaking powdery mildew infected plants above the test plants. After an incubation period of 6 days at 20°C / 18°C (day/night) and 60% relative humidity in a greenhouse the disease incidence is assessed. The fungicide interactions in the combinations are calculated according to COLBY method.

Example B-20: Action against Botrytis cinerea on tomatoes – protective treatment
4 week old tomato plants cv. Roter Gnom are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. Two days after application, the tomato plants are inoculated by spraying a spore suspension (1x10^5 conidia/ml) on the test plants. After an incubation period of 4 days at 20°C and 95% relative humidity in a growth chamber the disease incidence is assessed. The fungicide interactions in the combinations are calculated according to COLBY method.

**Example B-21: Action against Helminthosporium teres (Net blotch) on barley – protective treatment**

1 week old barley plants cv. Regina are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. Two days after application, the barley plants are inoculated by spraying a spore suspension (3x10^4 conidia/ml) on the test plants. After an incubation period of 4 days at 20°C and 95% relative humidity in a greenhouse the disease incidence is assessed. The fungicide interactions in the combinations are calculated according to COLBY method.

**Example B-22: Action against Ucinula necator (Powdery mildew) on grapes – protective treatment**

5 week old grape seedlings cv. Gutedel are treated with the formulated test compound (0.02% active ingredient) in a spray chamber. One day after application, the grape plants are inoculated by shaking plants infected with grape powdery mildew above the test plants. After an incubation period of 7 days at 26°C and 60% relative humidity under a light regime of 14/10 hours (light/dark) the disease incidence is assessed. The fungicide interactions in the combinations are calculated according to COLBY method.

The combinations according to the invention exhibit good activity in all of the above examples.

A further aspect of the instant invention is a method of controlling phytopathogenic diseases on useful plants or plant propagation material thereof, which comprises applying to said plant propagation material, preferably seeds, a fungicidally effective amount of a compound of formula I, especially a compound of formula Ia, Ib, Ic, Id or le, or a tautomer of such a compound.
WHAT IS CLAIMED IS:

1. A method of controlling phytopathogenic diseases on useful plants or on propagation material thereof, which comprises applying to the useful plants, the locus thereof or propagation material thereof a combination of components A) and B) in a synergistically effective amount, wherein component A) is a compound of formula I

\[
\begin{array}{c}
\text{N} \\
\text{R}_1 \\
\text{CH}_3 \\
\text{C}_6 \text{alkyl or trifluoromethyl and} \\
\text{R}_2 \\
\text{C}_1 - \text{C}_6 \text{alkyl, } \text{C}_1 - \text{C}_4 \text{alkoxy-} \text{C}_1 - \text{C}_6 \text{alkyl or } \text{C}_1 - \text{C}_6 \text{haloalkyl; or a tautomer of such a compound; and} \\
\text{component B) is a compound selected from the group consisting of a strobilurin fungicide; an azole fungicide; a phenyl pyrrole fungicide; an anilino-pyrimidine fungicide; a morpholine fungicide; a compound of formula F-1}
\end{array}
\]

a compound of formula F-2
a racemic compound of formula F-3 (syn)

(F-3);

a racemic compound of formula F-4 (anti)

(F-4);

a compound of formula F-5
which represents an epimeric mixture of the racemic compounds of formula F-3 (syn) and F-4 (anti), wherein the ratio of racemic compounds of formula F-3 (syn) to racemic compounds of formula F-4 (anti) is from 1000 : 1 to 1 : 1000;
a compound of formula F-6

a racemic compound of formula F-7 (trans)

a racemic compound of formula F-8 (cis)
a compound of formula F-9

which represents a mixture of racemic compounds of formula F-7 (trans) and F-8 (cis), wherein the ratio of racemic compounds of formula F-7 (trans) to racemic compounds of formula F-8 (cis) is from 2:1 to 100:1;

