

Institut für
Biologische Analytik und
Consulting IBACON GmbH

Final Report

**Laboratory Testing for Toxicity
(Acute Oral LD₅₀) of BNF 5119B
on Honey Bees (*Apis mellifera* L.)
(Hymenoptera, Apidae)
-Limit Test-**

(GLP compliant study based on EPPO 170 (1992))

Author: Dipl. Biol. [REDACTED]

Study Completion Date: September 21, 1999

Sponsor

Bayer AG
Crop Protection Division
Institute for Environmental Biology
Alfred-Nobel-Str. 50
40789 Monheim
Germany

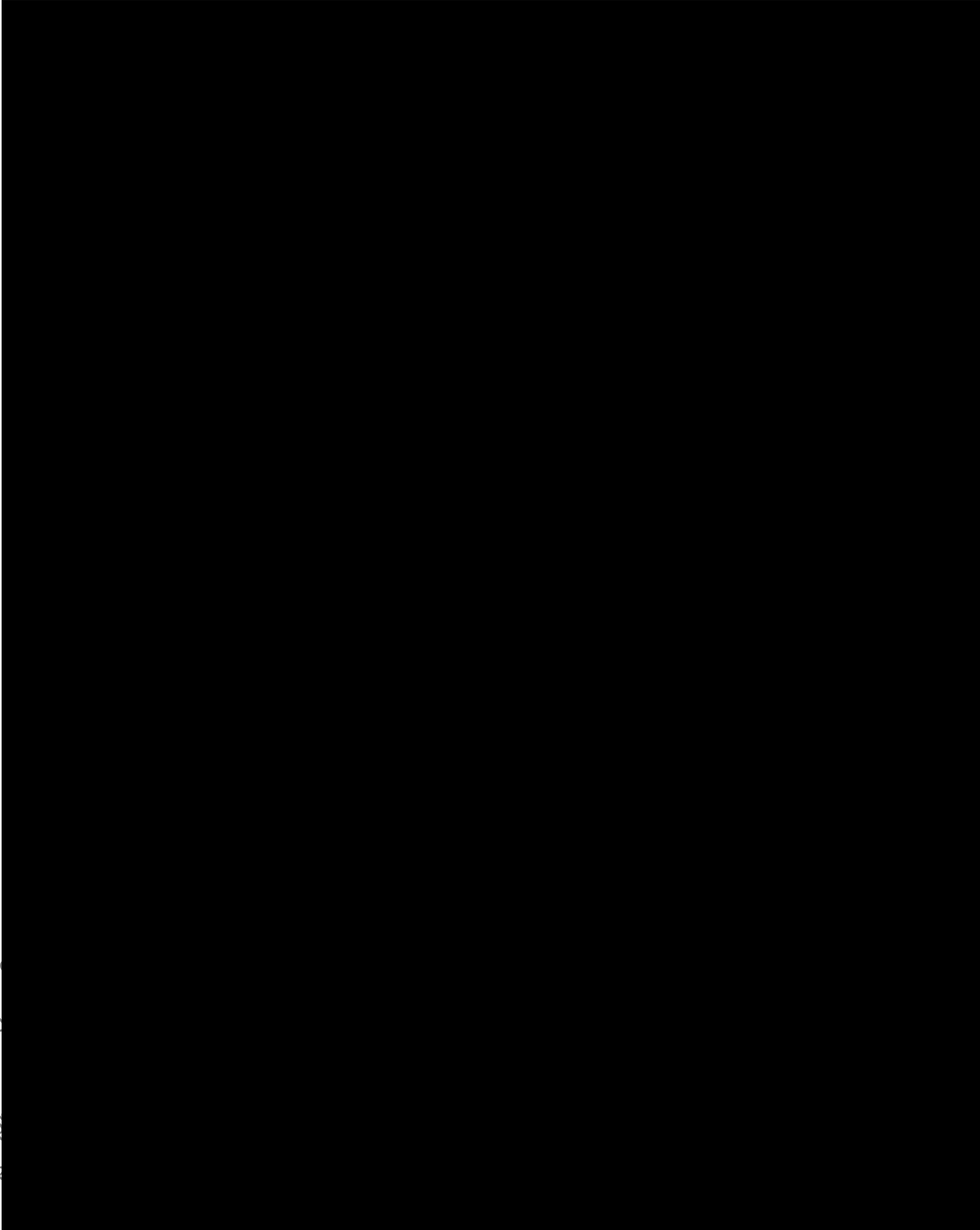
Test Facility

Institut für Biologische Analytik
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Project 6380036



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1. Summary

Report: [REDACTED] (1999): Laboratory Testing for Toxicity (Acute Oral LD₅₀) of BNF 5119B on Honey Bees (*Apis mellifera* L.) (Hymenoptera, Apidae) -Limit Test-
Source: IBACON, unpublished report No.: 6380036, September 21, 1999

