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(54) Title: CRYSTALLINE MODIFICATION OF FIPRONIL

(57) Abstract: The present invention relates to a crystalline modification of fipronil, to a process for the preparation of the same, to pesticidal and parasiticidal mixtures and compositions comprising said crystalline modification and to their use for combating pests and parasites.

CRYSTALLINE MODIFICATION OF FIPRONIL

The present invention relates to a novel crystalline modification of fipronil, to a process for the preparation of the same, to pesticidal and parasiticidal mixtures and compositions comprising said crystalline modification and to their use for combating pests and parasites.

Fipronil (formula I) is an active compound for controlling certain insect and acarid pests, and parasites.

Various processes for the preparation of fipronil have been described, generally and in detail. Documents which give detailed preparation procedures are e.g. EP 295 117; EP 460 940; EP 484 165; EP 668 269; EP 967 206; EP 1 331 222; EP 1 374 061; US 5 631 381; CN 1374298; or J. of Heibei University of Science and Technology, Vol. 25 (2), Sum 69 (2004), Dok. Serial No. 1008-1542 (2004) 02-0018-03.

Characterization of the fipronil material obtained by the processes described in the prior art is usually done by ¹H-NMR analysis and/ or measurement of the melting point. The described melting points are in the range of from 187°C to 203°C, mostly in the range of from 195°C to 203°C. In the Pesticidal Manual, 13th Edition (2003), British Crop Protection Council, p.433, fipronil is described as a white solid with a melting point of 200 to 201°C, with technical fipronil having a melting point of 195.5 °C to 203°C. Observations of different crystalline forms of fipronil have not been described, let alone any characterization of a certain crystalline modification or a preparation procedure for obtaining a certain crystalline modification.

For the large-scale preparation and formulation of a market compound such as fipronil, it is of crucial importance to know whether different crystalline modifications (also frequently referred to as polymoprhs) of a compound exist, how they can be obtained, and what their characteristic properties are. Crystalline modifications of one compound may have very different properties, for example with regard to solubility, rate of dissolution, suspension stability, stability during grinding, vapour pressure, optical and mechanical properties, hygroscopicity, crystal size, filtration properties, desiccation, density, melting point, degradation stability, stability against phase transformation into other crystalline modifications, colour, and even chemical reactivity.

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For example, different crystalline modifications frequently manifest themselves in different forms of the crystals, such as needles or plates. This is of relevance for e.g a filtration step in the preparation procedure. Plates typically will clog the pores of a filter leading to loss of time and product and tedious and expensive cleaning work. Also, a crystalline modification being present as plates and a crystalline modification being present as needles can have significantly different bulk densities which has implications for storage and packaging. Another relevant aspect, especially in the production of pesticides, is whether the crystalline modification is present as a fine powder which can produce hazardous dusts, or as dust-free larger crystals.

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Against this background, it has been an object of the present invention to find and characterize a novel crystalline modification of fipronil.

A further object has been to find preparation procedures for the novel crystalline modification which reproducibly give the crystalline modification.

Another object of the invention has been to find preparation procedures which give the novel crystalline modification II in high yield.

Yet another object of the invention has been to find preparation procedures which give the novel crystalline modification essentially excluding other crystalline modification forms (i.e. in over 80% by weight).

Accordingly, a novel crystalline modification of fipronil, a process for its preparation, pesticidal and parasiticidal mixtures and compositions comprising it and its use for combating pests and parasites has been found. The novel crystalline modification of fipronil is defined as "novel crystalline modification II" throughout this application. It is present as long needles which provides for its easy filtration.

Also, most suprisingly, 3 other crystalline modifications of fipronil have been found, which are subject to co-pending patent applications. Especially surprising was that the present crystalline modification II of fipronil does not melt but rather undergoes a phase transformation during heating into two thermodynamically more stable forms I and V and/or a mixture of them, and thus in a typical melting point measurement will give the melting points of these forms I and V or mixtures of them. Crystalline modification I has a very similar melting point as a second crystalline modification V (as described in copending patent applications), both melting points lying in the range of the melting points given in the prior art (i.e. 195 to 203°C). Moreover, one further crystalline modification IV of fipronil, as described in a co-pending patent application, also undergoes a phase transformation. The solid forms of fipronil thus are part of a very complex crystallization scenario. It can be concluded that the melting points given in the literature in no way

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can indicate which crystalline modification or crystalline modification mixtures were analyzed.

In T 605/02, the Technical Board of Appeal of the European Patent Authority ruled that, in the absence of a respective described preparation procedure, even the XRD pattern of a certain crystalline modification does not constitute prior art for lack of enablement. Thus, melting points given in documents published prior to the filing of this application cannot be regarded as prior art for the present invention as they do not enable the artisan to prepare the novel crystalline modification of fipronil.

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Crystalline modification II of fipronil, in an X-ray powder diffractogram at 25°C, shows at least 4, in particular at least 5, especially 7 and preferably all of the following reflexes:

	(1) d = 13.44 ± 0.2 Å	$2 \Theta = 6.6 \pm 0.2^{\circ}$
15	(2) $d = 7.84 \pm 0.1 \text{ Å}$	$2 \Theta = 11.3 \pm 0.2^{\circ}$
	(3) $d = 5.50 \pm 0.07 \text{ Å}$	$2 \Theta = 16.1 \pm 0.2^{\circ}$
	$(4) d = 5.14 \pm 0.05 Å$	$2 \Theta = 17.2 \pm 0.2^{\circ}$
	$(5) d = 4.95 \pm 0.05 Å$	$2 \Theta = 17.9 \pm 0.2^{\circ}$
	(6) $d = 3.95 \pm 0.05 \text{ Å}$	$2 \Theta = 22.4 \pm 0.2^{\circ}$
20	$(7) d = 3.77 \pm 0.05 Å.$	$2 \Theta = 23.5 \pm 0.2^{\circ}$
	(8) $d = 3.22 \pm 0.03 \text{ Å}$	$2 \Theta = 27.6 \pm 0.2^{\circ}$
	(9) $d = 2.91 \pm 0.03 \text{ Å}$	$2 \Theta = 30.8 \pm 0.2^{\circ}$.

In a particularly preferred embodiment, the crystalline modification II exhibits a powder X-ray diffraction pattern substantially the same as the pattern shown in Figure 1.

Studies of single crystals of the crystalline modification II have shown that the crystal system is monoclinic and has the space group P 2(1)/c. The characteristic data of the crystal structure of the crystalline modification II are shown in Table 1:

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Table 1: Crystallographic data and parameters of the crystalline modification II measured from

measured from		
Parameter	Modification II	
Class	Monoclinic	
Space group	P 2(1)/c	
a	8.606(1) Å	
b	26.919(2) Å	
С	16.086(1) Å	
α	90°	
β	102.066(1)°	
γ	90°	

Parameter	Modification II
Volume	3644.0(3) Å ³
Z	4
Temperature	-173.2°C
Density (calculated)	0.94 g/cm ³
R1, ωR2	0.081, 0.222

a,b,c = Length of the unit cell edges

 α, β, γ = Angles of the unit cell

Z = Number of molecules in the unit cell

^aDensity calculated without the incorporated solvent

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The single crystal structure further reveals, that modification II of fipronil has channel like tubes running through the crystal. These tubes are occupied by the corresponding crystallization solvent. This affects the powder diffraction patterns in a way that the X-ray reflections can be seen at slightly different 2Theta and d-spacing values (not more than \pm 0.34° 2Theta) for many of the solvates. The peak intensities are independent of the incorporated solvent[B1]. Modification II can therefore be regarded as a solvate of the corresponing co-crystallized solvent.

As the solvents trapped in the crystal are removed, the crystalline modification II of fipronil undergoes an endothermic phase transformation. In a differential scanning calorimetry, DSC, measurement (heating rate 5 K / min) this is observed typically at around 130°C, with an onset at 100 to 110°C and completion at 138°C to 145°C. This phase transformation leads to crystalline modifications I and V and/or depending on the heating rate to mixtures of them. We give here as an example a differential scanning calorimetry (DSC) thermogram of the crystalline modification II of fipronil, recorded with 10 K/ min. It further contains two endotherms with maxima at 196°C and 203°C, representing the melting points of the crystalline modifications I (mp. 196°C) and V (mp. 203°C) of fipronil. It is shown in figure 2.

Solvents in the process for preparation of modification II are inert and consist mainly of benzene derivatives, such as benzene, which may be substituted by one or more groups selected from halogen, cyano, C₁-C₆-alkyl, C₁-C₆-alkoxy, halogenmethyl, and nitro, and other solvents, such as tetrahydrofuran (THF), 1,2-dichloroethane, and acetonitrile. Preferred benzene derivatives are fluorobenzene, benzonitrile, anisole, p-xylene, o-xylene, m-xylene, CF₃-benzene, n-butylbenzene, t-butylbenzene, s-butylbenzene, i-butylbenzene, chlorobenzene, 2-chlorotoluene, 4-chlorotoluene, 1,2-dichlorobenzene, 1,3-dichlorobenzene, 1,4-dichlorobenzene, 1,4-diisopropylbenzene, mesitylene, nitrobenzene, 4-nitrotoluene, n-propylbenzene, and toluene, and more preferably selected from mono-, di- or tri(C₁-C₆-alkyl) benzenes[B2], which may be halogenated.

The crystalline modification II of fipronil can be achieved preferably from the following solvents; tetrahydrofuran, 1,2-dichloroethane, acetonitrile, toluene, mono chloro benzene, 1,2-dichlorobenzene, ethyl benzene, mesitylene, nitrobenzene and CF₃-benzene.

The amount of co-crystallized solvent depends of the properties, such as size of the corresponding solvent. For toluene the maximum amount of solvent in the crystals is in the range of 5 w-%. The solvent amount can be analysed for example via thermogravimetry. A TGA-trace of crystalline fipronil modification II from toluene is shown in figure 3.

In addition to the various solvent adduct versions of modification II, it was further discovered, that from all of the above mentioned solvents also at least one other crystalline modification can be achieved via solvent crystallization. Several conditions lead also to mixtures of forms, but also polymorph or modification pure material can be prepared by applying delicate and controlled, solvent dependant methods. The applicant has described preparation methods of modifications I and V in co-pending patent applications[B3].

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Also a further modification F-ST, a toluene solvate, is described in WO2007/069254. This solvate modification F-ST is different from the here described novel modification II which also contains co-crystallized toluene. This should be made clear by comparison of the powder diffraction patterns[B4] shown in Figure 1, and in WO2007/069254. It appears that F-ST is a stoichiometric solvate of fipronil and toluene in the ratio of about 2:1 respectively. In the case of toluene this means that there is about 8 w-% of toluene in the crystal lattice, which is about 3 w-% more than for crystalline modification II according to the current invention.

Further, structurally similar solvates to F-ST form can be prepared also with benzene, mono chloro benzene (MCB) as well as xylene (mixture).

Similar to F-ST, also crystalline modification II of fipronil can be used as starting material to prepare pure modification V via drying (tempering) above 100°C, preferably above 130°C. The drying time depends of the applied pressure. As modification II contains less toluene than F-ST, also less energy is needed in the drying (tempering) process.

In another embodiment, the present invention relates to the crystalline modification II having a fipronil content of at least 92 % by weight, particularly at least 96 % by weight and especially at least 98 % by weight.

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This invention also relates to solid (compositions of) fipronil comprising the crystalline modification II as defined hereinabove and a form of fipronil being different from said crystalline modification II (herein also referred to as "fipronil form"), e.g. amorphous fipronil or fipronil of a crystalline modification different from crystalline modification II. Preferably, the solid (compositions of) fipronil comprise the crystalline modification II in at least 85 % by weight, preferably in at least 90 % by weight, more preferably in at least 95 % by weight, and most preferably in at least 98 % by weight [B6].

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The crystalline modification II can be prepared using a process which comprises the 10 following steps:

- step i) preparing a solution of a solid form of fipronil being different from the crystalline modification II in a solvent S preferably selected from tetrahydrofurane (THF), 1,2-dichloroethane (DCE), acetonitrile, toluene, monochlorobenzene (MCB), 1,2-dichlorobenzene (DCB), ethyl benzene, mesitylene, nitrobenzene and CF₃benzene;
- step ii) effecting crystallization of fipronil; and step iii) isolating the resulting precipitate.
- 20 The preparation of modification II is favored by crystallization from concentrated solutions at elevated temperatures. Further, the resulting modification is dependent of especially the nucleation temperature followed by the formation of the first crystals. The beginning of the crystallization can be detected visually as clouding of the crystallization solution and/or with the aid of a turbidity probe. Usually the evaporation 25 or cooling crystallisation is effected not above 130°C, or close to the boiling point of the used solvent, and preferably not below 80°C.

Thus crystallization of modification II can be done by crystallization of fipronil from the aforesaid solvents at constant temperature by evaporation and/or by adding at least a polar solvent P to the above mentioned aromatic solvents. Such cooling cystallisation is advantageously started at a temperature close to the boiling point of the solvent, however, preferably not above 130°C or below 80°C. More precisely in the cases of THF, 1,2-dichloroethane and acetonitrile the crystallization temperature in an evaporation crystallization should be kept within a temperature window of 30 and 55°C, for toluene within a temperature window of 90°C to 110°C, for MCB and 1,2-dichlorobenzene in between 70°C and 130°C, nitrobenzene and mesitylene in between 90°C and 130°C and for ethyl benzene in between 100°C and 130°C and for CF₃-benzene in between 90°C and 103°C.

40 Modification II can also be crystallised from mixtures of the above mentioned aromatic solvents and polar solvents P, advantageously in amounts up to 20 vol.-%, preferably 5 to 15 vol.-% of P. Such solvent P is preferably selected from the group of methanol,

ethanol, propan-1-ol, propan-2-ol (isopropanol), butan-1-ol (n-butanol), butan-2-ol, tert-butanol, 2-methyl-propan-1-ol (iso-butanol), 2-methyl-propan-2-ol, pentan-3-ol, 2-methyl butan-1-ol, 3-methyl butan-1-ol, 1,2-ethanediol, 1,3-propandiol, 1,2-propandiol, cyclohexanol, acetonitrile, propionitrile, acetone, butanone (methyl ethyl ketone), pentan-2-one (methyl propyl ketone), pentan-3-one (diethylketone), 4-methyl-2-pentanone (isobutyl-methyl-ketone), 3-methyl-butan-2-one (iso-propyl-methyl-ketone), 3,3-dimethyl-2-butanone (tert-butyl-methyl-ketone), cyclohexanone, methylacetate, ethylacetate, isopropylacetate, N-butylacetate, isobutylacetate, diethylcarbonate, 2-butoxyethylacetate, dioxane, THF, diethylether, 2-methyl-THF, methyl-tert-butylether, dimethylformamide, dimethylacetamide, dimethylsulfoxide (DMSO), nitromethane, and nitroethane.

Modification II can also be crystallized by effecting the crystallization by cooling concentrated solutions of the corresponding solvent. In a cooling crystallization the crystallization begin needs to be within the aforesaid temperature ranges. Parameters effecting the nucleation temperature and beginning of crystallization are concentration, cooling rate and time of adding possible seeding crystals of the wanted modification to a supersaturated solution. The concentration depends naturally of the solubility of fipronil in the corresponding solvent. The cooling rate is typically in between 5 K/h and 20 K/h. The cooling rate may not exceed 1 K/minute.

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A person skilled in the art is able to perform a cooling crystallization in a corresponding solvent so that these conditions are met.

If the crystallization is not affected in a way that the nucleation begins in the aforesaid narrow temperature ranges, other modifications or mixtures of modifications result.

Modification II can be obtained from a solution in THF, or DCE by evaporation at from 40°C to the boiling point of the solvent used. Evaporation from a THF solution at 20-25°C yield a mixture of modifications I and II.

Alternatively modification II is obtainable from a solution in acetonitrile by cooling crystallization from 80 to 5°C (cf. example 4).

In a further embodiment of the invention modification II is obtainable from a solution in toluene by a cooling crystallization when nucleation and crystallization starts above 90°C (cf. example 5). In between 90 and 75°C mixtures of II and III (F-ST) result.

In a further embodiment of the invention modification II is obtainable from a solution in monochlorobenzene (MCB) also in a cooling crystallization (cf. example 6).

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In a further embodiment of the invention modification II is obtainable from a solution in mesitylene (cf. example 7).

In a further embodiment of the invention modification II is obtainable from a solution in dichlorobenzene (DCB) in two concentrations.

In a further embodiment of the invention modification II is obtainable from a solution in nitrobenzene (cf. example 8).

- In a further embodiment of the invention modification II is obtainable from a solution in ethyl benzene when crystallization took place above 100°C (cf. example 9). Preferably solutions with a fipronil content of 10 w.-% or more, more preferably of 15 w.-% or more are used.
- In a further embodiment of the invention modification II is obtainable from a solution in CF₃-benzene at 90°C or more, preferably 95°C or above.

A detailed description of these steps is as follows:

20 Step i)

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Suitable fipronil forms different from the crystalline modification II used in step i) are, for example, selected from amorphous fipronil or crystalline fipronil such as other triclinic or monoclinic forms, e.g. monoclinic fipronil of the space group C2/c, and also mixtures of crystalline modifications of fipronil.

The fipronil form used as starting material in step i) preferably has a purity of at least 85 % by weight, in particular at least 90 % by weight and especially at least 95 % by weight. "Purity" means the absence of chemical compounds other than fipronil.

The solvent S used in step i) consists of either acetonitrile, DCB, DCE, ethyl benzene, MCB, mesitylene, nitrobenzene, THF, toluene, or CF₃-benzene, each pure or in combination with a polar solvent P.

In a preferred embodiment, the solvent S used in step i) consists of DCB, mesitylene, and nitrobenzene, each pure or in combination with a polar solvent P

In another preferred embodiment, the solvent S used in step i) consists of ethyl benzene, MCB, and toluene, each pure or in combination with a polar solvent P

In step i), the fipronil form different from the crystalline modification II will usually be incorporated into the solvent S as a solid with mixing at a concentration and temperature where the solvent S is capable of completely dissolving the fipronil form.

The amount of fipronil form dissolved in the solvent S depends, on the nature of the solvent S and on the dissolution temperature. The person skilled in the art will be able to determine suitable conditions by standard experiments.

Step ii)

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In step ii) of the process of this invention, fipronil is then crystallized. Crystallization can be effected in a customary manner, for example by cooling the solution obtained in step i), by adding a solvent which reduces the solubility, or by concentrating the solution, or by a combination of the measures mentioned above.

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In a preferred embodiment, step ii) is carried out in the presence of seed crystals of the crystalline modification II.

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To achieve a conversion into the crystalline modification II which is as complete as possible, the crystallization is carried out over a period (duration of crystallization) of at least 1 h, in particular at least 3 h. Duration of crystallization is understood by the person skilled in the art as meaning the period of time between the beginning of the measure which initiates crystallization and the isolation of the fipronil by separating the crystalline material from the mother liquor.

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In general, the crystallization is allowed to proceed to a point where at least 60%, preferably at least 70%, in particular at least 90% by weight, for example from 80 to 90% by weight, of the fipronil employed has crystallized out.

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Concentration of the solution is effected by gradually removing the solvent S, such as by evaporation in vacuo, either at low temperature or at about 20°C to 25°C or at elevated temperature, and/or in the presence of a flow of an inert gas such as nitrogen or argon. The values of "low temperature" and "elevated temperature" depend, on the nature of the solvent S, however should not exceed 60°C.

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In a preferred embodiment, the crystallization is effected by concentrating the solution.

filtration, centrifugation or decanting. In general, the isolated precipitate will be washed,

Step iii)

In step iii) of the process of this invention, the crystalline modification II is isolated using customary techniques for separating solid components from liquids, for example by

for example with the solvent S used for the crystallization. The washing can be carried out in one or more steps. The washing is typically carried out at temperatures lower than 30°C and in particular lower than 25°C, to keep the loss of the product of value as low as possible. The resulting crystalline fipronil or modification II can then be dried and subjected to further processing.

The preparation process consisting of steps i) to step iii) can be repeated in order to achieve higher purities of fipronil.