Trinexapac-Ethyl; Chlormequat chloride; Ethephon; Prohexadione calcium; Mepiquat chloride; Proquinazid; Chlorothalonil; Famoxadone; Fenamidone; Benomyl; Carbendazim; Fuberidazole; Thiabendazole; Thiophanate; Thiophanate-methyl; Chlozolinate; Iprodione; Procymidine; Vinclozolin; Bitertanol; Fenarimol; Nuarimol; Pyrifoxon; Triforine; Benalaxyl; Furalaxy; Metalaxyl; Mefenoxam (Metalaxyl-M); Ofurace; Oxadixyl; Edifenphos; Ifoprobenfos (IBP); Isoprothiolane; Pyrazophos; Benodanil; Carboxin; Fenfuram; Flutolanil; Furametpyr; Mepronil; Oxycarboxin; Thifluзamide; Bupirimutate; Dimethirimol; Ethirimol; Diethofencarb; Famoxadone; Fenamidone; Quinoxyfen; Biphentyl; Chloroneb; Dicloran; Etridiazole; Quintozene (PCNB); Tecnazene (TCNB); Tolclofos-methyl; Dimethomorph; Carpropanid; Diclocymet; Fenoxanil; Fthalide; Pyroquilon; Tricyclazole; Fenhexamid; Polyoxyin; Pencycuron; Cyazofamid; Zoxamide; Basicidin-S; Kasugamycin; Streptomycin; Validamycin; Cymoxanil; Iodocarb; Propamocarb; Prothiocarb; Dinocap; Fluazinam; Fentin acetate; Fentin chloride; Fentin hydroxide; Oxolinic acid; Hymexazole; Ocithilone; Fosetyl-Aluminium; Phosphorsäure; Teclofluthalam (Bactericid); Triazoxide; Flusulfamide; Ferimzone; Diclomezine; Anilazine; Arsenates; Captaloil; Captan; Chlorothalonil; Copper (diverse salts); Copper Ammoniumcarbonate; Copper octanoate; Copper oleate; Copper sulphate; Copper hydroxide; Dichlofluanid; Dithianon; Dodine; Ferbam; Folpet; Guazatine; Iminoctadine; Mancozeb; Maneb; Mercury; Metiram; Propineb; Sulphur; Thiram; Tolyfluand; Zineb; Ziram; Acibenzolar-S-methyl; Probenazole; Bentiavalcarb; Iprovalicarb; Diflumetorim; Ethaboxam; Flusulfamide; Methasulfocarb; Silthiofam; Bacillus pumilus GB34; Bacillus pumilus strain QST 2808; Bacillus subtilis; Bacillus subtilis + PCNB + Metalaxyl; Cadmiumchlorid; Carbendisulfid; Bordeaux mixture; Cedar; Chlorine; Cinnamonaldehyde; Cycloheximide; Fenaminosulf; Fenamiphos; Dichloropropene; Dichlore; Formaldehyde; Gliocladium virens
- 113 -

