Cover Sheet for

ENVIRONMENTAL CHEMISTRY METHOD

Pestcide Name: Metolachlor

MRID #: 454996-01

Matrix: Water

Analysis: Immunoassay

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VOLUME 2 of 14 of submission

CGA-77102: METHOD

TITLE

ANALYTICAL METHOD FOR THE SEMI-QUANTITATIVE DETERMINATION OF CGA-77102 IN WATER BY ENZYME IMMUNOASSAY INCLUDING VALIDATION DATA

DATA REQUIREMENT

Not Applicable

AUTHORS

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COMPLETION DATE

June 23, 1999

PERFORMING LABORATORY

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Novartis Crop Protection, Inc.
Greensboro, NC 27419

LABORATORY STUDY IDENTIFICATION

Novartis Number 1004-98

SUBMITTER/SPONSOR

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VOLUME 1 OF 1 OF STUDY

PAGE <u>1</u> OF <u>40</u>

STATEMENT OF NO DATA CONFIDENTIALITY CLAIM

No claim of confidentiality is made for any information contained in this study on the basis of its falling within the scope of FIFRA Section 10 (d) (1) (A), (B) or (C).

Company: Novartis Crop Protection, Inc.

Company Representative: Karen Stumpf

Title: Senior Regulatory Manager

Signature: Kauen Stump Date: 6/23/99

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GOOD LABORATORY PRACTICE COMPLIANCE STATEMENT

This study was performed in accordance with Good Laboratory Practice Standards as required by the EPA-FIFRA Good Laboratory Practice Standards, 40 CFR part 160 with the following exception(s):

Alachlor and Dimethanamid-ESA analytical standards did not comply with the above guidelines.

Jody **L**. Needham.

Date

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CERTIFICATION

The analytical reports and experimental results included in Project H077102NAU950A, in support of analytical method 1004-98, are authentic accounts of the experiments.

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Date.

QUALITY ASSURANCE STATEMENT

Study Title:

Analytical Method for the Semi-Quantitative Determination of

CGA-77102 in Water by Enzyme Immunoassay Including

Validation Data

Study Director:

Needham, Jody L.

Method Number:

1004-98

Pursuant to Good Laboratory Practice Regulations, this statement verifies that the aforementioned study was inspected and/or audited and the findings reported to Management and to the Study Director by the Quality Assurance Unit on the dates listed below.

Inspection/Audit Type	Inspection/Audit Dates	Reporting Date
Audit Protocol	11/18/98 - 11/18/98	11/19/98
Inspect Analytical	05/05/99 - 05/05/99	05/06/99
Audit Final Report	05/18/99 - 05/20/99	05/24/99

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Staff Quality Assurance Auditor

Quality Assurance Unit

Novartis Crop Protection, Inc.

TABLE OF CONTENTS

				<u>P</u>	<u>age No</u>
TITLE P	AGE	•••••			1
STATE	ИΕΝΊ	OF N	O DATA	A CONFIDENTIALITY CLAIM	2
GOOD I	LABC	RATO	RY PRA	ACTICE COMPLIANCE STATEMENT	3
REPOR	TAP	PROV	AL		4
CERTIF	ICAT	10N			5
				ATEMENT	6
TABLE	OF C	ONTE	NTS		7.
GENER	AL IN	IFORM	MATION		10
1.	SU	MMAR	Y AND I	NTRODUCTION	12
	A.	SCO	PE		· 12
	B.	PRIN	CIPLE.		12
II.	MA	TERIA	LS AND	METHODS	12
: ,	A.	APP	ARATUS	S	12
٠	В.	REA	GENTS		. 13
	C.	ANAI	YTICAI	PROCEDURE	14
		1.0	Sampl	e Preparation	14
		2.0	Enzym	ne Immunoassay	15
			2.1	Inhibition of Enzyme Conjugate	15
			2.2	Wash	16
	•	•	2.3	Color Development	17

TABLE OF CONTENTS (Continued)

			<u>.</u>	Page No
	D.	İNST	RUMENTATION	18
•	,	1.0	Description and Operating Conditions	18
		2.0	Standardization	18
•	E.	INTER	RFERENCES	18
	F.	CONF	FIRMATORY TECHNIQUES	19
	G.	TIME	REQUIRED	19
	Н.	MODI	IFICATIONS AND POTENTIAL PROBLEMS	19
	I.	PREF	PARATION OF STANDARD SOLUTIONS	20
	J.	METH	HODS OF CALCULATION	20
111.	RES	SULTS	AND DISCUSSION	20
	A.	VALIE	DATION	20
		1.0.	ACCURACY AND PRECISION	21
		2.0.	SPECIFICITY	21
		3.0.	LIMITS OF DETECTION AND QUANTITATION	21
	В.	LIMIT	ATIONS	22
	C.	CIRC	UMSTANCES AFFECTING THE STUDY	22
IV.	COI	NCLUS	SION	22
V.	TAB	LES A	ND FIGURES	•
	TAB	BLE 1.	RESULTS OF STANDARD ADDITION EXPERIMENTS	23

TABLE OF CONTENTS

(Continued)

			<u>Pa</u>	ge ivo
	TABLE	≣ 2.	CROSS-REACTIVITY RESULTS FOR THE CGA-77102 ENZYME IMMUNOASSAY	24
	FIGUE	RE 1.	FLOW DIAGRAM FOR METHOD 1004-98	25
	FIGUF	RE 2.	PLATE LAYOUT CHART ON WHICH THE ANALYST WILL INDICATE THE POSITION OF SAMPLE AND STANDARD SOLUTIONS IN THE MICROWELL	00
		•	ASSAY PLATE	26
	FIGUF	RE 3.	A TYPICAL STANDARD CURVE OF THE CGA-77102 MICROWELL ASSAY	27
	FIGUF	RE 4.	STRUCTURES OF TEST SUBSTANCES EVALUATED FOR CROSS-REACTIVITY	28
VI. I	REFEREN	CE		32
APPE	ENDIX 1	RESID	UE TEST REPORT NO. 1	33
APPE	NDIX 2.	RESID	UE TEST REPORT NO. 2	37

GENERAL INFORMATION

Protocol Number: 1004-98 plus one amendment

Study Initiation Date: December 9, 1998

Study Personnel:

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Manager, Analytical Resources Group

Study Director and Analyst: Jody L. Needham, B.A.

