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⑤④ Virsraksts: **Ūdeni disperģējamas vai šķīdināmas pesticīdu granulas ar sildot aktivējamām saistvielām, granulēta kompozīcija**

⑤⑦ Kopsavilkums: Lētas, ūdenī ātri šķīdināmas vai disperģējamas granulētas kompozīcijas, kas satur ne mazāk kā 10% poru un sastāv no aglomerātiem, kuri satur pesticīdu granulas, kuras saistītas savā starpā ar sildot aktivējamas ūdenī šķīstošas saistvielas cietiem tiltniēm. Sildot aktivējamās ūdenī šķīstošas saistvielas kušanas temperatūra ir 40 - 120°C robežās; starpība starp mīkstināšanās un sacietēšanas temperatūrām ir mazāka par 5°C; hidrofilā/lipofilā attiecība ir 14 - 19; šķīšanas laiks ne lielāks par 50 minūtēm; kausējuma viskozitāte ne lielāka par 200 centipuāziem. Kā šādas saistvielas piemēri doti: polioksietilēns, dinonilfenols, etilēna oksīda un propilēna oksīda kopolimērs, vai to maisījumi.

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Izgudrojuma formula

1. Granulēta *kompozīcija*, kas *atšķiras* ar to, ka tā satur, % no kopējās kompozīcijas masas:

pesticīda daļiņas.....5-95

sildot aktivējamu ūdenī šķīstošu saistvielu, kuras kušanas temperatūra ir 40-120 °C robežās; starpība starp mīkstināšanās un sacietēšanas temperatūrām ir mazāka par 5 °C; hidrofilī/lipofilā attiecība ir 14-19; šķīšanas laiks ne lielāks par 50 minūtēm; kausējuma viskozitāte ne lielāka par 200 centipauziem.....5-40

un var papildus saturēt vēl vienu piedevu no rindas:

- 1) ūdeni iesūcoši, fiziski uzbriestoši vai gāzi izdaloši dezintegratori;
- 2) aģenti, kas novērš saķepšanu (sagulēšanos);
- 3) ķīmiskie stabilizatori;
- 4) virsmas aktīvās vielas (slapinošie un disperģējošie aģenti) un to maisījumi.

2. *Kompozīcija* pēc 1. punkta, kas *atšķiras* ar to, ka tā satur, % no kopējās kompozīcijas masas:

pesticīdus..... 20-80

sildot aktivējamu saistvielu..... 10-30.

3. *Kompozīcija* pēc 1. punkta, kas *atšķiras* ar to, ka starpība starp saistvielas mīkstināšanās un sacietēšanas temperatūrām ir mazāka par 3 °C.

4. *Kompozīcija* pēc 2. punkta, kas *atšķiras* ar to, ka starpība starp saistvielas mīkstināšanās un sacietēšanas temperatūrām ir mazāka par 3 °C.
5. *Kompozīcija* pēc 1. punkta, kas *atšķiras* ar to, ka piedevas kušanas temperatūra ir 45-100 °C robežās.
6. *Kompozīcija* pēc 2. punkta, kas *atšķiras* ar to, ka piedevas kušanas temperatūra ir 45-100 °C robežās.
7. *Kompozīcija* pēc 3. punkta, kas *atšķiras* ar to, ka piedevas kušanas temperatūra ir 45-100 °C robežās.
8. *Kompozīcija* pēc 2. punkta, kas *atšķiras* ar to, ka piedevas hidrofilī/lipofilā attiecība ir 16-19 robežās.
9. *Pesticīdu granulas*, kas ir disperģējamas vai šķīdinamas ūdenī, *atšķiras* ar to, ka tās satur ne mazāk kā 10% tukšumu un sastāv no aglomerātiem, kuru izmēri ir no 150 līdz 4000 mikroniem, pie kam šie aglomerāti satur pesticīdu daļiņas ar diametru no 1 līdz 50 mikroniem, kas saistītas savā starpā ar cietiem tiltiņiem no sildot aktivējamām ūdenī šķīstošām saistvielām pēc 1. punkta.
10. *Pesticīdu granulas* pēc 9. punkta, kas *atšķiras* ar to, ka tās satur ne mazāk kā 20% tukšumu.
11. *Pesticīdu granulas* pēc 10. punkta, kas *atšķiras* ar to, ka tās satur ne mazāk kā 20% tukšumu un saistvielu pēc 2. punkta.
12. *Pesticīdu granulas* pēc 11. punkta, kas *atšķiras* ar to, ka tās satur saistvielu pēc 8. punkta.
13. *Kompozīcija* pēc 1. punkta, kas *atšķiras* ar to, ka saistvielu ņem no rindas: polioksietilēts dinonilfenols, etilēna oksīda un propilēna oksīda kopolimērs, vai šo vielu maisījumi.
14. *Kompozīcija* pēc 8. punkta, kas *atšķiras* ar to, ka saistvielu ņem no rindas: polioksietilēts dinonilfenols, etilēna oksīda un propilēna oksīda kopolimērs, vai šo vielu maisījumi.
15. *Kompozīcija* pēc 9. punkta, kas *atšķiras* ar to, ka saistvielu ņem no rindas: polioksietilēts dinonilfenols, etilēna oksīda un propilēna oksīda kopolimērs, vai šo vielu maisījumi.
16. *Kompozīcija* pēc 12. punkta, kas *atšķiras* ar to, ka saistvielu ņem no rindas: polioksietilēts dinonilfenols, etilēna oksīda un propilēna oksīda kopolimērs, vai šo vielu maisījumi.

TITLE

5 WATER-DISPERSIBLE OR WATER-SOLUBLE
PESTICIDE GRANULES FROM HEAT-ACTIVATED BINDERS

BACKGROUND OF THE INVENTION

10 In general, water-dispersible or water-soluble
granular compositions are prepared by (1) processes
involving aqueous (or solvent) spraying and
subsequent drying such as pan or fluidized bed
granulation, high intensity mixing, granulation,
spray drying, or by spraying the active material (or
solution thereof) upon a preformed carrier or (2)
15 processes involving compaction such as briquetting,
tableting, and extrusion.

Japanese Patent Application No. 52/30577
discloses slow-release agrochemical-urea fertilizers
which are formed from liquid or organic solvent
20 solutions of agrochemicals, polyoxyethylene nonionic
surfactants, and urea.

U.S. 4,707,287 is directed to the protection of
certain enzymes from a peroxyacid bleach granulate
and discloses an improved granulate enzyme
25 composition comprising a core of enzyme material and
a protective coating comprising an alkaline buffer
salt. This patent broadly mentions the term
"alkylarylethoxylates" among many others as potential
waxy substances used as granulating agents, but there
30 is no teaching of any of the specific heat-activated
binders of the instant invention.