Guidelines: EPPO No. 170
Deviations: temperature: 29 °C; relative humidity: 64 -70 % instead of 25 ° C ± 2 ° C and relative humidity of 60 -70 % as indicated in the guideline

GLP: yes (certified laboratory)

Material and methods: test substance: BNF 5119B, purity: 99.6%, batch number: 870922ELB06; under laboratory conditions, starved honey bees (*Apis mellifera*, 3 groups of 10 bees per dose) received a single oral dose of either 121.5, 11.3 or 1.2 µg per bee in 20 mg sugar solution. Subsequently, honey bees were observed over a period of 48 hrs for behavioural impairments and survival rate. The reference treatment (0.2 µg dimethoate per bee) caused a 100 % mortality (the facility-specific LD₅₀ dose for dimethoate is typically between 0.10 and 0.14 µg/bee).

Findings: Toxicity to Honey Bees, Laboratory Tests

Test substance	BNF 5119B
Test object	<i>Apis mellifera</i>
Application rates µg product/bee	121.5*, 11.3* and 1.2*
Exposure	oral (sugar solution)
LD ₅₀ µg a.i./bee (24 and 48h)	> 121.5

* values based on actual intake of the test substance

Observations: One of 30 bees died after application of an oral dose with 121.5 µg BNF 5119B per bee. No bee died after application of an oral dose with 11.3 µg BNF 5119B per bee and two bees died after application of 1.2 µg BNF 5119B per bee. No behavioural abnormalities were observed for the 48 hours of the experimental time.

No bee died in neither the acetone and water, nor in the pure syrup control. All bees died after treatment with Dimethoate.

2. Survey of the Study

2.1 General Information

Title: Laboratory Testing for Toxicity (Acute Oral LD₅₀) of BNF 5119B on Honey Bees (*Apis mellifera* L.) (Hymenoptera, Apidae) - Limit Test -

Sponsor: Bayer AG
Crop Protection Division
Institute for Environmental Biology
Alfred-Nobel-Str. 50
40789 Monheim
Germany

Monitoring: [REDACTED]

Test Substance: BNF 5119B

Test Facility: Institut für Biologische Analytik
und Consulting IBACON GmbH
Industriestrasse 1
64380 Rossdorf
Germany

IBACON Project: 6380036

Project Staff:

Test Facility Management: Dr. [REDACTED]

Study Director: Dipl. Biol. [REDACTED]

Technical Coordination: [REDACTED]

Head of Quality Assurance Unit (QAU): Dipl. Biol. [REDACTED]

Quality Assurance Unit Manager: Dipl. Biol. [REDACTED]

Schedule:

Study Initiation Date: June 18, 1999
Experimental Starting Date: July 7, 1999
Experimental Completion Date: July 9, 1999
Date of 1st Study Protocol Amendment: August 25, 1999
Draft Report Date: August 25, 1999
Study Completion Date: September 21, 1999

2.2 Good Laboratory Practice

This study was performed in compliance with:

- The OECD Principles of Good Laboratory Practice (as revised in 1997) and the
- Chemikaliengesetz ('*Chemicals Act*') der Bundesrepublik Deutschland (ChemG), Anhang I ('*Annex 1*'), 1994/97.

This study was assessed in compliance with the study protocol and the IBACON Standard Operating Procedures. This study and/or test facility were periodically inspected by the Quality Assurance Unit (QAU) and the dates and the phases of the inspections are included in this final report. The data contained within this final report were audited in comparison to the raw data.

A quality assurance statement, signed by the Quality Assurance Unit, is included in this final report.

2.3 Archiving

The following data / sample(s) will be archived

for 15 years:

- all raw data
- the study protocol
- the study protocol amendment
- one certified copy of the final report

for at least 2 years:

- one sample of the test substance and of the toxic standard

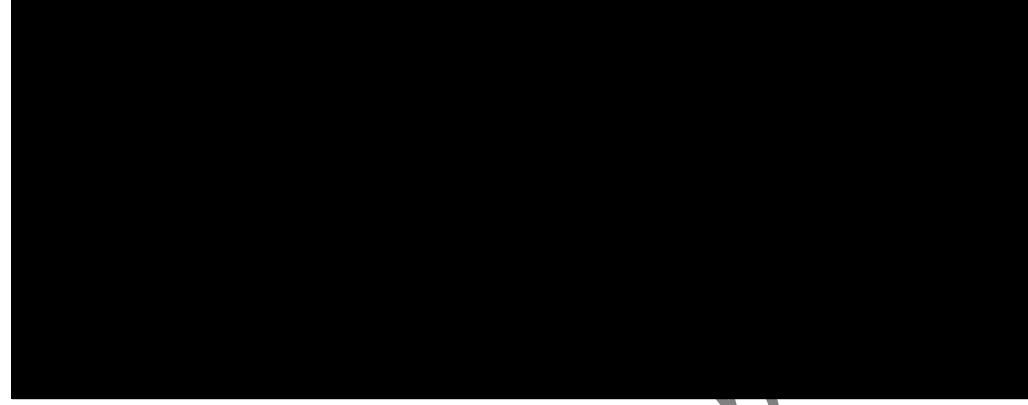
following the date on which the final report is audited by the Quality Assurance Unit at:

Institut für Biologische Analytik
und Consulting IBACON GmbH
Industriestrasse 1
64380 Rossdorf
Germany

No raw data or material relating to the study will be discarded without the sponsor's prior consent.