Chemist I, Analytical Resources Group

Analyst: James F. Brady, Ph.D.

Scientist III, Analytical Resources Group

Reference Substance Information:

Novartis Code Number	CAS Number	CAS Name	Lot Number	Purity	Reassay Date
Metolachlor	51218-45-2	Acetamide, 2-chloro-N- (2-ethyl-6-methylphenyl)-N- (2-methoxy-1-methylethyl)-	S94-1719	99.3 %	2/2000
. Alachlor	15972-60-8	Acetamide, 2-chloro-N-(2,6- diethylphenyl)-N- (methoxymethyl)-	S86-0926	98.4 %	NON-GLP
Alachlor-ESA (NOA-436644)	142363-53-9	Ethanesulfonic acid, 2-[(2,6- diethylphenyl)(methoxymethyl) amino}-2-oxo-	DAH-XXII-24	99.4 %	11/2000
Acetochlor	34256-82-1	Acetamide, 2-chloro-N- (ethoxymethyl)-N-(2-ethyl-6- methylphenyl)-	S97-2120	98 %	8/99
Acetochlor-ESA (NOA-436645)	187022-11-3	Ethanesulfonic acid, 2- [(ethoxymethyl)(2-ethyl-6- methylphenyl)amino]-2-oxo-	DAH-XXII-30	97.8 %	11/2000
Dimethanamid	87674-68-8	Acetamide, 2-chloro-N-(2,4- dimethyl-3-thienyl)-N- (2-methoxy-1-methylethyl)-	S97-2034	99.7 %	7/2000
Dimethanamid- ESA	205939-58-8	Ethanesulfonic acid, 2-[(2,4- dimethyl-3-thienyl)(2-methoxy- 1-methylethyl)amino]-2-oxo-	BPM-XVIII-19	Approx 87.5 %	NON-GLP
Metalaxyl	57837-19-1	DL-Alanine, N-[2,6- dimethylphenyl)-N- (methoxyacetyl)-,methyl ester	S94-1763	97.1 %	10/2000



GENERAL INFORMATION (continued)

Novartis CAS			Lot		Reassay
Code Number	Number	CAS Name	Number	Purity	Date
CGA-354743	Not Assigned	Ethanesulfonic acid, 2 [(2-ethyl-6-methylphenyl)(2methoxy-1-methylethyl)amino]-	DAH-XXVI-12	96.8 %	9/2000
		2-oxo-, sodium salt			
CGA-50720	152019-74-4	Acetic acid, [(2-ethyl-6- methylphenyl)amino]oxo-	DPS-I-74-2	>99.9%	6/99
CGA-40172	131068-72-9	Acetamide, N-(2-ethyl-6- methylphenyl)-2-hydroxy-N-(2- methoxy-1-methylethyl)-	, JAK-II-42(IV)	95.9 %	6/99
CGA-67869	66637-79-4	DL-Alanine, N-(2,6- dimethylphenyl)-N- (hydroxyacetyl)-, methyl ester	DAH-XV-35	>99.9%	6/99
CGA-67125	Not Assigned	Formamide, N-(2-ethyl-6- methylphenyl)-N-(2-methoxy-1- methylethyl)-	DPS-VII-41-1	99.9 %	12/99
CGA-37735	97055-05-5	Acetamide, N-(2-ethyl-6- methylphenyl)-2-hydroxy-	JAK-II-42(II)	99.2 %	3/2000
CGA-41638	65513-61-3	Acetamide, 2-chloro-N-(2- ethyl-6-methylphenyl)-N-(2- hydroxy-1-methylethyl)-	Acetamide, 2-chloro-N-(2- S94-1765 ethyl-6-methylphenyl)-N-(2-		9/99
CGA-51202	152019-73-3	Acetic acid, [(2-ethyl-6- methylphenyl)(2-methoxy-1- methylethyl)amino]oxo-	JAK-I-63	98.8 %	3/2000
CGA-62826	87764-37-2	DL-Alanine, N-(2,6- dimethylphenyl)-N- (methoxyacetyl)-	BPM-I-48	98.0 %	11/2000
CGA-46576	Not Assigned	Not Assigned	AW-I-43	97.4 %	6/99
CGA-380168	Not Assigned	Ethanesulfonic acid, 2-[(2- ethyl-6-methylphenyl)(2- methoxy-1-methylethyl)amino]- 2-oxo-, (S)-, sodium salt	thyl-6-methylphenyl)(2- loxy-1-methylethyl)amino]-		1/99
CGA-77102	87392-12-9	Acetamide, 2-chloro-N-(2- ethyl-6-methylphenyl)-N-(2- methoxy-1-methylethyl)-, (5)-	S95-1844	98.5 %	11/2000
CGA-37913	61520-53-4	1-Propanol, 2-[(2-ethyl-6- methylphenyl)amino]-	JAK-XVI-38	99.0 %	12/2000



SUMMARY AND INTRODUCTION

A. <u>SCOPE</u>

This method is to be used for the semiquantitative determination of CGA-77102 ([Acetamide, 2-chloro-*N*-(2-ethyl-6-methylphenyl)-*N*-(2-methoxy-1-methylethyl)-,(*S*)-], CAS # 87392-12-9) in water. The limit of detection (LOD, the smallest dose that yields a response that is statistically significant different than the response of the zero dose) is 0.05 ppb of CGA-77102. The limit of quantitation (LOQ, the lowest level of fortification with which an acceptable recovery can be obtained) for the method is 0.10 ppb of CGA-77102.

B. PRINCIPLE

A 100-µl aliquot of a representative water sample is added to a microwell coated with CGA-77102 antibody. The assay is carried out by sequential addition of enzyme conjugate, wash solution, enzyme substrate, and stop solution. The reaction is terminated by acidification. Quantification is performed spectrophotometrically at 450 nm. A flow diagram for the method is presented in Figure 1.