SUMMARY OF THE INVENTION

This invention comprises a low cost, rapidly
water-dispersible or water-soluble pesticidal
35 granular composition which is comprised of
agglomerates consisting essentially of solid

pesticidal particles bound together by solid bridges of a water-soluble heat activated binder (HAB). The granular composition contains at least about 10% voids (preferably 20% or greater) and comprises by weight based on the total weight of the composition:

- (1) 5 to 95% and preferably 20-80% of
pesticidal particles or a mixture of
pesticidal particles having a size in the
range of 1-50 microns or larger if the
pesticide is water-soluble; in
combination with or held together by
solid bridges of
- (2) 5 to 40% and preferably 10-30% of a heat
activated, water-soluble binder (HAB)
having one or more components wherein
said binder meets the following five
criteria:
 - (i) has a melting point range within
40-120°C, and preferably 45 to 100°C;
 - (ii) has a hydrophile/lipophile balance
(HLB) of about 14 to 19, preferably
16-19;
 - (iii) dissolves in mildly-agitated water in
60 min. or less, preferably 50 min. or
less;
 - (iv) has a melt viscosity of at least about
200 centipoise (cps); preferably 1,000
cps or greater, and most preferred
2,000 cps or greater; and
 - (v) has a difference of 5°C, and
preferably 3°C or less between the
softening point and onset of
solidification; and optionally
- (3) one or more additives selected from the
group consisting of

- (i) wicking, physically swelling, or gas-producing disintegrants;
5 (ii) anti-caking agents;
(iii) chemical stabilizers;
(iv) co-binders; and
(v) surfactants (wetting agents or dispersants)

10 such that said composition rapidly forms a high quality dispersion (or solution) in water, is attrition resistant (non-dusty), chemically stable, and non-caking. The agglomerates or granules are 150-4000 microns and preferably 250-1500 microns in
15 size.

DETAILED DESCRIPTION OF THE INVENTION

The most common method of applying agricultural pesticides involves their dilution in a solvent or non-solvent liquid in a mix tank followed by spraying
20 of the resulting solution or dispersion. Because of the increasing costs of non-aqueous solvents and the toxicity of some of them, formulations involving water-soluble or water-dispersible granules have become increasingly popular. In such formulations,
25 the dispersed particles formed on dilution should be no greater than 50 microns in their largest dimension to avoid nozzle pluggage or premature settling which results in uneven application of the pesticide. Consequently, it is necessary that all of the
30 components of the formulated product rapidly and completely disperse or dissolve in the dilution water.

Conventional methods for the preparation of water-soluble or water-dispersible granules involve
(1) solvent spraying such as fluidized bed or pan
35 granulation techniques or the impregnation of an active pesticide agent on preformed carrier granules or (2) compaction such as tableting or extrusion.

Granules prepared by bed or pan granulation are generally sprayable upon dilution with water, while
5 the impregnated or compacted compositions are usually applied dry and mechanically, for example, using spreaders. The solvent spraying processes can yield granules which are rapidly water-dispersible but are expensive, due to the drying step and extensive,
10 space-consuming equipment required. The granules produced from the compaction processes are generally slow in water dispersion. Furthermore, both of these processes often require specialized technology for their operation.

15 Often it is desirable to use mixtures of two or more pesticides of different functions, e.g., a mixture of a herbicide and an insecticide, to provide broad spectrum control over a variety of weeds and/or undesirable organisms. However, some of the
20 individual components are physically or chemically incompatible as mixtures, especially in long-term storage. For example, carbamate insecticides are generally unstable in the presence of alkaline components and sulfonylurea herbicides are known to
25 be unstable in the presence of acidic materials. The chemical incompatibility can be due to an impurity present in the complementary pesticide and not the bioactive component itself. For these reasons it would be desirable to have a sprayable, formulated
30 product consisting of particles or granules wherein potentially incompatible, active components are physically separated.

The present invention comprises low cost, rapidly water-dispersible or water-soluble granular
35 compositions consisting of agglomerates comprised of pesticidal particles bonded together by solid bridges of the heat-activated binder (HAB). The granules

have 10% voids or more, and have a preferred size of 150 to 4,000 microns. The preferred size of
5 pesticidal particles is 1 to 50 microns, especially
for pesticides with low water-solubility, to promote
water dispersion, avoid premature settling, and avoid
nozzle/screen pluggage during tank-mixing or
application in the field. Water-soluble pesticidal
10 particles may be larger.

The granular compositions of this invention
display a break-up time of three minutes or less in
water, good aqueous dispersion properties with a long
tube sedimentation value of 0.02 mL or less,
15 attrition of 33% or less, and are preferably
non-caking after 100 hours at 45°C under a pressure
of 3.5 Kg/cm².

The granules can be comprised of mixtures of
pesticidal particles which are ordinarily chemically
20 incompatible (e.g., in a conventional granule made by
water spraying, such as fluid bed or pan granulation)
because (1) the pesticidal particles can be
physically separated from each other via HAB bridges;
and (2) water is not required during
25 granulation/drying.

Advantages of the present granules include
potential incorporation of incompatible pesticides in
the same granule and low cost. The process used to
prepare these granules is simple and does not require
30 specialized technology. It utilizes readily
available, compact equipment. The process does not
require extensive dust collection systems nor a
space-consuming and expensive drying operation.

The compositions of this invention can be
35 prepared by several processes (either in a batch or
continuous mode) including the processes wherein (1)
the pesticidal particles, the HAB particles and

optional particulate additives are tumbled/mixed and heat is applied externally until the granules have grown to the desired size, following which the heat is shut off and the granules are allowed to cool while still tumbling or sitting in a separate container; or where (2) the pesticidal particles, HAB, and optional particulate additives are intensely sheared/mixed such that frictional heat melts the HAB thereby effecting granulation following which the aggregates are then cooled; or where (3) the pesticidal particles and optional particulate additives are tumbled/mixed and are sprayed with the heat-activated binder which has been pre-heated and is in a molten state following which the resulting agglomerates are cooled.

Processes (1) and (3), involving gentle tumbling/mixing, can be carried out, e.g. in a heated fluidized bed, a heated blender (e.g., paddle or ribbon type blenders, vee-blenders, zig-zag blenders, Lodige® blenders, Nauta® mixers) or a heated pan or drum granulator. Process (3) may not require additional heat other than that needed to melt the HAB for spraying. Subsequent cooling of the resulting agglomerates is done either in or outside of the processing vessel. Process (2) involving high intensity mixing/shearing can be carried out e.g., in Schugi® or turbulator-type vessels. In Process (1) a preferred method of preparing the initial mixture of particulates before granulation is to mill the pesticidal active plus additives and then mix (e.g., via tumbling) with HAB particulates (e.g., of a size 500-1,000 microns). Separation of pesticides can be enhanced and incompatibility then reduced (especially when one active is present in minor proportions) by forming granules from a particulate premix of the

major active component, HAB, and additives, followed
by introduction of the minor active component (and
5 optionally additional HAB), while the granules are
not so as to imbed the second active particulates in
a HAB layer on the surface of the first granules.

The term "pesticide" is intended to refer to
biologically active compositions containing chemicals
10 which are effective in killing pests or preventing or
controlling their growth. These chemicals are
commonly known as herbicides, fungicides,
insecticides, nematocides, acaricides, miticides,
virucides, algicides, bactericides, plant growth
15 regulants and their agriculturally suitable salts.
Preferred are those pesticides that have melting
points above 80°C; more preferred are pesticides that
melt above 100°C. The preferred size of the
pesticidal particles used in this invention is 1 to
20 50 microns. Examples of suitable pesticides are
listed below in Table 1.