2.4 Signatures

Study Director:

Dipl. Biol. 

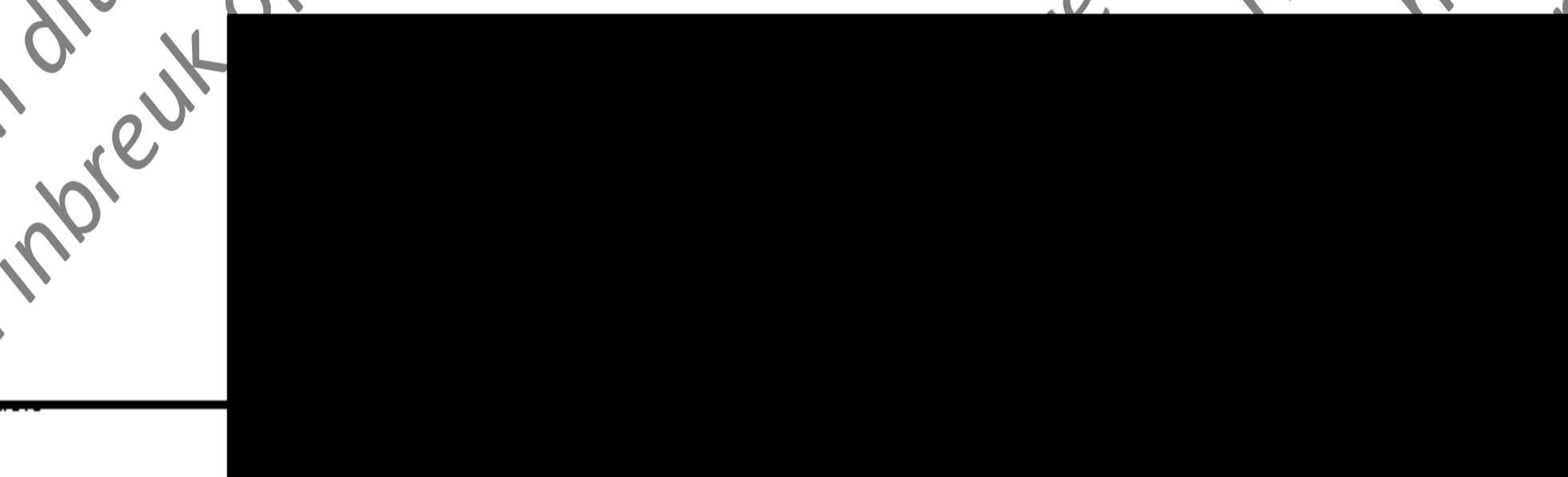


date:

September 21, 1999

Test Facility Management:

Dr. 