II. MATERIALS AND METHODS

A. <u>APPARATUS</u>

- 1.0 CGA-77102 Plate kit, enzyme immunoassay microwell assay, Beacon Analytical Systems, Scarborough, ME, catalog #CPP-005.
- 2.0 Microtiter plate reader, Titertek Multiskan MCC/340 MK II, ICN catalog #78-626-00, or equivalent.
- 3.0 AUTOmate™ microplate software and manual, ICN cat. #78-599-02 and 78-599-98, respectively, or equivalent.
- 4.0 Microtiter plate shaker, Lab-Line Instruments catalog #4625 or equivalent.
- 5.0 ELP-40 Microplate Strip Washer, Bio-Tek Instruments, Bio-Tek Part No. 4091000, or equivalent (use of equipment is optional).
- 6.0 Eight-channel pipette capable of dispensing 100-μl volumes, Elkay Labsystems catalog #4142-417 or equivalent.

- Adjustable microliter pipette, capable of dispensing 20- to 200-µl volumes, Rainin cat. #P-200 or equivalent.
- 8.0 Pipette tips for eight-channel pipette for volumes of 20- to 200-µl, packaged in an eight by twelve array, Costar cat. #4865 or equivalent.
- 9.0 Pipette tips for single-channel pipette for volumes of 100- to 1000-µl, packaged as 100 tips per rack, Costar cat. #4867 or equivalent.
- 10.0 Graduated pipette, 5- and 10-mL, Fisher cat. # 13-660E and 13-660F or equivalent.
- 11.0 Microtiter assay plate, 96 well, flat bottom, polystyrene, nonsterile, not treated for cell culture work, Flow Laboratories cat. #76-102-05 or equivalent.
- 12.0 Reagent reservoirs for multichannel pipettes, non-sterile, ICN Cat. #77-824-01 or equivalent.
- 13.0 pH meter, Corning M-90 pH Stick, VWR cat. #34102-081 or equivalent.
- 14.0 Magnetic stir bar, 0.5-1.0 inch in length, Fisher cat. #14-511-61 or 14-511-63 or equivalent.
- 15.0 Magnetic stirrer, Fisher cat. #11-496-21 or equivalent.
- 16.0 4-oz Amber Boston round bottles, Fisher cat. #03-320-4B or equivalent.

B. <u>REAGENTS</u>

- 1.0 Distilled, deionized water (H₂O).
- Acetonitrile (ACN), HPLC Grade, Fisher cat. #A998SK-4 or equivalent.
- 3.0 Sodium Chloride (NaCl) crystal, certified A.C.S., CAS No. 7647-14-5, Fisher cat. # S271-500 or equivalent.
- 4.0 Polyoxyethylene-sorbitan monolaurate (TWEEN 20). Sigma cat. #P-1379 or equivalent.

- 5.0 50% H₂O/TWEEN 20 Combined 1 mL of TWEEN 20 and 1 mL of H₂O.
- 6.0 H₂O/TWEEN 20 wash solution Dissolve 9 g of NaCl and 1 mL of 50% H₂O/TWEEN 20 in 1 liter of H₂O.
- 7.0 Enzyme conjugate, included in the CGA-77102 plate kit, Beacon Analytical Systems.
- 8.0 Substrate solution, included in the CGA-77102 plate kit, Beacon Analytical Systems. Alternatively, a commercially available horseradish peroxidase tetramethylene benzidine substrate, such as "K-Blue" (Neogen Corporation cat. #300177) may be used.
- 9.0 Stop solution, included in the CGA-77102 plate kit, Beacon Analytical Systems. Alternatively, 1.0 N HCl, prepared by the analyst, may be used.
- 10.0 CGA-77102 analytical standard, Novartis Crop Protection, P.O. Box 18300, Greensboro, NC 27419-8300. Storage conditions: frozen.

C. ANALYTICAL PROCEDURE

The antibody-coated microwells, all reagents and sample and standard solutions must be warmed to room temperature, approximately 22°C, prior to use. The reaction kinetics are temperature dependent. Be certain all solutions are warmed to room temperature before running the assay.

1.0 <u>Sample Preparation</u>

1.1 Preparation of Water Samples

- 1.1.1 Water samples are received and stored refrigerated until use.
- 1.1.2 The pH of each sample should be measured. Set the sample container on a magnetic stirrer and add a small stir bar. Allow the sample to stir for a few seconds prior to inserting the electrode. Record the value obtained. Samples having a pH value between 5.0 and 9.0 may be analyzed by this method. Samples having

a pH outside this range are unsuitable for this assay.

2.0 Enzyme Immunoassay

2.1 Inhibition of Enzyme Conjugate

- 2.1.1 Approximately 200 µl of each sample or standard solution is added individually to wells of the uncoated reservoir plate as indicated on the plate layout chart previously completed by the analyst. A sample chart is illustrated in Figure 2.
- 2.1.2 Decant the enzyme conjugate solution into a multichannel pipette reagent reservoir.
- 2.1.3 Adjust an eight-channel pipette to deliver 100 µl (refer to the pipette operation manual or applicable Novartis Crop Protection SOP for operation of the multichannel pipette). Snugly fit eight tips to the pipette and transfer a 100-µl aliquot from column 1, rows A through H, of the reservoir plate to the corresponding column on the antibody-coated assay plate. Eject the used tips to a waste receptacle. Attach eight new tips and proceed to transfer aliquots from columns 2 through 12 of the reservoir plate to the assay plate taking care to change tips after each transfer.

If an alternative, equivalent pipetting device is used in lieu of a hand-held multichannel pipette, the 100-µl aliquot from each well of the reservoir plate is transferred to the assay plate in a manner consistent with the operation of the device.

2.1.4 As soon as all sample and standard solutions have been transferred to the assay plate, successively pipette 100-µl aliquots of the enzyme conjugate solution from the reagent reservoir to each column of the assay plate from left to right as

described above. The pipette tips must be changed after each transfer.

Note: The enzyme conjugate solution has a surfactant-like character and tends to bubble and foam when pipetted. Through preliminary trials the analyst should ensure that manual or automated pipetting devices reproducibly deliver 100-µl aliquots without bubbling or foaming.