The crystalline modification II is especially suitable for efficiently combating the following pests:

millipedes (Diplopoda) such as Blaniulus or Narceus ssp;

15 insects (Insecta) such as:

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ants, bees, wasps, sawflies (Hymenoptera), e.g. Atta capiguara, Atta cephalotes, Atta laevigata, Atta robusta, Atta sexdens, Atta texana, Crematogaster spp., Hoplocampa minuta, Hoplocampa testudinea, Monomorium pharaonis, Solenopsis geminata,

Solenopsis invicta, Solenopsis richteri, Solenopsis xyloni, Pheidole megacephala, Pogonomyrmex species such as Pogonomyrmex barbatus and Pogonomyrmex californicus, Dasymutilla occidentalis, Bombus spp. Vespula squamosa, Paravespula vulgaris, Paravespula pennsylvanica, Paravespula germanica, Dolichovespula maculata, Vespa crabro, Polistes rubiginosa, Camponotus floridanus, and Linepithema humile,

beetles (Coleoptera), such as Agrilus sinuatus, Agriotes lineatus, Agriotes obscurus and other Agriotes species, Amphimallus solstitialis, Anisandrus dispar, Anthonomus grandis, Anthonomus pomorum, Aracanthus morei, Atomaria linearis, Blapstinus species, Blastophagus piniperda, Blitophaga undata, Bothynoderes punciventris, Bruchus rufimanus, Bruchus pisorum, Bruchus lentis, Byctiscus betulae, Cassida nebulosa, Cerotoma trifurcata, Ceuthorrhynchus assimilis, Ceuthorrhynchus napi, Chaetocnema tibialis, Conoderus vespertinus and other Conoderus species, Conorhynchus mendicus, Crioceris asparagi, Cylindrocopturus adspersus, Diabrotica (longicornis) barberi, Diabrotica semi-punctata, Diabrotica speciosa, Diabrotica undecimpunctata, Diabrotica virgifera and other Diabrotica species, Eleodes species, Epilachna varivestis, Epitrix hirtipennis, Eutinobothrus brasiliensis, Hylobius abietis, Hypera brunneipennis, Hypera postica, Ips typographus, Lema bilineata, Lema melanopus, Leptinotarsa decemlineata, Limonius californicus and other Limonius species, Lissorhoptrus oryzophilus, Listronotus bonariensis, Melanotus communis and other Melanotus species, Meligethes aeneus, Melolontha hippocastani, Melolontha melolontha, Oulema oryzae, Ortiorrhynchus sulcatus, Oryzophagus oryzae,

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Otiorrhynchus ovatus, Oulema oryzae, Phaedon cochleariae, Phyllotreta chrysocephala, Phyllophaga cuyabana and other Phyllophaga species, Phyllopertha horticola, Phyllotreta nemorum, Phyllotreta striolata, and other Phyllotreta species, Popillia japonica, Promecops carinicollis, Premnotrypes voraz, Psylliodes species, Sitona lineatus, Sitophilus granaria, Sternechus pinguis, Sternechus subsignatus, and Tanymechus palliatus and other Tanymechus species,

Centipedes (Chilopoda), e.g. Scutigera coleoptrata,

10 Cockroaches (Blattaria - Blattodea), e.g. Blattella germanica, Blattella asahinae, Periplaneta americana, Periplaneta japonica, Periplaneta brunnea, Periplaneta fuligginosa, Periplaneta australasiae, and Blatta orientalis,

Crickets, grasshoppers, locusts (Orthoptera), e.g. Acheta domestica, Gryllotalpa gryllotalpa, Locusta migratoria, Melanoplus bivittatus, Melanoplus femurrubrum, Melanoplus mexicanus, Melanoplus sanguinipes, Melanoplus spretus, Nomadacris septemfasciata, Schistocerca americana, Schistocerca gregaria, Dociostaurus maroccanus, Tachycines asynamorus, Oedaleus senegalensis, Zonozerus variegatus, Hieroglyphus daganensis, Kraussaria angulifera, Calliptamus italicus, Chortoicetes terminifera, and Locustana pardalina,

fleas (Siphonaptera), e.g. Ctenocephalides felis, Ctenocephalides canis, Xenopsylla cheopis, Pulex irritans, Tunga penetrans, and Nosopsyllus fasciatus,

- 25 Flies, mosquitoes (Diptera), e.g. Aedes aegypti, Aedes albopictus, Aedes vexans, Agromyza oryzea, Anastrepha ludens, Anopheles maculipennis, Anopheles crucians, Anopheles albimanus, Anopheles gambiae, Anopheles freeborni, Anopheles leucosphyrus, Anopheles minimus, Anopheles quadrimaculatus, Calliphora vicina, Chrysomya bezziana, Chrysomya hominivorax, Chrysomya macellaria, Chrysops 30 discalis, Chrysops silacea, Chrysops atlanticus, Cochliomyia hominivorax, Contarinia sorghicola, Cordylobia anthropophaga, Culicoides furens, Culex pipiens, Culex nigripalpus, Culex quinquefasciatus, Culex tarsalis, Culiseta inornata, Culiseta melanura, Dacus cucurbitae, Dacus oleae, Dasineura brassicae, Delia antique, Delia coarctata, Delia platura, Delia radicum, Dermatobia hominis, Fannia canicularis, 35 Gasterophilus intestinalis, Geomyza Tripunctata, Glossina morsitans, Glossina palpalis, Glossina fuscipes, Glossina tachinoides, Haematobia irritans, Haplodiplosis equestris, Hippelates spp., Hypoderma lineata, Leptoconops torrens, Liriomyza sativae, Liriomyza trifolii, Lucilia caprina, Lucilia cuprina, Lucilia sericata, Lycoria pectoralis, Mansonia spp., Mayetiola destructor, Musca domestica, Muscina stabulans,
- 40 Oestrus ovis, Oestrus ovis, Opomyza florum, Oscinella frit, Pegomya hysocyami, Phlebotomus argentipes, Phorbia antiqua, Phorbia brassicae, Phorbia coarctata, Progonya leyoscianii, Psila rosae, Psorophora columbiae, Psorophora discolor,

Prosimulium mixtum, Rhagoletis cerasi, Rhagoletis pomonella, Sarcophaga haemorrhoidalis, Sarcophaga sp., Simulium vittatum, Stomoxys calcitrans, Tabanus bovinus, Tabanus atratus, Tabanus lineola, Tabanus similis, Tetanops myopaeformis, Tipula olerace, and Tipula paludosa,

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Heteropterans (Heteroptera), such as Acrosternum hilare, Blissus leucopterus, Cicadellidae such as Empoasca fabae, Chrysomelidae, Cyrtopeltis notatus, Delpahcidae, Dysdercus cingulatus, Dysdercus intermedius, Eurygaster integriceps, Euschistus impictiventris, Leptoglossus phyllopus, Lygus lineolaris, Lygus pratensis, Nephotettix species, Nezara viridula, Pentatomidae, Piesma quadrata, Solubea insularis and Thyanta perditor,

Aphids and other homopterans (Homoptera), e.g. Acyrthosiphon onobrychis, Adelges laricis, Aphidula nasturtii, Aphis fabae, Aphis forbesi, Aphis glycines, Aphis gossypii, 15 Aphis grossulariae, Aphis pomi, Aphis schneideri, Aphis spiraecola, Aphis sambuci, Acyrthosiphon pisum, Aulacorthum solani, Brachycaudus cardui, Brachycaudus helichrysi, Brachycaudus persicae, Brachycaudus prunicola, Brevicoryne brassicae, Capitophorus horni, Cerosipha gossypii, Chaetosiphon fragaefolii, Cryptomyzus ribis, Dreyfusia nordmannianae, Dreyfusia piceae, Dysaphis radicola, Dysaulacorthum 20 pseudosolani, Dysaphis plantaginea, Dysaphis pyri, Empoasca fabae, Hyalopterus pruni, Hyperomyzus lactucae, Macrosiphum avenae, Macrosiphum euphorbiae, Macrosiphon rosae, Megoura viciae, Melanaphis pyrarius, Metopolophium dirhodum, Myzodes (Myzus) persicae, Myzus ascalonicus, Myzus cerasi, Myzus varians, Nasonovia ribis-nigri, Nilaparvata lugens, Pemphigus bursarius, Pemphigus 25 populivenae, and other Pemphigus species, Perkinsiella saccharicida, Phorodon humuli, Psyllidae such as Psylla mali, Psylla piri and other Psylla species, Rhopalomyzus ascalonicus, Rhopalosiphum maidis, Rhopalosiphum padi, Rhopalosiphum insertum, Sappaphis mala, Sappaphis mali, Schizaphis graminum, Schizoneura lanuginosa, Sitobion avenae, Trialeurodes vaporariorum, Toxoptera 30 aurantiiand, and Viteus vitifolii,

Lepidopterans (Lepidoptera), for example Agrotis ypsilon, Agrotis segetum and other Agrotis species, Alabama argillacea, Anticarsia gemmatalis, Argyresthia conjugella, Autographa gamma, Bupalus piniarius, Cacoecia murinana, Capua reticulana, Cheimatobia brumata, Chilo suppresalis and other Chilo species, Choristoneura fumiferana, Choristoneura occidentalis, Cirphis unipuncta, Cnaphlocrocis medinalis, Cydia pomonella, Dendrolimus pini, Diaphania nitidalis, Diatraea grandiosella, Earias insulana, Elasmopalpus lignosellus, Eupoecilia ambiguella, Euxoa species, Evetria bouliana, Feltia subterranea, Galleria mellonella, Grapholitha funebrana, Grapholitha molesta, Heliothis armigera, Heliothis virescens, Heliothis zea, Hellula undalis, Hibernia defoliaria, Hyphantria cunea, Hyponomeuta malinellus, Keiferia lycopersicella, Lambdina fiscellaria, Laphygma exigua, Lerodea eufala, Leucoptera coffeella,

Leucoptera scitella, Lithocolletis blancardella, Lobesia botrana, Loxostege sticticalis, Lymantria dispar, Lymantria monacha, Lyonetia clerkella, Malacosoma neustria, Mamestra brassicae, Momphidae, Orgyia pseudotsugata, Ostrinia nubilalis, Panolis flammea, Pectinophora gossypiella, Peridroma saucia, Phalera bucephala,

- Phthorimaea operculella, Phyllocnistis citrella, Pieris brassicae, Plathypena scabra,
 Plutella xylostella, Pseudoplusia includens, Rhyacionia frustrana, Scrobipalpula
 absoluta, Sesamia nonagrioides and other Sesamia species, Sitotroga cerealella,
 Sparganothis pilleriana, Spodoptera frugiperda, Spodoptera littoralis, Spodoptera litura,
 Thaumatopoea pityocampa, Tortrix viridana, Trichoplusia ni and Zeiraphera
 canadensis,
 - lice (Phthiraptera), e.g. Pediculus humanus capitis, Pediculus humanus corporis, Pthirus pubis, Haematopinus eurysternus, Haematopinus suis, Linognathus vituli, Bovicola bovis, Menopon gallinae, Menacanthus stramineus and Solenopotes capillatus,

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- orthopterans (Orthoptera), such as Acrididae, Acheta domestica, Forficula auricularia, Gryllotalpa gryllotalpa, Locusta migratoria, Melanoplus bivittatus, Melanoplus femurrubrum, Melanoplus mexicanus, Melanoplus sanguinipes, Melanoplus spretus, Nomadacris septemfasciata, Schistocerca americana, Schistocerca peregrina, Stauronotus maroccanus and Tachycines asynamorus,
- silverfish, firebrat (Thysanura), e.g. Lepisma saccharina and Thermobia domestica,
- termites (Isoptera), such as Calotermes flavicollis, Coptotermes ssp., Dalbulus maidis, Heterotermes aureus, Leucotermes flavipes, Macrotermes gilvus, Reticulitermes ssp., Termes natalensis, Coptotermes formosanus,
- thrips (Thysanoptera), such as Frankliniella fusca, Frankliniella occidentalis,

 Frankliniella tritici and other Frankliniella species, Scirtothrips citri, Thrips oryzae,

 Thrips palmi, Thrips simplex, and Thrips tabaci,
 - ticks and parasitic mites (Parasitiformes): ticks (Ixodida), e.g. Ixodes scapularis, Ixodes holocyclus, Ixodes pacificus, Rhiphicephalus sanguineus, Dermacentor andersoni, Dermacentor variabilis, Amblyomma americanum, Ambryomma maculatum, Ornithodorus hermsi, Ornithodorus turicata and parasitic mites (Mesostigmata), e.g. Ornithonyssus bacoti and Dermanyssus gallinae,
- true bugs (Hemiptera), e.g. Cimex lectularius, Cimex hemipterus, Reduvius senilis, 40 Triatoma spp., Rhodnius prolixus, and Arilus critatus,

Arachnoidea, such as arachnids (Acarina), for example of the families Argasidae, Ixodidae and Sarcoptidae, such as Amblyomma americanum, Amblyomma variegatum, Argas persicus, Boophilus annulatus, Boophilus decoloratus, Boophilus microplus, Dermacentor silvarum, Hyalomma truncatum, Ixodes ricinus, Ixodes rubicundus,

5 Latrodectus mactans, Loxosceles reclusa, Ornithodorus moubata, Otobius megnini, Dermanyssus gallinae, Psoroptes ovis, Rhipicephalus appendiculatus, Rhipicephalus evertsi, Sarcoptes scabiei, and Eriophyidae species such as Aculus schlechtendali, Phyllocoptrata oleivora and Eriophyes sheldoni; Tarsonemidae species such as Phytonemus pallidus and Polyphagotarsonemus latus; Tenuipalpidae species such as Brevipalpus phoenicis; Tetranychidae species such as Tetranychus cinnabarinus, Tetranychus kanzawai, Tetranychus pacificus, Tetranychus telarius and Tetranychus urticae, Panonychus ulmi, Panonychus citri, and Oligonychus pratensis,

Earwigs (Dermaptera), e.g. forficula auricularia; and

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Nematodes, including plant parasitic nematodes and nematodes living in the soil. Plant parasitic nematodes include, such as root knot nematodes, Meloidogyne hapla, Meloidogyne incognita, Meloidogyne javanica, and other Meloidogyne species; cystforming nematodes, Globodera rostochiensis and other Globodera species; Heterodera avenae, Heterodera glycines, Heterodera schachtii, Heterodera trifolii, and other Heterodera species; Seed gall nematodes, Anguina species; Stem and foliar nematodes, Aphelenchoides species; Sting nematodes, Belonolaimus longicaudatus and other Belonolaimus species; Pine nematodes, Bursaphelenchus xylophilus and other Bursaphelenchus species; Ring nematodes, Criconema species, Criconemella species, Criconemoides species, Mesocriconema species; Stem and bulb nematodes, Ditylenchus destructor, Ditylenchus dipsaci and other Ditylenchus species; Awl nematodes, Dolichodorus species; Spiral nematodes, Heliocotylenchus multicinctus and other Helicotylenchus species; Sheath and sheathoid nematodes, Hemicycliophora species and Hemicriconemoides species; Hirshmanniella species; Lance nematodes, Hoploaimus species; false rootknot nematodes, Nacobbus species; Needle nematodes, Longidorus elongatus and other Longidorus species; Pin nematodes, Paratylenchus species; Lesion nematodes, Pratylenchus neglectus, Pratylenchus penetrans, Pratylenchus curvitatus, Pratylenchus goodeyi and other Pratylenchus species; Burrowing nematodes, Radopholus similis and other Radopholus species; Reniform nematodes, Rotylenchus robustus and other Rotylenchus species; Scutellonema species; Stubby root nematodes, Trichodorus primitivus and other Trichodorus species, Paratrichodorus species; Stunt nematodes, Tylenchorhynchus claytoni, Tylenchorhynchus dubius and other Tylenchorhynchus species; Citrus nematodes, Tylenchulus species, Dagger nematodes, Xiphinema species and other plant parasitic nematode species.

Moreover, the crystalline modification II is especially useful for the control of crop pests, in particular of the Coleoptera, Lepidoptera and Acarina orders.

Moreover, the crystalline modification II is especially useful for the control of non-crop pests (household, turf, ornamental). Non-crop pests are pests of the classes Chilopoda and Diplopoda and of the orders Isoptera, Diptera, Blattaria (Blattodea), Dermaptera, Hemiptera, Hymenoptera, Orthoptera, Siphonaptera, Thysanura, Phthiraptera, and Acarina.

10 For use according to the present invention, the crystalline modification II can be converted into the customary formulations, for example solutions, emulsions, suspensions, dusts, powders, pastes and granules. The use form depends on the particular intended purpose; in each case, it should ensure a fine and even distribution of the compound according to the invention.

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The formulations are prepared in a known manner (see e.g. for review US 3,060,084, EP-A 707 445 (for liquid concentrates), Browning, "Agglomeration", Chemical Engineering, Dec. 4, 1967, 147-48, Perry's Chemical Engineer's Handbook, 4th Ed., McGraw-Hill, New York, 1963, pages 8-57 and et seq. WO 91/13546, US 4,172,714, 20 US 4,144,050, US 3,920,442, US 5,180,587, US 5,232,701, US 5,208,030, GB 2,095,558, US 3,299,566, Klingman, Weed Control as a Science, John Wiley and Sons, Inc., New York, 1961, Hance et al., Weed Control Handbook, 8th Ed., Blackwell Scientific Publications, Oxford, 1989 and Mollet, H., Grubemann, A., Formulation technology, Wiley VCH Verlag GmbH, Weinheim (Germany), 2001, 2. D. A. Knowles, 25 Chemistry and Technology of Agrochemical Formulations, Kluwer Academic Publishers, Dordrecht, 1998 (ISBN 0-7514-0443-8), for example by extending the active compound with auxiliaries suitable for the formulation of agrochemicals, such as solvents and/or carriers, if desired surfactants (e.g. adjuvans, emulsifiers, dispersing agents), preservatives, antifoaming agents, anti-freezing agents, for seed treatment 30 formulations also optionally colorants and/or binders and/or gelling agents.

Examples of suitable solvents are water, aromatic solvents (for example Solvesso products, xylene), paraffins (for example mineral oil fractions), alcohols (for example methanol, butanol, pentanol, benzyl alcohol), ketones (for example cyclohexanone, gamma-butyrolactone), pyrrolidones (NMP, NOP), acetates (glycol diacetate), glycols, fatty acid dimethylamides, fatty acids and fatty acid esters. In principle, solvent mixtures may also be used.

Examples of suitable carriers are ground natural minerals (for example kaolins, clays, talc, chalk) and ground synthetic minerals (for example highly disperse silica, silicates).

Suitable surfactants used are alkali metal, alkaline earth metal and ammonium salts of lignosulfonic acid, naphthalenesulfonic acid, phenolsulfonic acid, dibutylnaphthalenesulfonic acid, alkylarylsulfonates, alkyl sulfates, alkylsulfonates, fatty alcohol sulfates, fatty acids and sulfated fatty alcohol glycol ethers, furthermore condensates of sulfonated naphthalene and naphthalene derivatives with 5 formaldehyde, condensates of naphthalene or of naphthalenesulfonic acid with phenol and formaldehyde, polyoxyethylene octylphenol ether, ethoxylated isooctylphenol. octylphenol, nonylphenol, alkylphenol polyglycol ethers, tributylphenyl polyglycol ether, tristearylphenyl polyglycol ether, alkylaryl polyether alcohols, alcohol and fatty alcohol 10 ethylene oxide condensates, ethoxylated castor oil, polyoxyethylene alkyl ethers, ethoxylated polyoxypropylene, lauryl alcohol polyglycol ether acetal, sorbitol esters, lignosulfite waste liquors and methylcellulose.

Substances which are suitable for the preparation of directly sprayable solutions, 15 emulsions, pastes or oil dispersions are mineral oil fractions of medium to high boiling point, such as kerosene or diesel oil, furthermore coal tar oils and oils of vegetable or animal origin, aliphatic, cyclic and aromatic hydrocarbons, for example toluene, xylene, paraffin, tetrahydronaphthalene, alkylated naphthalenes or their derivatives, methanol, ethanol, propanol, butanol, cyclohexanol, cyclohexanone, isophorone, highly polar 20 solvents, for example dimethyl sulfoxide, N-methylpyrrolidone or water.

Also anti-freezing agents such as glycerin, ethylene glycol, propylene glycol and bactericides can be added to the formulation.

25 Suitable antifoaming agents are for example antifoaming agents based on silicon or magnesium stearate.

Suitable preservatives are for example Dichlorophen und enzylalkoholhemiformal.

30 Seed Treatment formulations may additionally comprise binders and optionally colorants.

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Binders can be added to improve the adhesion of the active materials on the seeds after treatment. Suitable binders are block copolymers EO/PO surfactants but also polyvinylalcoholsl, polyvinylpyrrolidones, polyacrylates, polymethacrylates, polybutenes, polyisobutylenes, polystyrene, polyethyleneamines, polyethyleneamides, polyethyleneimines (Lupasol®, Polymin®), polyethers, polyurethans, polyvinylacetate, tylose and copolymers derived from these polymers.

40 Optionally, also colorants can be included in the formulation. Suitable colorants or dyes for seed treatment formulations are Rhodamin B, C.I. Pigment Red 112, C.I. Solvent

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Red 1, pigment blue 15:4, pigment blue 15:3, pigment blue 15:2, pigment blue 15:1, pigment blue 80, pigment yellow 1, pigment yellow 13, pigment red 112, pigment red 48:2, pigment red 48:1, pigment red 57:1, pigment red 53:1, pigment orange 43, pigment orange 34, pigment orange 5, pigment green 36, pigment green 7, pigment white 6, pigment brown 25, basic violet 10, basic violet 49, acid red 51, acid red 52, acid red 14, acid blue 9, acid yellow 23, basic red 10, basic red 108.

Examples of a gelling agent is carrageen (Satiagel®).

10 Powders, materials for spreading and dustable products can be prepared by mixing or concomitantly grinding the active substances with a solid carrier.

Granules, for example coated granules, impregnated granules and homogeneous granules, can be prepared by binding the active compounds to solid carriers.

Examples of solid carriers are mineral earths such as silica gels, silicates, talc, kaolin, attaclay, limestone, lime, chalk, bole, loess, clay, dolomite, diatomaceous earth, calcium sulfate, magnesium sulfate, magnesium oxide, ground synthetic materials, fertilizers, such as, for example, ammonium sulfate, ammonium phosphate, ammonium nitrate, ureas, and products of vegetable origin, such as cereal meal, tree bark meal, wood meal and nutshell meal, cellulose powders and other solid carriers.

In general, the formulations comprise from 0.01 to 95% by weight, preferably from 0.1 to 90% by weight, of the active compound(s). In this case, the active compound(s) are employed in a purity of from 90% to 100% by weight, preferably 95% to 100% by weight (according to NMR spectrum).

For seed treatment purposes, the respective formulations can be diluted 2-10 fold leading to concentrations in the ready to use preparations of 0.01 to 60% by weight active compound by weight, preferably 0.1 to 40% by weight.

The crystalline modification II can be used as such, in the form of their formulations or the use forms prepared therefrom, for example in the form of directly sprayable solutions, powders, suspensions or dispersions, emulsions, oil dispersions, pastes, dustable products, materials for spreading, or granules, by means of spraying, atomizing, dusting, spreading or pouring. The use forms depend entirely on the intended purposes; they are intended to ensure in each case the finest possible distribution of the active compound(s) according to the invention.

40 Aqueous use forms can be prepared from emulsion concentrates, pastes or wettable powders (sprayable powders, oil dispersions) by adding water. To prepare emulsions, pastes or oil dispersions, the substances, as such or dissolved in an oil or solvent, can

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be homogenized in water by means of a wetter, tackifier, dispersant or emulsifier. However, it is also possible to prepare concentrates composed of active substance, wetter, tackifier, dispersant or emulsifier and, if appropriate, solvent or oil, and such concentrates are suitable for dilution with water.

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The active compound concentrations in the ready-to-use preparations can be varied within relatively wide ranges. In general, they are from 0.0001 to 10%, preferably from 0.01 to 1% per weight.