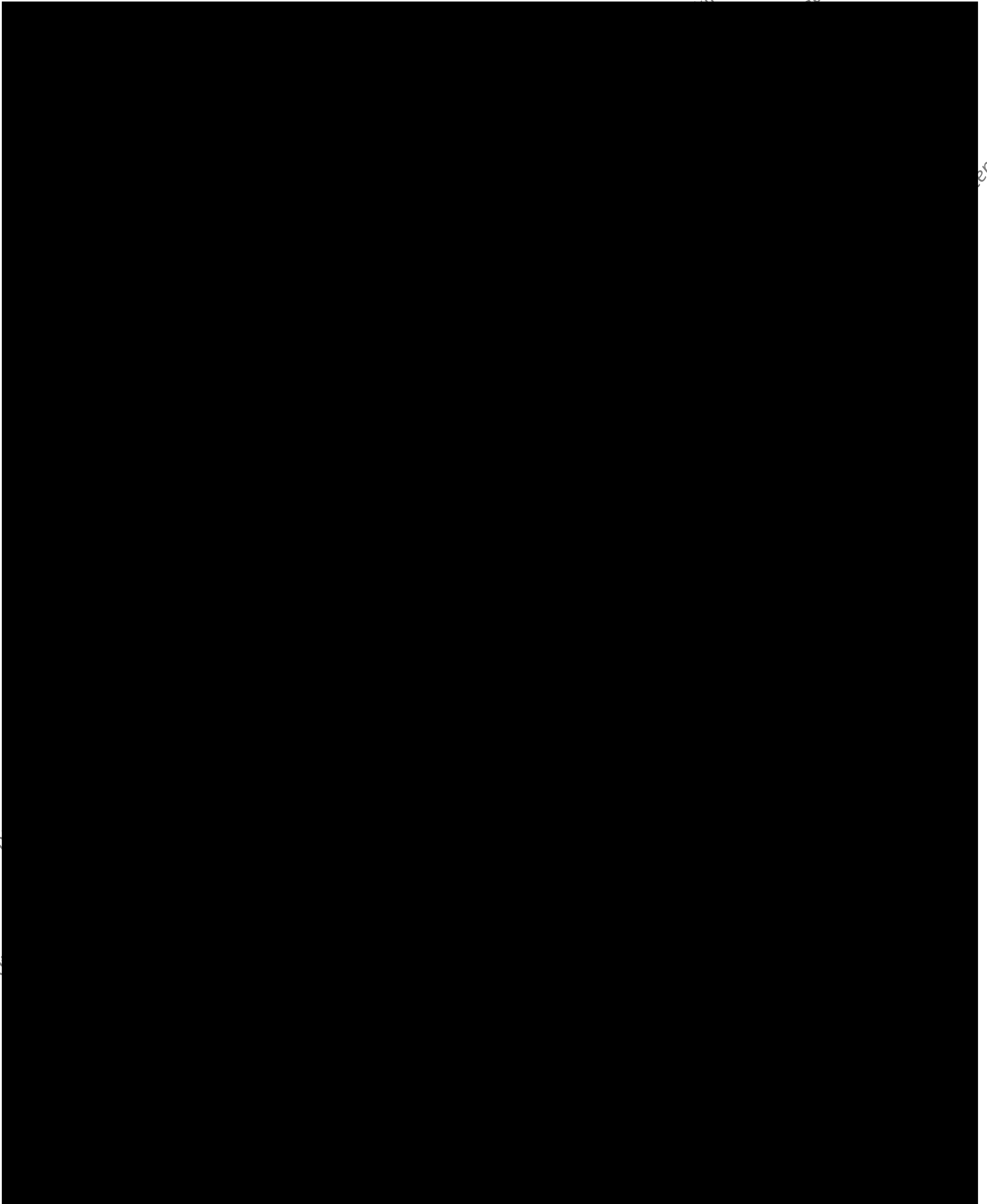


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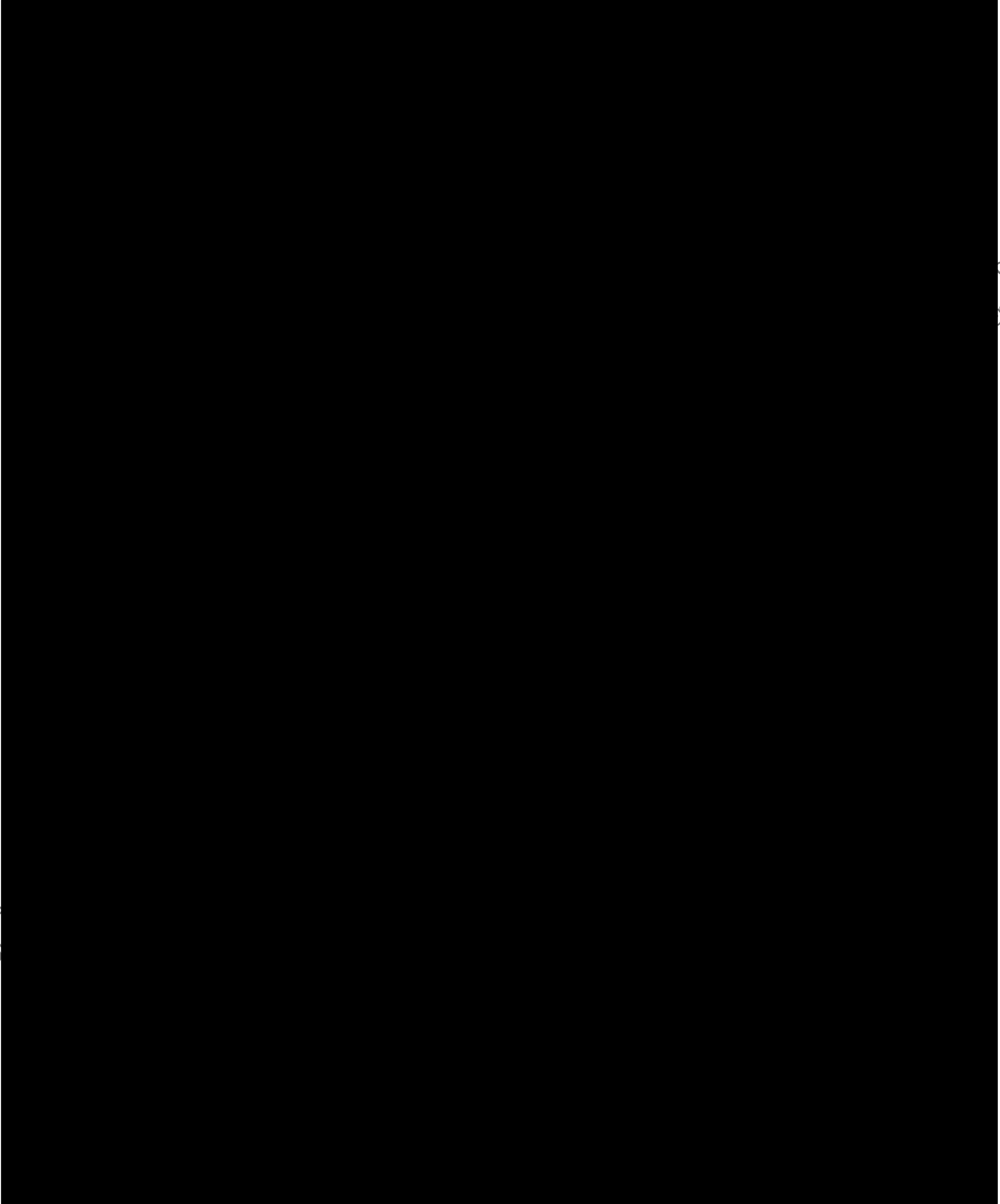
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3. Quality Assurance Unit Statement



