

TITLE PAGE

Residues of Imidacloprid and Imidacloprid Metabolites in Nectar, Blossoms, Pollen and Honey Bees Sampled from a Summer Rape Field in Sweden and Effects of These Residues on Foraging Honeybees

AUTHOR



TESTING FACILITY

BAYER AG

Group Protection-Development
Institute For Environmental Biology
D-51368 Leverkusen-Bayerwerk

GLP STUDY NUMBER

E3701360-0

REPORT NUMBER

SXR/Am 002

TOTAL NUMBER OF PAGES

28

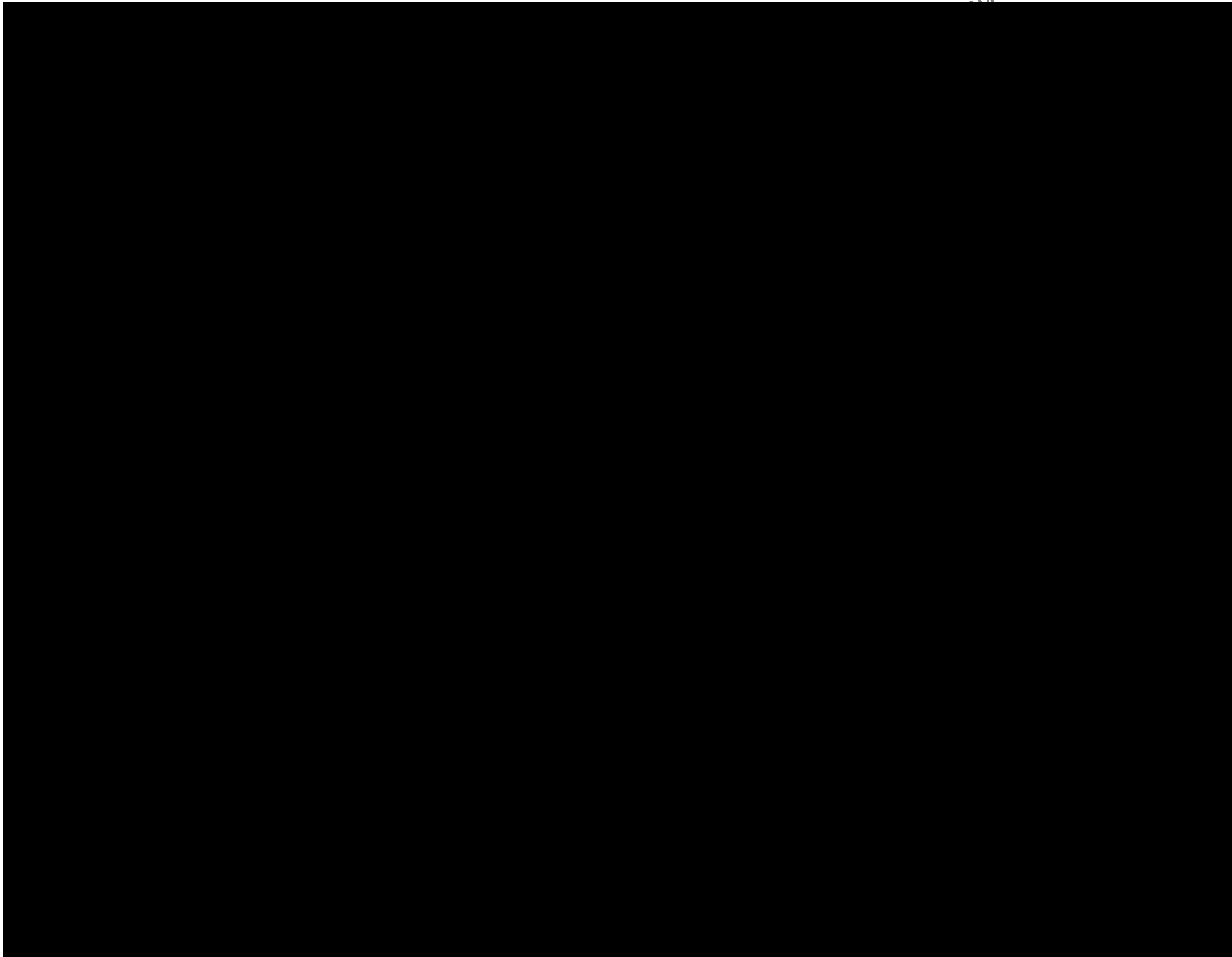
STUDY COMPLETION DATE:

21 Mai 1999



SXR/AM 002 / MO-99-008702

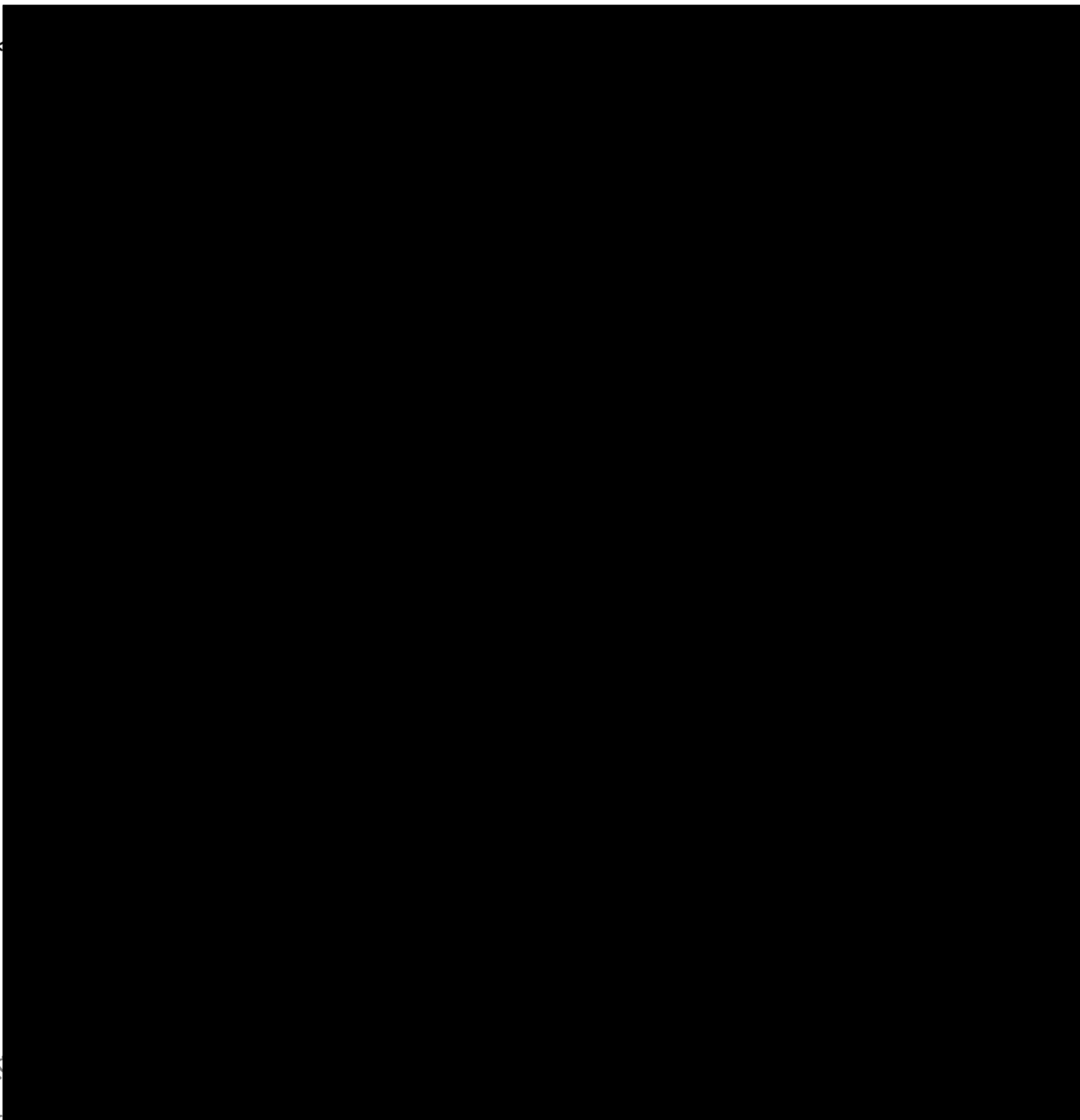
STATEMENT OF COMPLIANCE



Dit document is geen eigendom van het Ctgb
Op dit document kunnen rechten van
Voorts kan dit document
Publicatie, verspreiding, vermenigvuldiging, commerciële
rechthebbende van dit document kan derhalve
This document is not the property of the Ctgb and only pro
The document may be subject to rights such as in
Furthermore, this document may fall unde
Consequently, any publication, distribution, reproduction and/or publishing and
without the permission of the owner of this document may theref

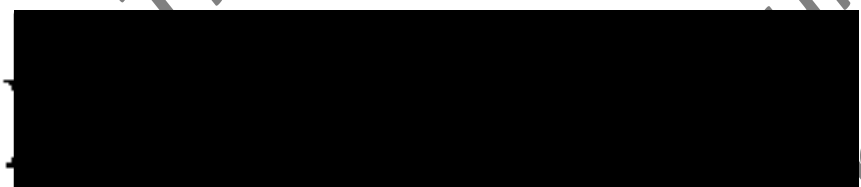
ment or its contents
owner.