The active compound(s) may also be used successfully in the ultra-low-volume process (ULV), it being possible to apply formulations comprising over 95% by weight of active compound, or even to apply the active compound without additives.

The following are examples of formulations: 1. Products for dilution with water for foliar applications. For seed treatment purposes, such products may be applied to the seed diluted or undiluted.

A) Water-soluble concentrates (SL, LS)

10 parts by weight of the active compound(s) are dissolved in 90 parts by weight of water or a water-soluble solvent. As an alternative, wetters or other auxiliaries are added. The active compound(s) dissolves upon dilution with water, whereby a formulation with 10 % (w/w) of active compound(s) is obtained.

B) Dispersible concentrates (DC)

20 parts by weight of the active compound(s) are dissolved in 70 parts by weight of cyclohexanone with addition of 10 parts by weight of a dispersant, for example polyvinylpyrrolidone. Dilution with water gives a dispersion, whereby a formulation with 20% (w/w) of active compound(s) is obtained.

30 C) Emulsifiable concentrates (EC)

15 parts by weight of the active compound(s) are dissolved in 80 parts by weight of xylene with addition of calcium dodecylbenzenesulfonate and castor oil ethoxylate (in each case 5 parts by weight). Dilution with water gives an emulsion, whereby a formulation with 15% (w/w) of active compound(s) is obtained.

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D) Emulsions (EW, EO, ES)

25 parts by weight of the active compound(s) are dissolved in 35 parts by weight of xylene with addition of calcium dodecylbenzenesulfonate and castor oil ethoxylate (in each case 5 parts by weight). This mixture is introduced into 30 parts by weight of water by means of an emulsifier machine (e.g. Ultraturrax) and made into a homogeneous emulsion. Dilution with water gives an emulsion, whereby a formulation with 25% (w/w) of active compound(s) is obtained.

E) Suspensions (SC, OD, FS)

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In an agitated ball mill, 20 parts by weight of the active compound(s) are comminuted with addition of 10 parts by weight of dispersants, wetters and 70 parts by weight of water or of an organic solvent to give a fine active compound(s) suspension. Dilution with water gives a stable suspension of the active compound(s), whereby a formulation with 20% (w/w) of active compound(s) is obtained.

- F) Water-dispersible granules and water-soluble granules (WG, SG)

 50 parts by weight of the active compound(s) are ground finely with addition of 50 parts by weight of dispersants and wetters and made as water-dispersible or water-soluble granules by means of technical appliances (for example extrusion, spray tower, fluidized bed). Dilution with water gives a stable dispersion or solution of the active compound(s), whereby a formulation with 50% (w/w) of active compound(s) is obtained.
- G) Water-dispersible powders and water-soluble powders (WP, SP, SS, WS)
 75 parts by weight of the active compound(s) are ground in a rotor-stator mill with addition of 25 parts by weight of dispersants, wetters and silica gel. Dilution with water
 gives a stable dispersion or solution of the active compound(s), whereby a formulation with 75% (w/w) of active compound(s) is obtained.
 - H) Gel-Formulation (GF) (for seed treatment purposes only)
- In an agitated ball mill, 20 parts by weight of the active compound(s) are comminuted with addition of 10 parts by weight of dispersants, 1 part by weight of a gelling agent/ wetters and 70 parts by weight of water or of an organic solvent to give a fine active compound(s) suspension. Dilution with water gives a stable suspension of the active compound(s), whereby a formulation with 20% (w/w) of active compound(s) is obtained.
 - 2. Products to be applied undiluted for foliar applications. For seed treatment purposes, such products may be applied to the seed diluted.
- 35 I) Dustable powders (DP, DS)
 5 parts by weight of the active compound(s) are ground finely and mixed intimately with
 95 parts by weight of finely divided kaolin. This gives a dustable product having 5%
 (w/w) of active compound(s)
- 40 J) Granules (GR, FG, GG, MG)
 0.5 part by weight of the active compound(s) is ground finely and associated with 95.5 parts by weight of carriers, whereby a formulation with 0.5% (w/w) of active

compound(s) is obtained. Current methods are extrusion, spray-drying or the fluidized bed. This gives granules to be applied undiluted for foliar use.

K) ULV solutions (UL)

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5 10 parts by weight of the active compound(s) are dissolved in 90 parts by weight of an organic solvent, for example xylene. This gives a product having 10% (w/w) of active compound(s), which is applied undiluted for foliar use.

Conventional seed treatment formulations include for example flowable concentrates

FS, solutions LS, powders for dry treatment DS, water dispersible powders for slurry treatment WS, water-soluble powders SS and emulsion ES and EC and gel formulation GF. These formulation can be applied to the seed diluted or undiluted. Application to the seeds is carried out before sowing, either directly on the seeds.

- In a preferred embodiment a FS formulation is used for seed treatment. Typcially, a FS formulation may comprise 1-800 g/l of active ingredient, 1-200 g/l surfactant, 0 to 200 g/l antifreezing agent, 0 to 400 g/l of binder, 0 to 200 g/l of a pigment and up to 1 liter of a solvent, preferably water.
- The invention relates in particular to pesticidal or parasiticidal compositions in the form of an aqueous suspension concentrate (SC). Such suspension concentrates comprise the crystalline modification II in a finely divided particulate form, where the particles of the crystalline modification II are suspended in an aqueous medium. The size of the active compound particles, i.e. the size which is not exceeded by 90% by weight of the active compound particles, is typically below 30 μm, in particular below 20 μm. Advantageously, at least 40% by weight and in particular at least 60% by weight of the particles in the SCs according to the invention have diameters below 2 μm.
- In addition to the active compound, suspension concentrates typically comprise surfactants, and also, if appropriate, antifoam agents, thickeners, antifreeze agents, stabilizers (biocides), agents for adjusting the pH and anticaking agents.

In such SCs, the amount of active compound, i.e. the total amount of the crystalline modification II and, if appropriate, further active compounds is usually in the range from 10 to 70% by weight, in particular in the range from 20 to 50% by weight, based on the total weight of the suspension concentrate.

Preferred surfactants are anionic and nonionic surfactants. The amount of surfactants will generally be from 0.5 to 20% by weight, in particular from 1 to 15% by weight and particularly preferably from 1 to 10% by weight, based on the total weight of the SCs according to the invention. Preferably, the surfactants comprise at least one anionic

surfactant and at least one nonionic surfactant, the ratio of anionic to nonionic surfactant typically being in the range from 10:1 to 1:10.

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Examples of anionic surfactants include alkylaryl sulfonates, phenyl sulfonates, alkyl sulfates, alkyl sulfonates, alkyl ether sulfates, alkylaryl ether sulfates, alkyl polyglycol ether phosphates, polyaryl phenyl ether phosphates, alkyl sulfosuccinates, olefin sulfonates, paraffin sulfonates, petroleum sulfonates, taurides, sarcosides, fatty acids, alkylnaphthalenesulfonic acids, naphthalenesulfonic acids, lignosulfonic acids, condensates of sulfonated naphthalenes with formaldehyde or with formaldehyde and phenol and, if appropriate, urea, and also condensates of phenolsulfonic acid, formaldehyde and urea, lignosulfite waste liquors and lignosulfonates, alkyl phosphates, alkylaryl phosphates, for example tristyryl phosphates, and also polycarboxylates, such as, for example, polyacrylates, maleic anhydride/olefin copolymers (for example Sokalan® CP9, BASF), including the alkali metal, alkaline earth metal, ammonium and amine salts of the substances mentioned above. Preferred anionic surfactants are those which carry at least one sulfonate group, and in particular their alkali metal and their ammonium salts.

Examples of nonionic surfactants comprise alkylphenol alkoxylates, alcohol 20 alkoxylates, fatty amine alkoxylates, polyoxyethylene glycerol fatty acid esters, castor oil alkoxylates, fatty acid alkoxylates, fatty amide alkoxylates, fatty polydiethanolamides, lanolin ethoxylates, fatty acid polyglycol esters, isotridecyl alcohol, fatty amides, methylcellulose, fatty acid esters, alkyl polyglycosides, glycerol fatty acid esters, polyethylene glycol, polypropylene glycol, polyethylene 25 glycol/polypropylene glycol block copolymers, polyethylene glycol alkyl ethers, polypropylene glycol alkyl ethers, polyethylene glycol/polypropylene glycol ether block copolymers (polyethylene oxide/polypropylene oxide block copolymers) and mixtures thereof. Preferred nonionic surfactants are fatty alcohol ethoxylates, alkyl polyglycosides, glycerol fatty acid esters, castor oil alkoxylates, fatty acid alkoxylates, 30 fatty amide alkoxylates, lanolin ethoxylates, fatty acid polyglycol esters and ethylene oxide/ propylene oxide block copolymers and mixtures thereof.

In particular, the SCs according to the invention comprise at least one surfactant which improves wetting of the plant parts by the aqueous application form (wetting agent) and at least one surfactant which stabilizes the dispersion of the active compound particles in the SC (dispersant). The amount of wetting agent is typically in the range from 0.5 to 10% by weight, in particular from 0.5 to 5% by weight and especially from 0.5 to 3% by weight, based on the total weight of the SC. The amount of dispersant is typically from 0.5 to 10% by weight and in particular from 0.5 to 5% by weight, based on the total weight of the SC.

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Preferred wetting agents are of anionic or nonionic nature and selected, for example, from naphthalenesulfonic acids including their alkali metal, alkaline earth metal, ammonium and amine salts, furthermore fatty alcohol ethoxylates, alkyl polyglycosides, glycerol fatty acid esters, castor oil alkoxylates, fatty acid alkoxylates, fatty amide alkoxylates, fatty polydiethanolamides, lanolin ethoxylates and fatty acid polyglycol esters.

Preferred dispersants are of anionic or nonionic nature and selected, for example, from polyethylene glycol/polypropylene glycol block copolymers, polyethylene glycol alkyl ethers, polyethylene glycol/polypropylene glycol ether block copolymers, alkylaryl phosphates, for example tristyryl phosphates, lignosulfonic acids, condensates of sulfonated naphthalenes with formaldehyde or with formaldehyde and phenol and, if appropriate, urea, and also condensates of phenolsulfonic acid, formaldehyde and urea, lignosulfite waste liquors and lignosulfonates, polycarboxylates, such as, for example, polyacrylates, maleic anhydride/olefin copolymers (for example Sokalan® CP9, BASF), including the alkali metal, alkaline earth metal, ammonium and amine salts of the substances mentioned above.

Viscosity-modifying additives (thickeners) suitable for the SCs according to the invention are in particular compounds which bestow upon the formulation pseudoplastic flow properties, i.e. high viscosity in the resting state and low viscosity in the agitated state. Suitable are, in principle, all compounds used for this purpose in suspension concentrates. Mention may be made, for example, of inorganic substances, such as bentonites or attapulgites (for example Attaclay® from Engelhardt), and organic substances, such as polysaccharides and heteropolysaccharides, such as xanthan gum such as sold under the trademarks Kelzan® from Kelco, Rhodopol® 23 from Rhone Poulenc or Veegum® from R.T. Vanderbilt, and preference is given to using xanthan gum. Frequently, the amount of viscosity-modifying additives is from 0.1 to 5% by weight, based on the total weight of the SC.

Antifoam agents suitable for the SCs according to the invention are, for example, silicone emulsions known for this purpose (Silikon® SRE, from Wacker, or Rhodorsil® from Rhodia), long-chain alcohols, fatty acids, defoamers of the type of aqueous wax dispersions, solid defoamers (so-called Compounds), organofluorine compounds and mixtures thereof. The amount of antifoam agent is typically from 0.1 to 1% by weight, based on the total weight of the SC.

Bactericides may be added for stabilizing the suspension concentrates according to the invention. Suitable bactericides are those based on isothiazolones, for example Proxel[®] from ICI or Acticide[®] RS from Thor Chemie or Kathon[®] MK from Rohm & Haas. The amount of bactericides is typically from 0.05 to 0.5% by weight, based on the total

weight of the SC.

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Suitable antifreeze agents are liquid polyols, for example ethylene glycol, propylene glycol or glycerol. The amount of antifreeze agents is generally from 1 to 20% by weight, in particular from 5 to 10% by weight, based on the total weight of the suspension concentrate.

If appropriate, the SCs according to the invention may comprise buffers for regulating the pH. Examples of buffers are alkali metal salts of weak inorganic or organic acids, such as, for example, phosphoric acid, boric acid, acetic acid, propionic acid, citric acid, fumaric acid, tartaric acid, oxalic acid and succinic acid.

The invention relates in particular to pesticidal or parasiticidal compositions in the form of water-dispersible granules (WG) or a water dispersible powder (WP). Such formulations comprise the crystalline modification II in a finely divided particulate form, where the particles of the crystalline modification II are homogenized in a solid or powder form. The size of the active compound particles, i.e. the size which is not exceeded by 90% by weight of the active compound particles, is typically below 30 μ m, in particular below 20 μ m. Advantageously, at least 40% by weight and in particular at least 60% by weight of the particles in the WGs or WPs according to the invention have diameters below 5 μ m.

In addition to the active compound, water-dispersible powders and water dispersible granules typically comprise surfactants, and also, if appropriate, antifoam agents, fillers, binders, and anticaking agents.

In such WGs and WPs, the amount of active compound, i.e. the total amount of the crystalline modification II and, if appropriate, further active compounds is usually in the range from 10 to 90% by weight, in particular in the range from 20 to 75% by weight, based on the total weight of the WG/WP.

Preferred surfactants are anionic and nonionic surfactants. The amount of surfactants will generally be from 0.5 to 20% by weight, in particular from 1 to 15% by weight and particularly preferably from 1 to 10% by weight, based on the total weight of the WGs or WPs according to the invention. Preferably, the surfactants comprise at least one anionic surfactant and at least one nonionic surfactant, the ratio of anionic to nonionic surfactant typically being in the range from 10:1 to 1:10.

Examples of anionic surfactants include alkylaryl sulfonates, phenyl sulfonates, alkyl sulfates, alkyl sulfonates, alkyl ether sulfates, alkylaryl ether sulfates, alkyl polyglycol ether phosphates, polyaryl phenyl ether phosphates, alkyl sulfosuccinates, olefin sulfonates, paraffin sulfonates, petroleum sulfonates, taurides, sarcosides, fatty acids,

alkylnaphthalenesulfonic acids, naphthalenesulfonic acids, lignosulfonic acids, condensates of sulfonated naphthalenes with formaldehyde or with formaldehyde and phenol and, if appropriate, urea, and also condensates of phenolsulfonic acid, formaldehyde and urea, lignosulfite waste liquors and lignosulfonates, alkyl phosphates, alkylaryl phosphates, for example tristyryl phosphates, and also polycarboxylates, such as, for example, polyacrylates, maleic anhydride/olefin copolymers (for example Sokalan® CP9, BASF), including the alkali metal, alkaline earth metal, ammonium and amine salts of the substances mentioned above. Preferred anionic surfactants are those which carry at least one sulfonate group, and in particular their alkali metal and their ammonium salts.

Examples of nonionic surfactants comprise alkylphenol alkoxylates, alcohol alkoxylates, fatty amine alkoxylates, polyoxyethylene glycerol fatty acid esters, castor oil alkoxylates, fatty acid alkoxylates, fatty amide alkoxylates, fatty polydiethanolamides, lanolin ethoxylates, fatty acid polyglycol esters, isotridecyl alcohol, fatty amides, methylcellulose, fatty acid esters, alkyl polyglycosides, glycerol fatty acid esters, polyethylene glycol, polypropylene glycol, polyethylene glycol alkyl ethers, polypropylene glycol alkyl ethers, polypropylene glycol alkyl ethers, polyethylene glycol/polypropylene glycol ether block copolymers (polyethylene oxide/polypropylene oxide block copolymers) and mixtures thereof. Preferred nonionic surfactants are fatty alcohol ethoxylates, alkyl polyglycosides, glycerol fatty acid esters, castor oil alkoxylates, fatty acid alkoxylates, fatty amide alkoxylates, lanolin ethoxylates, fatty acid polyglycol esters and ethylene oxide/ propylene oxide block copolymers and mixtures thereof.

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In particular, the WGs or WPs according to the invention comprise at least one surfactant which improves wetting of the formulation by the aqueous application form (wetting agent) and at least one surfactant which allows dispersion of the active compound particles in aqueous dilutions. The amount of wetting agent is typically in the range from 0.5 to 10% by weight, in particular from 0.5 to 5% by weight and especially from 0.5 to 3% by weight, based on the total weight of the WG/WP. The amount of dispersant is typically from 0.5 to 10% by weight and in particular from 2.0 to 8% by weight, based on the total weight of the WG/WP.

Preferred wetting agents are of anionic or nonionic nature and selected, for example, from naphthalenesulfonic acids including their alkali metal, alkaline earth metal, ammonium and amine salts, furthermore fatty alcohol ethoxylates, alkyl polyglycosides, glycerol fatty acid esters, castor oil alkoxylates, fatty acid alkoxylates, fatty amide alkoxylates, fatty polydiethanolamides, lanolin ethoxylates and fatty acid polyglycol esters.

Preferred dispersants are of anionic or nonionic nature and selected, for example, from polyethylene glycol/polypropylene glycol block copolymers, polyethylene glycol alkyl ethers, polyethylene glycol/polypropylene glycol ether block copolymers, alkylaryl phosphates, for example tristyryl phosphates, sodium phosphates, sodium lauryl sulphate, modified cellulose gum, polyvinylpyrrolidinone, lignosulfonic acids, condensates of sulfonated naphthalenes with formaldehyde or with formaldehyde and phenol and, if appropriate, urea, and also condensates of phenol-sulfonic acid, formaldehyde and urea, lignosulfite waste liquors and lignosulfonates, polycarboxylates, such as, for example, polyacrylates, maleic anhydride/olefin copolymers (for example Sokalan® CP9, BASF), including the alkali metal, alkaline earth metal, ammonium and amine salts of the substances mentioned above.

Antifoam agents suitable for the WGs or WPs according to the invention are, for example, tallow soap known for this purpose (Agnique Soap L, Foamaster Soap L), long-chain alcohols, fatty acids, organofluorine compounds and mixtures thereof. The amount of antifoam agent is typically from 0.1 to 1% by weight, based on the total weight of the WG/WP.

Fillers, binders, or additional dispersing aids suitable for the WGs and WPs according to the invention typically make up the remainer of the formulation. These typically are for example kaolin or attapulgite clay, fumed or precipitated silica, diatomateous earth, ammonium sulphate, or calcium silicate.

The crystalline modification II is effective through both contact and ingestion.

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According to a preferred embodiment of the invention, the crystalline modification II is employed via soil application. Soil application is especially favorable for use against ants, termites, crickets, or cockroaches.

According to another preferred embodiment of the invention, for use against non-crop pests such as ants, termites, wasps, flies, mosquitoes, crickets, locusts, or cockroaches the crystalline modification II is prepared into a bait preparation.

The bait can be a liquid, a solid or a semisolid preparation (e.g. a gel). Solid baits can be formed into various shapes and forms suitable to the respective application e.g. granules, blocks, sticks, disks. Liquid baits can be filled into various devices to ensure proper application, e.g. open containers, spray devices, droplet sources, or evaporation sources. Gels can be based on aqueous or oily matrices and can be formulated to particular necessities in terms of stickiness, moisture retention or aging characteristics.

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The bait employed in the composition is a product which is sufficiently attractive to incite insects such as ants, termites, wasps, flies, mosquitoes, crickets etc. or cockroaches to eat it. This attractant may be chosen from feeding stimulants or para and/or sex pheromones. Suitable feeding stimulants are chosen, for example, from animal and/or plant proteins (meat-, fish- or blood meal, insect parts, crickets powder, egg yolk), from fats and oils of animal and/or plant origin, or mono-, oligo- or polyorganosaccharides, especially from sucrose, lactose, fructose, dextrose, glucose, starch, pectin or even molasses or honey, or from salts such as ammonium sulfate, ammonium carbonate or ammonium acetate. Fresh or decaying parts of fruits, crops, plants, animals, insects or specific parts thereof can also serve as a feeding stimulant. Pheromones are known to be more insect specific. Specific pheromones are described in the literature and are known to those skilled in the art.

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Compositions of this invention may also contain other active ingredients, for example other pesticides, insecticides, fungicides, herbicides, fertilizers such as ammonium nitrate, urea, potash, and superphosphate, phytotoxicants and plant growth regulators, safeners and nematicides. These additional ingredients may be used sequentially or in combination with the above-described compositions, if appropriate also added only immediately prior to use (tank mix). For example, the plant(s) may be sprayed with a composition of this invention either before or after being treated with other active ingredients.