4. Statement of Compliance



5. Objectives of the Study

5.1 Title

Laboratory Testing for Toxicity (Acute Oral LD₅₀) of BNF 5119B on Honey Bees (*Apis mellifera* L.) (Hymenoptera, Apidae) - Limit Test -

5.2 Purpose

If honey bees can be exposed to residues of BNF 5119B, this study is useful for the registration of the pesticides in question. This study provides:

- the acute toxicity levels of the test substance to honey bees;
- toxicity information comparable to expected residues from standard rates, for assessment of the potential hazard to honey bees;
- informational support for precautionary label statements.

5.3 Guidelines/Recommendations

This study was designed to comply with the following method:

- EPPO 1992: Guideline on test methods for evaluating the side-effects of plant protection products on honey bees, Bulletin OEPP/EPPO Bulletin 22, 203-215 1992, No. 170

5.4 Justification for the Selection of the Test System

This test system is required by the Registration Directive 91/414/EEC and/or by the 'SETAC - Guidance document on regulatory testing procedures for pesticides with non-target arthropods' (Barrett et al. 1994) for the hazard assessment of pesticides.

6. Materials and Methods

6.1 Test Substance, Control and Toxic Standard

Test Substance

The test substance and the information concerning the test substance were provided by the sponsor:

Name:	BNF 5119B
Batch No.: (Lot No.):	870922ELB06
Active Ingredient(s)/Purity:	99.6 % according to certificate of analysis
Certificate of Analysis Ref. Code / Date:	Analysis Date: August 8, 1995
Indication:	insecticide
Aggregate State at Room Temperature:	solid
Colour:	white (according to IBACON personnel)
Solubility:	in water: not indicated in acetone: not indicated
Stability:	pure: see expiry date in water: test substance must be considered as stable under test conditions in acetone: not indicated
Expiry Date:	August 2000
Storage:	in original container, at room temperature, in the dark

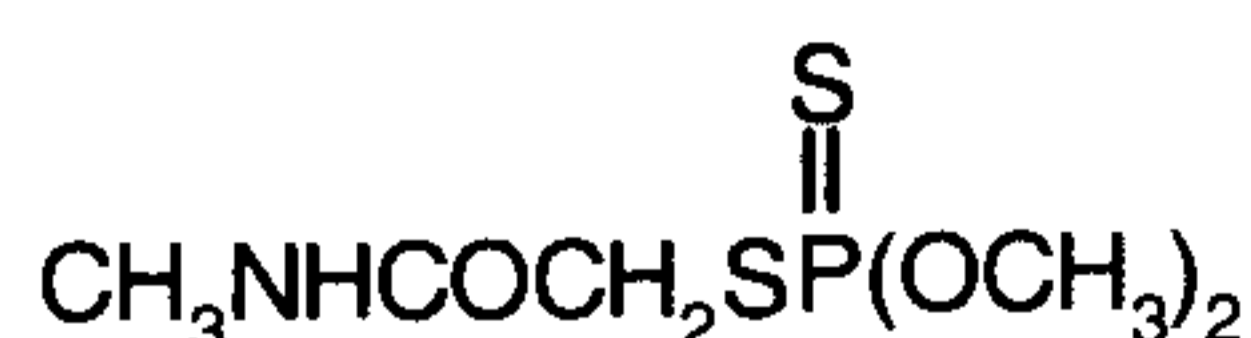
Controls

Controls:	1) pure syrup 2) acetone + syrup 3) tap water + syrup
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Toxic Standard

The information concerning the toxic standard according to the substance container label:

Name:	Perfekthion EC
Batch No.:	9801
Active Ingredient/Purity:	Dimethoate: 396 g/L
Chemical Structure of a.i.:	