CERTIFICATION OF AUTHENTICITY



INQUIRIES

Inquiries should be directed to:


BAYER AG
Crop Protection - Development
Institute for Environmental Biology
D-51368 Leverkusen - Bayerwerk, FRG

Phone No. (02173) / 38 - 3888

Dit document is geen eigendom van de afzender.
Op dit document is geen aanspraak te maken.
Publicatie, verspreiding, vermeerdering of anderszins openbaar maken van dit document is niet toegestaan.
This document is not the property of the sender.
The document may be used for internal purposes only.
Furthermore, the distribution, reproduction, or public use of this document is prohibited without the permission of the owner of the document.

document or its contents
its owner.

TABLE OF CONTENTS

	Page No.
TITLE PAGE.....	1
STATEMENT OF COMPLIANCE.....	2
CERTIFICATION OF AUTHENTICITY.....	3
TABLE OF CONTENTS.....	4
1.0 SUMMARY.....	5
2.0 INTRODUCTION.....	6
3.0 EXPERIMENTAL.....	6
3.1 Test Substance.....	6
3.2 Reference Substance.....	6
3.3 Execution of the Test.....	7
3.4 Origin of Honeybees.....	7
3.5 Location of the Trial Site and Description of the Study Plots.....	7
3.6 Drilling of the Rape Seed and Calibration of the Seed Machinery.....	7
3.7 Cultivation of the Plots.....	7
3.8 Sampling Procedure.....	8
3.9 Sample Processing and Residue Analysis.....	9
3.10 Climatic Conditions During the Study.....	9
3.11 Observations on Honeybees.....	9
4.0 FILING.....	9
5.0 RESULTS AND DISCUSSION.....	10
5.1 Analytical Findings.....	10
5.2 Observations on Foraging Honeybees.....	10
FIGURES	
Figure 1: Arrangement of the study plots on the study field.....	11
TABLES	
Table 1: Summary of the Analytical Findings.....	12
Table 2: Records on Flight and Foraging Activity of Honeybees During the Sampling Period.....	13
APPENDICES	
Appendix I: Analytical Report.....	14
Appendix II: Copy of the GLP Certificate.....	26
Appendix III: Quality Assurance Statement.....	28

1.0 SUMMARY

Report: [REDACTED] (1999): Residues of Imidacloprid and Imidacloprid Metabolites in Nectar, Blossoms, Pollen and Honey Bees Sampled from a Summer Rape Field in Sweden and Effects of These Residues on Foraging Honeybees
Bayer AG, unpublished report No: SXR/Am 002; 1999/05/21.
(Appendix I contains data from study RA-2043/98)

Guidelines: Internal Testing Method
Deviations: not applicable

GLP: yes (certified laboratory)

Material and methods: Poncho FS 500, a.i. content: 78.3 g/L Beta-Cyfluthrin & 428.2 g/L Imidacloprid; specification (formulation No.: 030 based on 06200/0029, developmental No.: 00195939); under field conditions small beehives (appr. 5,000 honeybees) were caged on flowering summer rape plots (drilling rate: 5 kg/ha) as a sampling device for rape nectar and rape pollen. Nectar was also directly sampled from flowers via micropipettes. In addition, flowers were sampled by hand. The honeybees used as samplers were observed for signs of behavioral impacts. All samples including the honeybees were subjected to a residue analysis for imidacloprid and its relevant metabolites.

Dates of biological work: July 2 - 6, 1998

Dates of analytical work: July 9 - 29, 1998

Findings: Residues in rape plant matrices and in the foraging honeybees

Type of Sample	Residue Level [mg/kg] *		
	Imidacloprid	Olefin-NTN	Hydroxy-NTN
<i>Control Samples</i>			
Honeybees before exposure	< 0.01	< 0.01	< 0.01
Honeybees after exposure	< 0.01	< 0.01	< 0.01
Rape nectar sampled by bees	< 0.01	< 0.01	< 0.01
Rape nectar sampled with micro-capillaries from the flowers	< 0.01	< 0.01	< 0.01
Rape blossoms	< 0.01	< 0.01	< 0.01
Rape pollen sampled by bees **	--	--	--
<i>Treatment Samples</i>			
Honeybees before exposure	< 0.01	< 0.01	< 0.01
Honeybees after exposure	< 0.01	< 0.01	< 0.01
Rape nectar sampled by bees	< 0.01	< 0.01	< 0.01
Rape nectar sampled with micro-capillaries from the flowers	< 0.01	< 0.01	< 0.01
Rape blossoms	< 0.01	< 0.01	< 0.01
Rape pollen sampled by bees **	--	--	--

* Limit of quantitation: 0.01 mg/kg. ** Amount insufficient for residue analysis

Observations: No behavioral impacts (e.g. apathy, exaggerated motility, discoordinated movements) or suspicious mortality was observed on the honeybees used for collecting rape nectar and rape pollen. At the time of sampling, aphids were observed on the rape plants.

2.0 INTRODUCTION

According to EU directive 91/414/EEG the impacts of pesticides on honeybees have to be examined. Besides the intrinsic toxicity of a pesticide the concentration to which a honeybee may be exposed under field conditions is an integral component for the hazard assessment. The present study aims to examine the exposure in greater detail for a refined risk assessment.

The rape samples were analysed for residues of imidacloprid and its olefin- and hydroxy metabolites. These metabolites were considered as relevant, since they have a chemical structure closely related to the parent molecule and were observed in plant metabolism studies in significant proportions (up to approx. 10 %).

3.0 EXPERIMENTAL

3.1 Test Substance

Test Substance:	Poncho FS 500
Active Ingredient(s):	(a) Beta-Cyfluthrin (FCR 4545) (b) Imidacloprid (NTN 33893)
Chemical Name(s) of AI(s):	(a) CYCLOPROPANECARBOXYLIC ACID, 3-(2,2-DICHLOROETHENYL)-2,2-DIMETHYL-,CYANO(4-FLUORO-3-PHENOXYPHENYL)METHYL ESTER (b) 1-(6-CHLORO-3-PYRIDINYL)-METHYL-N-NITRO-2-IMIDAZOLIDINIMINE
CAS Number of AI(s):	(a) 68359 - 37 - 5 (b) 138261 - 40 - 3
Indikation:	Seed dressing
Product Number	0195939
Formulation/Batch Number:	FL 0030 based on form no. 06200/0029
No. of Certificate:	FAR-No. 446-01
AI Content (acc. to Analysis):	(a) 78.3 g/l (b) 428.2 g/l
Analytical Method:	(a) GLC, int. Std. (b) HPLC, ext. Std.
Date of Analysis:	February 4, 1998
Expiry Date:	February 4, 1999
Physical Appearance:	dark blue suspension
Specific Density:	1.151 g/ml
Storage Conditions:	Room temperature
Seed Dressing Rate(s) Tested in the Study:	2.5 l Poncho 500 FS per 100 kg oilseed rape (= 1050 g/dt Imidacloprid & 200 g/dt beta-Cyfluthrin) (analytical findings*: 1026 g/dt Imidacloprid).
Seed Drilling Rate Tested in the Study:	5 kg seed per ha (68 g per 136 m ² plot) (seed variety: „Maskot“; summer rape)
Safety Precaution:	Routine hygienic precautions

3.2 Reference Substance

For this type of material and use pattern, a reference compound is not specified.