The following list of pesticidal or parasiticidal compounds which can be used together with the crystalline modification II according to the invention is intended to illustrate the possible combinations, but not to impose any limitation:

- A.1. Organo(thio)phosphates: acephate, azamethiphos, azinphos-methyl, chlorpyrifos, chlorpyrifos-methyl, chlorfenvinphos, diazinon, dichlorvos, dicrotophos, dimethoate, disulfoton, ethion, fenitrothion, fenthion, isoxathion, malathion, methamidophos, methidathion, methyl-parathion, mevinphos, monocrotophos, oxydemeton-methyl, paraoxon, parathion, phenthoate, phosalone, phosmet, phosphamidon, phorate, phoxim, pirimiphos-methyl, profenofos, prothiofos, sulprophos, tetrachlorvinphos, terbufos, triazophos, trichlorfon;
- A.2. Carbamates: alanycarb, aldicarb, bendiocarb, benfuracarb, carbaryl, carbofuran, carbosulfan, fenoxycarb, furathiocarb, methiocarb, methomyl, oxamyl, pirimicarb, propoxur, thiodicarb, triazamate;
- A.3. Pyrethroids: allethrin, bifenthrin, cyfluthrin, cyhalothrin, cyphenothrin,
 cypermethrin, alpha-cypermethrin, beta-cypermethrin, zeta-cypermethrin, deltamethrin,
 esfenvalerate, etofenprox, fenpropathrin, fenvalerate, imiprothrin, lambda-cyhalothrin,

permethrin, prallethrin, pyrethrin I and II, resmethrin, silafluofen, tau-fluvalinate, tefluthrin, tetramethrin, tralomethrin, transfluthrin, profluthrin, dimefluthrin;

A.4. Growth regulators: a) chitin synthesis inhibitors: benzoylureas: chlorfluazuron,
diflubenzuron, flucycloxuron, flufenoxuron, hexaflumuron, lufenuron, novaluron,
teflubenzuron, triflumuron; buprofezin, diofenolan, hexythiazox, etoxazole, clofentazine;
b) ecdysone antagonists: halofenozide, methoxyfenozide, tebufenozide, azadirachtin;
c) juvenoids: pyriproxyfen, methoprene, fenoxycarb; d) lipid biosynthesis inhibitors:
spirodiclofen, spiromesifen, spirotetramat;

A.5. Nicotinic receptor agonists/antagonists compounds: clothianidin, dinotefuran, imidacloprid, thiamethoxam, nitenpyram, acetamiprid, thiacloprid; the thiazol compound of formula Γ^1

A.6. GABA antagonist compounds: acetoprole, endosulfan, ethiprole, fipronil, vaniliprole, pyrafluprole, 5-Amino-1-(2,6-dichloro-4-trifluoromethyl-phenyl)-4-trifluoromethanesulfinyl-1H-pyrazole-3-carbothioic acid amide of formula Γ^2

$$S = NH_2$$
 CI CF_3 NH_2 CI CF_3 CF_3

- A.7. Macrocyclic lactone insecticides: abamectin, emamectin, milbemectin, lepimectin, spinosad;
 - A.8. METI I compounds: fenazaquin, pyridaben, tebufenpyrad, tolfenpyrad, flufenerim;
 - A.9. METI II and III compounds: acequinocyl, fluacyprim, hydramethylnon;
 - A.10. Uncoupler compounds: chlorfenapyr;

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- A.11. Oxidative phosphorylation inhibitor compounds: cyhexatin, diafenthiuron, fenbutatin oxide, propargite;
- A.12. Moulting disruptor compounds: cyromazine;
- A.13. Mixed Function Oxidase inhibitor compounds: piperonyl butoxide;

A.14. Sodium channel blocker compounds: indoxacarb, metaflumizone,

A.15. Various: benclothiaz, bifenazate, cartap, flonicamid, pyridalyl, pymetrozine, sulfur, thiocyclam, flubendiamide, cyenopyrafen, flupyrazofos, cyflumetofen, amidoflumet,

the anthranilamide compounds of formula Γ^3

wherein A¹ is CH₃, CI, Br, I, X is C-H, C-CI, C-F or N, Y' is F, CI, or Br, Y" is hydrogen, F, CI, CF₃, B¹ is hydrogen, CI, Br, I, CN, B² is CI, Br, CF₃, OCH₂CF₃, OCF₂H, and R^B is hydrogen, CH₃ or CH(CH₃)₂, and the malononitrile compounds as described in JP 2002 284608, WO 02/89579, WO 02/90320, WO 02/90321, WO 04/06677, WO 04/20399, JP 2004 99597, WO 05/68423, WO 05/68432, or WO 05/63694, especially the malononitrile compounds CF₂HCF₂CF₂CF₂CH₂C(CN)₂CH₂CH₂CF₃ (2-(2,2,3,3,4,4,5,5-octafluoropentyl)-2-(3,3,3-trifluoropropyl)malononitrile),

15 CF₃(CH₂)₂C(CN)₂CH₂(CF₂)₅CF₂H (2-(2,2,3,3,4,4,5,5,6,6,7,7-Dodecafluoro-heptyl)-2-(3,3,3-trifluoro-propyl)-malononitrile), CF₃(CH₂)₂C(CN)₂(CH₂)₂C(CF₃)₂F (2-(3,4,4,4-Tetrafluoro-3-trifluoromethyl-butyl)-2-(3,3,3-trifluoro-propyl)-malononitrile), CF₃(CH₂)₂C(CN)₂(CH₂)₂(CF₂)₃CF₃ (2-(3,3,4,4,5,5,6,6,6-Nonafluoro-hexyl)-2-(3,3,3-trifluoro-propyl)-malononitrile), CF₂H(CF₂)₃CH₂C(CN)₂CH₂(CF₂)₃CF₂H (2,2-Bis-(2,2,3,3,4,4,5,5-octafluoro-pentyl)-malononitrile), CF₃(CH₂)₂C(CN)₂CH₂(CF₂)₃CF₃ (2-(2,2,3,3,4,4,5,5-octafluoro-pentyl)-malononitrile).

 $(2,2,3,3,4,4,5,5\text{-octafluoro-pentyl})\text{-malononitrile}), CF_3(CH_2)_2C(CN)_2CH_2(CF_2)_3CF_3 \ (2-(2,2,3,3,4,4,5,5,5\text{-Nonafluoro-pentyl})-2-(3,3,3\text{-trifluoro-propyl})\text{-malononitrile}), \\ CF_3(CF_2)_2CH_2C(CN)_2CH_2(CF_2)_3CF_2H \ (2-(2,2,3,3,4,4,4\text{-Heptafluoro-butyl})-2-(2,2,3,3,4,4,5,5\text{-octafluoro-pentyl})\text{-malononitrile}) \ \text{and} \ CF_3CF_2CH_2C(CN)_2CH_2(CF_2)_3CF_2H \ (2-(2,2,3,3,4,4,5,5\text{-Octafluoro-pentyl})-2-(2,2,3,3,3\text{-pentafluoro-propyl})\text{-malononitrile}).$

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The commercially available compounds of the group A may be found in The Pesticide Manual, 13^{th} Edition, British Crop Protection Council (2003) among other publications. Thioamides of formula Γ^2 and their preparation have been described in WO 98/28279. Lepimectin is known from Agro Project, PJB Publications Ltd, November 2004.

30 Benclothiaz and its preparation have been described in EP-A1 454621. Methidathion and Paraoxon and their preparation have been described in Farm Chemicals Handbook, Volume 88, Meister Publishing Company, 2001. Acetoprole and its preparation have been described in WO 98/28277. Metaflumizone and its preparation have been described in EP-A1 462 456. Flupyrazofos has been described in Pesticide 35 Science 54, 1988, p.237-243 and in US 4822779. Pyrafluprole and its preparation have been described in JP 2002193709 and in WO 01/00614. Pyriprole and its preparation

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have been described in WO 98/45274 and in US 6335357. Amidoflumet and its preparation have been described in US 6221890 and in JP 21010907. Flufenerim and its preparation have been described in WO 03/007717 and in WO 03/007718. Cyflumetofen and its preparation have been described in WO 04/080180.

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- Anthranilamides of formula Γ³ and their preparation have been described in WO 01/70671; WO 02/48137; WO 03/24222, WO 03/15518, WO 04/67528; WO 04/33468; and WO 05/118552. The malononitrile compounds CF₂HCF₂CF₂CH₂C(CN)₂CH₂CH₂CF₃ (2-(2,2,3,3,4,4,5,5-octafluoropentyl)-2-(3,3,3-trifluoropropyl)malononitrile), CF₃(CH₂)₂C(CN)₂CH₂(CF₂)₅CF₂H (2-
- malononitrile), $CF_3(CH_2)_2C(CN)_2CH_2(CF_2)_3CF_3$ (2-(2,2,3,3,4,4,5,5,5-Nonafluoro-pentyl)-2-(3,3,3-trifluoro-propyl)-malononitrile), $CF_3(CF_2)_2CH_2C(CN)_2CH_2(CF_2)_3CF_2H$ (2-(2,2,3,3,4,4,4-Heptafluoro-butyl)-2-(2,2,3,3,4,4,5,5-octafluoro-pentyl)-malononitrile) and $CF_3CF_2CH_2C(CN)_2CH_2(CF_2)_3CF_2H$ (2-(2,2,3,3,4,4,5,5-Octafluoro-pentyl)-2-(2,2,3,3,3-pentafluoro-propyl)-malononitrile) have been described in WO 05/63694.

The following list of fungicidal compounds which can be used together with the crystalline modification II according to the invention is intended to illustrate the possible combinations, but not to impose any limitation:

25 Preferred are the binary mixtures containing modification II as compound I.

Preferred are the tertiary mixtures containing modification II as compound I, a compound IIA, and a compound IIB.

Preferred are the quaternary mixtures containing modification II as compound I, a compound IIA, and two compounds IIB1 and IIB2, resp.

Especially preferred are binary mixtures containing modification II as compound I and a fungicidal compound IIA selected from the list comprising azoles: cyproconazole, difenoconazole, epoxiconazole, fenbuconazole, fluquinconazole, flutriafol, hexaconazole, ipconazole, metconazole, propiconazole, prothioconazole, tebuconazole, tetraconazole, triadimenol, triadimefon, triticonazole, cyazofamid, imazalil, prochloraz, triflumizol, benomyl, carbendazim, thiabendazole, ethaboxam, and hymexazole.

40 Especially preferred are binary mixtures containing modification II as compound I and a fungicidal compound IIA selected from the list comprising strobilurins: azoxystrobin, dimoxystrobin, enestroburin, fluoxastrobin, kresoxim-methyl, metominostrobin, picoxy-

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strobin, pyraclostrobin, trifloxystrobin, methyl (2-chloro-5-[1-(3-methylbenzyloxyimino)ethyl]benzyl)carbamate, methyl (2-chloro-5-[1-(6-methylpyridin-2-ylmethoxyimino)ethyl]benzyl)carbamate, and methyl 2-(ortho-((2,5-dimethylphenyloxymethylene)phenyl)-3-methoxyacrylate;

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Especially preferred are binary mixtures containing modification II as compound I and a fungicidal compound IIA selected from the list comprising carboxamides: boscalid, carboxin, benalaxyl, fenhexamid, flutolanil, furametpyr, metalaxyl, mefenoxam (metalaxyl-M), ofurace, oxadixyl, oxycarboxin, penthiopyrad, thifluzamide, tiadinil, dimethomorph, fluopicolide (picobenzamid), diclocymet, N-(4'-bromobiphenyl-2-yl)-4-difluoromethyl-2-methylthiazole-5-carboxamide, N-(4'-trifluoromethylbiphenyl-2-yl)-4-difluoromethyl-2-methylthiazole-5-carboxamide, N-(4'-chloro-3'-fluorobiphenyl-2-yl)-4-difluoromethyl-2-methylthiazole-5-carboxamide, N-(3',4'-dichloro-4-fluorobiphenyl-2-yl)-3difluoromethyl-1-methylpyrazole-4-carboxamide, N-(3',4'-dichloro-5-fluorobiphenyl-2yl)-3-difluoromethyl-1-methylpyrazole-4-carboxamide; 3,4-dichloro-N-(2-cyanophenyl)isothiazol-5-carboxamide; N-(2',4'-difluorobiphenyl-2-yl)-1-methyl-3-trifluoromethyl-1H-pyrazole- 4-carboxamide; N-(2',4'-dichlorobiphenyl-2-yl)-1-methyl-3-trifluoromethyl-1H-pyrazole- 4-carboxamide; N-(2',4'-difluorobiphenyl-2-yl)-3-difluoromethyl-1-methyl-1H-pyrazole- 4-carboxamide; N-(2',4'-dichloro-biphenyl-2-yl)-3-difluoromethyl-1-methyl-1H-pyrazole-4-carboxamide; N-(2',5'-difluorobiphenyl-2-yl)-1-methyl-3-trifluoromethyl-1H-pyrazole-4-carboxamide; N-(2',5'-dichlorobiphenyl-2-yl)-1-methyl-3-trifluoromethyl-1H-pyrazole-4-carboxamide; N-(2',5'-difluorobiphenyl-2-yl)-3-difluoromethyl-1-methyl-1H-pyrazole-4-carboxamide; N-(2',5'-dichlorobiphenyl-2-yl)-3-di-fluoromethyl-1-methyl-1H-pyrazole- 4-carboxamide; N-(3',5'-difluoro-biphenyl-2-yl)-1-methyl-3-trifluoromethyl-1H-pyrazole- 4-carboxamide; N-(3',5'-dichlorobiphenyl-2-yl)-1-methyl-3-trifluoromethyl-1H-pyrazole-4-carboxamide, N-(3',5'-difluorobiphenyl-2-yl)-3-difluoromethyl-1-methyl-1H-pyrazole-4-carboxamide; N-(3',5'-dichlorobiphenyl-2-yl)-3-difluoromethyl-1-methyl-1H-pvrazole-4-carboxamide: N-(3'-fluorobiphenyl-2-yl)-1-methyl-3-trifluoro-methyl-1Hpyrazole-4-carboxamide, N-(3'-chlorobiphenyl-2-yl)-1-methyl-3-trifluoromethyl-1Hpyrazole-4-carboxamide; N-(3'-fluorobiphenyl-2-yl)-3-difluoromethyl-1-methyl-1Hpyrazole- 4-carboxamide, N-(3'-chlorobiphenyl-2-yl)-3-difluoromethyl-1-methyl-1Hpyrazole-4-carboxamide; N-(2'-fluorobiphenyl-2-yl)-1-methyl-3-trifluoromethyl-1Hpyrazole-4-carboxamide; N-(2'-chlorobiphenyl-2-yl)-1-methyl-3-trifluoro-methyl-1Hpyrazole-4-carboxamide; N-(2'-fluorobiphenyl-2-yl)-3-difluoro-methyl-1-methyl-1Hpyrazole-4-carboxamide; N-(2'-chlorbiphenyl-2-yl)-3-difluoromethyl-1-methyl-1Hpyrazole-4-carboxamide; N-(2'-fluoro-4'-chloro-5'-methylbiphenyl-2-yl)-1-methyl-3trifluoromethyl-1H-pyrazole-4-carbox-amide; N-(3',4',5'-trifluorobiphenyl-2-yl)-1-methyl-3-trifluoromethyl-1H-pyrazole-4-carboxamide; N-(3',4',5'-trifluorobiphenyl-2-yl)-1methyl-3-difluoromethyl-1H-pyrazole-4-carboxamide; N-(2',4',5'-trifluorobiphenyl-2-yl)-1-methyl-3-difluoromethyl-1H-pyrazole-4-carboxamide; N-(3',4',5'-tri-fluorobiphenyl-2yl)-3-chlorofluoromethyl-1-methyl-1H-pyrazole-4-carbox-amide; N-[2-(1,1,2,3,3,3-hexa-

fluoropropoxy)phenyl]-1-methyl-3-trifluoro-methyl-1H-pyrazole-4-carboxamide; N-[2-

(1,1,2,3,3,3-hexafluor-opropoxy)-phenyl]-3-difluoromethyl-1-methyl-1H-pyrazole-4-carboxamide, N-[2-(2-chloro-1,1,2-trifluoroethoxy)phenyl]-1-methyl-3-trifluoromethyl-1H-pyrazole-4-carboxamide; N-[2-(2-chlor-1,1,2-trifluoroethoxy)phenyl]-3-difluoromethyl-1-methyl-1H-pyrazole-4-carboxamide; N-[2-(1,1,2,2-tetra-fluoroethoxy)phenyl]-3-difluoromethyl-1-methyl-1H-pyrazole-4-carbox-amide; N-[2-(1,1,2,2-tetrafluoroethoxy)phenyl]-1-methyl-3-trifluoro-methyl-1H-pyrazole-4-carboxamide; N-(4'-(trifluoromethylthio)biphenyl-2-yl)-3-di-fluoromethyl-1-methyl-1H-pyrazole-4-carboxamide; N-(4'-(trifluoromethyl-thio)biphenyl-2-yl)-1-methyl-3-trifluoromethyl-1H-pyrazole-4-carboxamide; and 5-fluoro-1,3-dimethyl-1H-pyrazole-4-carboxylic acid [2-(1,2-dimethyl-propyl)-phenyl]-amide.

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Especially preferred are binary mixtures containing modification II as compound I and a fungicidal compound IIA selected from the list comprising heterocylic compounds: pyrimethanil, fenpiclonil, fludioxonil, aldimorph, dodemorph, fenpropimorph, tridemorph, 15 iprodione, procymidone, famoxadone, fenamidone, octhilinone, probenazole, diclomezine, pyroquilon, proquinazid, tricyclazole, captafol, captan, dazomet, fenoxanil, quinoxyfen, 5-chloro-7-(4-methylpiperidin-1-yl)-6-(2,4,6-trifluorophenyl)-[1,2,4]triazolo[1,5a]pyrimidine, 6-(3,4-dichloro-phenyl)-5-methyl-[1,2,4]triazolo[1,5-a]pyrimidine-7-ylamine, 6-(4-tert-butylphenyl)-5-methyl-[1,2,4]triazolo[1,5-a]pyrimidine-7-ylamine, 20 5-methyl-6-(3,5,5-trimethyl-hexyl)-[1,2,4]triazolo[1,5-a]pyrimidine-7-ylamine, 5-methyl-6-octyl-[1,2,4]triazolo[1,5-a]pyrimi-dine-7-ylamine, 6-methyl-5-octyl-[1,2,4]triazolo[1,5a]pyrimidine-7-ylamine, 6-ethyl-5-octyl-[1,2,4]triazolo[1,5-a]pyrimidine-7-ylamine, 5-ethyl-6-octyl-[1,2,4]triazolo[1,5-a]pyrimidine-7-ylamine, 5-ethyl-6-(3,5,5-trimethyl-hexyl)-[1,2,4]triazolo[1,5-a]pyrimidine-7-ylamine, 6-octyl-5-propyl-[1,2,4]tri-azolo[1,5-a]pyri-25 midine-7-ylamine, 5-methoxymethyl-6-octyl-[1,2,4]tri-azolo[1,5-a]pyrimidine-7-ylamine, 6-octyl-5-trifluoromethyl-[1,2,4]tri-azolo[1,5-a]pyrimidine-7-ylamine, and 5-trifluoromethyl-6-(3,5,5-trimethyl-hexyl)-[1,2,4]triazolo[1,5-a]pyrimidine-7-ylamine.

Especially preferred are binary mixtures containing modification II as compound I and a fungicidal compound IIA selected from the list comprising carbamates: mancozeb, maneb, metam, metiram, ferbam, propineb, thiram, zineb, ziram; diethofencarb, iprovalicarb, propamocarb, and methyl 3-(4-chlorophenyl)-3-(2-isopropoxycarbonyl-amino-3-methylbutyrylamino)propanoate.

Especially preferred are binary mixtures containing modification II as compound I and a fungicidal compound IIA selected from the list comprising: guazatine; streptomycin, validamycin A; binapacryl, dinocap, dinobuton; dithianon, isoprothiolane; fentin salts, such as fentin-acetate; edifenphos, iprobenfos, fosetyl, pyrazophos, chlorothalonil, dichlofluanid, flusulfamide, phthalide, quintozene, thiophanate-methyl, tolylfluanid;
 copper acetate, copper hydroxide, copper oxychloride, basic copper sulfate, sulfur; cyflufenamid, cymoxanil, dimethirimol, ethirimol, furalaxyl, metrafenone, and spiroxamine.

The active compounds IIA mentioned above, their preparation and their action against harmful fungi are generally known (cf.: http://www.hclrss.demon.co.uk/index.html); they are commercially available. The compounds named according to IUPAC, their preparation and their fungicidal activity are likewise known from EP-A 12 01 648; EP-A 226 917; WO 98/46608; WO 99/24413; WO 2004/049804; WO 2003/066609; WO 2003/053145; WO 2003/14103; EP-A 10 35 122; EP-A 10 28 125; EP-A 71 792; EP-A 141 317; WO 2003/009687; WO 05/087771; WO 2005/087772; WO 2005/087773; WO 2006/087325; WO 2006/087325; WO 2006/092428; WO 2006/092428; WO 2006/087343; WO 2001/42223; WO 2005/34628; WO 2005/123689; WO 2005/123690; WO 2006/120219; PCT/EP2006/064991; WO 2007/017450, and EP Application No. 06123463.9

With respect to their intended use, the following tertiary and quaternary mixtures of modification II as compound I are especially preferred:

Table1

Mixtures wherein compound IIA is trifloxystrobin, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table2

Mixtures wherein compound IIA is azoxystrobin, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

25 Table3

Mixtures wherein compound IIA is pyraclostrobin, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table4

Mixtures wherein compound IIA is boscalid, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table5

Mixtures wherein compound IIA is metalaxyl, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table6

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Mixtures wherein compound IIA is metalaxyl-M, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table7

Mixtures wherein compound IIA is cyproconazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

5 Table8

Mixtures wherein compound IIA is epoxiconazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table9

Mixtures wherein compound IIA is fenbuconazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table10

Mixtures wherein compound IIA is fluquinconazole, and the combination of compounds
15 IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table11

Mixtures wherein compound IIA is flutriafol, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table12

Mixtures wherein compound IIA is ipconazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

25 Table13

Mixtures wherein compound IIA is metconazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table14

Mixtures wherein compound IIA is propiconazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table15

Mixtures wherein compound IIA is prothioconazole, and the combination of compounds
IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table16

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Mixtures wherein compound IIA is tebuconazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table17

Mixtures wherein compound IIA is triadimenol, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

5 Table18

Mixtures wherein compound IIA is triticonazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table19

Mixtures wherein compound IIA is imazalil, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table20

Mixtures wherein compound IIA is prochloraz, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table21

Mixtures wherein compound IIA is carbendazim, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table22

Mixtures wherein compound IIA is thiabendazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

25 Table23

Mixtures wherein compound IIA is ethaboxam, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table24

Mixtures wherein compound IIA is hymexazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table25

Mixtures wherein compound IIA is pyrimethanil, and the combination of compounds
IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table26

Mixtures wherein compound IIA is fludioxonil, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table27

Mixtures wherein compound IIA is aldimorph, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

5 Table28

Mixtures wherein compound IIA is dodemorph, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table29

Mixtures wherein compound IIA is fenpropimorph, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table30

Mixtures wherein compound IIA is iprodione, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table31

Mixtures wherein compound IIA is captan, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table32

Mixtures wherein compound IIA is fenoxanil, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

25 Table33

Mixtures wherein compound IIA is probenazole, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table34

Mixtures wherein compound IIA is mancozeb, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table35

Mixtures wherein compound IIA is metiram, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table36

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Mixtures wherein compound IIA is thiram, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table37

Mixtures wherein compound IIA is ziram, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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5 Table38

Mixtures wherein compound IIA is guazatin, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table39

Mixtures wherein compound IIA is thiophanate-methyl, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table40

Mixtures wherein compound IIA is chlorothalonil, and the combination of compounds
15 IIB1 and IIB2 in each case corresponds to a row of Table Q.