Manufacturer:	BASF AG, Unternehmensbereich Pflanzenschutz, D-67056 Ludwigshafen
Expiry Date:	October/2000
Storage:	at room temperature, in the dark, in original container
Amount Applied in this Study:	0.2 µg active ingredient per bee

6.2 Test System

Taxonomic Group:	worker honey bees (Insecta, Hymenoptera)
Species:	adult <i>Apis mellifera carnica</i> L.
Age and Sex:	4 - 6 week old female bees
Origin:	honey bee colonies, disease-free and queen-right, bred by IBACON
Collection:	with glass tubes, from the flight board without anaesthetics
Bee Maintenance:	conformed to proper cultural practices

6.3 Test Units

Type:	stainless steel chambers
Size:	10 cm x 8.5 cm x 5.5 cm (length x width x height)
Front Side:	removable glass sheet
Bottom:	perforated with 98 ventilation holes; \varnothing 1 mm
Inner Walls:	lined with filter paper (Co. Macherey & Nagel, D-52355 Duren, Art. No. 68)
No. Of Individuals:	10 per test unit
Replicates:	3 per dosage/control
Identification:	test units were uniquely identified with study number, application date, treatment, concentration and replicate number
Bee Assignment:	assigned to test levels impartially

6.4 Test Conditions

Surrounding Type:	incubators
Temperature:	29 °C ¹
Relative Humidity:	64 - 70 %
Light:	darkness (except during observation)
Ventilation:	ventilation to avoid possible accumulation of pesticide vapour
Recording:	test conditions were recorded with suitable instruments and documented in the raw data

6.5 Food

Food:	commercial ready-to-use syrup (Apiinvert; 30 % Saccharose, 31 % Glucose, 39 % Fructose) <i>ad libitum</i>
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¹ in deviation from the guidelines which recommend a temperature of 25 ° C \pm 2 ° C and a relative humidity of 60 - 70 %. Experience of IBACON has shown that this deviation to the guideline will have no adverse effect on the study.

6.6 Application of the Test Substance, the Control and the Toxic Standard

The dosage applied was not adjusted to reflect the percentage a.i.

The procedure of preparation of the test substance solution is described in detail in the Attachment, page 21.

Application in the Oral Test:

ca. 20 mg of BNF 5119B contaminated food (for the 100 µg/bee nominal dose the test substance was distributed directly in the food, for the 10 and 1 µg/bee nominal dosages 1 part solvent (=water) including the test substance + 19 parts syrup were used) offered in syringes which were weighed before and after introduction into the cages (duration of uptake did not exceed 3 hours, except in the 10 µg/bee dosage group and in the control group with acetone as solvent; here, duration lasted up to 4 hours)

Dosages of the Test Substance in the Oral Test:

121.5, 11.3 and 1.2 µg/bee (values based on actual intake of the test substance)

Dosage of the Toxic Standard:

0.2 µg a.i. Dimethoate per bee

Controls:

- 1) BNF 5119B free sugar solution (used as the control for the 100µg/bee (nominal) dosage group)
- 2) BNF 5119B free sugar solution with acetone as solvent (used as the control for the 10µg/bee (nominal) dosage group)
- 3) BNF 5119B free sugar solution with water as solvent (used as the control for the 1µg/bee (nominal) dosage group)

6.7 Course of the Test

Treatment Groups:

controls, 3 dosages of test substance, toxic standard

Replicates:

3 per treatment group

Individuals:

10 per unit, 30 individuals per treatment group

Starvation Time:

60 minutes

Exposure Time:

48 hours

6.8 Test Parameters

Mortality:

number of dead bees after 60 minutes; 2, 4 hours (first day); 24 and 48 hours

Behavioural Abnormalities:

behavioural abnormalities (vomiting, apathy, intensive cleaning, discoordinated movements) after 60 minutes; 2, 4 hours (first day); 24 and 48 hours

6.9 Result Evaluation

Mortality:

results obtained from the bees treated with test substance are compared to those obtained from the toxic standard and the controls. Due to the results it was not necessary to conduct statistical analysis.

6.10 Validity Criteria of the Study

Control Mortality: no control mortality at experimental end (48 hours)
Toxic Standard Mortality: resulted in 100 % mortality

6.11 Deviations to the Study Protocol

Concerning: duration of uptake of the test substance
According to the Study Protocol: duration of uptake of the test substance should not exceed 3 hours
Deviation to the Study Protocol: uptake of test substance in the 10 µg/bee and in the acetone control took 4 hours and not 3 hours as indicated in the protocol
Reason for the Deviation: because the bees needed more than the expected 3 hours for the uptake of the contaminated food
Presumed Effect on the Study: none, because bees ingested enough amount of treated food and similar intake of solution in treated and control groups

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7. Results and Discussion

7.1 Oral Toxicity Test

After a starving period of 60 minutes, mortality occurred after ingestion of 121.5 (3.3 %) and 1.2 (6.7 %) μg BNF 5119B per bee. No mortality occurred after application of 11.3 μg BNF 5119B. Behavioural abnormalities were not observed (Table 1, Appendix Table 2 - 4).