* Dressed seeds were analysed for imidacloprid only.

3.3 Execution of the Test

The sampled study plots were drilled on April 28, 1998 (reserve plots drilled on May 13, 1998 but these plots were not used in this study). Sampling of nectar, flowers and honeybees and the behavioral observations were performed between July 3 and 6, 1998.

Sponsor:

BAYER AG
GB Plant Protection
Marketing - Seed Treatment (Dr. Krohn/Altmann)
D-40789 Monheim

Study Director:

Cultivar Manager:

Trials Officer:

Responsible Analyst:

Study Technicians:

Quality Assurance:

Laboratory Study Number:

SXR/Am 002

3.4 Origin of Honeybees

The honeybees used for pollen and nectar sampling were supplied by a commercial Swedish beekeeper [REDACTED]

The beehives used for the test were transported to the study site in the evening of July 1 and returned to the original place on July 7, 1998.

3.5 Location of the Trial Site and Description of the Study Plots

The trial site was located in the vicinity of Borunda-Skelinge, South of Eslov in Sweden. Before summer rape, the field was cultivated with sugarbeets in 1997.

There were two rape planted plots on the trial site (Fig. 1) which were separated by 2 m wide buffer strips. Each plot had a size of 8x17 m with a between-row distance of 12.5 cm. Each plot was staked out with marking sticks prior to the beginning of the experimental part.

3.6 Drilling of the Rape Seed and Calibration of the Seed Machinery

The control plot was drilled with 5 kg/ha (= 68 g per 136 m² plot) untreated rape seed whereas the treatment plot received 5 kg/ha rape seed coated with 2.5 l/dt Poncho FS 500.

Prior to sowing the proper functioning of the equipment was tested. The machinery control prior drilling showed a delivery rate between 4.2 and 4.3 g rape seed per application pipe (n = 16) over a 17 m drilling distance, i.e. the total seed delivery rate per plot was 16 x 4.2/4.3 g = 67.2/68.8 g per plot.

3.7 Cultivation of the Plots

Treated and untreated plots were cultivated in the same way according to the practice of the region. Before initiation of sampling, no protection treatments other than the seed treatment was necessary.

3.8 Sampling Procedure

Installment of bee hives

At the time of full rape blossom, tents of 4x4 m and 2 m height were installed on the control and the treatment plot (see Fig. 1). The tents consisted of an aluminium frame covered by gauze material (2x2 mm mesh size). For handling purposes, a walkway was created by removing all plants along a 50 cm transect between the tent entrance and the opposite end. One bee hive was placed at the end of the walkway opposite to the entrance in the treatment plot. The day after installment, hive entrances were disclosed and honeybees were allowed to forage on the study plot within the tent area. Before placing the beehives on the plots, appr. 100-200 honeybees were sampled to get blank samples of honeybees and honeybulbs for the residue work. The sampled honeybees were processed as described in the subsequent chapter.

Sampling of Nectar from the Honeybulbs

On the day after hive installment and the following two days a total of about 200 honeybees were sampled with tweezers directly from rape flowers after watching them for feeding over about 10-30 seconds. All sampled bees were killed by freezing (dry ice). Dead bees were stored on dry ice in the field and, at the end of each sampling day at the latest, transferred to a refrigerator (-19 to -20°C). At the end of the study the samples were shipped to Monheim on dry ice and further retained at -20°C until preparation of the honeybulbs. Honeybulbs were prepared by cutting the frozen bees into halves between the thorax and the abdomen and removing the filled honeybulbs by tweezers. All honeybulbs from one treatment group were pooled within an Eppendorf cap which was stored on dry ice. After all honeybees of a respective treatment were prepared (at the end of each preparation day at the latest), sampled honeybulbs were stored in a refrigerator at -20°C until residue analysis (see 3.9).

Sampling of Pollen from the Honeybees

From the prepared bees, pollen pockets were removed and stored in a refrigerator at -20°C until residue analysis (see 3.9). Unfortunately, not sufficient pollen could be obtained for residue analysis.

Sampling of Nectar from the Rape Flowers

The day before sampling, between 10 and 20 flowering rape plants outside the caged area were covered by plastic bags to prevent insects from foraging on that flowers. On the next day, nectar was directly drawn from that flowers by 5 µl micropipettes. After sampling, the micropipettes were emptied into a 1.5 ml Eppendorf tube which was stored on dry ice in the field. At the end of each sampling day at the latest, these samples were transferred into a refrigerator (-20°C) where they were retained until residue analysis (see 3.9).

As a check of possible contamination of the sampled material (from e.g. soil particles, dust), sampling people had to fill a second Eppendorf tube with uncontaminated tap water during sampling activities. Filling of this checking tube was done in several steps (5-10 pipetting events) from a water storage bottle. The analysis of these check samples revealed no residues indicating that the sampling people did not transfer inadvertently any residues from outside the flower into the nectar samples (see analytical report, page 25).

Sampling of Rape Flowers

About 10 g of rape flowers were sampled from plants outside the tent area. After sampling, the flowers were stored on dry ice in the field. At the end of each sampling day

at the latest, they were transferred into a refrigerator (-20°C) where they were retained until residue analysis (see 3.9).

Soil Samples

Some 200 g soil were sampled from the study field to characterize the soil at the study site. The soil at the study field can be classified as „sandy loam“. The organic carbon content was 1.8 % by weight. The water holding capacity was determined to be 63.1 g H₂O/100 g dry soil. The pH - value (1 KCL) was determined to be 6.0.

3.9 Sample Processing and Residue Analysis

Sample processing and analytical methods are described in detail in appendix I.

3.10 Climatic Conditions During the Study

During the study, temperature and precipitation events were recorded one to three times per day. The following records were made:

Date	Precipitation [mm]	Min. temperature [°C]	Max. temperature [°C]	Remarks
1 July	Not recorded	14	16	partly cloudy, rainy
2 July	0	13	24	stormy, cloudy
3 July	0	12	19	stormy, cloudy
4 July	0	12	14	cloudy, rain started
5 July	3	15	15	stormy, cloudy, rain
6 July	2	12	15	stormy, cloudy, rain

3.11 Observations on Honeybees

All behavioral anomalies of the honeybees were recorded together with the date of observation. In particular, the following behavioural aspects were observed:

Flight intensity:	Once per day, over a period of 5 minutes, the number of bees leaving the hive and returning to the hive was recorded.
Foraging intensity:	Once per day the number of bees foraging within a haphazardly assigned area of 1 m ² of flowering rape within the tent was recorded.
Behavioral anomalies:	Whenever observed, the following behavioral anomalies were recorded with the date and daytime of observation: - exaggerated motility - discoordinated movements (trembling, shaking, apathy)
Mortality:	Any suspicious numbers of dead bees in comparison to the controls during and after the test were recorded but no formal counts were made.

4.0 FILING

All raw data, the study protocol and the original of the report are filed in the Central GLP archive of PF/F, Crop Protection Center 40789 Monheim, FRG. Reserve samples of the test substance are stored in the pertinent archive of that test facility which provided or certified the test substance.

5.0 RESULTS AND DISCUSSION

5.1 Analytical Findings

Analytical findings are summarized in table 1 and given in detail in the analytical report (appendix I). No residues at or above the limit of quantitation were found in any of the examined matrices for either the parent compound or the relevant metabolites (olefin- and hydroxy-imidacloprid).

5.2 Observations on Foraging Honeybees

No behavioral impacts (e.g. apathy, exaggerated motility, disordinated movements) or suspicious mortality was observed on the honeybees used for collecting rape nectar and rape pollen. Due to the poor weather conditions prevailing during the sampling period, flight and foraging intensity of the honeybees was rather low. For this reason, the records made for these testing endpoints do not allow detailed conclusions (Table 2) other than that no marked differences between the control and the treatment plot were evident.