Table41

Mixtures wherein compound IIA is metrafenone, and the combination of compounds IIB1 and IIB2 in each case corresponds to a row of Table Q.

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Table Q

Table Q	T			T	T	
Mixture	Compound IIB1	Compound IIB2	Mixture	Compound IIB1	Compound IIB2	
No.	,	'	No.	,		
M-1	azoxystrobin	-	M-19	azoxystrobin	thiabendazole	
M-2	azoxystrobin	boscalid	M-20	azoxystrobin	ethaboxam	
M-3	azoxystrobin	metalaxyl	M-21	azoxystrobin	hymexazole	
M-4	azoxystrobin	cyproconazole	M-22	azoxystrobin	pyrimethanil	
M-5	azoxystrobin	epoxiconazole	M-23	azoxystrobin	fludioxonil	
M-6	azoxystrobin	fenbuconazole	M-24	azoxystrobin	aldimorph	
M-7	azoxystrobin	fluquinconazole	M-25	azoxystrobin	dodemorph	
M-8	azoxystrobin	flutriafol	M-26	azoxystrobin	fenpropimorph	
M-9	azoxystrobin	ipconazole	M-27	azoxystrobin	iprodione	
M-10	azoxystrobin	metconazole	M-28	azoxystrobin	captan	
M-11	azoxystrobin	propiconazole	M-29	azoxystrobin	fenoxanil	
M-12	azoxystrobin	prothioconazole	M-30	azoxystrobin	probenazol	
M-13	azoxystrobin	tebuconazole	M-31	azoxystrobin	mancozeb	
M-14	azoxystrobin	triadimenol	M-32	azoxystrobin	metiram	
M-15	azoxystrobin	triticonazole	M-33	azoxystrobin	thiram	
M-16	azoxystrobin	imazalil	M-34	azoxystrobin	ziram	
M-17	azoxystrobin	prochloraz	M-35	azoxystrobin	guazatin	
M-18	azoxystrobin	carbendazim	M-36	azoxystrobin	thiophanate-	

Mixture No.	Compound IIB1	Compound IIB2	Mixture No.	Compound IIB1	Compound IIB2
		methyl	M-74	trifloxystrobin	thiophanate-
M-37	azoxystrobin	chlorothalonil	101-74	unioxysuobin	methyl
M-38	azoxystrobin	metrafenone	M-75	trifloxystrobin	chlorothalonil
M-39	trifloxystrobin	-	M-76	trifloxystrobin	metrafenone
M-40	trifloxystrobin	boscalid	M-77	orysastrobin	-
M-41	trifloxystrobin	metalaxyl	M-78	orysastrobin	boscalid
M-42	trifloxystrobin	cyproconazole	M-79	orysastrobin	metalaxyl
M-43	trifloxystrobin	epoxiconazole	M-80	orysastrobin	cyproconazole
M-44	trifloxystrobin	fenbuconazole	M-81	orysastrobin	epoxiconazole
M-45	trifloxystrobin	fluquinconazole	M-82	orysastrobin	fenbuconazole
M-46	trifloxystrobin	flutriafol	M-83	orysastrobin	fluquinconazole
M-47	trifloxystrobin	ipconazole	M-84	orysastrobin	flutriafol
M-48	trifloxystrobin	metconazole	M-85	orysastrobin	ipconazole
M-49	trifloxystrobin	propiconazole	M-86	orysastrobin	metconazole
M-50	trifloxystrobin	prothioconazole	M-87	orysastrobin	propiconazole
M-51	trifloxystrobin	tebuconazole	M-88	orysastrobin	prothioconazole
M-52	trifloxystrobin	triadimenol	M-89	orysastrobin	tebuconazole
M-53	trifloxystrobin	triticonazole	M-90	orysastrobin	triadimenol
M-54	trifloxystrobin	imazalil	M-91	orysastrobin	triticonazole
M-55	trifloxystrobin	prochloraz	M-92	orysastrobin	imazalil
M-56	trifloxystrobin	carbendazim	M-93	orysastrobin	prochloraz
M-57	trifloxystrobin	thiabendazole	M-94	orysastrobin	carbendazim
M-58	trifloxystrobin	ethaboxam	M-95	orysastrobin	thiabendazole
M-59	trifloxystrobin	hymexazole	M-96	orysastrobin	ethaboxam
M-60	trifloxystrobin	pyrimethanil	M-97	orysastrobin	hymexazole
M-61	trifloxystrobin	fludioxonil	M-98	orysastrobin	pyrimethanil
M-62	trifloxystrobin	aldimorph	M-99	orysastrobin	fludioxonil
M-63	trifloxystrobin	dodemorph	M-100	orysastrobin	aldimorph
M-64	trifloxystrobin	fenpropimorph	M-101	orysastrobin	dodemorph
M-65	trifloxystrobin	iprodione	M-102	orysastrobin	fenpropimorph
M-66	trifloxystrobin	captan	M-103	orysastrobin	iprodione
M-67	trifloxystrobin	fenoxanil	M-104	orysastrobin	captan
M-68	trifloxystrobin	probenazol	M-105	orysastrobin	fenoxanil
M-69	trifloxystrobin	mancozeb	M-106	orysastrobin	probenazol
M-70	trifloxystrobin	metiram	M-107	orysastrobin	mancozeb
M-71	trifloxystrobin	thiram	M-108	orysastrobin	metiram
M-72	trifloxystrobin	ziram	M-109	orysastrobin	thiram
M-73	trifloxystrobin	guazatin	M-110	orysastrobin	ziram

Mixture	0 11104	0 11100
No.	Compound IIB1	Compound IIB2
M-111	orysastrobin	guazatin
M-112	orysastrobin	thiophanate- methyl
M-113	orysastrobin	chlorothalonil
M-114	orysastrobin	metrafenone
M-115	pyraclostrobin	-
M-116	pyraclostrobin	boscalid
M-117	pyraclostrobin	metalaxyl
M-118	pyraclostrobin	cyproconazole
M-119	pyraclostrobin	epoxiconazole
M-120	pyraclostrobin	fenbuconazole
M-121	pyraclostrobin	fluquinconazole
M-122	pyraclostrobin	flutriafol
M-123	pyraclostrobin	ipconazole
M-124	pyraclostrobin	metconazole
M-125	pyraclostrobin	propiconazole
M-126	pyraclostrobin	prothioconazole
M-127	pyraclostrobin	tebuconazole
M-128	pyraclostrobin	triadimenol
M-129	pyraclostrobin	triticonazole
M-130	pyraclostrobin	imazalil
M-131	pyraclostrobin	prochloraz
M-132	pyraclostrobin	carbendazim
M-133	pyraclostrobin	thiabendazole
M-134	pyraclostrobin	ethaboxam
M-135	pyraclostrobin	hymexazole
M-136	pyraclostrobin	pyrimethanil
M-137	pyraclostrobin	fludioxonil
M-138	pyraclostrobin	aldimorph
M-139	pyraclostrobin	dodemorph
M-140	pyraclostrobin	fenpropimorph
M-141	pyraclostrobin	iprodione
M-142	pyraclostrobin	captan
M-143	pyraclostrobin	fenoxanil
M-144	pyraclostrobin	probenazol
M-145	pyraclostrobin	mancozeb
M-146	pyraclostrobin	metiram
M-147	pyraclostrobin	thiram

Mixture		
No.	Compound IIB1	Compound IIB2
M-148	pyraclostrobin	ziram
M-149	pyraclostrobin	guazatin
M-150	ny ma alaatrahin	thiophanate-
IVI-150	pyraclostrobin	methyl
M-151	pyraclostrobin	chlorothalonil
M-152	pyraclostrobin	metrafenone
M-153	boscalid	-
M-154	boscalid	metalaxyl
M-155	boscalid	cyproconazole
M-156	boscalid	epoxiconazole
M-157	boscalid	fenbuconazole
M-158	boscalid	fluquinconazole
M-159	boscalid	flutriafol
M-160	boscalid	ipconazole
M-161	boscalid	metconazole
M-162	boscalid	propiconazole
M-163	boscalid	prothioconazole
M-164	boscalid	tebuconazole
M-165	boscalid	triadimenol
M-166	boscalid	triticonazole
M-167	boscalid	imazalil
M-168	boscalid	prochloraz
M-169	boscalid	carbendazim
M-170	boscalid	thiabendazole
M-171	boscalid	ethaboxam
M-172	boscalid	hymexazole
M-173	boscalid	pyrimethanil
M-174	boscalid	fludioxonil
M-175	boscalid	aldimorph
M-176	boscalid	dodemorph
M-177	boscalid	fenpropimorph
M-178	boscalid	iprodione
M-179	boscalid	captan
M-180	boscalid	fenoxanil
M-181	boscalid	probenazol
M-182	boscalid	mancozeb
M-183	boscalid	metiram
M-184	boscalid	thiram

Mixture	Compound IIB1	Compound IIB2	Mixture	Compound IIB1	Compound IIB2
No.	•	·	No.	·	
M-185	boscalid	ziram	M-222	metalaxyl	guazatin
M-186	boscalid	guazatin thiophanate-	M-223	metalaxyl	thiophanate- methyl
M-187	boscalid	methyl	M-224	metalaxyl	chlorothalonil
M-188	boscalid	chlorothalonil	M-225	metalaxyl	metrafenone
M-189	boscalid	metrafenone	M-226	cyproconazole	-
M-190	metalaxyl	-	M-227	cyproconazole	epoxiconazole
M-191	metalaxyl	cyproconazole	M-228	cyproconazole	fenbuconazole
M-192	metalaxyl	epoxiconazole	M-229	cyproconazole	fluquinconazole
M-193	metalaxyl	fenbuconazole	M-230	cyproconazole	flutriafol
M-194	metalaxyl	fluquinconazole	M-231	cyproconazole	ipconazole
M-195	metalaxyl	flutriafol	M-232	cyproconazole	metconazole
M-196	metalaxyl	ipconazole	M-233	cyproconazole	propiconazole
M-197	metalaxyl	metconazole	M-234	cyproconazole	prothioconazol
M-198	metalaxyl	propiconazole	M-235	cyproconazole	tebuconazole
M-199	metalaxyl	prothioconazole	M-236	cyproconazole	triadimenol
M-200	metalaxyl	tebuconazole	M-237	cyproconazole	triticonazole
M-201	metalaxyl	triadimenol	M-238	cyproconazole	imazalil
M-202	metalaxyl	triticonazole	M-239	cyproconazole	prochloraz
M-203	metalaxyl	imazalil	M-240	cyproconazole	carbendazim
M-204	metalaxyl	prochloraz	M-241	cyproconazole	thiabendazole
M-205	metalaxyl	carbendazim	M-242	cyproconazole	ethaboxam
M-206	metalaxyl	thiabendazole	M-243	cyproconazole	hymexazole
M-207	metalaxyl	ethaboxam	M-244	cyproconazole	pyrimethanil
M-208	metalaxyl	hymexazole	M-245	cyproconazole	fludioxonil
M-209	metalaxyl	pyrimethanil	M-246	cyproconazole	aldimorph
M-210	metalaxyl	fludioxonil	M-247	cyproconazole	dodemorph
M-211	metalaxyl	aldimorph	M-248	cyproconazole	fenpropimorph
M-212	metalaxyl	dodemorph	M-249	cyproconazole	iprodione
M-213	metalaxyl	fenpropimorph	M-250	cyproconazole	captan
M-214	metalaxyl	iprodione	M-251	cyproconazole	fenoxanil
M-215	metalaxyl	captan	M-252	cyproconazole	probenazol
M-216	metalaxyl	fenoxanil	M-253	cyproconazole	mancozeb
M-217	metalaxyl	probenazol	M-254	cyproconazole	metiram
M-218	metalaxyl	mancozeb	M-255	cyproconazole	thiram
M-219	metalaxyl	metiram	M-256	cyproconazole	ziram
M-220	metalaxyl	thiram	M-257	cyproconazole	guazatin
M-221	metalaxyl	ziram	M-258	cyproconazole	thiophanate-

Mixture No.	Compound IIB1	Compound IIB2	Mixture No.	Compound II
		methyl	M-295	fenbuconazo
M-259	cyproconazole	chlorothalonil	M-296	fenbuconazo
M-260	cyproconazole	metrafenone	M-297	fenbuconazo
M-261	epoxiconazole	-	M-298	fenbuconazo
M-262	epoxiconazole	fenbuconazole	M-299	fenbuconazo
M-263	epoxiconazole	fluquinconazole	M-300	fenbuconazo
M-264	epoxiconazole	flutriafol	M-301	fenbuconazo
M-265	epoxiconazole	ipconazole	M-302	fenbuconazo
M-266	epoxiconazole	metconazole	M-303	fenbuconazo
M-267	epoxiconazole	propiconazole	M-304	fenbuconazo
M-268	epoxiconazole	prothioconazole	M-305	fenbuconazo
M-269	epoxiconazole	tebuconazole	M-306	fenbuconazo
M-270	epoxiconazole	triadimenol	M-307	fenbuconazo
M-271	epoxiconazole	triticonazole	M-308	fenbuconazo
M-272	epoxiconazole	imazalil	M-309	fenbuconazo
M-273	epoxiconazole	prochloraz	M-310	fenbuconazo
M-274	epoxiconazole	carbendazim	M-311	fenbuconazo
M-275	epoxiconazole	thiabendazole	M-312	fenbuconazo
M-276	epoxiconazole	ethaboxam	M-313	fenbuconazo
M-277	epoxiconazole	hymexazole	M-314	fenbuconazo
M-278	epoxiconazole	pyrimethanil	M-315	fenbuconazo
M-279	epoxiconazole	fludioxonil	M-316	fenbuconazo
M-280	epoxiconazole	aldimorph	M-317	fenbuconazo
M-281	epoxiconazole	dodemorph	M-318	fenbuconazo
M-282	epoxiconazole	fenpropimorph	M-319	fenbuconazo
M-283	epoxiconazole	iprodione	M-320	fenbuconazo
M-284	epoxiconazole	captan	M-321	fenbuconazo
M-285	epoxiconazole	fenoxanil	M-322	fenbuconazo
M-286	epoxiconazole	probenazol	M-323	fenbuconazo
M-287	epoxiconazole	mancozeb	M-324	fenbuconazo
M-288	epoxiconazole	metiram	14.005	
M-289	epoxiconazole	thiram	M-325	fenbuconazo
M-290	epoxiconazole	ziram	M-326	fenbuconazo
M-291	epoxiconazole	guazatin	M-327	fenbuconazo
N4 000		thiophanate-	M-328	fluquinconaz
M-292	epoxiconazole	methyl	M-329	fluquinconaz
M-293	epoxiconazole	chlorothalonil	M-330	fluquinconaz
M-294	epoxiconazole	metrafenone	M-331	fluquinconaz

No. Compound IIB1 Compound IIB2 M-295 fenbuconazole - M-296 fenbuconazole fluquinconazole M-297 fenbuconazole ipconazole M-298 fenbuconazole ipconazole M-299 fenbuconazole propiconazole M-300 fenbuconazole propiconazole M-301 fenbuconazole prothioconazole M-302 fenbuconazole triadimenol M-303 fenbuconazole tridimenol M-304 fenbuconazole triticonazole M-305 fenbuconazole prochloraz M-306 fenbuconazole prochloraz M-307 fenbuconazole carbendazim M-308 fenbuconazole thiabendazole M-309 fenbuconazole pyrimethanil M-310 fenbuconazole pyrimethanil M-311 fenbuconazole pyrimethanil M-312 fenbuconazole fludioxonil M-313 fenbuconazole dodemorph M-314 fenbuconazole dodemorph M-315 fenbuconazole iprodione M-316 fenbuconazole probenazol M-317 fenbuconazole captan M-318 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole thiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole chlorothalonil M-325 fenbuconazole chlorothalonil M-327 fenbuconazole chlorothalonil M-328 fluquinconazole metrafenone M-328 fluquinconazole imetrafenone			I	
M-295fenbuconazole-M-296fenbuconazolefluquinconazoleM-297fenbuconazoleflutriafolM-298fenbuconazoleipconazoleM-299fenbuconazolemetconazoleM-300fenbuconazolepropiconazoleM-301fenbuconazoleprothioconazoleM-302fenbuconazoletriadimenolM-303fenbuconazoletriticonazoleM-304fenbuconazoleimazalilM-305fenbuconazoleprochlorazM-306fenbuconazoleprochlorazM-307fenbuconazolecarbendazimM-308fenbuconazolethiabendazoleM-309fenbuconazoleethaboxamM-310fenbuconazolehymexazoleM-311fenbuconazolepyrimethanilM-312fenbuconazolefludioxonilM-313fenbuconazolefludioxonilM-314fenbuconazoledodemorphM-315fenbuconazolefenpropimorphM-316fenbuconazoleiprodioneM-317fenbuconazolerenoxanilM-318fenbuconazoleprobenazolM-319fenbuconazolemetiramM-320fenbuconazolemetiramM-321fenbuconazolethiramM-322fenbuconazolethiramM-323fenbuconazolechlorothalonilM-324fenbuconazolechlorothalonilM-325fenbuconazolechlorothalonilM-326fenbuconazole <t< td=""><td>Mixture</td><td>Compound IIB1</td><td colspan="2">Compound IIB2</td></t<>	Mixture	Compound IIB1	Compound IIB2	
M-296fenbuconazolefluquinconazoleM-297fenbuconazoleflutriafolM-298fenbuconazoleipconazoleM-299fenbuconazolemetconazoleM-300fenbuconazolepropiconazoleM-301fenbuconazoleprothioconazoleM-302fenbuconazoletebuconazoleM-303fenbuconazoletridimenolM-304fenbuconazoletriticonazoleM-305fenbuconazoleprochlorazM-306fenbuconazolecarbendazimM-307fenbuconazolethiabendazoleM-308fenbuconazoleethaboxamM-309fenbuconazolehymexazoleM-310fenbuconazolepyrimethanilM-311fenbuconazolefludioxonilM-312fenbuconazolefludioxonilM-313fenbuconazoledodemorphM-314fenbuconazoledodemorphM-315fenbuconazolefenpropimorphM-316fenbuconazolecaptanM-317fenbuconazoleprobenazolM-318fenbuconazoleprobenazolM-319fenbuconazolemetiramM-320fenbuconazolemetiramM-321fenbuconazolethiramM-322fenbuconazolethiramM-323fenbuconazolechlorothalonilM-324fenbuconazolechlorothalonilM-325fenbuconazolechlorothalonilM-326fenbuconazolechlorothalonilM-328fluquin		6 1		
M-297fenbuconazoleflutriafolM-298fenbuconazoleipconazoleM-299fenbuconazolemetconazoleM-300fenbuconazolepropiconazoleM-301fenbuconazoleprothioconazoleM-302fenbuconazoletebuconazoleM-303fenbuconazoletriadimenolM-304fenbuconazoletriticonazoleM-305fenbuconazoleimazalilM-306fenbuconazoleprochlorazM-307fenbuconazolecarbendazimM-308fenbuconazolethiabendazoleM-309fenbuconazolethiabendazoleM-310fenbuconazolepyrimethanilM-311fenbuconazolepyrimethanilM-312fenbuconazolefludioxonilM-313fenbuconazoledodemorphM-314fenbuconazoledodemorphM-315fenbuconazolefenpropimorphM-316fenbuconazolefenoxanilM-317fenbuconazoleprobenazolM-318fenbuconazolemancozebM-319fenbuconazolemetiramM-320fenbuconazolemetiramM-321fenbuconazolethiramM-323fenbuconazoleziramM-324fenbuconazolechlorothalonilM-325fenbuconazolechlorothalonilM-326fenbuconazolechlorothalonilM-327fenbuconazolemetrafenoneM-328fluquinconazole-			-	
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M-299 fenbuconazole metconazole M-300 fenbuconazole propiconazole M-301 fenbuconazole prothioconazole M-302 fenbuconazole tebuconazole M-303 fenbuconazole triadimenol M-304 fenbuconazole triticonazole M-305 fenbuconazole imazalil M-306 fenbuconazole prochloraz M-307 fenbuconazole carbendazim M-308 fenbuconazole thiabendazole M-309 fenbuconazole ethaboxam M-310 fenbuconazole pyrimethanil M-312 fenbuconazole pyrimethanil M-313 fenbuconazole fludioxonil M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole probenazol M-318 fenbuconazole probenazol M-320 fenbuconazole metiram M-320 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole metrafenone M-328 fluquinconazole metrafenone				
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M-302 fenbuconazole tebuconazole M-303 fenbuconazole triadimenol M-304 fenbuconazole triticonazole M-305 fenbuconazole imazalil M-306 fenbuconazole prochloraz M-307 fenbuconazole carbendazim M-308 fenbuconazole thiabendazole M-309 fenbuconazole ethaboxam M-310 fenbuconazole hymexazole M-311 fenbuconazole pyrimethanil M-312 fenbuconazole fludioxonil M-313 fenbuconazole aldimorph M-314 fenbuconazole fenpropimorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole fenoxanil M-318 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole thiram M-322 fenbuconazole guazatin M-323 fenbuconazole guazatin M-324 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole metrafenone	M-300	fenbuconazole	propiconazole	
M-303 fenbuconazole triadimenol M-304 fenbuconazole triticonazole M-305 fenbuconazole imazalil M-306 fenbuconazole prochloraz M-307 fenbuconazole carbendazim M-308 fenbuconazole thiabendazole M-309 fenbuconazole ethaboxam M-310 fenbuconazole hymexazole M-311 fenbuconazole pyrimethanil M-312 fenbuconazole fludioxonil M-313 fenbuconazole dodemorph M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole mancozeb M-321 fenbuconazole thiram M-322 fenbuconazole ziram M-323 fenbuconazole guazatin M-324 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole metrafenone	M-301	fenbuconazole	prothioconazole	
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M-305fenbuconazoleimazalilM-306fenbuconazoleprochlorazM-307fenbuconazolecarbendazimM-308fenbuconazolethiabendazoleM-309fenbuconazoleethaboxamM-310fenbuconazolehymexazoleM-311fenbuconazolepyrimethanilM-312fenbuconazolefludioxonilM-313fenbuconazolealdimorphM-314fenbuconazoledodemorphM-315fenbuconazolefenpropimorphM-316fenbuconazoleiprodioneM-317fenbuconazolecaptanM-318fenbuconazolefenoxanilM-319fenbuconazoleprobenazolM-320fenbuconazolemetiramM-321fenbuconazolemetiramM-322fenbuconazolethiramM-323fenbuconazoleziramM-324fenbuconazoleguazatinM-325fenbuconazolechlorothalonilM-326fenbuconazolemetrafenoneM-327fenbuconazolemetrafenoneM-328fluquinconazole-	M-303	fenbuconazole	triadimenol	
M-306fenbuconazoleprochlorazM-307fenbuconazolecarbendazimM-308fenbuconazolethiabendazoleM-309fenbuconazoleethaboxamM-310fenbuconazolehymexazoleM-311fenbuconazolepyrimethanilM-312fenbuconazolefludioxonilM-313fenbuconazolealdimorphM-314fenbuconazoledodemorphM-315fenbuconazolefenpropimorphM-316fenbuconazoleiprodioneM-317fenbuconazolecaptanM-318fenbuconazoleprobenazolM-319fenbuconazolemancozebM-320fenbuconazolemetiramM-321fenbuconazolethiramM-322fenbuconazoleguazatinM-323fenbuconazoleguazatinM-324fenbuconazolethiophanatemethylM-325fenbuconazolechlorothalonilM-326fenbuconazolechlorothalonilM-327fenbuconazolemetrafenoneM-328fluquinconazole-	M-304	fenbuconazole	triticonazole	
M-307 fenbuconazole thiabendazole M-308 fenbuconazole thiabendazole M-309 fenbuconazole ethaboxam M-310 fenbuconazole hymexazole M-311 fenbuconazole pyrimethanil M-312 fenbuconazole fludioxonil M-313 fenbuconazole aldimorph M-314 fenbuconazole dodemorph M-315 fenbuconazole iprodione M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole mancozeb M-320 fenbuconazole metiram M-321 fenbuconazole thiram M-322 fenbuconazole guazatin M-323 fenbuconazole guazatin M-324 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-305	fenbuconazole	imazalil	
M-308 fenbuconazole thiabendazole M-309 fenbuconazole ethaboxam M-310 fenbuconazole hymexazole M-311 fenbuconazole pyrimethanil M-312 fenbuconazole fludioxonil M-313 fenbuconazole aldimorph M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole thiram M-322 fenbuconazole thiram M-323 fenbuconazole guazatin M-324 fenbuconazole guazatin M-325 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-306	fenbuconazole	prochloraz	
M-309 fenbuconazole ethaboxam M-310 fenbuconazole hymexazole M-311 fenbuconazole pyrimethanil M-312 fenbuconazole fludioxonil M-313 fenbuconazole aldimorph M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole guazatin M-324 fenbuconazole guazatin M-325 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-307	fenbuconazole	carbendazim	
M-310 fenbuconazole hymexazole M-311 fenbuconazole pyrimethanil M-312 fenbuconazole fludioxonil M-313 fenbuconazole aldimorph M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-308	fenbuconazole	thiabendazole	
M-311 fenbuconazole pyrimethanil M-312 fenbuconazole fludioxonil M-313 fenbuconazole aldimorph M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole guazatin M-324 fenbuconazole guazatin M-325 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-309	fenbuconazole	ethaboxam	
M-312 fenbuconazole fludioxonil M-313 fenbuconazole aldimorph M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole M-328 fluquinconazole -	M-310	fenbuconazole	hymexazole	
M-313 fenbuconazole aldimorph M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-311	fenbuconazole	pyrimethanil	
M-314 fenbuconazole dodemorph M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole guazatin M-324 fenbuconazole guazatin M-325 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-312	fenbuconazole	fludioxonil	
M-315 fenbuconazole fenpropimorph M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-313	fenbuconazole	aldimorph	
M-316 fenbuconazole iprodione M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-314	fenbuconazole	dodemorph	
M-317 fenbuconazole captan M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-315	fenbuconazole	fenpropimorph	
M-318 fenbuconazole fenoxanil M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-316	fenbuconazole	iprodione	
M-319 fenbuconazole probenazol M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-317	fenbuconazole	captan	
M-320 fenbuconazole mancozeb M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-318	fenbuconazole	fenoxanil	
M-321 fenbuconazole metiram M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-319	fenbuconazole	probenazol	
M-322 fenbuconazole thiram M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanate- methyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-320	fenbuconazole	mancozeb	
M-323 fenbuconazole ziram M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanatemethyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-321	fenbuconazole	metiram	
M-324 fenbuconazole guazatin M-325 fenbuconazole thiophanate- methyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-322	fenbuconazole	thiram	
M-325 fenbuconazole thiophanate- methyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-323	fenbuconazole	ziram	
M-325 fenbuconazole methyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-324	fenbuconazole	guazatin	
M-325 fenbuconazole methyl M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -			thiophanate-	
M-326 fenbuconazole chlorothalonil M-327 fenbuconazole metrafenone M-328 fluquinconazole -	M-325	fenbuconazole		
M-328 fluquinconazole -	M-326	fenbuconazole	· · · · · · · · · · · · · · · · · · ·	
M-328 fluquinconazole -	M-327	fenbuconazole	metrafenone	
<u>'</u>			-	
		fluquinconazole	flutriafol	
M-330 fluquinconazole ipconazole		-		
M-331 fluquinconazole metconazole		· ·	· ·	