25

30

35

TABLE 1

HERBICIDES

5

Cmpd.

10

15

20

25

30

35

No.Common Namem.p. (°C)Chemical Name

1

acifluorfen

142-160

5-[2-chloro-4-(trifluoro
methyl)phenoxy]-2-nitro-
benzoic acid

2

asulam

142-144

methyl [(4-aminophenyl)-
sulfonyl]carbamate

3

atrazine

175-177

6-chloro-N-ethyl-N'-(1-
methylethyl)-1,3,5-
triazine-2,4-diamine

4

bensulfuron
methyl

185-188

2-[[[(4,6-dimethoxy-2-
pyrimidinyl)amino]-
carbonyl]amino]sulfonyl]-
methyl]benzoic acid,
methyl ester

5

bentazon

137-139

3-(1-methylethyl)-(1H)-2,1,3-
benzothiadiazin-4(3H)-one,
2,2-dioxide

6

bromacil

158-159

5-bromo-6-methyl-3-(1-methyl-
propyl)-2,4(1H,3H)pyrimi-
dinedione

7

bromoxynil

194-195

3,5-dibromo-4-hydroxybenzo-
nitrile

Cmpd.				
	No.	Common Name	m.p. (°C)	Chemical Name
5	8	chloramben	200-201	3-amino-2,5-dichlorobenzoic acid
10	9	chlorimuron ethyl	>100	2-[[[(4-chloro-6-methoxy-2-pyrimidinyl)amino]carbonyl amino]sulfonyl]benzoic acid, ethyl ester
15	10	chloroxuron	151-152	N'-[4-(4-chlorophenoxy)-phenyl]N,N-dimethylurea
20	11	chlorsulfuron	174-178	2-chloro-N-[[[(4-methoxy-6-methyl-1,3,5-triazin-2-yl)-amino]carbonyl]benzene-sulfonamide
	12	chlortoluron	147-148	N'-(3-chloro-4-methylphenyl)-N,N-dimethylurea
25	13	clomazone	oil	2-[(2-chlorophenyl)methyl]-4,4-dimethyl-3-isoxazolidinone
30	14	cyanazine	166-167	2-[[[4-chloro-6-(ethylamino)-1,3,5-triazin-2-yl]amino]-2-methylpropanenitrile
35	15	dazomet	104-105	tetrahydro-3,5-dimethyl-2H-1,3,5-thiadiazine-2-thione

	Cmpd.		m.p. (°C)	Chemical Name
	No.	Common Name		
5	16	desmediphan	120	ethyl [3-[[[(phenylamino)- carbonyl]oxy]phenyl]- carbamate
10	17	dicamba	114-116	3,6-dichloro-2-methoxybenzoic acid
	18	dichlobenil	139-145	2,6-dichlorobenzonitrile
15	19	dichlorprop	117-118	(±)-2-(2,4-dichlorophenoxy)- propanoic acid
	20	diphenamid	134-135	N,N-dimethyl-α-phenylbenzene acetamide
20	21	dipropetryn	104-106	6-(ethylthio)-N,N'-bis(1- methylethyl)-1,3,5- triazine-2,4-diamine
25	22	diuron	158-159	N'-(3,4-dichlorophenyl)-N,N- dimethylurea
30	23	thiameturon	>100	3-[[[[(4-methoxy-6-methyl- 1,3,5-triazin-2-yl)amino]- carbonyl]amino]sulfonyl]- 2-thiophenecarboxylic acid. methyl ester

Cmpd.				
	<u>No.</u>	<u>Common Name</u>	<u>m.p. (°C)</u>	<u>Chemical Name</u>
5	24	----	>100	2-[[[N-(4-methoxy-6-methyl-1,3,5-triazine-2-yl)-N-methylamino]carbonyl]-amino]sulfonyl]benzoic acid, methyl ester
10	25	fenac	156	2,3,6-trichlorobenzeneacetic acid
15	26	fenuron	133-134	N,N-dimethyl-N'-phenylurea
	27	fluometuron	163-164	N,N-dimethyl-N'-[3-(trifluoromethyl)phenyl]urea
20	28	fluridone	151-154	1-methyl-3-phenyl-5-[3-(trifluoromethyl)phenyl]-4(1H)-pyridinone
25	29	fomesafen	220-221	5-[2-chloro-4-(trifluoromethyl)phenoxy]-N-(methylsulfonyl)-2-nitrobenzamide
	30	glyphosate	200	N-(phosphonomethyl)glycine
30	31	hexazinone	115-117	3-cyclohexyl-6-(dimethylamino)-1-methyl-1,3,5-triazine-2,4(1H,3H)-dione

Cmpd.				
	<u>No.</u>	<u>Common Name</u>	<u>m.p. (°C)</u>	<u>Chemical Name</u>
5	32	imazamethabenz	>100	6-(4-isopropyl-4-methyl-5-oxo-2-imidazolin-2-yl)- m-toluic acid, methyl ester and 6-(4-isopropyl-4-methyl-5-oxo-2-imidazolin-2-yl)- p-toluic acid, methyl ester
10	33	imazaquin	219-222	2-[4,5-dihydro-4-methyl-4-(1-methylethyl)-5-oxo-1H-imidazol-2-yl]-3-quinoline-carboxylic acid
15	34	imazethapyr	172-175	(±)-2-[4,5-dihydro-4-methyl-4-(1-methylethyl)-5-oxo-1H-imidazol-2-yl]-5-ethyl-3-pyridinecarboxylic acid
20	35	ioxynil	209	4-hydroxy-3,5-diiodobenzo-nitrile
25	36	isoproturon	155-156	N-(4-isopropylphenyl)-N',N'-dimethylurea
30	37	isouron	119-120	N'-[5-(1,1-dimethylethyl)-3-isoxazolyl]-N,N-dimethylurea
35	38	isoxaben	176-179	N-[3-(1-ethyl-1-methylpropyl)-5-isoxazolyl]-2,6-dimethoxybenzamide

Cmpd.				
No.	Common Name	m.p. (°C)	Chemical Name	
5				
	39 karbutilate	176-178	3-[[[(dimethylamino)carbonyl]-amino]phenyl-(1,1-dimethylethyl)carbamate	
10	40 lenacil	316-317	3-cyclohexyl-6,7-dihydro-1H-cyclopentapyrimidine-2,4-(3H,5H)dione	
15	41 MCPA	100-115	(4-chloro-2-methylphenoxy)-acetic acid	
	42 MCPB	100	4-(4-chloro-2-methylphenoxy)-butanoic acid	
20	43 mefluidide	183-185	N-[2,4-dimethyl-5-[[[(trifluoromethyl)sulfonyl]-amino]phenyl]acetamide	
25	44 methabenzthiazuron	119-120	1,3-dimethyl-3-(2-benzothiazolyl)urea	
	45 methazole	123-124	2-(3,4-dichlorophenyl)-4-methyl-1,2,4-oxadiazolidine-3,5-dione	
30	46 metribuzin	125-126	4-amino-6-(1,1-dimethylethyl)-3-(methylthio)-1,2,4-triazin-5(4H)-one	