2.1.5 Place the plate on the plate shaker. Start the shaker with the power control set to the "constant" position. The speed setting should be adjusted to approximately ninety oscillations per minute or slightly less than "2" on the Lab Line shaker. Allow the plate to shake for one hour.

Note: The reaction kinetics of this assay are temperature dependent. To avoid exposing the plate to harsh fluctuations in temperature such as strong drafts, the plate shaker should be placed inside an opaque closed chamber. Placing a cardboard box over the shaker will suffice if a more elaborate apparatus, such as a darkened Plexiglas chamber, is not available.

2.2 <u>Wash</u>

- 2.2.1 Remove the plate from the shaker. Using a hand-held multichannel pipette, remove the reactants from each column left to right across the plate.
- 2.2.2 Starting with column one, add approximately 200 µl of wash solution to the each well, gently agitate the plate and immediately remove the wash solution. Proceed across the plate from column two to twelve washing each column in a similar fashion. Wash the plate two additional times.

2.2.3 After the final wash, shake the plate vigorously to remove most of the remaining liquid. The plate may be inverted and blotted with dry paper towels. Blot only the exterior of the plate. Do not insert the blotting paper into the wells. Visually examine the plate to ensure little wash remains.

Should only part of the microtiter plate be used, the analyst should exercise care not to contaminate unused wells.

If an alternative washing device, such as the ELP-40 microplate strip washer, is used, be certain the reactants are removed in a manner consistent with their addition.

2.3 Color Development

- 2.3.1 Add 100 µl of substrate to each well across the plate in the same fashion as the previous solutions were dispensed. Tips need not be changed between additions.
- 2.3.2 Place the plate on the plate shaker and shake for 0.5 hour with the controls set at the same settings as described in Section II.C.2.1.5. Individual wells will gradually turn varying shades of blue.
- 2.3.3 Terminate color development by adding 100 µl of stop solution to each well in the same fashion as the color reagent was added. Mix the acidified solution well by repeated pipetting and dispensing of each well's contents, taking care not to form bubbles or to leave liquid behind in the pipette tips. Should bubbles form, they can be broken by manipulation with a pipette tip. Tapping the side of the strip holder gently may aid mixing of an acidified solution. A well-mixed solution will appear yellow to the eye with no traces of blue remaining. Be sure to change tips

between each addition to avoid contaminating a column of wells with residue from a previous column.

The absorbance of the final reaction product within each microwell should be analyzed within 30 minutes of addition of stop solution.

D. INSTRUMENTATION

1.0 Description and Operating Conditions

1.1 The Titertek Multiskan MCC/340 MK II eight-channel filter spectrophotometer is used to measure the absorbance of the final reaction solutions in each well. The operator should select filter 4 (450 nm) for this method. The instrument should be warmed up for ten minutes prior to use. Refer to the instrument operating instructions for further details on the operation of this instrument.

2.0 Standardization

- 2.1 Each assay consists of standards and samples run concurrently on the same plate. The absorbance values obtained from sample solutions may only be compared to the absorbance of standards run in the same analytical set. CGA-77102 standards range from 4.0 to 0.05 ng/mL in addition to a H₂O blank (zero dose).
- 2.2 Spectrophotometric analysis of the colored reaction products obtained from standard solutions will yield absorbance values measured at 450 nm (A_{450}). With a calculator or computer, use these data to generate a log/linear regression function. This curve is in the form of $y = m \log(x) + b$. The concentrations of the CGA-77102 standards are plotted on a logarithmic scale on the horizontal (x) axis and the absorbance values are plotted on a linear scale on the vertical (y) axis. An example of a typical standard curve is shown in Figure 3.

E. INTERFERENCES

1.0 The antibodies used in this assay bind primarily to CGA-77102 and metolachlor. Refer to Section III.A.2.0.



SPECIFICITY for a detailed discussion of the cross-reactivity parameters of this assay.

2.0 Slight inhibition was observed when glassware was not rinsed before use. Therefore, to obtain best results, all glassware must be pre-rinsed before use. This includes volumetric flasks, bottles, and scintillation vials used in the preparation of standards, beakers used to contain distilled water, and any other glassware that may be used. Pre-rinsing involves triple rinsing with Acetone and blowing dry with Nitrogen air. Vessels used to contain water should then be rinsed three times with de-ionized water before filling with distilled water.

F. CONFIRMATORY TECHNIQUES

1.0 The ability of the microwell assay to respond to CGA-77102 in water may be assessed by the analysis of samples fortified with a known amount of CGA-77102.

G. <u>TIME REQUIRED</u>

An analyst can analyze forty-two samples and six standards in duplicate in approximately three hours.

H. MODIFICATIONS AND POTENTIAL PROBLEMS

- 1.0 As previously stated in the introduction to the ANALYTICAL PROCEDURE, the microwell assay plates, reagents and sample solutions must be warmed to room temperature, approximately 22°C, before use. Placing the plate (in its plastic bag) and the reagents in a hood with the sash drawn low for approximately 30 minutes will bring the reagents to temperature efficiently.
- 2.0 The pH of the water sample, if extreme, may affect antibody binding. The pH of each sample should be examined. If a sample is found to have a pH outside the range of 5.0 9.0, it is unsuitable for analysis by this method.
- 3.0 The bottom surface of the microwell strips is the optical surface through which the absorbance of the final reaction product will be measured. The analyst should exercise care to prevent damage to this surface. The strip holder is designed to prevent the bottom of the strips from contacting flat surfaces, such as a bench top, upon which the strip

holder may be placed. Nevertheless, the analyst should maintain a clean work area as a preventative measure.

- 4.0 The analyst should take care to be certain all sample and standard solutions are positioned properly in the reservoir plate. The large sample load requires that special attention to detail be maintained throughout the analysis.
- 5.0 The analyst may observe absorbance values which are less than those of the highest standard, 4.0 ng/mL. In this event, the concentration of the corresponding sample cannot be calculated since its absorbance readings do not fall within the range of the standard curve. The sample should be diluted and re-assayed to obtain absorbance readings which lie within the bounds of the standard curve. The concentration of the undiluted sample can be calculated by multiplying the concentration of the diluted sample by the dilution factor.
- 6.0 The transfer of solutions by multichannel pipettes requires the analyst to constantly monitor his or her technique.