GL-21; Glyodin; Hexachlorobenzene; Iprovalicarb; Manganese dimethylidithiocarbamate; Mercuric chloride; Nabam; Neem oil (hydrophobic extract); Oxytetracycline; Oxythioquinox; Paraformaldehyde; Pentachloronitrobenzene; Pentachlorophenol; paraffin oil; Polyoxin D; zinc salt; sodium bicarbonate; potassium bicarbonate; sodium diacetate; sodium propionate; TCMTB; Benalaxyl-M; Boscalid; Hexaconazole; Metrafenone; Oxine Copper; Penthiopyrad; Perfurazolate; Tolyfluanid; Terramycin; Trichoderma; Trichoderma harzianum; Triphenyltin hydroxide and Xanthomonas campestris subsp. Vescicatoria; Paclorbutrazol; 1,1-bis(4-chlorophenyl)-2-ethoxyethanol; 2,4-dichlorophenyl benzenesulfonate; 2-fluoro-N-methyl-N-1-naphthylacetamide; 4-chlorophenyl phenyl sulfone; abamectin; acequinocyl; acetoprole; acrinathrin; aldicarb; aldoxycarb; alpha-cypermethrin; amidithion; amidoflumet; amidothioate; amiton; amiton hydrogen oxalate; amitraz; aramite; arsenous oxide; AVI 382; AZ 60541; azinphos-ethyl; azinphos-methyl; azobenzene; azocyclotin; azothoate; benomy; benoxafos; benzoximate; benzyl benzoate; bifenthrin; bifenthrin; binapacryl; brofenvalerate; bromocyclen; bromophos; bromophos-ethyl; bromopropylate; buprofezin; butocarboxim; butoxycarboxim; butylpyridaben; calcium polysulfide; camphechlor; carbanolate; carbaryl; carbofuran; carbofenothion; CGA 50’439; chlorimethionat; chlorbenside; chlordimeform; chlordimeform hydrochloride; chlorfenapyr; chlorfenathol; chlorfenoson; chlorfenosulphide; chlorfenvinphos; chlorfenizolate; chloromebuform; chloromethuron; chloropropylate; chlorpyrifos; chlorpyrifos-methyl; chlorthiophos; cinerin I; cinerin II; cinerins; clofentazine; closantel; coumaphos; crotamiton; crotroxyphos; cufraneb; cyanthoate; cyhalothrin; cyhexatin; cypermethrin; DCPM; DDT; demephion; demephion-O; demephion-S; demeton; demeton-methyl; demeton-O; demeton-O-methyl; demeton-S; demeton-S-methyl; demeton-S-methylsulphon; diafenthion; dialifos; diazinon; dichlofluanid; dichlorvos; dicliphos; dicofol; dicrotophos; dienochlor; dimefox; dimethoate; dinactin; dinex; dinex-dicloxine; dinobuten; dinocap; dinocap-4; dinocap-6; dinocap; dinopenton; dinosulfon; dinoterbon; dioxathion; diphenyl sulfone; disulfiram; disulfoton; DNOC; dofenapyn; doramectin; endosulfan; endothion; EPN; eprinomectin; ethion; ethoate-methyl; etoxazol; etrimfos; fenazaflor; fenazaquin; fenbutatin oxide; fenothiocarb; fenpropathrin; fenpyrad; fenpyroximate; fenson; fentifanil; fenvalerate; fipronil; fluacrypyrim; fluazuron; flubenzimine; flucycloxuron; flucytthrinate; fluenetil; flufenoxuron; flumethrin; fluorbenside; fluvialate; FMC 1137; fortemanate; fortemanate hydrochloride; formation; formparanate; gamma-HCH; glyodin; halifenprox; heptenophos; hexadeacyl cyclopropenecarboxylate; hexythiazox; iodomethane; isocarbophos; isopropyl O-(methoxyaminothiophosphoryl)salicylate; ivermectin; jasmolin I; jasmolin II; jodfenphos; lindane; lufenuron; malathion; malonoben; mecarbam; mephosfolan;
mesulfen; methacrifos; methamidophos; methidathion; methiocarb; methomyl; methyl bromide; metolcarb; mevinphos; mexacarbate; milbemectin; milbemycin oxime; mipafox; monocrotophos; morphothion; moxidectin; naled; NC-184; nifuradine; nikkomycins; nitrilacarb; nitrilacarb 1:1 zinc chloride complex; NNI-0101; NNI-0250; omethoate; oxamyl; oxydeprofos; oxydisulfoton; pp'-DDT; parathion; permethrin; petroleum oils; phenkapton; phenthoate; phorate; phosalon; phosfalan; phosmet; phosphamidon; phoxim; pirimiphos-methyl; polychloroterpene; polynactins; proclonol; profenofos; promacyl; propargite; propetamphos; propoxur; prothidithion; prothoate; pyrethrin I; pyrethrin