Cmpd.			
No.	Common Name	m.p. (°C)	Chemical Name
5			
47	met-sulfuron methyl	163-166	2-[[[(4-methoxy-6-methyl-1,3,5-triazin-2-yl)amino]-carbonyl]amino]sulfonyl]-benzoic acid, methyl ester
10			
48	monuron	174-175	N'-(4-chlorophenyl)-N,N-dimethylurea
15			
49	naptalam	185	2-[(1-naphthalenylamino)-carbonyl]benzoic acid
50	neburon	102-103	1-butyl-3-(3,4-dichlorophenyl)-1-methylurea
20			
51	nitralin	151-152	4-(methylsulfonyl)-2,6-dinitro-N,N-dipropyl-aniline
25			
52	norflurazon	174-180	4-chloro-5-(methylamino)-2-[3-(trifluoromethyl)phenyl]-3(2H)-pyridazinone
53	oryzalin	141-142	4-(dipropylamino)-3,5-dinitro-benzenesulfonamide
30			
54	perfluidone	142-144	1,1,1-trifluoro-N-[2-methyl-4-(phenylsulfonyl)phenyl]-methanesulfonamide
35			
55	phenmedipham	143-144	3-[(methoxycarbonyl)amino]-phenyl (3-methylphenyl)-carbamate

Cmpd.			
No.	Common Name	m.p. (°C)	Chemical Name
5	56 picloram	>215 (DEC)	4-amino-3,5,6-trichloro-2-pyridinecarboxylic acid
10	57 prometryn	118-120	N,N'-bis(1-methylethyl)-6-(methylthio)-1,3,5-triazine-2,4-diamine
15	58 pronamide	155-156	3,5-dichloro-N-(1,1-dimethyl-2-propynyl)benzamide
	59 propazine	212-214	6-chloro-N,N'-bis(1-methylethyl)-1,3,5-triazine-2,4-diamine
20	60 pyrazon	205-206	5-amino-4-chloro-2-phenyl-3(2H)pyridazinone
	61 siduron	133-138	N-(2-methylcyclohexyl)-N'-phenylurea
25	62 simazine	225-227	6-chloro-N,N'-diethyl-1,3,5-triazine-2,4-diamine
30	63 sulfometuron methyl	182-189	2-[[[(4,6-dimethyl-2-pyrimidinyl)amino]carbonyl]-amino]sulfonyl]benzoic acid, methyl ester
35	64 tebuthiuron	161-164	N-[5-(1,1-dimethylethyl)-1,3,4-thiadiazol-2-yl]-N,N'-dimethylurea

Cmpd.		Common Name	m.p. (°C)	Chemical Name
No.				
5	65	terbacil	175-177	5-chloro-3-(1,1-dimethyl-ethyl)-6-methyl-2,4(1H,3H)-pyrimidinedione
10	66	terbuthyl- azine	177-179	2-(tert -butylamino)-4-chloro-6-(ethyl-amino)- <u>s</u> -triazine
	67	terbutryn	104-105	N-(1,1-dimethylethyl)-N'-ethyl-6-(methylthio)-1,3,5-triazine-2,4-diamine
15	68	triclopyr	148-150	[(3,5,6-trichloro-2-pyridinyl)oxy]acetic acid
20	69	2,4-D	140	(2,4-dichlorophenoxy)acetic acid
	70	2,4-DB	119-120	4-(2,4-dichlorophenoxy)-butanoic acid
25	71	triasulfuron	>100	(3-(6-methoxy-4-methyl-1,3,5-triazin-2-yl)-1-[2-(2-chloroethoxy)phenylsulfonyl]urea
30	72	primisulfuron	>100	[2-/3-(4,6-bis(difluoromethoxypyrimidin-2-yl-ureidosulfonyl)benzoic acid methylester]
35				

	Cmpd.		m.p. (°C)	Chemical Name
	No.	Common Name		
5	73	----	>100	[2-/3-(4,6-bis(difluoro-methoxy)-pyrimidin-2-yl)-ureidosulfonyl]-benzoic acid methylester]
10	74	NC-311	170-172	[5-pyrazolesulfonamide, N-[(4-methoxy-6-methyl-pyrimidine-2-yl)-amino-carbonyl]-4-methoxy-carbonyl-1-methyl-]
15	75	----	160-162	N-[[[(4,6-dimethoxy-2-pyrimidinyl)amino]carbonyl]-3-(ethylsulfonyl)-2-pyridinesulfonamide
20	76	----	152-159	2-[[[[(4,6-dimethoxy-2-pyrimidinyl)amino]carbonyl]-amino]sulfonyl]-N,N-dimethyl-3-pyridine-carboxamide
25	77	----	204-206	Methyl 2-[[[[(4-ethoxy-6-(methylamino)-1,3,5-triazin-2-yl)amino]carbonyl]amino]-sulfonyl]benzoate
30				
<u>FUNGICIDES</u>				
35	78	carbendazim	302-307	methyl 2-benzimidazole-carbamate

Cmpd.			
No.	Common Name	m.p. (°C)	Chemical Name
5	79 thiuram	146	tetramethylthiuram disulfide
	80 dodine	136	n-dodecylguanidine acetate
10	81 chloroneb	133-135	1,4-dichloro-2,5-dimethoxy- benzene
	82 cymoxanil	160-161	2-cyano-N-ethylcarbamoyl- 2-methoxyiminoacetamide
15	83 captan	178	N-trichloromethylthiotetra- hydrophthalamide
	84 folpet	177	N-trichloromethylthio- phthalimide
20	85 thiophanate- methyl	195	dimethyl 4,4'-(o-phenylene)- bis(3-thioallophanate)
	86 thiabendazole	304-305	2-(thiazol-4-yl)benzimidazole
25	87 chlorothalonil	240-241	tetrachloroisophthalonitrile
	88 dichloran	195	2,6-dichloro-4-nitroaniline
30	89 captafol	160-161	cis-N-[1,1,2,2-tetrachloro- ethylthio]cyclohex-4- ene-1,2-dicarboximide
35			

Cmpd.				
No.	Common Name	m.p. (°C)	Chemical Name	
5				
	90	iprodione	133-136	3-(3,5-dichlorophenyl)-N-(1-methylethyl)-2,4-dioxo-1-imidazolidine carboxamide
10				
	91	vinclozolin	108	3-(3,5-dichlorophenyl)-5-ethenyl-5-methyl-2,4-oxazolidinedione
15				
	92	kasugamycin	202-204 (DEC)	kasugamycin
	93	triadimenol	121-127	beta-(4-chlorophenoxy)-alpha-(1,1-dimethylethyl)-1-H-1,2,4-triazol-1-ethanol
20				
	94	flutriafol	130	+alpha-(2-fluorophenyl)-alpha-(4-fluorophenyl)-1H-1,2,4-triazole-1-ethanol
25				
	95	flusilazol	52-53 HCl 201-203	1-[[bis(4-fluorophenyl)-methylsilyl)methyl]-1H-1,2,4-triazole
30				
	96	hexaconazole	111	(+/-)-alpha-butyl-alpha-(2,4-dichlorophenyl)-1H-1,2,4-triazole-1-ethanol
	97	fenarimol	117-119	alpha-(2-chlorophenyl)-alpha(4-chlorophenyl)-5-pyridinemethanol
35				