 Pipetting errors are the major source of error in immunoassay methodology.

I. PREPARATION OF STANDARD SOLUTIONS

Dissolve 5.0 mg of CGA-77102 in a minimal amount of ACN (approximately 15 mL), sonicate, and bring volume up to 100 mL with H_2O . This will make 100 mL of a 50 μ g/mL solution. Serially dilute this solution with H_2O to make 5000-, 1000-, 10-, 4.0-, 1.0-, 0.3-, 0.1-, and 0.05-ng/mL standards. Prepare a blank consisting solely of H_2O . All standard solutions should be stored in amber Boston round bottles. Store these solutions at 4°C when not in use.

J. <u>METHODS OF CALCULATION</u>

CGA-77102 residues in sample solutions are determined by inserting the absorbance value of a given sample into the log/linear regression function generated by methods described in Section II.D.2.2. These calculations may be made on a computer or hand-held calculator.

III. RESULTS AND DISCUSSION

A. <u>VALID</u>ATION

The objective of this study was to validate "draft" analytical method 1004-98 by analysis of freshly fortified water samples.

1.0 ACCURACY AND PRECISION

The accuracy and precision of this method were assessed by conducting standard addition experiments using laboratory reagent water. Two groups of ten samples each were fortified with 0.10 and 1.0 ppb of CGA-77102, respectively, and analyzed pre- and post-fortification. The results of these experiments are shown in Table 1. The mean percent recovered (95% at 0.10 ppb and 123% at 1.0 ppb) indicate the method has acceptable accuracy. The standard deviations of these measurements (0.01 ppb at 0.10 ppb and 0.09 ppb at 1.0 ppb) indicate Method 1004-98 has acceptable precision.

2.0. SPECIFICITY

The specificity of Method 1004-98 was evaluated by determining the cross reactivity parameters of twenty-one structurally related test substances. Results of these experiments are shown in Table 2. These data indicate the antibodies used in Method 1004-98 are primarily reactive to CGA-77102. Metolachlor (CGA-24705) is strongly recognized (55%) because one-half of this material is, in fact, CGA-77102 (S-CGA-24705). Dimethanamid and CGA-380168, the ethanesulfonic acid metabolite of CGA-77102, are far less reactive (12.9 and 2.1%, respectively). The remaining test substances are less than 1% reactive or not reactive at all. Thus, this method has little or no reactivity with degradates of chloroacetanalide compounds. Refer to Figure 4 for the chemical structures of the test substances screened for cross reactivity.

3.0. LIMITS OF DETECTION AND QUANTITATION

The limits of detection for test substances screened were calculated using Brady's modification of Rodbard's method (1). This method was found to have an LOD of 0.05 ppb for CGA-77102, 0.05 ppb for Metolachlor, 0.20 ppb for Dimethanamid, and 0.23 ppb for CGA-380168. All other test substances were found to have less than one percent reactivity to the assay relative to CGA-77102 so additional limits of detection were not determined. An LOQ was determined only for CGA-77102 because this was the only test substance used to conduct standard addition experiments. Based on the data generated in these

experiments, CGA-77102 was determined to have an LOQ of 0.10 ppb.

B. LIMITATIONS

- 1.0 This method is intended to complement, not replace, chromatographic analyses. It can be used as an inexpensive screening technique to remove samples that yield responses below a pre-selected level of concern from further, more expensive analytical procedures.
- 2.0 As stated in Section II.H.2.0, the pH of a sample may affect antibody binding. Consequently, the pH of all samples must be checked prior to analysis. If the pH of a sample is outside the range of 5.0 9.0, that sample is unsuitable for analysis by Method 1004-98.
- The cross reactivity data presented in Table 2 is not meant to be an exhaustive evaluation of the antibodies used in Method 1004-98. The twenty-one test substances screened represent a limited attempt to investigate the cross reactivity of the assay. Therefore, analysts should confirm residues of concern determined by this method by an additional analytical method.

C. CIRCUMSTANCES AFFECTING THE STUDY

There were no adverse circumstances affecting the quality or integrity of the data.

IV. <u>CONCLUSION</u>

This method has been found to be a rapid, reliable and accurate method for the semi-quantitative determination of CGA-77102 in water.

٧. TABLES AND FIGURES

TABLE 1. ¹RESULTS OF STANDARD ADDITION EXPERIMENTS

	CGA-77102					
Distilled Water						Fortification
Samples	Pre-fort	tification	Post-fort	<u>ification</u>	² Net	<u>Level (ppb)</u>
Α	< 0.05	<0.05	0.09	0.09	0.09	0.10
В	<0.05	<0.05	0.08	0.09	80.0	0.10
С	<0.05	<0.05	· 0.10	0.09	0.10	0.10
D	<0.05	<0.05	0.09	0.09	0.09	0.10
E	<0.05	<0.05	0.10	0.16	0.13	0.10
F	< 0.05	<0.05	0.09	0.10	0.10	0.10
G	<0.05	<0.05	0.08	0.10	0.09	0.10
Н	< 0.05	< 0.05	0.10	0.10	0.10	0.10
1	< 0.05	<0.05	0.09	0.10	0.09	0.10
J	<0.05	< 0.05	0.08	0.09	0.09	0.10

Mean \pm SD = 0.10 \pm 0.01 N = 10% CV = 13.4

	ppb	CGA-77102	Equivalents F	ound		CGA-77102
Distilled Water						Fortification
Samples	Pre-fort	ification	Post-for	tification	² Net	Level (ppb)
A .	<0.05	<0.05	1.03	1.22	1.12	1.0
В	<0.05	<0.05	1.10	1.23	1.17	1.0
Ç	<0.05	<0.05	1.36	1.39	1.37	1.0
D	<0.05	<0.05	1.24	1.24	1.24	1.0
E	<0.05	<0.05	1.21	- 1.28	1.25	1.0
F	<0.05	<0.05	1.34	1.37	1.36	1.0
G	<0.05	<0.05	1.15	1.32	1.24	1.0
Н	<0.05	<0.05	1.16	1.25	1.20	1.0
i	< 0.05	<0.05	1.21	1.28	1.25	1.0
J	<0.05	<0.05	1.09	1.08	1.09	1.0

Mean \pm SD = 1.23 \pm 0.09 N = 10 % CV = 7.3



¹Refer to Appendix 2. ²Net values were calculated by subtracting the mean pre-fortification result from the mean post-fortification result.