Dit document is geen eigendom van het Ctgb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot openbaarmaking.
Op dit document kunnen rechten van derden rusten, waaronder intellectuele eigendomsrechten of auteursrechten.
Voorts kan dit document onder een regeling omtrent gegevensbescherming vallen.
Publicatie, verspreiding, vermenigvuldiging, commerciële exploitatie en gebruik van dit document of de inhoud hiervan zonder de toestemming van de rechthebbende van dit document kan derhalve verboden zijn en een inbreuk opleveren van de rechten van de rechthebbende.
This document is not the property of the Ctgb and only provided based on mandatory freedom of information requirements.
The document may be subject to rights such as intellectual property and copy rights of third parties.
Furthermore, this document may fall under a regulatory data protection and any commercial exploitation and use of this document or its contents without the permission of the owner of this document may therefore be prohibited and violate the rights of its owner.

FIGURES

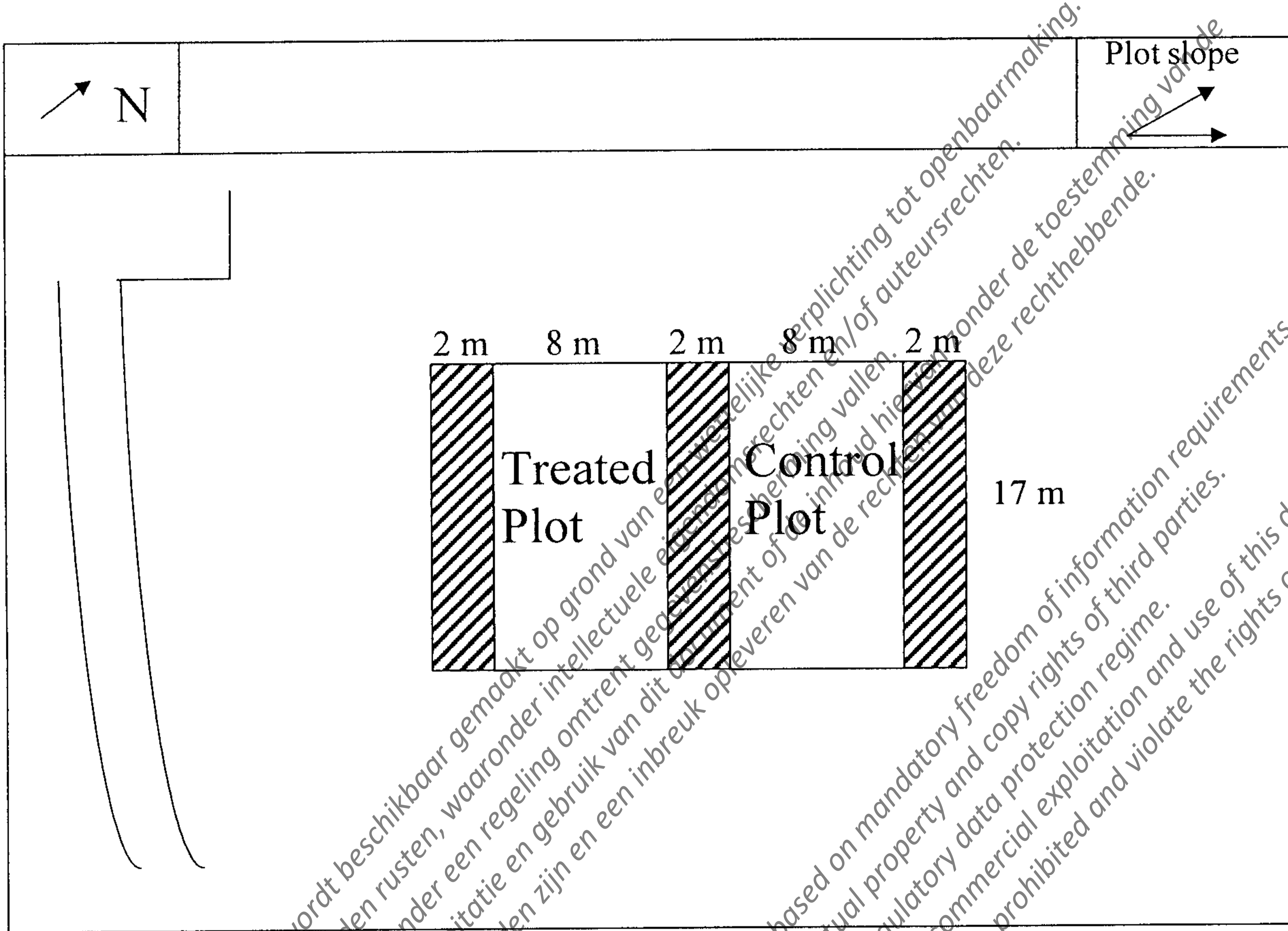


Figure 1: Arrangement of the study plots on the study field.

There were two plots of rape plants separated from each other by 2 m buffer strips. Each plot had a size of 8x17 m with a distance of 12.5 cm between rows. Plot management is reported in detail in the raw data to study RA-2043/98.

Dit document is geen eigendom van het Ctgb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot openbaarmaking. Op dit document kunnen rechten van derden rusten, waaronder intellectuele eigendomsrechten en/of auteursrechten. Voorts kan dit document onder een regeling omtrent intellectuele eigendomsrechten vallen. Het gebruik van dit document is onder de toestemming van de rechthebbende van dit document voor de afgeleide verwerking verboden. Dit document is niet de eigendom van de Ctgb en wordt uitsluitend als intellectuele eigendom van de rechthebbende van dit document ter beschikking gesteld. Het gebruik van dit document is onder de toestemming van de rechthebbende van dit document voor de afgeleide verwerking verboden. Consequently, any publication, distribution, reproduction and/or publishing and any commercial exploitation and use of this document or its contents without the permission of the owner of this document may therefore be prohibited and violate the rights of its owner.

TABLES

Table 1: Summary of the Analytical Findings.

Type of Sample	Residue Level [mg/kg] *		
	Imidacloprid	Olefin-NTN	Hydroxy-NTN
<i>Control Samples</i>			
Honeybees before exposure	< 0.01	< 0.01	< 0.01
Honeybees after exposure	< 0.01	< 0.01	< 0.01
Rape nectar sampled by bees	< 0.01	< 0.01	< 0.01
Rape nectar sampled with micro-capillaries from the flowers	< 0.01	< 0.01	< 0.01
Rape blossoms	< 0.01	< 0.01	< 0.01
Rape pollen sampled by bees **	--	--	--
<i>Treatment Samples</i>			
Honeybees before exposure	< 0.01	< 0.01	< 0.01
Honeybees after exposure	< 0.01	< 0.01	< 0.01
Rape nectar sampled by bees	< 0.01	< 0.01	< 0.01
Rape nectar sampled with micro-capillaries from the flowers	< 0.01	< 0.01	< 0.01
Rape blossoms	< 0.01	< 0.01	< 0.01
Rape pollen sampled by bees **	--	--	--

* Limit of quantitation: 0.01 mg/kg

** Amount harvested from dissected bees was not sufficient for residue analysis

Dit document is geen eigendom van het Ggb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot openbaar maken.
 Op dit document rusten rechten van de Ggb en/of auteursrechten.
 Publicatie, verspreiding, vermenigvuldiging, commerciële exploitatie en gebruik van dit document of de inhoud hiervan zonder de toestemming van de rechthebbende van dit document zijn en blijven verboden.
 This document is not the property of the Ggb and only provided based on mandatory freedom of information requirements.
 The document may be subject to rights such as intellectual property and copy rights of third parties.
 Consequently, any publication, distribution, reproduction and/or publishing of this document may therefore be prohibited and violate the rights of its owner.

Table 2: Records on Flight and Foraging Activity of Honeybees During the Sampling Period.

Day after hive installment	No. of bees which left the hive during the 5 min. observation period	No. of bees which returned to the hive during the 5 min. observation period	No. of bees which foraged on the flowering rape during the check
<i>Control Plots</i>			
+ 1	84	65	12
+ 2	74	91	6
+ 4	4	13	1
<i>Treatment Plots</i>			
+ 1	96	130	4
+ 2	55	42	2
+ 4	5	4	0

Dit document is geen eigendom van het Ctgb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot openbaarmaking. Op dit document kunnen rechten van derden rusten, waaronder intellectuele eigendomsrechten en/of auteursrechten. Voorts kan dit document onder een regeling omtrent gegevensbescherming vallen. Publicatie, verspreiding, vermenigvuldiging, commerciële exploitatie en gebruik van dit document of de inhoud hiervan zonder de toestemming van de rechthebbende van dit document kan derhalve verboden zijn en een inbreuk opleveren van de rechten van deze rechthebbende.