Mixture No.	Compound IIB1	Compound IIB2	Mixture No.	(
M-332	fluquinconazole	propiconazole	M-369	1
M-333	fluquinconazole	prothioconazole	M-370	1
M-334	fluquinconazole	tebuconazole	M-371	1
M-335	fluquinconazole	triadimenol	M-372	1
M-336	fluquinconazole	triticonazole	M-373	1
M-337	fluquinconazole	imazalil	M-374	1
M-338	fluquinconazole	prochloraz	M-375	1
M-339	fluquinconazole	carbendazim	M-376	
M-340	fluquinconazole	thiabendazole	M-377	
M-341	fluquinconazole	ethaboxam	M-378	
M-342	fluquinconazole	hymexazole	M-379	
M-343	fluquinconazole	pyrimethanil	M-380	
M-344	fluquinconazole	fludioxonil	M-381	
M-345	fluquinconazole	aldimorph	M-382	
M-346	fluquinconazole	dodemorph	M-383	
M-347	fluquinconazole	fenpropimorph	M-384	
M-348	fluquinconazole	iprodione	M-385	
M-349	fluquinconazole	captan	M-386	
M-350	fluquinconazole	fenoxanil	M-387	1
M-351	fluquinconazole	probenazol	M-388	
M-352	fluquinconazole	mancozeb	101-300	
M-353	fluquinconazole	metiram	M-389	Ŀ
M-354	fluquinconazole	thiram	M-390	ŀ
M-355	fluquinconazole	ziram	M-391	
M-356	fluquinconazole	guazatin	M-392	
M-357	fluquinconazole	thiophanate-	M-393	
101-007	naquinconazoie	methyl	M-394	
M-358	fluquinconazole	chlorothalonil	M-395	
M-359	fluquinconazole	metrafenone	M-396	
M-360	flutriafol	-	M-397	
M-361	flutriafol	ipconazole	M-398	
M-362	flutriafol	metconazole	M-399	
M-363	flutriafol	propiconazole	M-400	
M-364	flutriafol	prothioconazole	M-401	
M-365	flutriafol	tebuconazole	M-402	
M-366	flutriafol	triadimenol	M-403	
M-367	flutriafol	triticonazole	M-404	
M-368	flutriafol	imazalil	M-405	i

Mixturo		Ι
Mixture No.	Compound IIB1	Compound IIB2
M-369	flutriafol	prochloraz
M-370	flutriafol	carbendazim
M-371	flutriafol	thiabendazole
M-372	flutriafol	ethaboxam
M-373	flutriafol	hymexazole
M-374	flutriafol	pyrimethanil
M-375	flutriafol	fludioxonil
M-376	flutriafol	aldimorph
M-377	flutriafol	dodemorph
M-378	flutriafol	fenpropimorph
M-379	flutriafol	iprodione
M-380	flutriafol	captan
M-381	flutriafol	fenoxanil
M-382	flutriafol	probenazol
M-383	flutriafol	mancozeb
M-384	flutriafol	metiram
M-385	flutriafol	thiram
M-386	flutriafol	ziram
M-387	flutriafol	guazatin
101 007	natiaioi	thiophanate-
M-388	flutriafol	methyl
M-389	flutriafol	chlorothalonil
M-390	flutriafol	metrafenone
M-391	ipconazole	-
M-392	ipconazole	metconazole
M-393	ipconazole	propiconazole
M-394	ipconazole	prothioconazole
M-395	ipconazole	tebuconazole
M-396	ipconazole	triadimenol
M-397	ipconazole	triticonazole
M-398	ipconazole	imazalil
M-399	ipconazole	prochloraz
M-400	ipconazole	carbendazim
M-401	ipconazole	thiabendazole
M-402	ipconazole	ethaboxam
M-403	ipconazole	hymexazole
M-404	ipconazole	pyrimethanil
M-405	ipconazole	fludioxonil

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Mixture No.	Compound IIB1	Compound IIB2
M-406	ipconazole	aldimorph
M-407	ipconazole	dodemorph
M-408	ipconazole	fenpropimorph
M-409	ipconazole	iprodione
M-410	ipconazole	captan
M-411	ipconazole	fenoxanil
M-412	ipconazole	probenazol
M-413	ipconazole	mancozeb
M-414	ipconazole	metiram
M-415	ipconazole	thiram
M-416	ipconazole	ziram
M-417	ipconazole	guazatin
M-418	inconazala	thiophanate-
101-410	ipconazole	methyl
M-419	ipconazole	chlorothalonil
M-420	ipconazole	metrafenone
M-421	metconazole	-
M-422	metconazole	propiconazole
M-423	metconazole	prothioconazole
M-424	metconazole	tebuconazole
M-425	metconazole	triadimenol
M-426	metconazole	triticonazole
M-427	metconazole	imazalil
M-428	metconazole	prochloraz
M-429	metconazole	carbendazim
M-430	metconazole	thiabendazole
M-431	metconazole	ethaboxam
M-432	metconazole	hymexazole
M-433	metconazole	pyrimethanil
M-434	metconazole	fludioxonil
M-435	metconazole	aldimorph
M-436	metconazole	dodemorph
M-437	metconazole	fenpropimorph
M-438	metconazole	iprodione
M-439	metconazole	captan
M-440	metconazole	fenoxanil
M-441	metconazole	probenazol
M-442	metconazole	mancozeb

Mixture No.	Compound IIB1	Compound IIB2
M-443	metconazole	metiram
M-444	metconazole	thiram
M-445	metconazole	ziram
M-446	metconazole	guazatin
	motoonazoro	thiophanate-
M-447	metconazole	methyl
M-448	metconazole	chlorothalonil
M-449	metconazole	metrafenone
M-450	propiconazole	-
M-451	propiconazole	prothioconazole
M-452	propiconazole	tebuconazole
M-453	propiconazole	triadimenol
M-454	propiconazole	triticonazole
M-455	propiconazole	imazalil
M-456	propiconazole	prochloraz
M-457	propiconazole	carbendazim
M-458	propiconazole	thiabendazole
M-459	propiconazole	ethaboxam
M-460	propiconazole	hymexazole
M-461	propiconazole	pyrimethanil
M-462	propiconazole	fludioxonil
M-463	propiconazole	aldimorph
M-464	propiconazole	dodemorph
M-465	propiconazole	fenpropimorph
M-466	propiconazole	iprodione
M-467	propiconazole	captan
M-468	propiconazole	fenoxanil
M-469	propiconazole	probenazol
M-470	propiconazole	mancozeb
M-471	propiconazole	metiram
M-472	propiconazole	thiram
M-473	propiconazole	ziram
M-474	propiconazole	guazatin
M-475	propiconazole	thiophanate-
		methyl
M-476	propiconazole	chlorothalonil
M-477	propiconazole	metrafenone
M-478	prothioconazole	-

Mixture No.	Compound IIB1	Compound IIB2	Mixtu No.
M-479	prothioconazole	tebuconazole	M-51
M-480	prothioconazole	triadimenol	M-51
M-481	prothioconazole	triticonazole	M-51
M-482	prothioconazole	imazalil	M-51
M-483	prothioconazole	prochloraz	M-52
M-484	prothioconazole	carbendazim	M-52
M-485	prothioconazole	thiabendazole	M-52
M-486	prothioconazole	ethaboxam	M-52
M-487	prothioconazole	hymexazole	M-52
M-488	prothioconazole	pyrimethanil	M-52
M-489	prothioconazole	fludioxonil	M-52
M-490	prothioconazole	aldimorph	M-52
M-491	prothioconazole	dodemorph	N4.50
M-492	prothioconazole	fenpropimorph	M-52
M-493	prothioconazole	iprodione	M-52
M-494	prothioconazole	captan	M-53
M-495	prothioconazole	fenoxanil	M-53
M-496	prothioconazole	probenazol	M-53
M-497	prothioconazole	mancozeb	M-53
M-498	prothioconazole	metiram	M-53
M-499	prothioconazole	thiram	M-53
M-500	prothioconazole	ziram	M-53
M-501	prothioconazole	guazatin	M-53
M-502	prothioconazole	thiophanate- methyl	M-53
M-503	prothioconazole	chlorothalonil	M-54
M-504	prothioconazole	metrafenone	M-54
M-505	tebuconazole	-	M-54
M-506	tebuconazole	triadimenol	M-54
M-507	tebuconazole	triticonazole	M-54
M-508	tebuconazole	imazalil	M-54
M-509	tebuconazole	prochloraz	M-54
M-510	tebuconazole	carbendazim	M-54
M-511	tebuconazole	thiabendazole	M-54
M-512	tebuconazole	ethaboxam	M-54
M-513	tebuconazole	hymexazole	M-55
M-514	tebuconazole	pyrimethanil	M-55
M-515	tebuconazole	fludioxonil	M-55

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Mixture No.	Compound IIB1	Compound IIB2
M-516	tebuconazole	aldimorph
M-517	tebuconazole	dodemorph
M-518	tebuconazole	fenpropimorph
M-519	tebuconazole	iprodione
M-520	tebuconazole	captan
M-521	tebuconazole	fenoxanil
M-522	tebuconazole	probenazol
M-523	tebuconazole	mancozeb
M-524	tebuconazole	metiram
M-525	tebuconazole	thiram
M-526	tebuconazole	
M-527	tebuconazole	ziram
IVI-527	tebuconazoie	guazatin
M-528	tebuconazole	thiophanate-
M-529	tebuconazole	methyl chlorothalonil
M-530	tebuconazole	metrafenone
M-531	triadimenol	-
M-532	triadimenol	triticonazole
M-533	triadimenol	imazalil
M-534	triadimenol	prochloraz
M-535	triadimenol	carbendazim
M-536	triadimenol	thiabendazole
M-537	triadimenol	ethaboxam
M-538		
M-539	triadimenol triadimenol	hymexazole
		pyrimethanil
M-540	triadimenol	fludioxonil
M-541	triadimenol	aldimorph
M-542	triadimenol	dodemorph
M-543	triadimenol	fenpropimorph
M-544	triadimenol	iprodione
M-545	triadimenol	captan
M-546	triadimenol	fenoxanil
M-547	triadimenol	probenazol
M-548	triadimenol	mancozeb
M-549	triadimenol	metiram
M-550	triadimenol	thiram
M-551	triadimenol	ziram
M-552	triadimenol	guazatin

Mixture	0 11154	C LUDO
No.	Compound IIB1	Compound IIB2
M-553	triadimenol	thiophanate-
101-333	triadimento	methyl
M-554	triadimenol	chlorothalonil
M-555	triadimenol	metrafenone
M-556	triticonazole	-
M-557	triticonazole	imazalil
M-558	triticonazole	prochloraz
M-559	triticonazole	carbendazim
M-560	triticonazole	thiabendazole
M-561	triticonazole	ethaboxam
M-562	triticonazole	hymexazole
M-563	triticonazole	pyrimethanil
M-564	triticonazole	fludioxonil
M-565	triticonazole	aldimorph
M-566	triticonazole	dodemorph
M-567	triticonazole	fenpropimorph
M-568	triticonazole	iprodione
M-569	triticonazole	captan
M-570	triticonazole	fenoxanil
M-571	triticonazole	probenazol
M-572	triticonazole	mancozeb
M-573	triticonazole	metiram
M-574	triticonazole	thiram
M-575	triticonazole	ziram
M-576	triticonazole	guazatin
M-577	4	thiophanate-
101-577	triticonazole	methyl
M-578	triticonazole	chlorothalonil
M-579	triticonazole	metrafenone
M-580	imazalil	_
M-581	imazalil	prochloraz
M-582	imazalil	carbendazim
M-583	imazalil	thiabendazole
M-584	imazalil	ethaboxam
M-585	imazalil	hymexazole
M-586	imazalil	pyrimethanil
M-587	imazalil	fludioxonil
M-588	imazalil	aldimorph

Mixture No. Compound IIB1 Compound IIB	2
No.	
M-589 imazalil dodemorph	
M-590 imazalil fenpropimorph	
M-591 imazalil iprodione	
M-592 imazalil captan	
M-593 imazalil fenoxanil	
M-594 imazalil probenazol	
M-595 imazalil mancozeb	
M-596 imazalil metiram	
M-597 imazalil thiram	
M-598 imazalil ziram	
M-599 imazalil guazatin	
M COO imposibility thiophanate-	
M-600 imazalil methyl	
M-601 imazalil chlorothalonil	
M-602 imazalil metrafenone	
M-603 prochloraz -	
M-604 prochloraz carbendazim	
M-605 prochloraz thiabendazole	
M-606 prochloraz ethaboxam	
M-607 prochloraz hymexazole	
M-608 prochloraz pyrimethanil	
M-609 prochloraz fludioxonil	
M-610 prochloraz aldimorph	
M-611 prochloraz dodemorph	
M-612 prochloraz fenpropimorph	
M-613 prochloraz iprodione	
M-614 prochloraz captan	
M-615 prochloraz fenoxanil	
M-616 prochloraz probenazol	
M-617 prochloraz mancozeb	
M-618 prochloraz metiram	
M-619 prochloraz thiram	
M-620 prochloraz ziram	
M-621 prochloraz guazatin	
thiophanate-	
M-622 prochloraz methyl	
M-623 prochloraz chlorothalonil	
M-624 prochloraz metrafenone	

Mixture	0 11154	C LUDO
No.	Compound IIB1	Compound IIB2
M-625	carbendazim	-
M-626	carbendazim	thiabendazole
M-627	carbendazim	ethaboxam
M-628	carbendazim	hymexazole
M-629	carbendazim	pyrimethanil
M-630	carbendazim	fludioxonil
M-631	carbendazim	aldimorph
M-632	carbendazim	dodemorph
M-633	carbendazim	fenpropimorph
M-634	carbendazim	iprodione
M-635	carbendazim	captan
M-636	carbendazim	fenoxanil
M-637	carbendazim	probenazol
M-638	carbendazim	mancozeb
M-639	carbendazim	metiram
M-640	carbendazim	thiram
M-641	carbendazim	ziram
M-642	carbendazim	guazatin
M-643	carbendazim	thiophanate-
1010	oarboriaaziiri	methyl
M-644	carbendazim	chlorothalonil
M-645	carbendazim	metrafenone
M-646	thiabendazole	-
M-647	thiabendazole	ethaboxam
M-648	thiabendazole	hymexazole
M-649	thiabendazole	pyrimethanil
M-650	thiabendazole	fludioxonil
M-651	thiabendazole	aldimorph
M-652	thiabendazole	dodemorph
M-653	thiabendazole	fenpropimorph
M-654	thiabendazole	iprodione
M-655	thiabendazole	captan
M-656	thiabendazole	fenoxanil
M-657	thiabendazole	probenazol
M-658	thiabendazole	mancozeb
M-659	thiabendazole	metiram
M-660	thiabendazole	thiram
M-661	thiabendazole	ziram

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Mixture	Compound IIB1	Compound IIB2
No. M-662	thichandarala	au anatia
IVI-002	thiabendazole	guazatin
M-663	thiabendazole	thiophanate-
NA 004	45:-541-	methyl
M-664	thiabendazole	chlorothalonil
M-665	thiabendazole	metrafenone
M-666	ethaboxam	-
M-667	ethaboxam	hymexazole
M-668	ethaboxam	pyrimethanil
M-669	ethaboxam	fludioxonil
M-670	ethaboxam	aldimorph
M-671	ethaboxam	dodemorph
M-672	ethaboxam	fenpropimorph
M-673	ethaboxam	iprodione
M-674	ethaboxam	captan
M-675	ethaboxam	fenoxanil
M-676	ethaboxam	probenazol
M-677	ethaboxam	mancozeb
M-678	ethaboxam	metiram
M-679	ethaboxam	thiram
M-680	ethaboxam	ziram
M-681	ethaboxam	guazatin
NA COO	- 41 1	thiophanate-
M-682	ethaboxam	methyl
M-683	ethaboxam	chlorothalonil
M-684	ethaboxam	metrafenone
M-685	hymexazole	-
M-686	hymexazole	pyrimethanil
M-687	hymexazole	fludioxonil
M-688	hymexazole	aldimorph
M-689	hymexazole	dodemorph
M-690	hymexazole	fenpropimorph
M-691	hymexazole	iprodione
M-692	hymexazole	captan
M-693	hymexazole	fenoxanil
M-694	hymexazole	probenazol
M-695	hymexazole	mancozeb
M-696	hymexazole	metiram
M-697	hymexazole	thiram
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Mixture		
No.	Compound IIB1	Compound IIB2
M-698	hymexazole	ziram
M-699	hymexazole	guazatin
M-700	hymexazole	thiophanate-
101-7 00	Trymexazore	methyl
M-701	hymexazole	chlorothalonil
M-702	hymexazole	metrafenone
M-703	pyrimethanil	-
M-704	pyrimethanil	fludioxonil
M-705	pyrimethanil	aldimorph
M-706	pyrimethanil	dodemorph
M-707	pyrimethanil	fenpropimorph
M-708	pyrimethanil	iprodione
M-709	pyrimethanil	captan
M-710	pyrimethanil	fenoxanil
M-711	pyrimethanil	probenazol
M-712	pyrimethanil	mancozeb
M-713	pyrimethanil	metiram
M-714	pyrimethanil	thiram
M-715	pyrimethanil	ziram
M-716	pyrimethanil	guazatin
M-717	no unimo a the a mil	thiophanate-
IVI-7 17	pyrimethanil	methyl
M-718	pyrimethanil	chlorothalonil
M-719	pyrimethanil	metrafenone
M-720	fludioxonil	-
M-721	fludioxonil	aldimorph
M-722	fludioxonil	dodemorph
M-723	fludioxonil	fenpropimorph
M-724	fludioxonil	iprodione
M-725	fludioxonil	captan
M-726	fludioxonil	fenoxanil
M-727	fludioxonil	probenazol
M-728	fludioxonil	mancozeb
M-729	fludioxonil	metiram
M-730	fludioxonil	thiram
M-731	fludioxonil	ziram
M-732	fludioxonil	guazatin
M-733	fludioxonil	thiophanate-