TABLE 2.

1 CROSS-REACTIVITY RESULTS FOR THE CGA-77102 ENZYME IMMUNOASSAY

TEST SUBSTANCE	LOD	² LOQ	³ /50	PERCENT REACTIVITY RELATIVE TO CGA-77102	SET#	ANALYSIS Dates	NUMBER OF ANALYSES
4CGA-77102	. 0.05	0.10	0.48	100	990104A	1/4/99	. 24
COM-11102	. 0.05	0.10	0.40	100	990105A	1/5/99	24
•	•				990107A	1/7/99	
					990128A	1/28/99	
Metolachlor	0.05		1.0	55	990104A	1/4/99	6
Dimethanamid	0.20		4.2	12.9	990104A	1/4/99	6
CGA-380168	0.23		13.5	2.1	990105A	1/5/99	6
CGA-380108 CGA-41638	0.23		46	<1.0	990105A	1/5/99	6
Acetochlor			68	<1.0	990103A	1/4/99	6
CGA-40172				<1.0 <1.0	990104A	1/7/99	6
	•-		84				
Alachlor Alachlor-ESA		••	470	<1,0 NR	990104A	1/4/99 1/4/99	6
			· 5NR		990104A		6
Acetochlor-ESA	••	**	NR	NR	990104A	1/4/99	6
Metalaxyl			NR	NR	990107A	1/7/99	6
CGA-354743			NR	NR	990107A	1/7/99	6
CGA-50720			NR	NR	990107A	1/7/99	6
CGA-67869		••	NR	, NR	990107A	1/7/99	6
CGA-67125		**	NR	NR	990107A	1/7/99	6
CGA-37735	••		NR	NR	990104A	1/4/99	6.
CGA-51202		'	NR	NR	990105A	1/5/99	6
CGA-62826	••		NR	NR	990105A	1/5/99	6
CGA-46576			NR	NR	990105A	1/5/99	6 -
Dimethanamid-ESA	. 		NR	NR	990128A	1/28/99	6
CGA-37913		••	NR	NR	990128A	1/28/99	6

¹ Refer to Appendix 1.

⁵ NR - not reactive.

Limit of Quantitation (LOQ), the lowest level of fortification (in units of ng/mL) with which an acceptable recovery can be obtained. An LOQ was determined for CGA-77102 only.

³I₅₀, the concentration of test substance (in units of ng/mL) that yields half the response of the zero dose.

⁴ Parameters shown for CGA-77102 are the average of four determinations.

FIGURE 1. FLOW DIAGRAM FOR METHOD 1004-98

Measure sample pH

L

Add sample aliquot to microwell plate

.

Add enzyme conjugate to microwell plate

J.

Incubate one hour with shaking

1

Wash well three times

l

Add substrate to microwell plate

1

Incubate 30 minutes with shaking

↓

Stop color development by acidification

Ľ

Measure absorbance at 450 nm

FIGURE 2. PLATE LAYOUT CHART ON WHICH THE ANALYST WILL INDICATE THE POSITION OF SAMPLE AND STANDARD SOLUTIONS IN THE MICROWELL ASSAY PLATE

PLATE LAYOUT SHEET

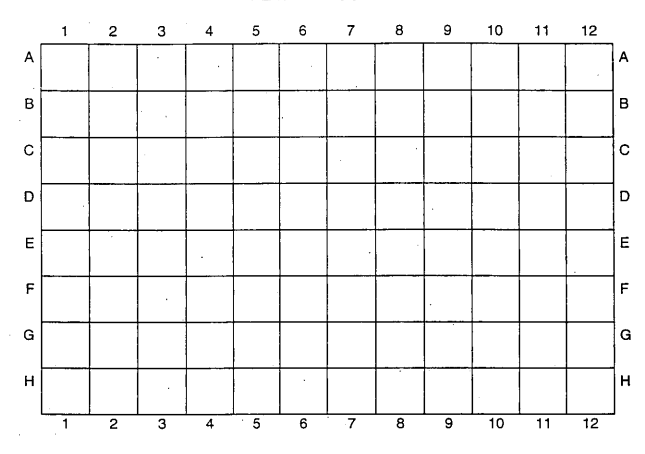


Plate ID Date Analyst Notebook/Page Comments



FIGURE 3. <u>A TYPICAL STANDARD CURVE OF THE CGA-77102</u> MICROWELL ASSAY

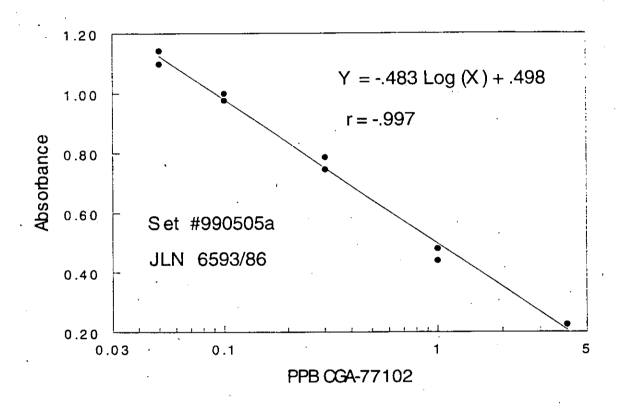


FIGURE 4. STRUCTURES OF TEST SUBSTANCES EVALUATED FOR CROSS-REACTIVITY

Metolachlor (CGA-24705)

FIGURE 4. STRUCTURES OF TEST SUBSTANCES EVALUATED FOR CROSS-REACTIVITY (continued)

CGA-354743

CGA-40172

FIGURE 4. <u>STRUCTURES OF TEST SUBSTANCES EVALUATED FOR CROSS-REACTIVITY</u> (continued)

CGA-67869

CGA-51202

CGA-62826

FIGURE 4. STRUCTURES OF TEST SUBSTANCES EVALUATED FOR CROSS-REACTIVITY (continued)

VI. REFERENCE

1. Brady, J. F. Interpretation of immunoassay data. In *Immunoanalysis of Agrochemicals: Emerging Technologies*. Karu, A., Nelson, J. Wong, R., Eds.; ACS Symposium Series 586; American Chemical Society: Washington, DC, 1995; pp 266-287.