This document is not the property of the Ctgb and only provided based on mandatory freedom of information requirements. The document may be subject to rights such as intellectual property and copy rights of third parties. Consequently, any publication, distribution, reproduction and/or publishing and any commercial exploitation and use of this document or its contents without the permission of the owner of this document may therefore be prohibited and violate the rights of its owner.

APPENDICES

APPENDIX I: Analytical Report.



Study No.: E 370 1360-0
(contains data from study no. RA-2043/98)

STUDY TITLE

Analysis of Rape Nectar and Rape Blossoms for Residues of Imidacloprid and Imidacloprid Metabolites and Preliminary Observations of Effects on Domestic Honeybees in Sweden

Residue Analytical Method for the Determination of Imidacloprid, Hydroxy- and Olefin-Metabolite in Rape Flower, Rape Pollen, Bee and Nectar Samples by HPLC-MS/MS

Dit document is geen eigendom van het Ctgb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot openbaarmaking. Op dit document kunnen rechten van derden rusten, waaronder intellectuele eigendomsrechten en/of auteursrechten. Voorts kan dit document onder een regeling omtrent gegevensbescherming vallen. Het gebruik van dit document of de inhoud hiervan zonder de toestemming van de rechthebbende van dit document kan derhalve verboden zijn en een inbreuk opleveren van de rechten van deze rechthebbende.

This document is not the property of the Ctgb and only provided based on mandatory freedom of information requirements. The document may be subject to rights such as intellectual property and copy rights of third parties. Furthermore, the document may fall under a regulatory data protection regime. Consequently, any publication, distribution, reproduction and/or publishing and any commercial exploitation and use of this document or its contents without the permission of the owner of this document may therefore be prohibited and violate the rights of its owner.

1) Nectar-Samples:

Extraction and sample clean-up:

1. Place for e.g. 1.0 g of the sample material in a 150-ml beaker.
2. Add 10 ml of water and place the sample for 2 min into a Ultrasonic Bath.
3. Add 20 ml of methanol.
4. Blend the sample using an ultra-turrax blender (or equivalent) for approximately 1 min.
5. Vacuum filter the suspension through 2.5 g of Celite filter aid using Schwarzband filter paper supported on a Büchner funnel into a 250-ml vacuum filter flask.
6. Wash the filtered solids with a total of 20 ml of methanol/water (3/1, v/v). Press residual solvent from the solids using rubber damming. Discard the filtered solids.
7. Transfer the filtrate into a 250-ml brown glass round-bottom flask.
8. Concentrate the filtrate to an aqueous remainder of 5 to 10 ml using a rotary evaporator with a max. bath temperature of 50 °C.
9. Add 5 to 10 ml water to the aqueous solution from step 8 to bring the total volume of the extracts to approx. 20 ml.
10. Place the aqueous solution on the top of the ChemElut® CE 1020 (20 ml volume) column fitted with a disposable stainless steel needle and wait for approx. 15 minutes to achieve an uniform distribution of the liquid on the column.
11. Elute the residues from the column with 140 ml of CH₂Cl₂. Collect the eluate in a 250-ml brown glass round-bottom flask.
12. Evaporate the eluate from step 11 to dryness using a vacuum rotary evaporator and a max. bath temperature of 40 °C.
13. Dissolve the residues in 1.00 ml of acetonitrile/water (2/8, v/v) and determine the residues with HPLC-MS/MS.

2.) Bee-Samples, Rape Flowers, Rape Pollen:

Extraction and sample clean-up:

1. Place for e.g. 2.0 g of the sample material in a 150-ml beaker. Add 30 ml of methanol/water (3/1,v/v) and allow the sample to soak for 30 min.
2. Blend the sample using an ultra-turrax blender (or equivalent) for approximately 1 min.
3. Vacuum filter the suspension through 2.5 g of Celite filter aid using Schwarzband filter paper supported on a Büchner funnel into a 250-ml vacuum filter flask.
4. Wash the filtered solids with a total of 30 ml of methanol/water (3/1, v/v). Press residual solvent from the solids using rubber damming. Discard the filtered solids.
5. Transfer the filtrate to a 100-ml graduated cylinder. Determine the total volume of the extracts. Mix the solution well, and transfer the half (e.g. 1.0 g sample equivalent) to a 250-ml brown glass round-bottom flask.
6. Concentrate the aliquot to an aqueous remainder of 5 to 10 ml using a rotary evaporator with a max. bath temperature of 50 °C.
7. Add 5 to 10 ml water to the aqueous solution from step 6 to bring the total volume of the extracts to approx. 20 ml.
8. Place the aqueous solution on the top of the ChemElut® CE 1020 (20 ml volume) column fitted with a disposable stainless steel needle and wait for approx. 15 minutes to achieve an uniform distribution of the liquid on the column.
9. Elute the residues from the column with 140 ml of CH₂Cl₂. Collect the eluate in a 250-ml brown glass round-bottom flask.
10. Evaporate the eluate from step 9 to dryness using a vacuum rotary evaporator and a max. bath temperature of 40 °C.
11. Dissolve the residues in 2 ml of toluene/ethyl acetate (85/15, v/v).
12. Apply the organic solution from step 11 onto a 0.5 g (3 ml) silica gel (SiOH) column (e.g. Varian).
13. Allow the solution to pass through the column at a flow rate of 1 ml/min.
14. Rinse the 250-ml brown glass round-bottom flask with 10 ml of toluene/ethyl acetate (70/30, v/v) and apply the solution onto the column, too.
15. Elute the residues with 5 ml of acetonitrile at a flow rate of 1 ml/min. Collect the eluate in a 25-ml brown glass pear-shaped flask.
16. Evaporate the eluate from step 15 to dryness using a vacuum rotary evaporator and a max. bath temperature of 40 °C. Dissolve the residues in e.g. 1.00 ml of acetonitrile/water (2/8, v/v) and determine the residues with HPLC-MS/MS.

NOTE

1. The volumes to be used for flushing the column with toluene/ethyl acetate and for elution with acetonitrile must be newly determined for each batch of SiOH-column!
2. The flow rate should not be too high, since otherwise losses of the residues in may occur with recoveries below 70 % and the clean-up is less effective.

3.) HPLC-MS/MS determination of Imidacloprid, Hydroxy and Olefin Metabolite:

A) Measuring equipment and HPLC conditions:

Instrument: Hewlett Packard 1100
 Column: e.g.: Phenomenex, Luna C18 (2), 5 µm, 15 x 0.46 cm i.D. or Merck, Superspher, RP select-B, 4 µm, 12.5 x 0.4 cm i.D.
 Solvent A: Water + 0.1 ml Acetic acid/L
 Solvent B: ACN + 0.1 ml Acetic acid/L
 Oventemperature: 40 °C
 Inject.volume: 50 µL
 Flow: 1.0 mL/min
 Split: 150 µL into MS from 1000 µL

Time Table	0 min	20 % B
	10 min.	20 % B
	11 min	90 % B
	15 min	90 % B
	16 min	20 % B
	19 min	20 % B
	Stoptime	19 min

Retention Times: Olefin-Imidacloprid approx. 4.5 min
 Hydroxy-Imidacloprid approx. 5.5 min
 Imidacloprid approx. 8.5 min

B) Mass Spectroscopy

The experiments were performed on a triple-quadrupole mass spectrometer fitted with an electrospray interface operated in the positive ion mode under MRM conditions.

The mass spectrometer was tuned by infusing a standard solution of 0.5 mg/l Imidacloprid, Hydroxy-Metabolite and Olefin-Metabolite (dissolved in acetonitrile/water (2/8, v/v) + 0.1 ml acetic acid per litre) at a flow rate of 5-10 µl/min.

Mass axis calibration was done by infusing a polypropylene glycol 3000 solution. Unit mass resolution was established and maintained in each mass resolving quadrupole by maintaining a full width at half-maximum of between 0.8 and 1.0 DA. After tuning and calibration, optimal collision-activated dissociation (CAD) conditions for fragmentation of Imidacloprid, Hydroxy-Metabolite and Olefin-Metabolite were determined. These experiments were performed with nitrogen as collision gas with a collision offset of +20 eV for Imidacloprid,

-23 eV for Hydroxy-Metabolite and -13 eV for Olefin-Metabolite at an approximate collision gas thickness of 1.56×10^{15} atoms/cm².