Due to the results of this study (no mortality > 50 % in the highest dosage group) the LD_{50} must be > 121.5 $\mu\text{g}/\text{bee}$.

Three different controls were used for this test:

The pure syrup control was used for the 121.5 $\mu\text{g}/\text{bee}$ group, because the test substance was mixed directly in the syrup. In the 11.3 and 1.2 $\mu\text{g}/\text{bee}$ dosage groups, acetone and water was used as solvent, respectively.

No mortality occurred in the control groups within the whole experiment.

100 % mortality occurred after ingestion of 0.2 μg Dimethoate per bee in the toxic standard group.

Table 1. Mortality^a and behavioural abnormalities^a of the bees in the oral toxicity test^b

ingested test substance $\mu\text{g}/\text{bee}$	after 1 hour		after 4 hours		after 24 hours		after 48 hours	
	mortality	behav. abnorm.	mortality	behav. abnorm.	mortality	behav. abnorm.	mortality	behav. abnorm.
	mean %	mean %	mean %	mean %	mean %	mean %	mean %	mean %
121.5	3.3	0.0	3.3	0.0	3.3	0.0	3.3	0.0
11.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
1.2	6.7	0.0	6.7	0.0	6.7	0.0	6.7	0.0
syrup	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
acetone	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
water	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
toxic st.	0.0	0.0	96.7	3.3	100.0	0.0	100.0	0.0

^a results are averages from three replicates (ten bees each) per dosage/control

^b see Appendix for details; behav. abnorm = behavioural abnormalities

acetone = control with acetone; water = control with water

syrup = pure syrup control; toxic st. = toxic standard

7.2 Conclusions

Mortality: Oral LD₅₀ (24 and 48h) of BNF 5119B: > 121.5 µg/bee
Behavioural Abnormalities: No behavioural abnormalities occurred in the test substance treatment groups.

8. References

Barrett K.L., Grandy N., Harrison E.G., Hassan S.A. & Oomen P. 1994: SETAC Guidance document on regulatory testing procedures for pesticides with non-target arthropods. 28-30 March 1994, IAC Wageningen, The Netherlands

Chemikaliengesetz der Bundesrepublik Deutschland (ChemG). Anhang 1, in der Fassung der Bekanntmachung vom 25. Juli 1994 (BGBl. I S. 1703) mit Änderungen vom 27. September 1994 (BGBl. I S. 2705) und 14. Mai 1997 (BGBl. I S. 1060)

EC Agrochemical Registration Directive (DS65) (Directive 91/414/EEC). ANNEX III. Requirements for the dossier to be submitted for the authorization of a plant production product

EPPO 1992: Guideline on test methods for evaluating the side-effects of plant protection products on honey bees, Bulletin OEPP/EPPO Bulletin 22, 203-215 1992, No. 170

OECD Principles of Good Laboratory Practice, adopted by Council on 26th November 1997 [C(97)186/Final], Environment Directorate, Organisation for Economic Co-operation and Development, Paris 1998

9. Distribution of the Final Report

Sponsor: 1x (the original final report)

IBACON: 1x (one certified copy of the original final report)

Appendix

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Table 2. (Exact Data). Definitive oral toxicity test; mortality and behavioural abnormalities of the bees

test substance		after 1 hour			after 2 hours			after 4 hours			after 24 hours			after 48 hours		
unit #	dosage µg/bee	dead #	beh. abnor. #	symp.	dead #	beh. abnor. #	symp.	dead #	beh. abnor. #	symp.	dead #	beh. abnor. #	symp.	dead #	beh. abnor. #	symp.
1	122.0	0	0		0	0		0	0		0	0		0	0	
2	120.5	1	0		1	0		1	0		1	0		1	0	
3	122.0	0	0		0	0		0	0		0	0		0	0	
1	12.3	0	0		0	0		0	0		0	0		0	0	
2	9.8	0	0		0	0		0	0		0	0		0	0	
3	11.8	0	0		0	0		0	0		0	0		0	0	
1	1.2	0	0		0	0		0	0		0	0		0	0	
2	1.2	2	0		2	0		2	0		2	0		2	0	
3	1.2	0	0		0	0		0	0		0	0		0	0	
syrup																
1	control	0	0		0	0		0	0		0	0		0	0	
2	control	0	0		0	0		0	0		0	0		0	0	
3	control	0	0		0	0		0	0		0	0		0	0	
acetone																
1	control	0	0		0	0		0	0		0	0		0	0	
2	control	0	0		0	0		0	0		0	0		0	0	
3	control	0	0		0	0		0	0		0	0		0	0	
water																
1	control	0	0		0	0		0	0		0	0		0	0	
2	control	0	0		0	0		0	0		0	0		0	0	
3	control	0	0		0	0		0	0		0	0		0	0	
toxic																
1	standard	0	0		0	8	c	9	f	e	10	0		10	0	
2	standard	0	0		1	6	b,c	10	0		10	0		10	0	
3	standard	0	0		0	10	b,c	10	0		10	0		10	0	