Cmpd.				
	<u>No.</u>	<u>Common Name</u>	<u>m.p. (°C)</u>	<u>Chemical Name</u>
5				
	<u>BACTERICIDES</u>			
10	98	oxytetracycline dihydrate	181-182 (DEC)	oxytetracycline dihydrate
	<u>ACARICIDES</u>			
15	99	hexathiazox	108-109	trans-5-(4-chlorophenyl)-N-cyclohexyl-4-methyl-2-oxo-3-thiazolidinecarboxamide
20	100	oxythioquinox	169-170	6-methyl-1,3-dithiolo-[2,3-B]quinonolin-2-one
	101	dienochlor	122-123	bis(pentachloro-2,4-cyclopentadien-1-yl)
25	102	cyhexatin	245	tricyclohexyltin hydroxide
	<u>INSECTICIDES</u>			
30	103	carbofuran	150-152	methylcarbamic acid, ester with 2,3-dihydro-2,2-dimethyl-7-benzofuranol
35	104	carbaryl	142	methylcarbamic acid, ester with α -naphthol

Cmpd.			
	<u>No.</u>	<u>Common Name</u>	<u>m.p. (°C)</u>
5			<u>Chemical Name</u>
	105	thiodicarb	173-174
			dimethyl N,N'-[thiobis-(N-methylimmo)carbonyl-oxy]]-bis[ethanimido-thioate]
10			
	106	deltamethrin	98-101
			α -cyano-3-phenoxybenzyl-cis-3-(2,2-dibromovinyl)-2,2-dimethylcyclopropane carboxylate
15			
20			
25			
30			
35			

The term "heat-activated binder" refers to any surface active material comprised of one or more components which dissolve rapidly in water, have some viscosity near the melting point for tackiness, and are thus capable of acting as a binder when heat is applied. At some elevated temperature, the binder softens and melts, thereby becoming sticky enough to bind the pesticidal particles into granules. A more preferred amount of binder used in this invention is 10-30% by weight based on the total weight of the composition. A more preferred melting point range for the binders of this invention is 45°C to 100°C. Examples of suitable heat-activated binders, which are not intended to be limiting, are ethylene oxide/propylene oxide copolymers and polyethoxylated dinonylphenol.

The HAB can be a single component or multi-components which are mixed in the solid state, co-melted or co-dissolved. Preferred single component HAB's are ethylene oxide/propylene oxide copolymers and polyethoxylated dinonylphenol. Specifically preferred single components are block copolymers of ethylene oxide/propylene oxide, where 80% is ethylene oxide, and 20% is propylene oxide, and polyethoxylated dinonylphenol with 150 ethylene oxide units. The preferred copolymer has an HLB of 16 and a melting point of about 45 to 61°C. The preferred dinonylphenol derivative has an HLB of about 19 and a melting point of about 48 to 63°C.

The HAB must meet the following five criteria:

- (1) have a melting point range within 40 to 120°C;
- (2) be water-soluble with a hydrophile/lipophile balance (HLB) of about 14 to 19;

(3) dissolve in mildly-agitated water in 50 minutes or less;

5 (4) have a melt viscosity of at least 200 cps; and

(5) have a difference of 5°C or less between the softening point and onset of solidification.

10 The use of a HAB having a very low melting point can lead to caking of the granules, while use of a HAB having a very high melting point can require a temperature sufficiently high so that decomposition of the pesticide or other components can occur during granulation.

15 Surface activity as measured by the critical HLB range is necessary to provide good bonding of the HAB to pesticidal particles and rapid wetting at the onset of bridge dissolution when the granules are placed in water. Materials which have too low an HLB
20 are not completely water-soluble.

The dissolution rate in water is very important, since factors other than HLB affect dissolution, e.g., viscosity of the hydrated HAB and its tendency to form a gel-like layer when in contact
25 with mildly or non-agitated water.

The use of a HAB having the specified melt viscosity and minimum difference between softening and solidification temperatures is necessary so that it will be tacky enough to effect agglomeration of
30 pesticidal particles near the melting point of the HAB.

Additives, many of which are commonly used in conventional granules, may optionally be used in HAB granules. Examples include:

35 (1) disintegrants which wick in water, physically expand, or produce gas to aid break-up of

the granule. Non-limiting examples of suitable disintegrants include cross-linked polyvinyl
5 pyrrolidone, microcrystalline cellulose, cross-linked sodium carboxymethyl cellulose, salts of polyacrylates of methacrylates, and the combination of sodium or potassium bicarbonates or carbonates
10 with acids such as citric or fumaric acid, used alone or in combination, at levels of up to 30% by weight based on the total weight of the composition;

(2) anticaking agents to prevent clumping of granules when stored under hot warehouse conditions. Non-limiting examples of suitable anticaking agents
15 include sodium or ammonium phosphates, sodium carbonate or bicarbonate, sodium acetate, sodium metasilicate, magnesium or zinc sulfates, magnesium hydroxide (all optionally as hydrates), and sodium alkylsulfosuccinates;

20 (3) chemical stabilizers to prevent decomposition of the active(s) during storage. Non-limiting examples of suitable chemical stabilizers include alkaline earth or transition metal sulfates such as magnesium, zinc, aluminum, and
25 iron (optionally as hydrates) used at levels of 1-9% by weight based on the total weight of the composition;

(4) co-binders to achieve optimized properties such as increased granulation efficiency
30 or improved anticaking. Up to 50% co-binders such as polyethylene glycols, polyethylene oxide, polyethoxylated fatty acids or alcohols, hydrated inorganics such as sodium silicate, sorbitol, or urea may be used; and

35 (5) surfactants to improve the speed and quality of wetting and dispersion of the granule upon

mixing with water. Often dispersing agents are most useful, since the HAB itself has wetting characteristics.

Examples of preferred dispersants include sodium or ammonium salts of sulfonated naphthalene (or methyl naphthalene)-formaldehyde condensates, sodium, calcium, or ammonium salts of ligninsulfonates (optionally polyethoxylated); dialkyl; diolalkynes; sodium taurates; and sodium or ammonium salts of maleic anhydride copolymers.