Nebulization gas is set at 1.48 l/min, curtain gas is set at 0.95 l/min and turbo gas is set at 6 l/min.

Detector: e.g. Triple Quadrupole MS/MS Mass Spectrometer

Perkin-Elmer Sciex Instruments

API 300, Apple™ Macintosh® System 8.0

Interface: Electrospray, TurboIon Spray

Potential: +4900 V

Temperature: 300 °C

Nebulizer gas: Nitrogen 5.0 (99.999% purity), 1.48 l/min

Scan type: MRM (Multiple Reaction Monitoring Mode)

Polarity: Positive

Acquisition mode: Profile

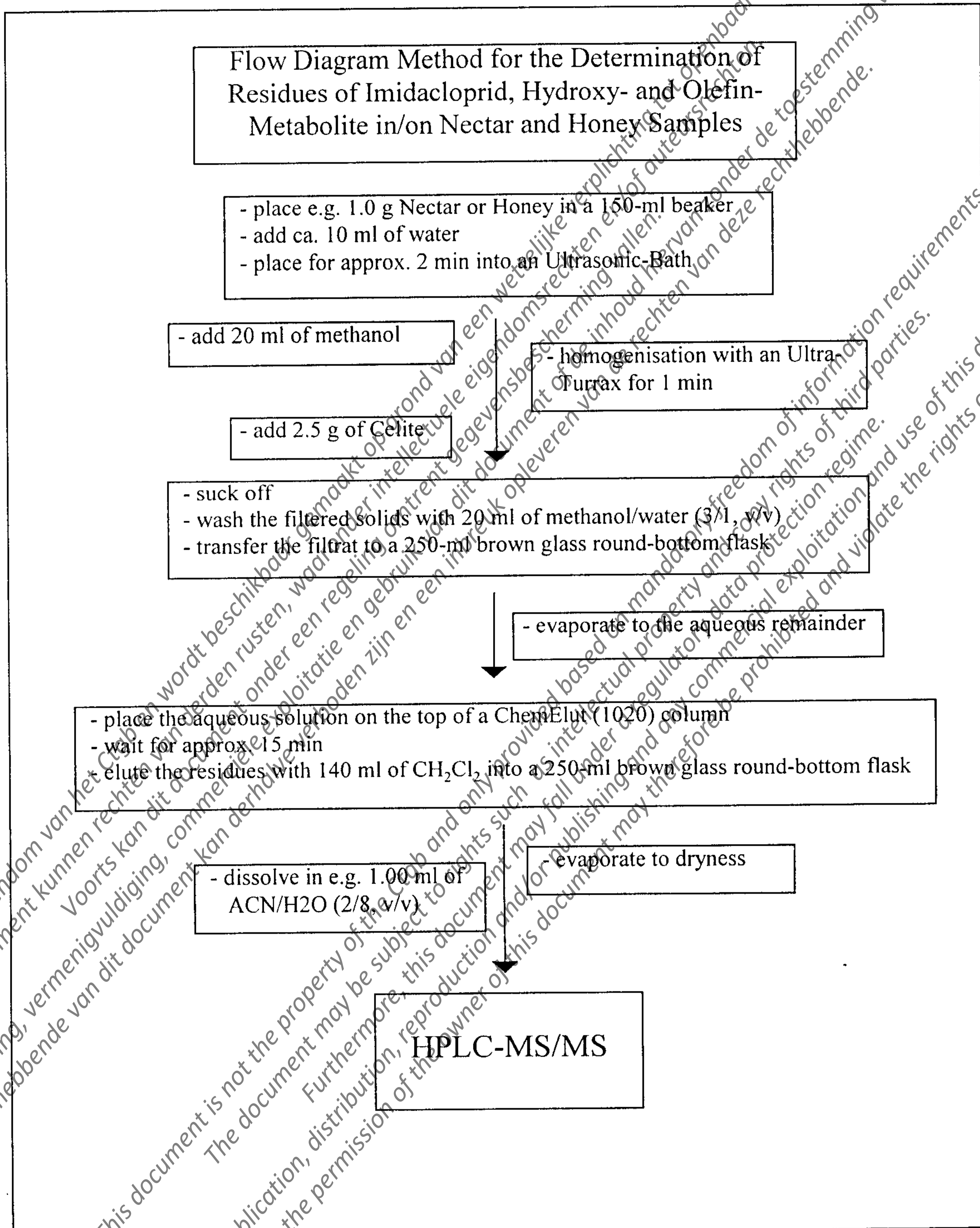
Mass spectrometer operating parameters

Compound	Precursor Ion Q1 Mass (amu)	Product Ion Q3 Mass (amu)	Dwell Time (msec)	Collision Energy (eV)
Imidacloprid (Cl 37)#	258.0	210.9	500	-20
Imidacloprid (Cl 35)	256.0	208.9	500	-20
Hydroxy-Metabolite (Cl 37)#	274.0	190.8	250	-23
Hydroxy-Metabolite (Cl 35)	272.0	190.8	250	-23
Olefin-Metabolite (Cl 37)#	256.0	237.8	250	-13
Olefin-Metabolite (Cl 35)	254.0	235.8	250	-13

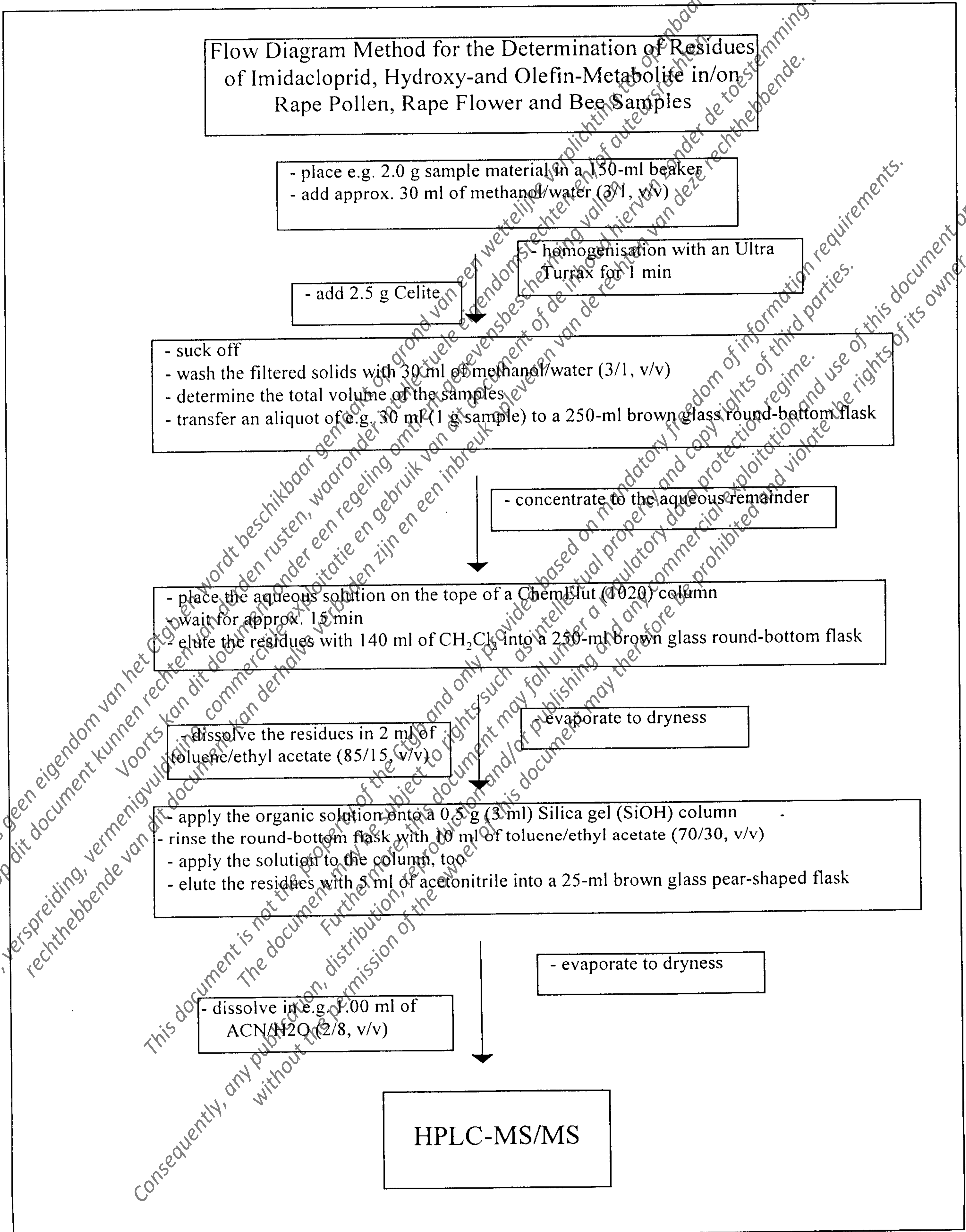
#= ³⁷Cl isotope of all substances were detected to use as qualifiers