II; pyrethrins; pyridaben; pyridaphenthion; pyrimidifen; pyrimate; quinalphos; quintetofos; R-1492; RA-17; rotenone; schradan; sebufos; selamectin; SI-0009; sophamide; spirodiclofen; spiromesifen; SSI-121; sulfiram; sulfuranid; sulfotep; sulfur; SZI-121; tau-fluvalinate; tebufenpyrad; TEPP; terbam; tetrachlorvinphos; tetradifon; tetrachlor; tetrasul; thiafenox; thiocarboxime; thioflanox; thiometon; thioquinox; thuringiensi; triamiphos; triarathene; triazophos; triazuron; trichlorfon; trifenofos; trinactin; vamidothion; vaniliprole; YI-5302; bethoxazin; copper dioctanoate; copper sulfate; cybutryne; dichlone; dichlorphen; endothal; fentin; hydrated lime; nabam; quinoclamine; quinoamid; simazine; triphenyltin acetate; triphenyltin hydroxide; abamectin; crufoate; doramectin; emamectin; emamectin benzoate; eprinomectin; ivermectin; milbemycin oxime; moxidectin; piperazine; selamectin; spinosad; thiophanate; chloralose; endrin; fenthion; pyridin-4-amine; strychnine; 1-hydroxy-1H-pyridine-2-thione; 4-(quinoxalin-2-ylamino)benzenesulfonamide; 8-hydroxyquinoline sulfate; bronopol; copper dioctanoate; copper hydroxide; cresol; dichlorphen; dipyrithione; dodicin; fenaminosulf; formaldehyde; hydrargaphen; kasugamycin; kasugamycin hydrochloride hydrate; nickel bis(dimethylthiocarbamate); nitrapyrin; ochthilone; oxolinic acid; oxytetracycline; potassium hydroxyquinoline sulfate; probenazole; streptomycin; streptomycin sesquisulfate; teclofamil; thiomersal; iodomethane; methyl bromide; apholate; bisazir; busulfan; diflubenzuron; dimatit; hemel; hempha; metepa; methiotepa; methyl apholate; morzid; penfluron; tepa; thiocarboxime; thiota; tretamine; urepeta; (E)-dec-5-en-1-yl acetate with (E)-dec-5-en-1-ol; (E)-tridec-4-en-1-yl acetate; (E)-6-methylhept-2-en-4-ol; (E,Z)-tetradeca-4,10-dien-1-yl acetate; (Z)-dodec-7-en-1-yl acetate; (Z)-hexadec-11-enal; (Z)-hexadec-11-en-1-yl acetate; (Z)-hexadec-13-en-11-yn-1-yl acetate; (Z)-icos-13-en-10-one; (Z)-tetradec-7-en-1-ol; (Z)-tetradec-9-en-1-ol; (Z)-tetradec-9-en-1-yl acetate; (9Z,12Z)-dodeca-7,9-dien-1-yl acetate; (9Z,11E)-tetradeca-9,11-dien-1-yl acetate; (9Z,12E)-tetradeca-9,12-dien-1-yl acetate; 14-methyloctadec-1-ene; 4-methylnonan-5-ol with 4-methylnonan-5-one; alpha-multistriatin; brévicom; codleure; codleusene; cuelure; disparlure; dodec-8-ene-
1-yl acetate; dodec-9-en-1-yl acetate; dodeca-8,10-dien-1-yl acetate; dominicalure; ethyl 4-methyloctanoate; eugenol; frontalin; gossypure; grandlure; grandlure I; grandlure II; grandlure III; grandlure IV; hexalure; ipsdienol; ipsenol; japonilure; lineatin; litlure; looplure; medlure; megatomoic acid; methyl eugenol; muscalure; octadeca-2,13-dien-1-yl acetate; octadeca-3,13-dien-1-yl acetate; orfralure; oryctalure; ostramone; siglure; sordidin; sulcatol; tetradec-11-en-1-yl acetate; trimedlure; trimedlure A; trimedlure B₁; trimedlure B₂; trimedlure C; trunc-call; 2-(octylthio)ethanol; butopyronoxyl; butoxy(polypropylene glycol); dibutyl adipate; dibutyl phthalate; dibutyl succinate; diethyltoluamide; dimethyl carbonate; dimethyl phthalate; ethyl hexanediol; hexamide; methoquin-buty1; methylneodecanamide; oxamate; picaridin; 1,1-dichloro-1-nitroethane; 1,1-dichloro-2,2-bis(4-ethylphenyl)ethane; 1,2-dichloropropene; 1,2-dichloropropene with 1,3-dichloropropene; 1-bromo-2-chloroethane; 2,2,2-trichloro-1-(3,4-dichlorophenyl)ethyl acetate; 2,2-dichlorovinyl 2-ethylsulfinylethyl methyl phosphate; 2-(1,3-dithiolan-2-yl)phenyl dimethylcarbamate; 2-(2-butoxyethoxy)ethyl thiocyanate; 2-(4,5-dimethyl-1,3-dioxolan-2-yl)phenyl methylcarbamate; 2-(4-chloro-3,5-xylyloxy)ethanol; 2-chlorovinyl diethyl phosphate; 2-imidazolidone; 2-isovalerylindan-1,3-dione; 2-methyl(prop-2-ynyl)aminophenyl methylcarbamate; 2-thiocyanatoethyl laurate; 3-bromo-1-chloroprop-1-ene; 