HAB candidates may be identified by the following tests:

(1) the melting point is measured by DSC (Differential Scanning Calorimetry) at a 5°C/minute heating rate. The onset of the melt should be no lower than 40°C;

(2) The hydrophile/lipophile balance with a total possible range of 1 to 20 is determined by the method outlined in McCutcheon's "Detergents and Emulsifiers", 1971 annual, page 223;

(3) The rate of dissolution in water is determined by the following procedure:

- (a) a sample of the test material (0.15 g) is placed in the bottom of a glass graduated cylinder with an inside diameter of 2.8 cm,
- (b) the cylinder is placed on a steam bath (alternately heated externally with a stream of hot air near the bottom) until the sample is fully melted,
- (c) the cylinder is placed on a level surface and the sample allowed to

solidify upon cooling to 25°C,
yielding an even layer in the
bottom,

(d) water is added (100 mL at 25°C) to
the cylinder and is stirred at 110
rpm with a rectangular metal or
plastic paddle having a thickness
of 1.5 mm, a width of 18 mm, and a
height of 16.5 mm so that the
bottom of the paddle is 48 cm above
the surface of the solidified
sample, and

(e) the time for complete dissolution
of the sample is noted;

(4) The viscosity at the softening point is
calculated using an Arrhenius plot ($\ln$ viscosity vs
 $1/T$).

The plot is derived from experimental viscosity
measurements vs temperature using a rotational
viscometer operated at a shear rate of 1.16 sec^{-1} .
Viscosity measurements are taken over at least a 30°C
temperature range whose minimum temperature is within
1°C of the softening point as measured by DSC.

Another requirement of the melt behavior of HAB
candidates is that the onset of softening in the
heating curve differs by 5°C or less from the onset
of solidification in the consecutive cooling curve.

This parameter is measured using a differential
scanning calorimeter (e.g., Du Pont Instruments 1090
Thermal Analyser with model 910 DSC module). One to
three milligrams of sample is typically used in a
hermetically-sealed coated aluminum pan. The heating
curve endotherm is observed at 5°C/minute, while the
cooling curve exotherm is observed at 1°C/minute.

Typically a sample is heated from 25°C to 100°C to 120°C and then allowed to cool back to 25°C. It
5 should be noted that a given HAB can exhibit a broad melting behavior (usually 12 to 16°C from softening to full melt).

Granules of this invention have at least 10% voids and preferably at least 20%. The upper limit
10 of voids is set by the fragility (high attrition) of the granule. Determination of voidage is accomplished by pycnometer measurements of the starting premix powder and the final HAB granules, using a paraffin oil. Alternately, helium
15 porosimetry may be used. Voids are important to speed penetration of water into the granule and thus aid break-up in the mix tank.

The granules also exhibit break-up times in water of less than 3 minutes and preferably less than
20 2. Break-up time is measured by adding a sample of the granules (0.5 g, 250 to 1410 microns) to a 100 mL graduated cylinder [internal height after stoppering is 22.5 cm; I.D. is 28 mm] containing 90 mL of distilled water at 25°C, following which the cylinder
25 is clamped in the center, stoppered, and rotated about the center at 8 rpm until the sample is completely broken up in the water.

Formation of a high quality aqueous dispersion is also a desirable property and is determined by the
30 long tube sedimentation test in U.S. 3,920,442 (Col. 9, lines 1 to 39). Acceptable values correspond to 0.02 mL, preferably 0.01 mL of solids after 5 minutes of settling.

The granules should exhibit low attrition
35 characteristics which can be determined by the attrition test in U.S. 3,920,442 (Col. 8, lines 5 to 48). The test is modified to use test samples of the

commercial granule size (e.g., 250 to 1410 microns). Attrition values of less than 40% and preferably less than 30% are acceptable.

The granules should also resist caking. This property is determined by taping a stainless steel disc (0.9 mm thick x 51 mm diameter) flush with the bottom of a glass cylinder (46.5 mm i.d. x 75 mm length x 51 mm thickness) following which the granular sample (20 g) is delivered to the cylinder assembly and leveled, and a second stainless steel disc (0.9 mm thick x 44.5 mm diameter) is placed on the top of the granules.

A 400 g weight (45 mm diameter or less) is then placed on top of the inner disc, and the entire assembly is placed in an oven for 100 hours at 45°C (preferably 55°C) following which the assembly is removed from the oven, the weight removed, and the sample allowed to cool to room temperature. The bottom disc is then detached and if the sample flows out of the cylinder, the resistance to caking is excellent, and if the sample remains in the cylinder, the cake is removed, placed onto a flat surface and a penetrometer is used with a single-edged razor to measure the minimum force necessary to cleave the cake.

Cakes requiring a force of less than 100 g, and preferably less than 5 g are acceptable.

The following examples are presented to illustrate, but not to restrict, this invention.

Definitions of Ingredients Used in Examples

	<u>Name</u>	<u>Identity</u>
5	Macol® DNP 150 (Mazer Chemicals)	polyethylated dinonylphenol (150 ethylene oxide units)
		<u>Melt behavior:</u>
10		• melting point - softening point 48°C; finish 63°C
		• difference between softening point and onset of solidification = 2°C
15		• melt viscosity - about 1,900 cps at softening point
		<u>Dissolution Rate:</u> 19 minutes
20		<u>HLB:</u> 19
	Pluronic® F108 (BASF)	ethylene oxide/propylene oxide block copolymer with 80% ethylene oxide and 20% propylene oxide units
25		<u>Melt behavior:</u>
		• melting point - softening point 45°C; finish 61°C
30		• difference between softening point and onset of solidification = 0°C
35		

	<u>Name</u>	<u>Identity</u>
5		• melt viscosity: at softening point about 26,500 centipoises (cps)
10		<u>Dissolution Rate:</u> 50 minutes <u>HLB:</u> 16
15	Hodag® E100 (Hodag Chemical Corp.)	a 100 mole ethoxylate of nonylphenol
20		<u>Melt behavior:</u> • melting point - softening point 40°C; finish point 64°C • difference between softening point and onset of solidification = 0°C • melt viscosity at softening point = 1,100 cps
25		<u>Dissolution Rate:</u> 20 minutes
30		<u>HLB:</u> 19
35		

31

	<u>Name</u>	<u>Identity</u>
5	Iconol® OP-40 (BASF)	a 40 mole ethoxylate of octylphenol
10		<u>Melt behavior:</u> • melting point - softening point = 40°C; finish = 55°C • difference between softening point and onset of solidification = 3°C
15		• viscosity at softening point about 700 cps
20		<u>Dissolution Rate:</u> 18 minutes
		<u>HLB:</u> 18
	Polyplasdone® XL-10 (GAF)	Cross-linked polyvinyl pyrrolidone
25	Avicel® PH-105 (FMC)	microcrystalline cellulose
30	AC-DI-SOL® (FMC)	Cross-linked sodium carboxymethyl cellulose
	Morwet® D425 (Desoto)	sodium naphthalene sulfonate formaldehyde condensate
35	Morwet® EFW (Desoto)	sodium alkyl naphthalene sulfonate

	<u>Name</u>	<u>Identity</u>
5	Lignosol® TSF (Reed)	ammonium lignosulfonate
	Monawet® MB100 (Mona)	sodium dibutylsulfosuccinate
10	Aerosol® A196 (Amer. Cy.)	sodium dicyclohexylsulfo- succinate (+ 15% sodium benzoate)
15	Explotab® (Edward Mendell Co.)	sodium starch glycolate
	Triton® AG-120 (R & H)	polyethoxylated nonyl phenol absorbed on silica
20	Triton® X-120 (R & H)	polyethoxylaed nonyl phenol absorbed in MgCo ₃

EXAMPLE 1

25 All ingredients below (with the exception of
the Macol®) were mixed and then milled on a high
intensity rotary shearing mill. The resultant
mixture was then mixed with Macol® DNP 150 (<840
microns) to form a premix for granulation. A 150 g
30 portion of the premix was placed in a fluidized bed
and heated air was applied to the granules. When the
temperature of the granules reached 70°C (about 12
minutes) the heat was shut off and the granules
allowed to cool while still fluidized by unheated
35 air. A 70% yield of a 250 to 1410 micron spherical
granules was realized. The premix formulation and
resulting properties of the granules are given below.