Mixture No.	Compound IIB1	Compound IIB2
		methyl
M-734	fludioxonil	chlorothalonil
M-735	fludioxonil	metrafenone
M-736	aldimorph	-
M-737	aldimorph	dodemorph
M-738	aldimorph	fenpropimorph
M-739	aldimorph	iprodione
M-740	aldimorph	captan
M-741	aldimorph	fenoxanil
M-742	aldimorph	probenazol
M-743	aldimorph	mancozeb
M-744	aldimorph	metiram
M-745	aldimorph	thiram
M-746	aldimorph	ziram
M-747	aldimorph	guazatin
M-748	aldimorph	thiophanate- methyl
M-749	aldimorph	chlorothalonil
M-750	aldimorph	metrafenone
M-751	dodemorph	-
M-752	dodemorph	fenpropimorph
M-753	dodemorph	iprodione
M-754	dodemorph	captan
M-755	dodemorph	fenoxanil
M-756	dodemorph	probenazol
M-757	dodemorph	mancozeb
M-758	dodemorph	metiram
M-759	dodemorph	thiram
M-760	dodemorph	ziram
M-761	dodemorph	guazatin
NA 700	d - d l-	thiophanate-
M-762	dodemorph	methyl
M-763	dodemorph	chlorothalonil
M-764	dodemorph	metrafenone
M-765	fenpropimorph	-
M-766	fenpropimorph	iprodione
M-767	fenpropimorph	captan
M-768	fenpropimorph	fenoxanil

Mixture		C LUDO
No.	Compound IIB1	Compound IIB2
M-769	fenpropimorph	probenazol
M-770	fenpropimorph	mancozeb
M-771	fenpropimorph	metiram
M-772	fenpropimorph	thiram
M-773	fenpropimorph	ziram
M-774	fenpropimorph	guazatin
M-775	fenpropimorph	thiophanate- methyl
M-776	fenpropimorph	chlorothalonil
M-777	fenpropimorph	metrafenone
M-778	iprodione	-
M-779	iprodione	captan
M-780	iprodione	fenoxanil
M-781	iprodione	probenazol
M-782	iprodione	mancozeb
M-783	iprodione	metiram
M-784	iprodione	thiram
M-785	iprodione	ziram
M-786	iprodione	guazatin
M-787	iprodione	thiophanate-
101-707	prodione	methyl
M-788	iprodione	chlorothalonil
M-789	iprodione	metrafenone
M-790	captan	-
M-791	captan	fenoxanil
M-792	captan	probenazol
M-793	captan	mancozeb
M-794	captan	metiram
M-795	captan	thiram
M-796	captan	ziram
M-797	captan	guazatin
M-798	cantan	thiophanate-
IVI-7 90	captan	methyl
M-799	captan	chlorothalonil
M-800	captan	metrafenone
M-801	fenoxanil	-
M-802	fenoxanil	probenazol
M-803	fenoxanil	mancozeb

Г	Γ	T
Mixture	Compound IIB1	Compound IIB2
No.	•	·
M-804	fenoxanil	metiram
M-805	fenoxanil	thiram
M-806	fenoxanil	ziram
M-807	fenoxanil	guazatin
M-808	fenoxanil	thiophanate- methyl
M-809	fenoxanil	chlorothalonil
M-810	fenoxanil	metrafenone
M-811	probenazol	-
M-812	probenazol	mancozeb
M-813	probenazol	metiram
M-814	probenazol	thiram
M-815	probenazol	ziram
M-816	probenazol	guazatin
1404		thiophanate-
M-817	probenazol	methyl
M-818	probenazol	chlorothalonil
M-819	probenazol	metrafenone
M-820	mancozeb	-
M-821	mancozeb	metiram
M-822	mancozeb	thiram
M-823	mancozeb	ziram
M-824	mancozeb	guazatin
NA 005		thiophanate-
M-825	mancozeb	methyl
M-826	mancozeb	chlorothalonil
M-827	mancozeb	metrafenone
M-828	metiram	-
M-829	metiram	thiram
M-830	metiram	ziram
M-831	metiram	guazatin
NA 000		thiophanate-
M-832	metiram	methyl
M-833	metiram	chlorothalonil
M-834	metiram	metrafenone
M-835	thiram	-
M-836	thiram	ziram
M-837	thiram	guazatin
L	L	

Mixture No.	Compound IIB1	Compound IIB2
M-838	thiram	thiophanate- methyl
M-839	thiram	chlorothalonil
M-840	thiram	metrafenone
M-841	ziram	-
M-842	ziram	guazatin
M-843	ziram	thiophanate- methyl
M-844	ziram	chlorothalonil
M-845	ziram	metrafenone
M-846	guazatin	-
M-847	guazatin	thiophanate-

Mixture	Compound IIB1	Compound IIB2
No.	Compound his i	Compound ribz
		methyl
M-848	guazatin	chlorothalonil
M-849	guazatin	metrafenone
M-850	thiophanate-	-
IVI-050	methyl	
M-851	thiophanate-	oblorotholonil
IVI-05 I	methyl	chlorothalonil
M-852	thiophanate-	metrafenone
IVI-052	methyl	
M-853	chlorothalonil	-
M-854	chlorothalonil	metrafenone
M-855	metrafenone	-

The crystalline modification II and the one or more compound(s) of groups A.1 – A.15 are usually applied in a weight ratio of from 500:1 to 1:100, preferably from 20:1 to 1:50, in particular from 5:1 to 1:20.

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The afore-mentioned applies also to the ratios of combinations of modification II with fungicidal compounds IIA. Compounds IIB are usually combined with modification II in ratios from 100:1 to 1:100.

- Depending on the desired effect, the application rates of the mixtures according to the invention are from 5 g/ha to 2000 g/ha, preferably from 50 to 1500 g/ha, in particular from 50 to 750 g/ha.
- The crystalline modification II, the mixtures and the compositions according to the invention can be applied to any and all developmental stages, such as egg, larva, pupa, and adult. The pests may be controlled by contacting the target pest, its food supply, habitat, breeding ground or its locus with a pesticidally effective amount of the crystalline modification II, the mixtures or the compositions according to the invention.
- 20 "Locus" means a plant, seed, soil, area, material or environment in which a pest is growing or may grow.
 - In general, "pesticidally effective amount" means the amount of the crystalline modification II, the mixtures and the compositions according to the invention needed to achieve an observable effect on growth, including the effects of necrosis, death, retardation, prevention, and removal, destruction, or otherwise diminishing the occurrence and activity of the target organism. The pesticidally effective amount can

vary for the various mixtures / compositions used in the invention. A pesticidally effective amount of the mixtures / compositions will also vary according to the prevailing conditions such as desired pesticidal effect and duration, weather, target species, locus, mode of application, and the like.

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The crystalline modification II, the mixtures and the compositions according to the invention can also be employed for protecting plants from attack or infestation by insects, acarids or nematodes comprising contacting a plant, or soil or water in which the plant is growing.

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In the context of the present invention, the term plant refers to an entire plant, a part of the plant or the propagation material of the plant, that is, the seed or the seedling.

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Plants which can be treated with the crystalline modification II, the mixtures and the compositions according to the invention include all genetically modified plants or transgenic plants, e.g. crops which tolerate the action of herbicides or fungicides or insecticides owing to breeding, including genetic engineering methods, or plants which have modified characteristics in comparison with existing plants, which can be generated for example by traditional breeding methods and/or the generation of mutants, or by recombinant procedures.

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Some of the inventive mixtures and compositions have systemic action and can therefore be used for the protection of the plant shoot against foliar pests as well as for the treatment of the seed and roots against soil pests. The term seed treatment comprises all suitable seed treatment techniques known in the art, such as, but not limited to, seed dressing, seed coating, seed dusting, seed soaking, seed film coating, seed multilayer coating, seed encrusting, seed dripping, and seed pelleting.

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The present invention also comprises seeds coated with or containing the crystalline modification II or the mixtures or the compositions according to the invention.

The term seed embraces seeds and plant propagules of all kinds including but not limited to true seeds, seed pieces, suckers, corms, bulbs, fruit, tubers, grains, cuttings, cut shoots and the like and means in a preferred embodiment true seeds.

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Suitable seed is seed of cereals, root crops, oil crops, vegetables, spices, ornamentals, for example seed of durum and other wheat, barley, oats, rye, maize (fodder maize and sugar maize / sweet and field corn), soybeans, oil crops, crucifers, cotton, sunflowers, bananas, rice, oilseed rape, turnip rape, sugarbeet, fodder beet, eggplants, potatoes, grass, lawn, turf, fodder grass, tomatoes, leeks, pumpkin/squash, cabbage, iceberg lettuce, pepper, cucumbers, melons, Brassica species, melons, beans, peas, garlic,

onions, carrots, tuberous plants such as potatoes, sugar cane, tobacco, grapes, petunias, geranium/pelargoniums, pansies and impatiens.

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In addition, the crystalline modification II, the mixtures and the compositions according to the invention may also be used for the treatment seeds from plants, which tolerate the action of herbicides or fungicides or insecticides or nematicides owing to breeding, mutation and/or genetic engineering methods.

For example, the crystalline modification II, the mixtures and the compositions 10 according to the invention can be employed in transgenic crops which are resistant to herbicides from the group consisting of the sulfonylureas (EP-A-0257993, U.S. Pat. No. 5,013,659), imidazolinones (see for example US 6222100, WO0182685, WO0026390, WO9741218, WO9802526, WO9802527, WO 04/106529, WO 05/20673, WO 03/14357, WO 03/13225, WO 03/14356, WO 04/16073), glufosinate-type (see for 15 example EP-A-0242236, EP-A-242246) or glyphosate-type (see for example WO 92/00377) or in plants resistant towards herbicides selected from the group of cyclohexadienone/aryloxyphenoxypropionic acid herbicides (US 5,162,602, US 5,290,696, US 5,498,544, US 5,428,001, US 6,069,298, US 6,268,550, US 6,146,867, US 6,222,099, US 6,414,222) or in transgenic crop plants, for example 20 cotton, with the capability of producing Bacillus thuringiensis toxins (Bt toxins) which make the plants resistant to certain pests (EP-A-0142924, EP-A-0193259).

Furthermore, the crystalline modification II, the mixtures and the compositions according to the invention can be used also for the treatment of seeds from plants, which have modified characteristics in comparison with existing plants consist, which can be generated, for example by traditional breeding methods and/or the generation of mutants, or by recombinant procedures). For example, a number of cases have been described of recombinant modifications of crop plants for the purpose of modifying the starch synthesized in the plants (e.g. WO 92/11376, WO 92/14827, WO 91/19806) or of transgenic crop plants having a modified fatty acid composition (WO 91/13972).

The seed treatment application of the crystalline modification II, the mixtures and the compositions according to the invention is carried out by spraying or dusting the seeds before sowing of the plants and before emergence of the plants.

In the treatment of seeds the corresponding formulations are applied by treating the seeds with an effective amount of the crystalline modification II, the mixtures or the compositions according to the invention. Herein, the application rates of the crystalline modification II are generally from 0.1 g to 10 kg per 100 kg of seed, preferably from 1 g

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to 5 kg per 100 kg of seed, in particular from 1 g to 2.5 kg per 100 kg of seed. For specific crops such as lettuce and onions the rates can be higher.

The mixtures and the compositions according to the invention are effective through both contact (via soil, glass, wall, bed net, carpet, plant parts or animal parts), and ingestion (bait, or plant part) and through trophallaxis and transfer.

Preferred application methods are into water bodies, via soil, cracks and crevices, pastures, manure piles, sewers, into water, on floor, wall, or by perimeter spray application and bait.

According to another preferred embodiment of the invention, for use against non-crop pests such as ants, termites, wasps, flies, mosquitoes, crickets, locusts, or cockroaches the mixtures and the compositions according to the invention are prepared into a bait preparation.

The bait can be a liquid, a solid or a semisolid preparation (e.g. a gel). The bait employed in the mixtures/compositions is a product which is sufficiently attractive to incite insects such as ants, termites, wasps, flies, mosquitoes, crickets etc. or cockroaches to eat it. This attractant may be chosen from feeding stimulants or para and / or sex pheromones readily known in the art.

Methods to control infectious diseases transmitted by insects (e.g. malaria, dengue and yellow fever, lymphatic filariasis, and leishmaniasis) with the inventive mixtures and their respective compositions also comprise treating surfaces of huts and houses, air spraying and impregnation of curtains, tents, clothing items, bed nets, tsetse-fly trap or the like. Insecticidal compositions for application to fibers, fabric, knitgoods, nonwovens, netting material or foils and tarpaulins preferably comprise a composition including the inventive mixtures, optionally a repellent and at least one binder.

The crystalline modification II, the mixtures and the compositions according to the invention can be used for protecting wooden materials such as trees, board fences, sleepers, etc. and buildings such as houses, outhouses, factories, but also construction materials, furniture, leathers, fibers, vinyl articles, electric wires and cables etc. from ants and/or termites, and for controlling ants and termites from doing harm to crops or human being (e.g. when the pests invade into houses and public facilities).

In the case of soil treatment or of application to the pests dwelling place or nest, the quantity of active ingredient ranges from 0.0001 to 500 g per 100 m², preferably from 0.001 to 20 g per 100 m².

Customary application rates in the protection of materials are, for example, from 0.01 g to 1000 g of active compound per m² treated material, desirably from 0.1 g to 50 g per m².

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- Insecticidal compositions for use in the impregnation of materials typically contain from 0.001 to 95 weight %, preferably from 0.1 to 45 weight %, and more preferably from 1 to 25 weight % of at least one repellent and / or insecticide.
- For use in bait compositions, the typical content of active ingredient(s) is from 0.0001 weight % to 15 weight %, desirably from 0.001 weight % to 5% weight % of active compound. The composition used may also comprise other additives such as a solvent of the active material, a flavoring agent, a preserving agent, a dye or a bitter agent. Its attractiveness may also be enhanced by a special color, shape or texture.
- For use in spray compositions, the content of the active ingredient(s) is from 0.001 to 80 weights %, preferably from 0.01 to 50 weight % and most preferably from 0.01 to 15 weight %.
- For use in treating crop plants, the rate of application of the active ingredient(s) may be in the range of 0.1 g to 4000 g per hectare, desirably from 25 g to 600 g per hectare, more desirably from 50 g to 500 g per hectare.
 - It was also an object of the present invention to provide mixtures suitable for treating, controlling, preventing and protecting warm-blooded animals, including humans, and fish against infestation and infection by pests. Problems that may be encountered with pest control on or in animals and/or humans are similar to those described at the outset, namely the need for reduced dosage rates, and / or enhanced spectrum of activity and / or combination of knock-down activity with prolonged control and / or resistance management.

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- This invention also provides a method for treating, controlling, preventing and protecting warm-blooded animals, including humans, and fish against infestation and infection by pests of the orders Siphonaptera, Hymenoptera, Hemiptera, Orthoptera, Acarina, Phthiraptera, and Diptera, which comprises orally, topically or parenterally administering or applying to said animals a pesticidally effective amount of the crystalline modification II, the mixtures and the compositions according to the invention.
- The invention also provides a process for the preparation of a composition for treating, controlling, preventing or protecting a warm-blooded animal or a fish against infestation or infection by pests of the Siphonaptera, Hymenoptera, Hemiptera, Orthoptera, Acarina, Phthiraptera, and Diptera orders which comprises a pesticidally effective

amount of the crystalline modification II, the mixtures and the compositions according to the invention.

The above method is particularly useful for controlling and preventing infestations and infections in warm-blooded animals such as cattle, sheep, swine, camels, deer, horses, poultry, goats, dogs and cats as well as humans.

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Infestations in warm-blooded animals and fish including, but not limited to, lice, biting lice, ticks, nasal bots, keds, biting flies, muscoid flies, flies, myiasitic fly larvae, chiggers, gnats, mosquitoes and fleas may be controlled, prevented or eliminated by the crystalline modification II, the mixtures and the compositions according to the invention.

For oral administration to warm-blooded animals, the crystalline modification II, the mixtures and the compositions according to the invention may be formulated as animal feeds, animal feed premixes, animal feed concentrates, pills, solutions, pastes, suspensions, drenches, gels, tablets, boluses and capsules. In addition, the crystalline modification II, the mixtures and the compositions according to the invention may be administered to the animals in their drinking water. For oral administration, the dosage form chosen should provide the animal with 0.01 mg/kg to 100 mg/kg of animal body weight per day of the crystalline modification II, the mixtures and the compositions according to the invention.

Alternatively, the crystalline modification II, the mixtures and the compositions according to the invention may be administered to animals parenterally, for example, by intraruminal, intramuscular, intravenous or subcutaneous injection. The crystalline modification II, the mixtures and the compositions according to the invention may be dispersed or dissolved in a physiologically acceptable carrier for subcutaneous injection. Alternatively, the crystalline modification II, the mixtures and the compositions according to the invention may be formulated into an implant for subcutaneous administration. In addition, the crystalline modification II, the mixtures and the compositions according to the invention may be transdermally administered to animals. For parenteral administration, the dosage form chosen should provide the animal with 0.01 mg/kg to 100 mg/kg of animal body weight per day of the crystalline modification II, the mixtures and the compositions according to the invention.

The crystalline modification II, the mixtures and the compositions according to the invention may also be applied topically to the animals in the form of dips, dusts, powders, collars, medallions, sprays, spot-on and pour-on formulations. For topical application, dips and sprays usually contain 0.5 ppm to 5000 ppm and preferably 1 ppm to 3000 ppm of the crystalline modification II. In addition, the crystalline

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modification II may be formulated as ear tags for animals, particularly quadrupeds such as cattle and sheep.

The figure and examples below serve to illustrate the invention and are not to be understood as limiting it.

Figure 1: X-ray powder diffractogram of modification II

Figure 2: Differential scanning calorimetry thermogram of modification II

Figure 3: Thermogravimetric analysis of modification II

10 Figure 4: X-ray powder diffractograms of mixtures of modifications I and V

Preparation examples

reflexes listed in Table 2 below.

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Example 1: Characterization of a solid form of fipronil as starting material 15 All preparation procedures below were conducted with two samples of solid fipronil as starting materials which were obtained according to procedures as described in WO 2001/30760, with final crystallization of the product from a solvent mixture of MCB/ethanole (% by weight of ethanol at crystallization start: 13 %) at temperatures of 70°C to 35°C. This solid form in X-ray powder diffractogram studies proved to be 20 crystalline fipronil of a mixture of several crystalline modifications. This mixture has been characterized to consist of crystalline modification I and crystalline modification V, as for the first time identified and described in a co-pending patent application. A least squares refinement with the Topas program with simulated X-ray powder diffractogram patterns from single crystal data of form I and form V shows that in these two example 25 samples, the percentage of form I varies from 30% to 70 %. X-ray powder diffractograms of the two samples are shown in figure 4.

Irrespective of the sample of solid fipronil used as starting material, the crystallization examples given below yielded in modification II.

Example 2: Preparation of modification II by crystallization from THF 1,0 g of crystalline fipronil having a chemical purity of about 96 % by weight was dissolved in 25 ml of THF at 45 to 50°C[B7] in a round bottomed flask. The solution was kept at this temperature while the solvent was slowly evaporated with a gentle flow of inert N₂ gas. The solvent was left to evaporate for about 15 hours, after which the obtained crystalline material was filtered from some residual solvent on a paper filter. The material obtained has the X-ray powder diffractogram shown in Figure 1 with the

Example 3: Preparation of modification II by crystallization from 1,2-dichloroethane (DCE)

1,0 g of crystalline fipronil having a chemical purity of about 96 % by weight was dissolved in 25 ml of DCE at 45°C to 50°C in a round bottomed flask. Then the heating was switched off and the oil bath was left to cool to 20°C to 25°C. The solvent was left to evaporate for about 15 hours. The material obtained has the X-ray powder diffractogram shown in Figure 1 with the reflexes listed in Table 2 below.

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Example 4: Preparation of modification II by crystallization from acetonitrile
 2,0 g of crystalline fipronil having a chemical purity of about 96 % by weight was dissolved in 4 ml of acetonitrile at 75°C to 80°C in a 10 ml test tube to give a clear solution. The solution was cooled down to 5°C in approximately 8h and left at 5°C over night. The supernatant was removed by using a pipet. The solid was dried on a filter
 paper for a few hours and then analyzed to give within a 0.3° 2θ error marginal the X-ray powder diffractogram shown in Figure 1.