3-methyl-1-phenylpyrazol-5-yl dimethylcarbamate; 4-methyl(prop-2-ynyl)amino-3,5-xylyl methylcarbamate; 5,5-dimethyl-3-oxocyclohex-1-enyl dimethylcarbamate; abamectin; acephate; acetamiprid; acethion; acetoprole; acrinathrin; acrylonitrile; alanycarb; aldicarb; aldoxycarb; aldrin; allethrin; allosamidin; allyxycarb; alphacypermethrin; alpha-ec dysone; aluminium phosphate; amidithion; amidothioate; aminocarb; amiton; amiton hydrogen oxalate; amitraz; anabasine; aethidion; AVI 382; AZ 60541; azadirachtin; azamethiphos; azinphos-ethyl; azinphos-methyl; azothoate; Bacillus thuringiensis delta endotoxins; barium hexafluorosilicate; barium polysulfide; barthrin; BAS 320 I; Bayer 22/190; Bayer 22408; bendiocarb; benfuracarb; bensultap; beta-cyfluthrin; beta-cypermethrin; bifenthrin; bioallethrin; bioallethrin S-cyclopentenyl isomer; bioethanomethrin; biopermethrin; bioresmethrin; bis(2-chloroethyl) ether; bistrifluron; borax; brofenvalerate; bromfenvinfos; bromocyclen; bromo-DDT; bromophos; bromophos-ethyl; bufencarb; buprofezin; butacarb; butathiofos; butocarboxim; butonate; butoxycarboxim; butylpyridaben; cadusafos; calcium arsenate; calcium cyanide; calcium polysulfide; camphechlor; carbonolate; carbaryl; carbofuran; carbon disulfide; carbon tetrachloride; carbothothonth; carbosulfan; cartap; cartap hydrochloride; cevadine; chlorbicyclen; chlordane; chlodecone; chlormethyform; chlordimeform hydrochloride; chlorethoxyfos; chlorfenapry; chlorfenvinphos; chlorfluazuron; chlorimephos; chlorform; chloropicrin; chlorphoxim; chlorprazophos;
chlorpyrifos; chlorpyrifos-methyl; chlorthiofos; chromafenozide; cinerin I; cinerin II; cinerins; cis-resmethrin; cismethrin; clocythrin; cloethocarb; closantel; clothianidin; copper acetoxarsenate; copper arsenate; copper oleate; coumaphos; coumithoate; crotamiton; crotopyrophos; crufomate; cryolite; CS 708; cyanofenphos; cyanothoate; cyclethrin; cycloprothrin; cyfluthrin; cyhalothrin; cypermethrin; cyphenothrin; cyromazine; cythioate; d-limonene; d-tetramethrin; DAEP; dazomet; DDT; decarbofuran; deltamethrin; demephion; demephion-O; demephion-S; demeton; demeton-methyl; demeton-O; demeton-O-methyl; demeton-S; demeton-S-methyl; demeton-S-methylsulphon; diafenthion; dialifos; dianidafos; diazinon; dicapthon; dichlorfenthion; dichlorvos; dicliphos; dicresyl; dicrotophos; dicyclanil; dielidrin; diethyl 5-methylpyrazol-3-yl phosphate; diflubenzuron; dilor; dimefluthrin; dimethox; dimetan; dimethoate; dimethrin; dimethylvinphos; dimetilan; dinex; dinex-diclexine; dinoprop; dinosam; disocarb; disofuran; diofenolan; diosabenzofos; dioxacarb; dioxathion; disulfoton; dithiocrofos; DNOC; doramectin; DSP; ecdysonester; EI 1642; emamectin; emamectin benzoate; EMPC; empenthrin; endosulfan; endothion; endrin; EPBP; EPN; epofenonane; epirinocetin; esfenvalerate; etaphos; ethiofencarb; ethion; ethiprole; ethoate-methyl; ethoprophos; ethyl formate; ethyl-DDD; ethylene dibromide; ethylene dichloride; ethylene oxide; etofenprox; etrimfos; EXD; famphur; fenamiphos; fenazaflor; fenchlorphos; fenethacarb; fenfluthrin; fenitrothion; fenobucarb; fenoxacrin; fenoxycarb; fenpirithrin; fenpropathrin; fenpyrad; fensulfothion; fenthion; fenthion-ethyl; fenvalerate; fipronil; flinonicamid; flucofuron; flucycloxuron; flucythrinate; flufenetil; flufenoxuron; flufenprox; flumethrin; fluvalinate; FMC 1137; fonofos; formetanate; formetanate hydrochloride; formothion; formparanate; fosmethian; fospirate; fosthiazate; fosthietan; furathiocarb; furethrin; gamma-cyhalothrin; gamma-HCH; guazatine; guazatine acetates; GY-81; halifenprox; halofenozide; HCH; HEOD; heptachlor; heptenophos; heterophos; hexafiumuron; HHDN; hydramethylnon; hydrogen cyanide; hydroprone; hyquin