Premix Formulation

5		<u>Weight Percent</u>
	Chlorsulfuron	75.0
	Macol® DNP150	10.0
	Anhydrous MgSO ₄	6.0
	Polyplasdone® XL-10 +	9.0
10	Impurities	

Properties of Granules (250 to 1410 microns)

	% Attrition	21.0%
	25°C Break-up Time	
15	in water	75.0 sec
	OC Break-up Time	
	in water	102.0 sec
	25°C Break-up Time -	
	in 28-0-0 Liquid	
20	Fertilizer	214.0 sec
	55°C Caking	0.0 g
	Long Tube	
	Sedimentation	Trace
	Long Tube	
25	Sedimentation	
	(1 wk/55°C)	.003
	Assay (% chlor-	
	sulfuron) Control	72.9%
	1 wk/55°C	70.0%
30		

EXAMPLE 2

Example 1 was repeated except that the premix without binder was hammer-milled.

	<u>Premix Formulation</u>	<u>Example 2</u>
5	Chlorsulfuron Tech (%)	77
	Macol® DNP150 (%)	10
	MgSO ₄ ·7H ₂ O (%)	3
	ZnSO ₄ ·7H ₂ O (%)	3
	Polyplasdone® XL-10 (%)	7
10	Avicel® PH-105 (%)	-
	AC-DI-SOL® (%)	-
	Premix Charge (GM)	1934
	Conversion (%)	73
	Attrition (%)	29
15	Long tube sedimentation	0.002
	Long tube sedimentation (1 wk/55°C)	0.002
	25°C H ₂ O Break-up (sec)	83
	0°C H ₂ O Break-up (sec)	107
20	25°C 28-0- Break-up (sec)	250
	55°C Caking (GM Force)	0

EXAMPLE 3

Approximately 73.84 g of 2,4-D, Na salt and

25 1.16 g of 2-[[N-(4-methoxy-6-methyl-1,3,5-triazin-2-yl)-N-methylamino]carbonyl]amino]sulfonyl]benzoic acid, methyl ester, Na salt were milled together as in Example 1. This material was then blended with 25 g of Macol® DNP-150 (<840 microns). The mixture was

30 then added to a laboratory double cone blender and heated with a heat gun to 77°C, whereupon granulation was observed. The heat was removed and the granules allowed to cool to 50°C then removed from the blender. Approximately 97.3 g were recovered with

35 88.3% being between 250 and 1410 microns in size. The physical properties of the granules were: long tube sedimentation (5 minute reading) 0 ml,

attrition-33.7%, break-up time in 25°C water-2.17 minutes and bulk density-0.50 g/ml.

5 The granules exhibited good chemical stability upon aging, also with no loss of the above physical properties.

EXAMPLE 4

10 Approximately 72.86 g 2,4-D, Na salt, 1.14 g of 2-[[N-(4-methoxy-6-methyl-1,3,5-triazin-2-yl)-N-methylamino]carbonyl]amino[sulfonyl]benzoic acid, methyl ester, Na salt and 1 g NaHCO₃ were milled together as in Example 1. This mixture was then blended with 25 g Macol® DNP-150 (<840 microns). The
15 procedure of Example 3 was then followed to produce granules. Approximately 93.4 g were recovered with 90.9% being between 250 and 1410 microns. The physical properties of the as made granules were: long tube sedimentation (5 minute reading)-trace,
20 bulk density-0.5 g/ml, attrition 37.5%, and break-up time in 25°C water-2.18 minutes. The properties after aging at 45°C for 3 weeks were: long tube sedimentation (5 minute reading)-trace, bulk
25 density-0.5 g/ml, attrition-36.8%, break-up time in 25°C water-2.19 minutes. As in Example 3, these granules also maintained good chemical stability on aging.

EXAMPLE 5

30 Approximately 1480 g methabenzthiazuron, 9.80 g ammonium salt of chlorsulfuron (technical), 5.16 g Sellogen® HR, 6.88 g Petro® D425, 12.90 g diammonium hydrogen phosphate, 137.26 g kaolin clay, 40 g MgSO₄, and 48 g Polypladone® XL-10 were milled in an ACM
35 mill at 90 g/min feed rate, rotor speed of 11,000 rpms, classifier speed of 6,000 rpms and as air flow of 50 cfm. Approximately 1643 g of milled material

was recovered from the mill. Three 600 g batches were granulated in a laboratory 2.2 liter vee blender by combining 522 g of the milled material and 78 g of less than 500 micron Pluronic® F108. This material was blended and heated as in Example 3 to 70°C when granulation was observed. The heat was removed and the granules cooled to 45°C before removing from the blender. Approximately 1787 g of granules were recovered from the blender with 88.7% being between 250 to 1410 microns in size. The physical properties of the granules were: long tube sedimentation (5 minute reading)-0.015 ml, break-up time in 25°C water-1.70 minutes, bulk density-0.5 g/ml, and attrition-11.9%. This material passed both the 45°C and 55°C caking test.

EXAMPLE 6

The following were hammer-milled:
86.9% Na 2,4-D tech (83% assay as acid)
1.3% 2-[[N-(4-methoxy-6-methyl-1,3,5-triazine-2-yl)-N-methylamino]carbonyl]-amino]sulfonyl]benzoic acid, methyl ester,
Na salt (92% assay as the free sulfonylurea)
5.9% Morwet® D425
5.9% Morwet® EFW

The resulting premix was continuously auger fed [62 g/min.] to a 35.6 cm diameter disc agglomerator (56° angle with horizontal) rotating at 30 rpm. Molten Macol® DNP-150 (95°C) was sprayed continuously (23 g/minute) onto the premix in the agglomerator using an external mix, air-atomizing, spray nozzle. The Macol® comprised 25-30% of the total mass of resulting granules. A yield of 61% of 1410 to 1680 micron granules was obtained. The granules had an attrition of 39%, a long tube sedimentation of

0 mL, and a break-up time in 25°C water of 150 seconds.

5

EXAMPLE 7

The premix of Example 1 was metered continuously to a 2 liter capacity stainless steel drum (10 cm high x 12 cm diameter) which was rotated at 34 rpm at a 30° angle with the horizontal. The bed of premix on the drum was maintained at 70-77°C by heating the external wall of the drum with an infrared lamp. Approximately 89% of granules exiting the drum were 250 to 1410 microns in size. These granules had a long tube sedimentation of 0.01 ml, an attrition of 40%, and a break-up time of 64 seconds in 25°C water.