Example 5: Preparation of modification II by crystallization from toluene 0,7 g of fipronil having chemical purity of >98 % by weight was suspended in 10 ml of toluene in a 30 ml glass reactor. Stirring was applied and the suspension was heated up to 110 °C where the solid was completely dissolved. The solution was then cooled down to 25 °C with a cooling rate of -1 K/min. The crystallization was observed to begin at 86 °C. The solid product was filtrated and dried on filtration paper over night and then analyzed to give within a 0.3° 20 error marginal the X-ray powder diffractogram shown in Figure 1. The content of fipronil in this experiment was 8.2 w-%.

Example 6: Preparation of modification II by crystallization from monochlorobenzene (MCB)

0,60 g of fipronil having chemical purity of >98 % by weight was suspended in 15 ml of MCB. Stirring was applied and the suspension was heated up to 137 °C where the solid was completely dissolved. The solution was then cooled down to 25 °C with a cooling rate of -1 K/min. The crystallization was observed to begin at 76 °C. The solid product was filtrated and dried on filtration paper over night and then analyzed to give within a 0.3° 2 θ error marginal the X-ray powder diffractogram shown in Figure 1.

Example 7: Preparation of modification II by crystallization from mesitylene 2g of fipronil having chemical purity of ~96 % by weight was dissolved at 15 ml of mesitylene. The solution was cooled down within 5 hours to $20\text{-}25^{\circ}\text{C}$ and left to stand over night. The crystallized solid was filtered and dried in a vacuum oven (at 10 mbar) for ~1.5 h at 40°C . The sample was analyzed to give within a 0.3° 2θ error marginal the X-ray powder diffractogram shown in Figure 1[B8].

Example 8: Preparation of modification II by crystallization from nitrobenzene 2g of fipronil having chemical purity of about 96 % by weight was dissolved at 5 ml of nitrobenzene. The solution was cooled down within 5 hours to 20-25°C and left to stand over night. The crystallized solid was filtered and dried in a vacuum oven (at 10 mbar) for about 1.5 h at 40°C. The sample was analyzed to give within a 0.3° 2θ error marginal the X-ray powder diffractogram shown in Figure 1[B9].

Example 9: Preparation of modification II by crystallization from ethyl benzene 1.5 g of fipronil having chemical purity of >98 % by weight was suspended in 10 ml of ethyl benzene. Stirring was applied and the suspension was heated up to 136°C giving a clear solution. The solution was cooled down to 25°C with a cooling rate of -1 K/min. The crystallization was detected to begin at 118°C. The solid product was filtrated and dried on filtration paper over night and then analyzed to give within a 0.3° 20 error marginal the X-ray powder diffractogram shown in Figure 1.

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Example 10: Preparation of modification II by crystallization from CF_3 -benzene 1.5 g of fipronil having chemical purity of >98 % by weight was suspended in 18.5 ml of CF_3 -benzene. Stirring was applied and the suspension was heated up to 103 °C giving a clear solution. The solution was cooled down to 25°C with a cooling rate of -1 K/min. The crystallization was detected to begin at 97°C. The solid product was filtrated and dried on filtration paper over night at 20-25°C. The sample was analyzed to be within 0.3° 20 error marginal the X-ray powder diffractogram shown in Figure 1.

Table 2: d-spacings and 2θ-angles of modification II

d (Å)	2θ
d = 13.44 ± 0.2 Å	6.6 ± 0.2°
d = 7.84 ± 0.1 Å	11.3 ± 0.2°
$d = 5.50 \pm 0.07 \text{Å}$	16.1 ± 0.2°
d = 5.14 ± 0.05 Å	17.2 ± 0.2°
d = 4.95 ± 0.05 Å	17.9 ± 0.2°
$d = 3.95 \pm 0.05 \text{Å}$	22.4 ± 0.2°
d = 3.77± 0.05 Å.	23.5 ± 0.2°
d = 3.22 ± 0.03 Å	27.6 ± 0.2°
d = 2.91 ± 0.03 Å	30.8 ± 0.2°

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Analysis:

The X-ray powder diffractogram displayed in Figure 1 was recorded using a Siemens D-5000 diffractometer (manufacturer: Bruker AXS) in reflection geometry in the range from 2θ = 2° - 60° with increments of 0.02° using Cu-K α radiation at 25°C. The 2θ values found were used to calculate the stated interplanar spacing d. In Figure 1, the

intensity of the peaks (y-axis: linear intensity in counts) is plotted versus the 2θ angle (x-axis in degrees 2θ).

The single crystal X-ray diffraction data was collected on a Bruker AXS CCD Detector using graphite $Cu_{K\alpha}$ radiation. The structure was solved by using direct methods, refined, and expanded by using Fourier techniques with the SHELX software package (G.M. Sheldrick, SHELX-97, Universität Göttingen, 1997). Absorption correction was performed with SADABS software.

Melting points indicated herein refer to values determined on a Mettler hot stage microscope and represent equilibrium melting points.

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DSC was performed on a Mettler Toledo DSC 823 module in air atmosphere. Crystals taken from the mother liquor were blotted dry on filter paper and place in crimped but vented aluminum sample pans for the DCS experiment. The sample size in each case was 5 to 10 mg. The temperature range was typically 30°C to 230°C at a heating rate of 5°C/min.

TGA measurements were performed on a SEIKO Instrument in nitrogen atmosphere in platinum pans. The sample size in each case was ~ 8-10 mg. The temperature range was 30°C to 600°C at a heating rate 10°C/min.

Investigations of the effect of the temperature of nucleation and beginning of the crystallization to the modification of the crystalline end product carried out with

Polyblock by HEL Ltd. The multi reactor crystallization system allows the monitoring of the crystallization process and change in turbidity with special reflectance turbidity probes by HEL. The heating/cooling mantle and thermostat "Julabo FP 50" as well as the turbidity probes were controlled with a PC.

Claims:

1. A crystalline modification II of fipronil which has an X-ray powder diffractogram showing, at 25°C, at least 5 of the following reflexes:

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$$(1) d = 13,44 \pm 0.2 \text{ Å}$$

$$(2) d = 7.84 \pm 0.1 \text{ Å}$$

(3)
$$d = 5.50 \pm 0.07 \text{ Å}$$

$$(4) d = 5.14 \pm 0.05 \text{ Å}$$

(5)
$$d = 4,95 \pm 0,05 \text{ Å}$$

(6)
$$d = 3.95 \pm 0.05 \text{ Å}$$

$$(7) d = 3,77 \pm 0,05 Å$$

(8)
$$d = 3.22 \pm 0.03 \text{ Å}$$

(9)
$$d = 2.91 \pm 0.03 \text{ Å}$$
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2. The crystalline modification II according to claim 1 undergoing a phase transformation at 105°C to 145°C into two other crystalline modifications I and V of fipronil having melting points at 196 to 197°C and at 202 to 203°C, respectively.

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- 3. The crystalline modification II according to claims 1 or 2 having a fipronil content of at least 98% by weight.
- 4. Solid fipronil comprising the crystalline modification II as defined in any of claims
 25 1 to 3 and a form of fipronil being different from crystalline modification II.
 - 5. Solid fipronil according to claim 4 comprising the crystalline modification II as defined in any of claims 1 to 3 in at least 85% by weight.
- 30 6. A process for preparing the crystalline modification II as defined in any of claims 1 to 3, comprising the steps of:
 - step i) preparing a solution of a solid form of fipronil being different from the crystalline modification II in a solvent S selected from tetrahydrofurane, 1,2-dichloroethane, acetonitrile, mono-, di- or tri(C₁-C₆-alkyl) benzenes[B1], which may be halogenated;

- step ii) effecting crystallization of fipronil; and step iii) isolating the resulting precipitate.
- 7. The process according to claim 6, wherein the solvent S is selected from tetrahydrofuran, monochlorobenzene and 1,2-dichloroethane.
 - 8. The process according to claim 6 or 7, wherein the solvent S is tetrahydrofuran.

- 9. The process according to claim 6 or 7, wherein the solvent S is 1,2-dichloroethane.
- 5 10. The process according to claim 6 or 7, wherein the solvent S is acetonitrile.
 - 11. The process according to claim 6 or 7, wherein the solvent S is monochlorobenzene.
- 10 12. The process according to claim 6, wherein the solvent S is selected from toluene, 1,2-dichlorobenzene, ethyl benzene,-mesitylene, nitrobenzene and CF₃-benzene.
- The process according to claims 11 or 12, wherein, in step ii), the crystallization 13. 15 of fipronil is effected by adding a polar solvent P to the solvent S selected from the group of methanol, ethanol, propan-1-ol, propan-2-ol, butan-1-ol, butan-2-ol, tert-butanol, 2-methyl-propan-1-ol, 2-methyl-propan-2-ol, pentan-3-ol, 2-methylbutan-1-ol, 3-methyl butan-1-ol, 1,2-ethanediol, 1,3-propandiol, 1,2-propandiol, cyclohexanol, acetonitrile, propionitrile, acetone, butanone, pentan-2-one, pen-20 tan-3-one, 4-methyl-2-pentanone, 3-methyl-butan-2-one, 3,3-dimethyl-2-butanone, cyclohexanone, methylacetate, ethylacetate, isopropylacetate, N-butylacetate, isobutylacetate, diethylcarbonate, 2-butoxyethylacetate, dioxane, tetrahydrofuran, diethylether, 2-methyl-tetrahydrofuran, methyl-tert-butylether, dimethylformamide, dimethylacetamide, dimethylsulfoxide, nitromethane, and 25 nitroethane.
 - 14. The process according to any of claims 6 to 12, wherein, in step ii), the crystallization of fipronil is effected by concentration of the solution obtained in step i).
 - 15. The process according to any of claims 6 to 14, wherein, in step ii), the crystallization of fipronil is effected by adding a solvent which reduces the solubility.

- 35 16. The process according to any of claims 6 to 15, wherein step ii) is carried out in the presence of seed crystals of the crystalline modification II as defined in any of claims 1 to 3.
- 17. A synergistic pesticidal or parasiticidal mixture comprising, as active components,
 40 the crystalline modification II as defined in any of claims 1 to 3 and one or more pesticidal or parasiticidal compounds.

18. A pesticidal or parasiticidal composition, comprising the crystalline modification II as defined in any of claims 1 to 3 or the mixture as defined in claim 17 and pesticidally or parasiticidally acceptable carriers and/or auxiliaries.

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5 19. The composition according to claim 18 in the form of an aqueous suspension concentrate.

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- 20. The composition according to claim 18 in the form of water-dispersible granules.
- 10 21. The composition according to claim 18 in the form of a water-dispersible powder.
 - 22. Use of the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21 for controlling pests.
 - 23. A method for controlling pests which comprises contacting the pests or their food supply, habitat, breeding grounds or their locus with a pesticidally effective amount of the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21.
 - 24. A method for protecting a plant from infestation and attack by pests which comprises applying to the foliage or stem of said plant a pesticidally effective amount of the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21.
 - 25. A method as claimed in any of claims 23 or 24, wherein the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21 are applied in an amount of from 5 g/ha to 2000 g/ha.
- 26. The use according to claim 22 or the method according to any of claims 23 to 25 wherein the pests are insects, arachnids or plant nematodes.
 - 27. A method of the protection of seed comprising contacting the seeds with the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21 in pesticidally effective amounts.

28. A method as claimed in claim 27 wherein the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21 is applied in an amount of from 0.1 g to 10 kg per 100 kg of seeds.

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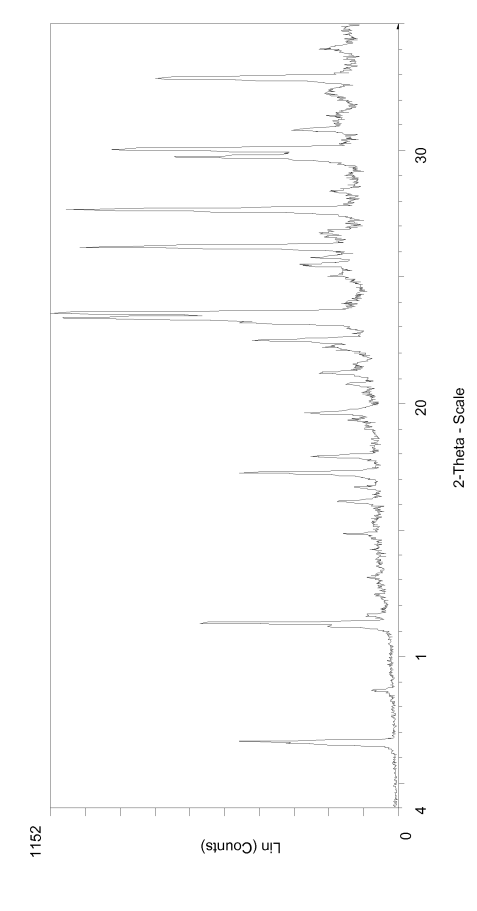
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- 29. Seed comprising the crystalline modification II as defined in any of claims 1 to 3 or the solid fipronil as defined in claims 5 or 6 or the mixture as defined in claim 17 in an amount of from 0.1 g to 10 kg per 100 kg of seeds.
- 10 30. Use of the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21 for combating parasites in and on animals.
- 15 31. A method for treating, controlling, preventing or protecting animals against infestation or infection by parasites which comprises orally, topically or parenterally administering or applying to the animals a parasiticidally effective amount of the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21.
 - 32. A process for the preparation of a composition for treating, controlling, preventing or protecting animals against infestation or infection by parasites which comprises a parasiticidally effective amount of the crystalline modification II as defined in any of claims 1 to 3 or of the solid fipronil as defined in claims 4 or 5 or of the mixture as defined in claim 17 or of the composition as defined in any of claims 18 to 21.

Figure 1. Powder diffractogram of modification II



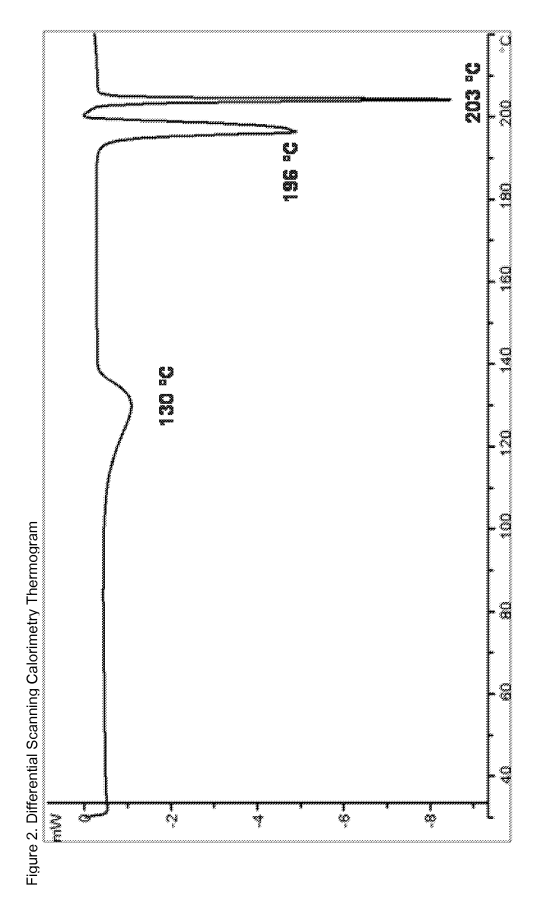
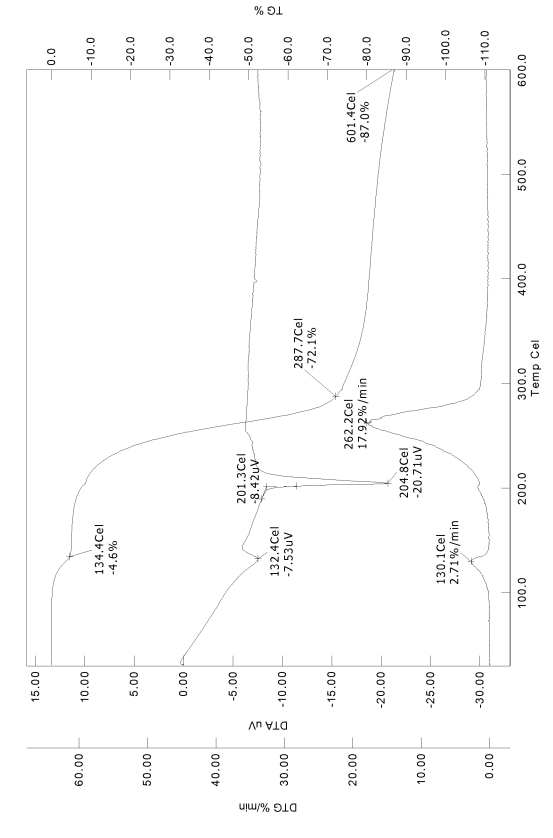
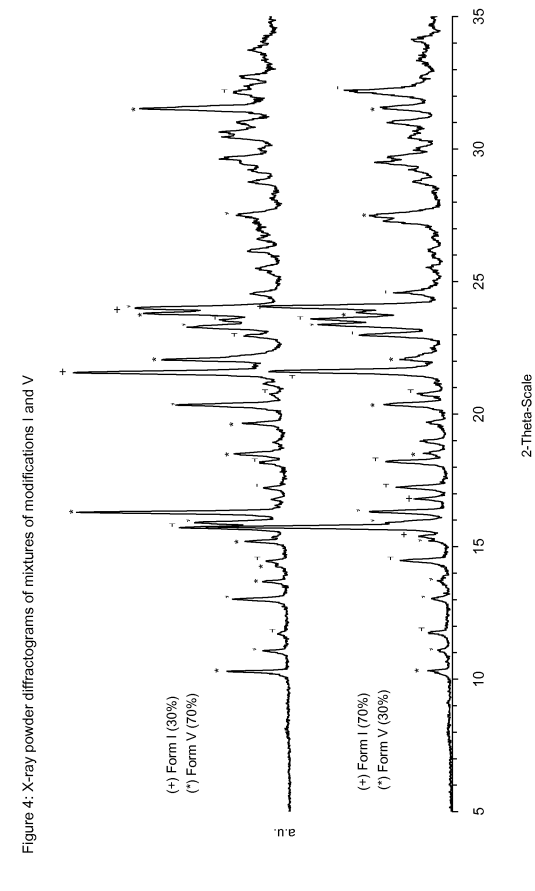


Figure 3. Differential Scanning Calorimetry Thermogram





INTERNATIONAL SEARCH REPORT

International application No PCT/EP2007/061896

A. CLASSIFICATION OF SUBJECT MATTER INV. A01N47/02 A01N2 A01N25/00 A01P7/00 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) AO1N Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) EPO-Internal, CHEM ABS Data, BIOSIS, WPI Data C. DOCUMENTS CONSIDERED TO BE RELEVANT Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X TANG, RI YUAN; ZHONG, PING; LIN, QIU LIAN; 1-6. HU, MAO LIN; SHI, QIAN: 14 - 16"5-Amino-1-[2,6-dichloro-4-(trifluoromethy 1)phenyl]-4-(trifluoromethylsulfanyl)-1H-p yrazole-3-carbonitrile" ACTA CRYSTALLOGRAPHICA, SECTION E: STRUCTURE REPORTS ONLINE. vol. E61, no. 12, 2005, pages 04374-04375, XP002418262 the whole document Α WO 00/62616 A (AVENTIS CROPSCIENCE SA [FR]; HUBER SCOT KEVIN [US]) 26 October 2000 (2000-10-26) the whole document -/-χ Further documents are listed in the continuation of Box C. See patent family annex. Special categories of cited documents: later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. "O" document referring to an oral disclosure, use, exhibition or other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 23 January 2008 05/02/2008 Name and mailing address of the ISA/ Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Bertrand, Franck Fax: (+31-70) 340-3016

INTERNATIONAL SEARCH REPORT

International application No
PCT/EP2007/061896

		PC1/EF200//001890		
C(Continua	ation). DOCUMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.		
		Relevant to claim No. 1-7,11, 14-16		

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No
PCT/EP2007/061896

cited in search report		date		member(s)	date
WO 0062616	Α	26-10-2000	AT	242968 T	15-07-2003
			AU	774162 B2	17-06-2004
			AU	5212000 A	02-11-2000
			BR	0011174 A 2369753 A1	19-02-2002
			CA		26-10-2000
			CN	1351468 A	29-05-2002 13-03-2002
			CZ	20013670 A3	24-07-2003
			DE	60003405 D1	13-05-2004
			DE DK	60003405 T2 1168920 T3	06-10-2003
			EG	22187 A	31-10-2003
			EP	1168920 A1	09-01-2002
			ES	2202135 T3	01-04-2004
			HU	0201728 A2	28-09-2002
			JP	2002542173 T	10-12-2002
			KR	20020009590 A	01-02-2002
			MX	PA01010424 A	30-07-2002
			NO	20014995 A	14-12-2001
			PL	351518 A1	22-04-2003
			PT	1168920 T	31-10-2003
			RÜ	2241331 C2	10-12-2004
			SK	14572001 A3	04-06-2002
			TW	273891 B	21-02-2007
WO 0130760	A	03-05-2001	AT	273961 T	15-09-2004
MO 0120/00	Α.	03-03-2001	ΑÜ	783139 B2	29-09-2005
			AU	1270700 A	08-05-2001
			BG	106622 A	29-12-2002
			BR	PI9917518 A	18-06-2002
			CA	2384283 A1	03-05-200
			CN	1332730 A	23-01-2002
			CZ	20021384 A3	14-08-2002
			DE	69919599 D1	23-09-2004
			DE	69919599 T2	11-08-200
			DK	1222173 T3	20-09-2004
			EΑ	5077 B1	28-10-2004
			ΕP	1222173 A1	17-07-2002
			ES	2222743 T3	01-02-200
			HR	20020438 A2	31-08-2004
			HU	0203206 A2	28-03-2003
			JP	2003512456 T	02-04-200
			MΧ	PA02003944 A	15-10-2003
			PL	354611 A1	09-02-200
			PT	1222173 T	30-11-200
			RO	121209 B1	30-01-200
			TR	200201072 T2	22-07-200
			TR	200202806 T2	21-03-2003
			TW	553936 B	21-09-200
			UA	73752 C2	15-08-200
		•	US	6620943 B1	16-09-2003