carb; imidacloriprid; imiprothrin; indoxacarb; iodomethane; IPSP; isazofos; isobenzan; isocarbo phos; isodrin; isofenphos; isolane; isopropcarb; isopropyl O-(methoxyaminothiophosphoryl)-salicylate; isoprothiolane; isothioate; isoxathion; ivermectin; jasmolin I; jasmolin II; jodfenphos; juvenile hormone I; juvenile hormone II; juvenile hormone III kelevan; kinoprene; lambda-cyhalothrin; lead arsenate; leptophos; lindane; lirimfos; lufenuron; lyhidathion; m-cumeryl methylcarbamate; magnesium phosphate; malathion; malonobenz; mazidox; mecarbam; mecarphon; menazon; mephostalan; mercurous chloride; mesulfenfos; metam; metam-potassium; metan-sodium; methacrifos; methamidophos; methanesulfonyl fluoride; methidathion; methiocarb; methocrotophos; methomyl; methoprene; methoquin-butyl;
methothrin; methoxychlor; methoxyfenoizide; methyl bromide; methyl isothiocyanate; methylchloroform; methylene chloride; metofluthrin; metolcarb; metoxadiazone; mevinphos; mexacarbate; milbemectin; milbemycin oxime; mipafox; mirex; monocrotophos; morphothion; moxidectin; naftalofos; naled; naphthalene; NC-170; NC-184; nicotine; nicotine sulfate; nifluridine; nitenupyram; nithiazine; nitrilcarb; nitrilcarb 1:1 zinc chloride complex; NNI-0101; NNI-0250; nornicotine; novaluron; nufamfluron; O-2,5-dichloro-4-iodophenyl O-ethyl ethylphosphonothioate; O,O-diethyl O-4-methyl-2-oxo-2H-chromen-7-yl phosphorothioate; O,O-diethyl O-6-methyl-2-propylpyrimidin-4-yl phosphorothioate; O,O,O',O'-tetrapropyl dithiopyrophosphate; oleic acid; omethoate; oxamyl; oxydemeton-methyl; oxydeprofos; oxydisulfoton; pp'DDT; para-dichlorobenzene; parathion; parathion-methyl; penfluron; pentachlorophenol; pentachlorophenyl laurate; permethrin; petroleum oil; PH 60-38; phenkapton; phenothrin; phenthoate; phorate; phosalone; phosfolan; phosmet; phosnichlor; phosphamidon; phosphine; phoxim; phoxim-methyl; pirimetaphos; pirimicar; pirimiphos-ethyl; pirimiphos-methyl; polychlordicyclopentadiene isomers; polychloroterpenes; potassium arsenite; potassium thiocyanate; prallethrin; precocene I; precocene II; precocene III; primidophos; profenofos; profluthrin; promacyl; promecarb; propaphos; propetamphos; propoxur; prothidathion; prothiofos; prothoate; protrifenbute; pymetrozine; pyraclofos; pyrazophos; pyresmethrin; pyrethrin I; pyrethrin II; pyrethrins; pyridaben; pyridalyl; pyridaphenthion; pyrimidifen; pyrimidinate; pyriproxyfen; quassia; quinalphos; quinalphos-methyl; quinothion; quintiofos; R-1492; rafosanide; resmethrin; rotenone; RU 15525; RU 25475; ryania; ryanodine; sabadilla; schradan; sebufos; selamectin; SI-0009; silafluofen; SN 72129; sodium arsenite; sodium cyanide; sodium fluoride; sodium hexafluorosilicate; sodium pentachlorophenoxyde; sodium selenate; sodium thiocyanate; sophamide; spinosad; spiromesifen; sulcofuron; sulcofuron-sodium; sulfuramid; sulfotep; sulfuryl fluoride; sulprofos; tar oils; tau-fluvalinate; tazimcarb; TDE; tebufenoizde; tebufenpyrad; tebupirimfos; teflubenzuron; tefluthrin; temephos; TEPP; terallethrin; terbam; terbufos; tetrachloroethane; tetrachlorvinphos; tetramethrin; theta-cypermethrin; thiacloprid; thiafenox; thiamethoxam; thicrofos; thiocarboxime; thiocyclam; thiocyclam hydrogen oxalate; thiodicarb; thiofanox; thiometon; thionazin; thiosultap; thiosultap-sodium; thuringiensin; tolfenpyrad; tralomethrin; transfluthrin; transpermethrin; triamiphos; triazamate; triazophos; triazuron; trichlorfon; trichlorometaphos-3; trichloronat; trifenofos; triflumuron; trimethacarb; triprene; vamidothion; vaniliprole; veratridine; veratrine; XMC; xylylcarb; YI-5302; zeta-cypermethrin; zetamethrin; zinc phosphide; zolaprofos und ZXI 8901; a compound of formula A-1
a compound of formula A-2