10

15

EXAMPLE 8

A premix of 20 g of sodium 2,4D (84% assay), 0.5 g 2-[[N-(4-methoxy-6-methyl-1,3,5-triazin-2-yl)-N-methylamino]carbonyl]amino]sulfonyl]benzoic acid, methyl ester, Na salt (91% assay), and 3.6 g of Macol® DNP-150 were milled as in Example 1 for two minutes. The dustless contents were then screened, revealing that 61% of the granules produced were in the 149 to 840 micron range and 89% in the 74 to 840 micron range. The long tube sedimentation of the granules in the latter size range was 0 ml, the break-up time in 25°C water was 90 seconds, and the attrition was 40%. There was 0% decomposition of either active after aging 1 week and 3% decomposition of the sulfonylurea after 2 weeks at 55°C.

25

30

35

EXAMPLE 9

Approximately 100 g of premix was made by
 5 combining the following ingredients:

	Chlorsulfuron technical	78.5 g
	Macol® DNP-150	12.0 g
	ZnSO ₄ ·7H ₂ O	2.0 g
	MgSO ₄ ·7H ₂ O	2.0 g
10	Ac-Di-Sol®	2.75 g
	Avicel® PH-105	2.75 g

The premix was milled as in Example 1 to a
 powder which was then placed in a fluid bed
 15 granulator and fluidized with hot air. The bed was
 gradually heated to 70°C (9-10 minutes). The
 granules formed as the binder softened. The heat was
 removed and the granules were allowed to cool while
 fluidization was continued. After cooling, the
 20 granulated product was sieved. Approximately 76 g of
 granules were recovered in the 250 to 1410 micron
 size range which exhibited the following properties.

	Break-up Time in 25°C	81 seconds
	water	
25	Caking (1 day/55°C/3.5 Kg/cm ²)	none
	Assay on sample stored 1 week	74.4%
	at -6°C	
	Assay on sample stored 1 week	75.8%
	at 55°C	
30	Long Tube Sedimentation	
	(before and after aging)	0.005 mL
	Attrition	12%

EXAMPLE 10

100 g of a premix was prepared by combining:

5	Chlorsulfuron technical	77.0 g
	Hodag® E-100	10.0 g
	Anhydrous sodium carbonate	2.0 g
	Polyplasdone® XL-10	2.0 g
	Sodium acetate trihydrate	2.75 g

10

The premix was milled and granulated as described in Example 9. Approximately 60 g of 250 to 1410 micron granules were recovered. Break-up times in 25°C water averaged 91 seconds. There was no caking after 4 days at 55°C.

15

EXAMPLE 11

100 g of a premix was prepared by combining:

20	Chlorsulfuron technical	77.0 g
	Iconol® OP-40	10.0 g
	Anhydrous sodium carbonate	6.0 g
	Polyplasdone® XL-10	7.0 g
	Sodium acetate trihydrate	2.75 g

25

The premix was milled and granulated as described in Example 9. Approximately 57 g of 250 to 1410 micron granules were obtained. The break-up time in 25°C water was 69 seconds. There was no caking after 4 days at 55°C. Attrition was 34%.

30

EXAMPLE 12

100 g of a premix was prepared by combining:

35

5	Benzoic Acid, 2-[[[(4-ethoxy-6-methylamino-1,3,5-triazin-2-yl)aminocarbonyl]amino-sulfonyl]methyl ester	77.0 g
	Hodag® E-100	10.0 g
	Anhydrous MgSO ₄	6.0 g
	Polyplasdone® XL-10	7.0 g

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The premix was milled and granulated as described in Example 9. Approximately 59 g of 250 to 1410 micron granules were obtained. The break-up time in 25°C water was 90 seconds. The granules did not cake after 4 days at 55°C. Attrition was 28% and long tube sedimentation was 0.005 ml. Chemical stability was excellent.

EXAMPLE 13

The granulation procedure of Example 5 was repeated, using the following ingredients in the premix:

25	methabenzthiazuron	1460 g
	Chlorsulfuron tech.	10 g
	Pluronic® F108	240 g
	MgSO ₄	10 g
	ZnSO ₄ ·7H ₂ O	90 g
	Morwet® D-425	50 g
	Avicel® PH-105	140 g

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The resulting granules (250 to 1410 microns) were produced in 83% yield and had the following properties: long tube sedimentation-0.015 ml, caking-100 g at 45°C, break-up time in 25°C water-90 seconds, and attrition-10%.

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CLAIMS

What is claimed is:

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1. A granular composition which comprises by weight based on the total weight of the composition: 5 to 95% of pesticidal particles in combination with 5 to 40% of a water-soluble heat activated binder having a melting point range within 40° to 120°C, a difference of less than 5°C between the softening point and the onset of solidification, a hydrophile/lipophile balance of about 14 to 19, a dissolution time of not greater than about 50 minutes; and a melt viscosity of at least about 200 cps; and optionally at least one additive selected from the group consisting of

- 10
- (i) wicking, physically swelling, or gas-producing disintegrants;
 - 20 (ii) anti-caking agents; and
 - (iii) chemical stabilizers; and
 - (iv) surfactants (wetting or dispersants) agents and mixtures of the foregoing.

25 2. A composition of Claim 1 comprising by total weight 20 to 80% of pesticidal particles, and 10 to 30% of the heat activated binder.

30 3. The composition of Claim 1 wherein the difference between the softening point and the onset of solidification is less than 3°C.

35 4. The composition of Claim 2 wherein the difference between the softening point and the onset of solidification is less than 3°C.

5. The composition of Claim 1 wherein the melting point range is 45-100°C.

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6. The composition of Claim 2 wherein the melting point range is 45-100°C.

7. The composition of Claim 3 wherein the melting point range is 45-100°C.

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8. The composition of Claim 2 wherein the hydrophile/lipophile balance is in the range of 16-19.

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9. Water-dispersible or water-soluble pesticidal granules which contain at least about 10% voids and comprise agglomerates having a size in the range 150 to 4,000 microns which agglomerates are comprised of pesticidal particles having a size in the range of 1 to 50 microns in diameter bonded together by solid bridges of a water-soluble heat-activated binder as described in Claim 1.

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10. The granules of Claim 9 which contain at least about 20% voids.

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11. The granules of Claim 10 which contain at least about 20% voids and the binder of Claim 2.

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12. The granules of Claim 11 which contain the binder of Claim 8.

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13. The composition of Claim 1 wherein the binder is selected from the class consisting of
5 polyethoxylated dinonylphenol, ethylene oxide/-propylene oxide copolymer and mixtures of the foregoing.

14. The composition of Claim 8 wherein the
10 binder is selected from the class consisting of polyethoxylated dinonylphenol, ethylene oxide/-propylene oxide copolymer and mixtures of the foregoing.

15 15. The composition of Claim 9 wherein the binder is selected from the class consisting of polyethoxylated dinonylphenol, ethylene oxide/-propylene oxide copolymer and mixtures of the
20 foregoing.

16. The composition of Claim 12 wherein the binder is selected from the class consisting of polyethoxylated dinonylphenol, ethylene oxide/-propylene oxide copolymer and mixtures of the
25 foregoing.

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