= number of individuals; beh. abnor. = behavioural abnormalities
 symp. = observed symptoms according to the following key: a = food refusal/vomiting; b = moving coordination problems;
 c = apathy; d = intensive cleaning; e = nervous; f = sitting in one corner of the chamber

Table 3. (Relative Data). Definitive oral toxicity test; mortality and behavioural abnormalities of the bees

test substance dosage µg/bee	after 1 hour				after 4 hours				after 24 hours				after 48 hours			
	mortality		beh.abnor. ^a		mortality		beh.abnor. ^a		mortality		beh.abnor. ^a		mortality		beh.abnor. ^a	
	mean	±SD	mean	±SD	mean	±SD	mean	±SD	mean	±SD	mean	±SD	mean	±SD	mean	±SD
	%	%	%	%	%	%	%	%	%	%	%	%	%	%	%	%
121.5	3.3	5.8	0.0	0.0	3.3	5.8	0.0	0.0	3.3	5.8	0.0	0.0	3.3	5.8	0.0	0.0
11.3	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
1.2	6.7	11.5	0.0	0.0	6.7	11.5	0.0	0.0	6.7	11.5	0.0	0.0	6.7	11.5	0.0	0.0
syrup	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
acetone	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
water	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
toxic st.	0.0	0.0	0.0	0.0	96.7	5.8	3.3	5.8	100.0	0.0	0.0	0.0	100.0	0.0	0.0	0.0

^abeh. abnor. = behavioural abnormalities; mean = mean of three replicates; ±SD = standard deviation from three replications
acetone = control with acetone; water = control with water; syrup = pure syrup control; toxic st. = toxic standard

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Table 4. Definitive oral intake

substance dosage nominal	substance concentrations		weight of syringes			ingested solution	ingested substance	
	in food µg / mg	start ^a mg	end ^b mg	difference mg	mg / bee	µg/bee	µg/bee	Ø ^c µg/bee
100µg/bee								
1	5	11158	10914	244	24	122.0		
2	5	11146	10905	241	24	120.5	121.5	
3	5	11146	10902	244	24	122.0		
10µg/bee								
1	0.5	11220	10974	246	25	12.3		
2	0.5	11219	11024	195	20	9.8	11.3	
3	0.5	11230	10994	236	24	11.8		
1µg/bee								
1	0.05	11160	10913	247	25	1.2		
2	0.05	11148	10902	246	25	1.2	1.2	
3	0.05	11176	10934	242	24	1.2		
syrup control								
1	0	11177	10944	233	23	0.0		
2	0	11161	10899	262	26	0.0	0.0	
3	0	11162	10891	271	27	0.0		0.0
acetone control								
1	0	11095	10863	232	23	0.0		
2	0	11146	10894	252	25	0.0	0.0	0.0
3	0	11242	11004	238	24	0.0		
water control								
1	0	11125	10870	255	26	0.0		
2	0	11186	10942	244	24	0.0	0.0	0.0
3	0	11180	10937	243	24	0.0		
toxic standard								
Dimethoate								
0.2 µg/bee								
1	0.01	11084	10844	240	24	0.2		
2	0.01	11197	10957	240	24	0.2	0.2	
3	0.01	11128	10889	239	24	0.2		

^a weight of syringes at the start of the experiment, ^b after removing from the test cages;

^c ingested solution as calculated average, ^d results are rounded results, calculated from the exact data

Attachment: Preparation of the Test Substance Solutions

Oral Test

100 µg/bee dosage (nominal):

20 mg of the test substance was mixed with 4 g syrup and treated with ultra sonic for 45 minutes and overnight stored in a refrigerator until the application.

Control: pure syrup.

10 µg/bee dosage (nominal):

25 mg of the test substance was dissolved ad 2.5 g acetone (stock solution). The test substance was completely dissolved in water. The stock solution was clearly visible.

500 mg of the stock solution was added to 9.5 g syrup (1:20).

Control: 500 mg acetone and 9.5 g syrup.

1 µg/bee dosage (nominal):

50 mg of the test substance was dissolved ad 50 g water (stock solution) overnight stored in a refrigerator until the application. The test substance was completely dissolved in water. The stock solution was clearly visible.

500 mg of the stock solution was added to 9.5 g syrup (1:20).

Control: 500 mg water and 9.5 g syrup.

100, 10 and 1 µg/bee should be obtained when 20 mg of the contaminated sugar solution per bee were administered.

121.5, 11.3 and 1.2 µg/bee were obtained, because the bees ingested between 20 and 25 mg contaminated food per bee.