a compound of formula A-3

a compound of formula A-4
a compound of formula A-5

\[ \text{A-5} \]

a compound of formula A-6

\[ \text{A-6} \]

a compound of formula A-7

\[ \text{A-7} \]

a compound of formula A-8
a compound of formula A-9

a compound of formula A-10

a compound of formula A-11
a compound of formula A-12

![Chemical Structure A-12](image)

(A-12);

a compound of formula A-13

![Chemical Structure A-13](image)

(A-13);

a compound of formula A-14

![Chemical Structure A-14](image)

(A-14);

a compound of formula A-15

![Chemical Structure A-15](image)

(A-15);
a compound of formula A-15A

\[
\begin{align*}
\text{H}_2\text{N} & \text{C(CH}_3\text{)}_2\text{CH}_2\text{SCH}_3 \\
\text{H}_3\text{C} & \text{C} \text{CF(CF}_3\text{)}_2
\end{align*}
\]

(A-15A);

a compound of formula (A-16)

\[
\begin{align*}
\text{H}_3\text{C} & \text{O} \text{N} \text{N} \text{H} \text{N} \text{CH}_3 \\
\text{H}_3\text{C} & \text{O} \text{N} \text{N} \text{H} \text{N} \text{CH}_3
\end{align*}
\]

(A-16);

a compound of formula (A-17)

\[
\begin{align*}
\text{H}_3\text{C} & \text{O} \text{N} \text{N} \text{H} \text{N} \text{CH}_3 \\
\text{H}_3\text{C} & \text{O} \text{N} \text{N} \text{H} \text{N} \text{CH}_3
\end{align*}
\]

(A-17);

a compound of formula (A-18)
a compound of formula (A-19)

(a-18);

a compound of formula (A-20)

(a-19);

a compound of formula (A-20)

(a-20);

a compound of formula (A-21)
a compound of formula (A-21);

a compound of formula (A-22);

a compound of formula (A-23);

a compound of formula (A-24)
a compound of formula (A-25)

a compound of formula (A-26)

bis(tributyltin) oxide; bromoacetamide; calcium arsenate; cloethocarb; copper acetoarsenite; copper sulfate; fentin; ferric phosphate; metaldehyde; methiocarb; niclosamide; niclosamide-olamine; pentachlorophenol; sodium pentachlorophenoxide; tazimcarb; thiodicarb; tributyltin
oxide; trifenmorph; trimethacarb; triphenylin acetate; triphenylin hydroxide; 1,2-dibromo-3-chloropropane; 1,2-dichloropropane; 1,2-dichloropropane with 1,3-dichloropropene; 1,3-dichloropropene; 3,4-dichlorotetrahydrothiophene 1,1-dioxide; 3-(4-chlorophenyl)-5-methylrhodanine; 5-methyl-6-thioxo-1,3,5-thiadiazinan-3-yacetic acid; 6-isopentenylaminopurine; abamectin; acetoprole; alanycarb; aldicarb; aldoxycarb; AZ 60541; benclothiaz; benomyl; butylpyridaben; cadusafos; carbofuran; carbon disulfide; carbosulfan; chloropicrin; chlorpyrifos; cloethocarb; cytokinins; dazomet; DBCP; DCIP; diamidafos; dichloflenthion; dicliphos; dimethoate; doramectin; emamectin; emamectin benzoate; eprinomectin; ethoprophos; ethylene dibromide; fenamiphos; fenpyrad; fensulfothion; fosthiazate; fosthietan; furfural; GY-81; heterophos; iodomethane; isamidofos; isazofos; ivermectin; kinetin; mecarphon; metam; metam-potassium; metam-sodium; methyl bromide; methyl isothiocyanate; milbemycin oxime; moxidectin; Myrothecium verrucaria composition; NC-184; oxamyl; phorate; phosphamidon; phosphocarb; sebufos; selamectin; spinosad; terban; terbufos; tetrachlorothiophene; thiafenox; thionazin; triazophos; triazuron; xlenols; YI-5302; zeatin; potassium ethylxanthate; nitrpyrin; acibenzolar; acibenzolar-S-methyl; probenazole; Reynoutria sachalinensis extract; 2-isovalerylindan-1,3-dione; 4-(quinoxalin-2-ylamino)benzenesulfonamide; alpha-chlorohydrin; aluminium phosphide; antu; arsenous oxide; barium carbonate; bisthiosemi; brodifacoum; bromadiolone; bromethalin; calcium cyanide; chloralose; chlorophacinone; cholecalciferol; coumachlor; coumafuryl; coumatetralyl; crimidine; difenacoum; difethialone; diphenacine; ergocalciferol; flocoumafen; fluoroacetamide; flupropadine; flupropadine hydrochloride; gamma-HCH; HCH; hydrogen cyanide; iodomethane; lindane; magnesium phosphide; methyl bromide; norbormide; phosacetim; phosphine; phosphorus; pindone; potassium arsenite; pyrinuron; scilliroside; sodium arsenite; sodium cyanide; sodium fluoroacetate; strychnine; thallium sulfate; warfarin; zinc phosphide; 2-(2-butoxyethoxy)ethyl piperonylate; 5-(1,3-benzodioxol-5-yl)-3-hexylcyclohex-2-enone farnesol with nerolidol; MB-599; MGK 264; piperonyl butoxide; piprotal; propyl isome; S421; sesamex; sesamolin; sulfoxide; anthraquinone; chloralose; copper naphthenate; copper oxychloride; diazinon; dicyclopentadiene; guazatine; guazatine acetates; methiocarb; pyridin-4-amine; thiram; trimethacarb; zinc naphthenate; ziram; imanin; ribavirin; mercuric oxide; octxillinone; thiophanate-methyl; a compound of formula B-1A
wherein \( R' \) is hydrogen, \( C_{1-4} \) alkyl or \( C_{1-4} \) haloalkyl;
a compound of formula B-2