Dit document is geen eigendom van het Ctgb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot openbaarmaking. Op dit document kunnen rechten van derden rusten, waaronder intellectuele eigendomsrechten en/of auteursrechten. Voorts kan dit document onder een regeling omtrent gegevensbescherming vallen en/of de inhoud hiervan kan deze rechten van de rechthebbende. Publicatie, verspreiding, vermenigvuldiging, commerciële exploitatie en gebruik van dit document of de inhoud hiervan kan derhalve verboden zijn en een inbreuk opleveren van de rechten van de rechthebbende. Dit document is not the property of the Ctgb and only provided based on mandatory freedom of information requirements. The document may be subject to rights such as intellectual property and copy rights of third parties. Furthermore, this document may fall under a regulatory data protection regime. Consequently, any publication, distribution, reproduction and/or publishing and any commercial exploitation and use of this document without the permission of the owner of this document may therefore be prohibited and violate the rights of its owner.

4.) Flow Diagram Method for the Determination of Residues of Imidacloprid and Metabolites in/on Nectar and Honey Samples



5.) Flow Diagram Method for the Determination of Residues of Imidacloprid and Metabolites in/on Rape Pollen, Rape Flower and Bee Samples:



6.) Results of Bee Samples, Nectar Samples of Bees, Rape Flower and Rape Pollen Samples.

A) Bee Samples:

Sample name	Sample description	Sample weight	Residues		
			Hydroxy-NTN [mg/kg]	Olefin-NTN [mg/kg]	Imidacloprid [mg/kg]

Samples taken before exposure					
D Bees Control 07/02/98	D Sweden	10.2 g	< 0.01	< 0.01	< 0.01
C Bees Poncho 07/02/98	C Sweden	12.9 g	< 0.01	< 0.01	< 0.01

Samples taken during exposure					
A Bees Control 07/03,04,06/98	A Sweden	7.4 g	< 0.01	< 0.01	< 0.01
B Bees Poncho 07/03,04,06/98	B Sweden	8.1 g	< 0.01	< 0.01	< 0.01

Dit document is geen eigendom van het Ctgb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot openbaarmaking. Voorts kan dit document onder een regeling omtrent gegevensbescherming vallen. Het gebruik van dit document of de inhoud hiervan zonder de toestemming van de rechthebbende is niet toegestaan.

Publicatie, verspreiding, vermenigvuldiging, commerciële exploitatie en gebruik van dit document kan derhalve verboden zijn en een inbreuk opleveren van de rechten van deze rechthebbende.

This document is not the property of the Ctgb and only provided based on mandatory freedom of information requirements. The document may be subject to rights such as intellectual property and copy rights of third parties. Furthermore, this document may fall under a regulatory data protection regime. Consequently, any publication, distribution, reproduction and/or publishing and any commercial exploitation and use of this document or its contents without the permission of the owner of this document may therefore be prohibited and violate the rights of its owner.

B) Nectar Samples:

Sample Name	Sample description	Sample weight	Residues		
			Hydroxy-NTN [mg/kg]	Olefin-NTN [mg/kg]	Imidacloprid [mg/kg]

Nectar harvested from the honeybulb of bees which were collected before exposure

D Nectar Control 07/02/98	D Sweden N	0.976 g	< 0.01	< 0.01	< 0.01
C Nectar Poncho 07/02/98	C Sweden N	1.665 g	< 0.01	< 0.01	< 0.01

Nectar harvested from the honeybulb of bees which were collected during exposure

A Nectar Control 07/03/04/98	A Sweden N	0.06 g	< 0.01	< 0.01	< 0.01
B Nectar Poncho 07/03/04/98	B Sweden	1.019 g	< 0.01	< 0.01	< 0.01

Nectar harvested directly from the plants via micro-capillaries

Nectar Control 07/02/98	Nectar ca. 1 ml Eppendorfgefäße 02.07.98	ca 1 ml	< 0.01	< 0.01	< 0.01
Nectar Control 07/03/98	Nectar ca. 1 ml Eppendorfgefäße 03.07.98	ca 1 ml	< 0.01	< 0.01	< 0.01
Nectar Poncho 07/02/98	Nectar ca. 1 ml Eppendorfgefäße 02.07.98	ca 1 ml	< 0.01	< 0.01	< 0.01
Nectar Poncho 07/03/98	Nectar ca. 1 ml Eppendorfgefäße 03.07.98	ca 1 ml	< 0.01	< 0.01	< 0.01

C) Rape Flower Samples:

Sample Name	Sample description	Sample weight	Residues		
			Hydroxy-NTN [mg/kg]	Olefin-NTN [mg/kg]	Imidacloprid [mg/kg]
Rape Flower Control	Blüten in 1 Probengefäß ca. 20 g 02.07.98	ca. 20 g	< 0.01	< 0.01	< 0.01
Rape Flower Poncho	Blüten in 1 Probengefäß ca. 20 g 02.07.98	ca. 20 g	< 0.01	< 0.01	< 0.01

D) Rape Pollen Samples:

There was not a sufficient amount of pollen for a reasonable residue analysis

Dit document is geen eigendom van het Ctgb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot informatieverschaffing. Op dit document kunnen rechten van derden rusten, waaronder intellectuele eigendomsrechten en/of auteursrechten. Voorts kan dit document onder een regeling omtrent gegevensbescherming vallen. Het gebruik van dit document of de inhoud hiervan zonder de toestemming van de rechthebbende van dit document kan derhalve verboden zijn en een inbreuk opleveren van de rechten van de rechthebbende van deze informatie.

This document is not the property of the Ctgb and only provided based on mandatory freedom of information requirements. The document may be subject to rights such as intellectual property and copy rights of third parties. Furthermore, this document may fall under a regulatory data protection regime. Consequently, any publication, distribution, reproduction and/or publishing and any commercial exploitation and use of this document or its contents without the permission of the owner of this document may therefore be prohibited and violate the rights of its owner.