APPENDIX 1

RESIDUE TEST REPORT NO. 1

Development Resources/Chemical Support Department Novartis Crop Protection, Inc. Greensboro, North Carolina

RESIDUE TEST REPORT TEST NUMBER: RI-MV-002-99 INVENTORY NUMBER: 18425.1 REPORT NO.: 1

PROJECT NUMBER:

H077102NAU950A

PROTOCOL NUMBER:

1004-98

TEST SUBSTANCES:

CGA-77102, Metolachior, Alachior, Alachior-ESA, Acetochlor, Acetochlor-ESA, Dimethanamid, Dimethanamid-ESA, Metalaxyl, CGA-354743, CGA-50720, CGA-40172, CGA-67869, CGA-67125, CGA-37735, CGA-41638, CGA-51202, CGA-62826, CGA-46576.

CGA-380168, CGA-37913

TEST SYSTEM:

Water

NO. OF ANALYSES:

144

LABORATORY:

Development Resources/

Chemical Support Department

DESCRIPTION: Distilled water samples were fortified with a range of concentrations of each test substance listed above to assess the reactivity of the CGA-77102 enzyme immunoassay as part of the method validation of "Draft" Analytical Method 1004-98.

STUDY DIRECTOR: Jody L. Needham, B.A.

Jody L. needham

DATE: 6/23/99

MANAGEMENT REPRESENTATIVE: Max W. Cheung, Ph.D.

SIGNATURE:

M, M, M

DATE: 06/23/99

SUBMITTED BY: Jody L. Needham and James F. Brady

QUALITY ASSURANCE AUDITOR:

SIGNATURE: Versa SCX for Yathlun Comphell DATE:

DISTRIBUTION:

J. Needham

M. W. Cheung

J. Brady

Main File

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RESIDUE TEST REPORT TEST NUMBER: RI-MV-002-99 INVENTORY NO.: 18425.1

REPORT NO.: 1

PROTOCOL NUMBER: 1004-98

METHOD OF ANALYSIS: "Draft" 1004-98

PROJECT NUMBER: H077102NAU950A

TEST SUBSTANCES: See below TEST SYSTEM: Water

TABLE 1. CROSS-REACTIVITY RESULTS FOR THE CGA-77102 ENZYME IMMUNOASSAY

·				PERCENT			
	•			REACTIVITY	•		NUMBER
TEST		· .	•	RELATIVE TO		ANALYSIS	OF
SUBSTANCE	1LOD	² LOQ	³ <i>I₅₀</i>	CGA-77102	SET#	DATE	ANALYSES
4.	•						_
⁴CGA-77102	0.05	0.10	0.48	100	990104A	1/4/99	24
					990105A	1/5/99	
,					990107A	1/7/99	
		•			990128A	1/28/99	
Metolachlor	0.05		1.0	55	990104A	1/4/99	. 6
Dimethanamid	0.20	 ,	4.2	12.9	990104A	1/4/99	6
CGA-380168	0.23	,	13.5	2.1	990105A	1/5/99	. 6
CGA-41638			46	<1.0 ·	990105A	1/5/99	6
Acetochlor			68	<1.0	990104A	1/4/99	6
CGA-40172			84	<1.0	990107A	1/7/99	6
Alachlor			470	<1.0	990104A	1/4/99	. 6
CGA-354743			⁵NR	NR	990107A	1/7/99	6
Metalaxyl			NR	NR	990107A	1/7/99	6
Acetochlor-ESA			NR	NR	990104A	1/4/99	6
Alachlor-ESA			NR	NR	990104A	1/4/99	6
Dimethanamid-							
ESA '		••	NR	NR	990128A	1/28/99	6
CGA-50720			NR	NR	990107A	1/7/99	6
CGA-67869			NR	NR	990107A	1/7/99	6
CGA-67125			NR	NR	990107A	1/7/99	6
CGA-37735			NR	NR	990104A	1/4/99	6
CGA-51202			NR	NR	990105A	1/5/99	6
CGA-62826			NR	NR	990105A`	1/5/99	6
CGA-46576			NR	NR	990105A	1/5/99	6
CGA-37913			NR	NR	990128A	1/28/99	. 6
				, ** *			_

Limit of Detection (LOD), the smallest dose (in units of ng/ml) that yields a response that is significantly different from the response of the zero dose.

NR - not reactive.

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Limit of Quantitation (LOQ), the lowest level of fortification (in units of ng/ml) that gives an acceptable recovery. An LOQ was determined for CGA-77102 only.

 l_{50} , the concentration of test substance (in units of ng/ml) that yields half the response of the zero dose.

Parameters shown for CGA-77102 are the average of four determinations.