\[
\text{(B-2)};
\]
a compound of formula B-3

\[
\text{(B-3)};
\]
a compound of formula B-4
a compound of formula B-5

a compound of formula B-8

and a compound of formula B-9
2. A method according to claim 1, wherein component A) is a compound of formula I, wherein R₁ is difluoromethyl.

3. A method according to claim 1, wherein component A) is a compound of the formula Ia

4. A method according to claim 1, wherein component A) is a compound of the formula Ib
and one component B) as described above.

5. A method according to claim 1, wherein component A) is a compound of the formula (Ic):

and one component B) as described above.

6. A method according to claim 1, wherein component A) is a compound of the formula (Id):
and one component B) as described above.

7. A method according to claim 1, wherein component A) is a compound of the formula \( \text{Ie} \)

and one component B) as described above.

8. A method according to claim 1, wherein component B) is selected from the group consisting of a strobilurin fungicide, selected from the group consisting of Azoxystrobin, Dimoxystrobin, Fluoxastrobins, Kresoxim-methyl, Metominostrobin, Oryxastrobins, Picoxystrobin, Pyraclostrobin; Trifloxystrobin and a compound of formula B-6.
an azole fungicide, selected from the group consisting of Azaconazole, Bromuconazole, Cyproconazole, Difenoconazole, Diniconazole, Diniconazole-M, Epoxiconazole, Fenbuconazole, Fluquinconazole, Flusilazole, Flutriafol, Hexaconazole, Imazalil, Imibenconazole, Ipconazole, Metconazole, Myclobutanil, Oxoconazole, Pefurazoxide, Penconazole, Prochloraz, Propiconazole, Prothioconazole, Simeconazole, Tebuconazole, Tetraconazole, Triadimefon, Triadimenol, Triflumizole, Triflumizole, Diclobutrazol, Etaconazole, Furconazole, Furconazole-cis and Quinconazole;
a phenyl pyrrole fungicide, selected from the group consisting of Fenpiclonil and Fludioxonil;
an anilino-pyrimidine fungicide, selected from the group consisting of Cyprodinil, Mepanipyrim and Pyrimethanil;
a morpholine fungicide, selected from the group consisting of Aldimorph, Dodemorph, Fenpropimorph, Tridemorph, Fenpropidin, Spiroxamine; Piperalin and a compound of formula B-7

a compound of formula F-1
9. A method according to claim 1, wherein component B) is selected from the group consisting of Azoxystrobin; Picoxystrobin; Cyproconazole; Difenconazole; Propiconazole; Fludioxonil; Cyprodinil; Fenpropimorph; Fenpropidin; a compound of formula F-1
and Chlorothalonil.

10. A method according to claim 1, wherein component B) is selected from the group consisting of Azoxystrobin; Cyproconazole; Propiconazole; Cyprodinil; Chlorothalonil; Fludioxonil and Fenpropidin.

11. A fungicidal composition comprising a combination of components A) and B) according to claim 1 in a synergistically effective amount, together with an agriculturally acceptable carrier, and optionally a surfactant.

12. A fungicidal composition comprising a combination of components A) and B) according to claim 1 together with an agricultural acceptable carrier, and optionally a surfactant, wherein the weight ratio of A) to B) is between 2000 : 1 and 1 : 1000.

13. A method of protecting natural substances of plant and/or animal origin, which have been taken from the natural life cycle, and/or their processed forms, which comprises applying to said natural substances of plant and/or animal origin or their processed forms a combination of components A) and B) according to claim 1 in a synergistically effective amount.