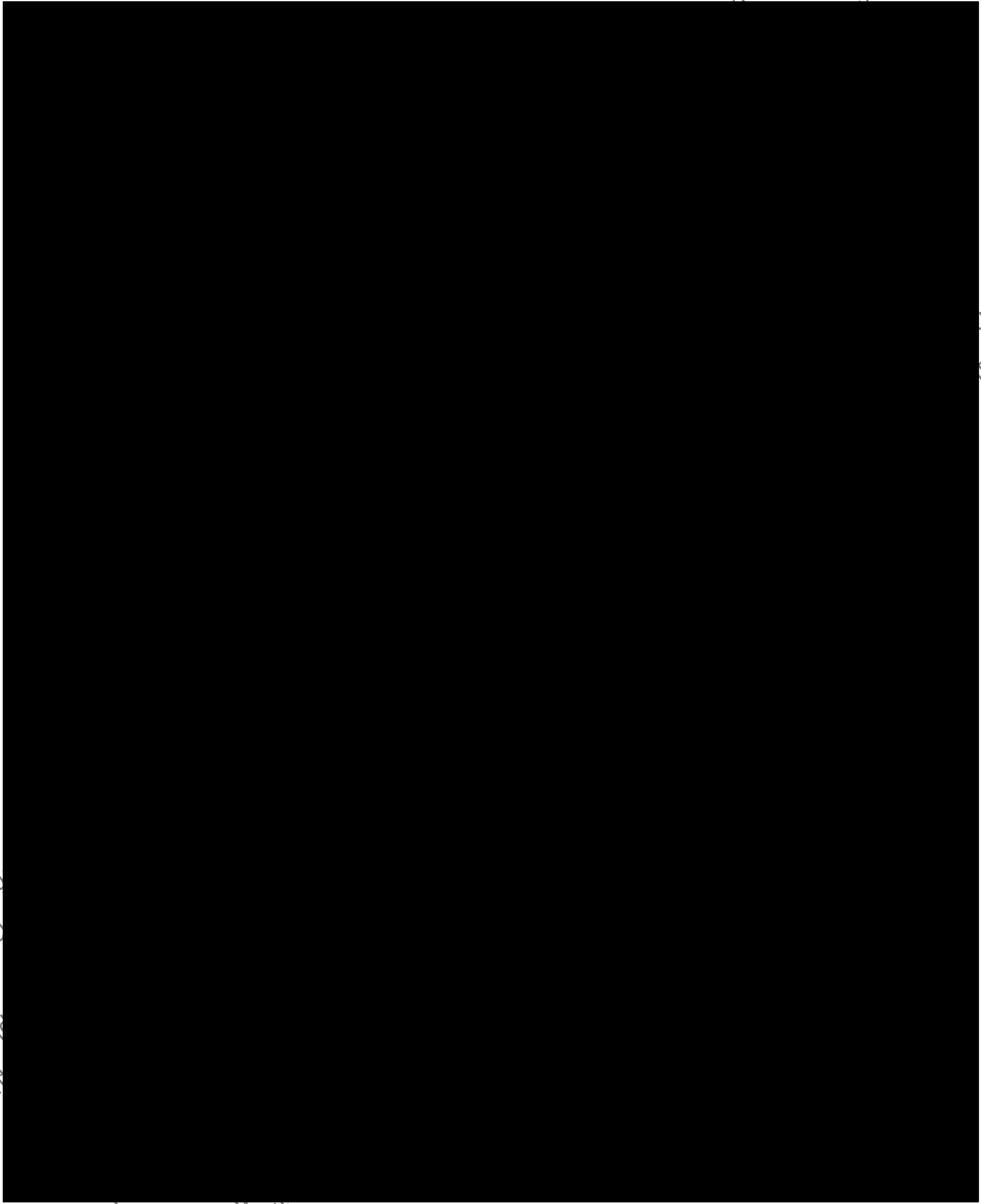
7.) Results of Water Control Samples

Sample Name	Sample description	Sample weight	Residues		
			Hydroxy-NTN [mg/kg]	Olefin-NTN [mg/kg]	Imidacloprid [mg/kg]
Water SWE E 3701360-0 C Blank Control Sample	Sweden	1 g	< 0.01	< 0.01	< 0.01
Water SWE E 3701360-0 N Blank Control Sample	Sweden	1 g	< 0.01	< 0.01	< 0.01
Water SWE E 3701360-0 C Poncho FS 500 Blank Sample	Sweden	1 g	< 0.01	< 0.01	< 0.01
Water SWE E 3701360-0 N Poncho FS 500 Blank Sample	Sweden	1 g	< 0.01	< 0.01	< 0.01

Limit of Quantitation = 0.01 mg/kg

Dit document is geen eigendom van het Ctgb en wordt beschikbaar gemaakt op grond van een wettelijke verplichting tot openbaarmaking.
 Op dit document kunnen rechten van derden rusten, waaronder intellectuele eigendomsrechten en/of auteursrechten.
 Voorts kan dit document onder een regeling omtrent gegevensbescherming vallen.
 Publicatie, verspreiding, vermenigvuldiging, commerciële exploitatie en gebruik van dit document of de inhoud hiervan zonder de toestemming van de rechthebbende van dit document kan derhalve verboden zijn en een inbreuk opleveren van de rechten van deze rechthebbende van de informatievereisen.
 This document is not the property of the Ctgb and only provided based on mandatory freedom of information requirements.
 The document may be subject to rights such as intellectual property and copy rights of third parties.
 Furthermore, this document may fall under a regulatory data protection and use of this document and/or publishing and any commercial exploitation and use of this document may therefore be prohibited and violate the rights of its owner.
 Consequently, any publication, distribution, reproduction and/or publishing and any commercial exploitation and use of this document may therefore be prohibited and violate the rights of its owner.

Appendix II: Copy of the GLP Certificate



making.

van de

document or its contents
its owner.

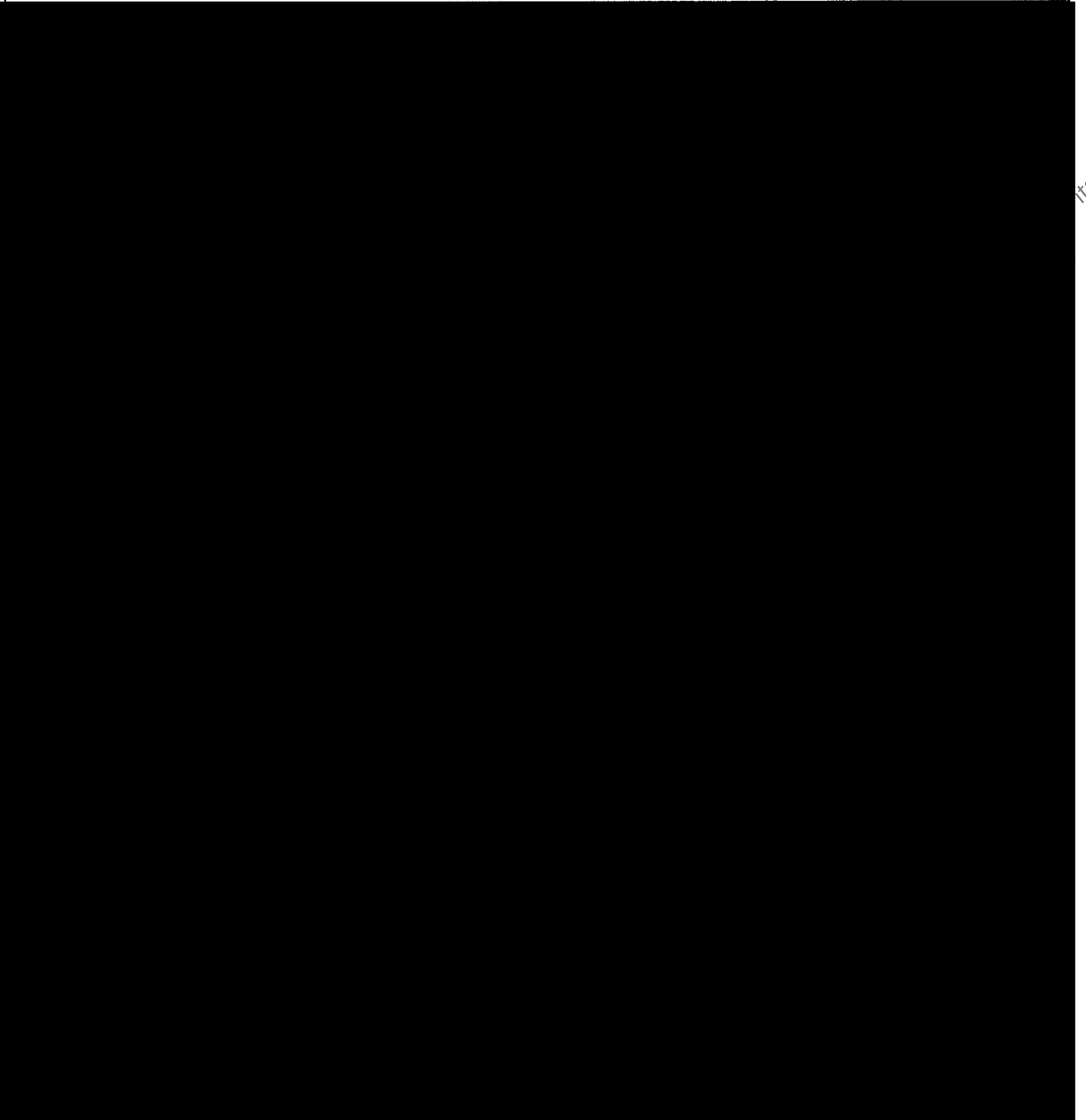
Dit document is geen eigendom
Op dit document
Publicatie, verspreiding
recht

Consequently, any publication
without

Appendix III: Quality Assurance Statement

Referat GLP

Quality Assurance Statement



garmaking.
ng van de

its contents

Dit docu

Publ

Consequently, any
witho