RESIDUE TEST REPORT TEST NUMBER: <u>RI-MV-002-99</u> INVENTORY NO.: <u>18425.1</u> REPORT NO.: <u>1</u>

PROTOCOL NUMBER: 1004-98

METHOD OF ANALYSIS: "Draft" 1004-98

PROJECT NUMBER: H077102NAU950A

TEST SUBSTANCES: See below

TEST SYSTEM: Water

TABLE 2. TEST SUBSTANCES USED IN PROTOCOL 1004-98

NAME	CODE NUMBER	% PURITY	EXPIRATION DATE
CGA-77102	S95-1844	98.5	11/2000
Metolachlor	S94-1719	99.3	2/2000
² Alachlor	S86-0926	98.4	NON-GLP
² Alachior-ESA	DAH-XXII-24	99.4	11/2000
² Acetochlor	S97-2120	98	. 8/99
² Acetochlor-ESA	DAH-XXII-30	97.8	11/2000
² Dimethanamid	S97-2034	99.7	7/2000
² Dimethanamid-ESA	BPM-XVIII-19	approx. 87.5	NON-GLP
Metalaxyl	S94-1763	97.1	10/2000
¹ CGA-354743	DAH-XXVI-12	96.8	9/2000
¹ CGA-50720	DPS-I-74-2	>99.9	6/99
¹ CGA-40172	JAK-II-42 (IV)	95.9	6/99
¹ CGA-67869	DAH-XV-35	>99.9	6/99
¹CGA-67125	DPS-VII-41-1	99.9	12/99
¹ CGA-37735	JAK-II-42 (II)	99.2	3/2000
¹CGA-41638	S94-1765	95	9/99
¹CGA-51202	JAK-I-63	98.8	3/2000
¹ CGA-62826	BPM-I-48	98.0	11/2000
¹ CGA-46576	AW-I-43	97.4	6/99
¹ CGA-380168	DAH-XXIII-62	98.8	1/99
¹ CGA-37913	JAK-XVI-38	99.0	12/2000

¹These compounds lack common names.

²These compounds, or their metabolites, are not Novartis products.

APPENDIX 2

RESIDUE TEST REPORT NO. 2

Development Resources/Chemical Support Department Novartis Crop Protection, Inc. Greensboro, North Carolina

RESIDUE TEST REPORT

TEST NUMBER: RI-MV-002-99 INVENTORY NUMBER: 18425.1 REPORT NO.: 2

PROJECT NUMBER:

H077102NAU950A

PROTOCOL NUMBER:

1004-98

TEST SUBSTANCE:

CGA-77102

TEST SYSTEM:

Water

NO. OF ANALYSES:

22

LABORATORY: Development Resources/

Chemical Support Department

DESCRIPTION: Distilled water samples were fortified with CGA-77102 and analyzed pre- and post-fortification as part of the method validation of "Draft" Analytical Method 1004-98.

STUDY DIRECTOR: Jody L. Needham, B.A.

SIGNATURE: Ody L. needham

DATE: 6/23/99

MANAGEMENT REPRESENTATIVE: Max W. Cheung, Ph.D.

SIGNATURE:

m. W. Champ

DATE: 06/23/99

SUBMITTED BY: Jody L. Needham and James F. Brady

QUALITY ASSURANCE AUDITOR:

SIGNATURE: Stere SA SCIX for Mathieux Constsell

DATE: 6/23/99

DISTRIBUTION:

J. Needham

J. Brady

M. W. Cheung

Main File

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Novartis Number 1004-98

Page 38 of 40

Residue Test Report

Test Number: RI-MV-002-99 Inventory No.: 18425.1 Report No.: 2 Test Substance: Test System:

CGA-77102 Water

TABLE 1. RESULTS OF WATER STANDARD ADDITION EXPERIMENTS IN THE METHOD VALIDATION OF "DRAFT" 1004-98

	NB ref.	6593/64	6593/64	6593/64	6593/64	6593/64	6593/64	6593/64	6593/64	6593/64	6593/64
٠	Date	1/26/99	1/26/99	1/26/99	1/26/99	1/26/99	1/26/99	1/26/99	1/26/99	1/26/99	1/26/99
Analysis	Set Name	990126A	990126A	990126A	990126A	990126A	990126A	990126A	990126A	990126A	990126A
CGA-77102	Level (ppb)	0.10	0.10	0.10	0.10	0.10	0.10	0.10	0.10	0.10	0.10
	Net	0.09	0.08	0.10	0.09	0.13	0.10	0.09	0.10	0.09	0.09
ppb CGA-77102 Equivalents Found	fication	0.09	0.09	0.09	0.09	0.16	0.10	0.10	0.10	0.10	0.09
	Post-fortification	60.0	0.08	0.10	0.09	0.10	60.0	0.08	0.10	0.09	0.08
	ppb CGA-77102 Equ Pre-fortification	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05
	Pre-forti	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05
Distilled Water	Samples	∢	6 0	ر ن	۵	ш	Ľ.	IJ	エ	· .	¬

¹Net values were calculated by subtracting the mean pre-fortification result from the mean post-fortification result.

Mean \pm SD = 0.10 \pm 0.01

= 0.13

z >

Protocol Number: Project Number:

1004-98 H077102NAU950A "Draft" 1004-98

Method of Analysis:

Residue Test Report

Test Number: RI-MV-002-99 Inventory No.: 18425.1 Report No.: 2

CGA-77102 Water Test Substance: Test System:

TABLE 1. RESULTS OF WATER STANDARD ADDITION EXPERIMENTS IN THE METHOD VALIDATION OF "DRAFT" 1004-98 (continued) 1004-98 H077102NAU950A "Draft" 1004-98 Method of Analysis:

		NB ref.	6593/86	6593/86	6593/86	6593/86	6593/86	6593/86	6593/86	6593/86	6593/86	6593/86
CGA-77102 02 Fortivalents Found Fortification		Date	2/5/99	5/5/99	5/5/99	5/2/99	5/5/99	5/2/99	2/2/99	5/2/99	2/5/99	5/5/99
	Analysis	Set Name	990505A									
	Fortification	Level (ppb)	1.0	1.0.	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
		Net	1.12	1.17	1.37	1.24	1.25	1.36	1.24	1.20	1.25	1.09
	s Found	fication	1.22	1.23	1.39	1.24	1.28	1.37	1.32	1.25	1.28	1.08
	2 Equivalents	Post-fortification	1.03	1.10	1.36	1.24	1.21	1.34	1.15	1.16	1.21	1.09
	CGA-7710	ppb CGA-77103 Pre-fortification	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05
	qdd	Pre-forti	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05	<0.05
	Distilled Water	Samples	¥	B	ပ		ш	u.	ഗ	I	_	_

Mean \pm SD = 1.23 \pm 0.09 10 0.07 z > ¹Net values were calculated by subtracting the mean pre-fortification result from the mean post-fortification result.

Protocol Number: Project Number: