TEXT SEARCHABLE DOCUMENT

Common Name	Flumioxazin					
PC Code	129034					
Chemical Name of the Test S	ubstance					
IUPAC	N-(7-fluoro-3,4-dil cyclohex-1-ene-1,2			p-2-ynyl-2H	-1,4-benzoxaz	in-6-yl)
CAS	2-[7-Fluoro-3,4-dil 4,5,6,7-tetrahydro-				I-1,4-benzoxa	zin-6-y1]-
CAS Number	103361-09-7					
Chemical Structure	See back					
Empirical Formula	-					
Molecular Formula	C ₁₉ H ₁₅ FN ₂ O ₄					
Molecular Weight (g/mole)	354.3					
SMILES Code	-					
	Temperature (° C)	Value		Source/Con	ments
Solubility (mg/L)	•		_			
Water						
Solvent A						
Solvent B						
Solvent C						
Specific Gravity						
Kow						
Henry's Law Constant (atm-r	n3/mol)					
Measured					•	
Estimated						
Vapor Pressure (mmHg)						
Dissociation Constant						
pKa1						
pKa2				·		
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pKb1			
pKb2			
Boiling Point (° C)			
Melting Point (° C)			

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Anaerobic Aquatic Metabolisi	n (162-3)	
Acceptability of the Study	Acceptable	· · · · · · · · · · · · · · · · · · ·
Purpose of the Study	Registration (ne	w use)
Common Name for the Test Substance	Flumioxazin	
PC Code	129034	
If test substance is a degradate, provide parent's common name	-	
If test substance is a degradate, is it also a registered pesticide (give the common name)?	-	
Chemical Name for the Test Sul	ostance	
IUPAC		dihydro-3-oxo-4-prop-2-ynyl-2H-1,4-benzoxazin-6- ne-1,2-dicarboxamide
CAS	-	-dihydro-3-oxo-4-(2-propynyl)-2H-1,4-benzoxazin-6- nydro-1H-isoindole-1,3(2H)-dione
CAS Number	103361-09-7	
Chemical Structure	See back	
Study Temperature (°C)	25	
Solid Phase (if solid and aqueor	us phases were no	t reported separately, report the Combined Phase)
Half-life (days)	48.5	
Soil Texture	Clay	
Sand (%)	14	
Silt (%)	28	
Clay (%)	58	
Organic Matter (%)	5	
Organic Carbon (%)	3	
рН	6.4	
Soil Mapping Unit	-	
Soil Taxonomy Classification	-	

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Initial Concentration (mg/kg)	-				
Major Degradate(s)	Common Name	Chemical Structure	Chemical Name IUPAC	CAS	CAS Number
Degradate #1	APF	See back	6-Amino-7-fluoro-4-(2- propenyl)-2H-1,4- benzoxazin-3(4H)-one		
Degradate #2	DAPF	See back		-	-
Major Degradate #1 (APF):	Days to Maxin	num Concent	tration	21	
-	Maximum Cor mg/kg	ncentration	· · · · ·	0.664	
	Maximum Cor % of ap			19.3	
	Half-life (days))		-	
Major Degradate #2 (DAPF):	Days to Maxin	num Concent	tration	360	
	Maximum Cor mg/L	ncentration		0.462	
	Maximum Concentration % of applied			13.5	
	Half-life (days)			-	-
Aqueous Phase	1				
Half-life (days)	79.7			_	
Initial Concentration (mg/L)	0.858				
Major Degradate(s)	Common Name	Chemical Structure	Chemical Name	CAS	CAS Number
Degradate #1	APF	See back	IUPAC 6-Amino-7-fluoro-4-(2- propenyl)-2H-1,4- benzoxazin-3(4H)-one	CAS -	
Major Degradate #1 (APF):	Days to Maximum Concentration			2	
	Maximum Cor mg/L	centration	· · · · · · · · · · · · · · · · · · ·	0.421	
	Maximum Cor % of ap			49.1	
	Half-life (days))		-	

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MRID	45914602	
Source	DER	
Date DER Reviewed	July 15, 2003 b	y Dynamac Corporation.
Comments	sediment collect potentials were	label. Study conducted for 360 days. Pond water and ted from Greenville, Mississippi. In water, redox -231.5 to -160.8 mV and oxygen content was 0.00 ppm. C-482-HA" was maximum of 16.9% of applied at 182 tem.

Anaerobic Aquatic Metabolisr	n (162-3)
Acceptability of the Study	Acceptable
Purpose of the Study	Registration (new use)
Common Name for the Test Substance	Flumioxazin
PC Code	129034
If test substance is a degradate, provide parent's common name	
If test substance is a degradate, is it also a registered pesticide (give the common name)?	
Chemical Name for the Test Sub	ostance
IUPAC	N-(7-fluoro-3,4-dihydro-3-oxo-4-prop-2-ynyl-2H-1,4-benzoxazin-6- yl)cyclohex-1-ene-1,2-dicarboxamide
CAS	2-[7-Fluoro-3,4-dihydro-3-oxo-4-(2-propynyl)-2H-1,4-benzoxazin-6- yl]-4,5,6,7-tetrahydro-1H-isoindole-1,3(2H)-dione
CAS Number	103361-09-7
Chemical Structure	See back
Study Temperature (°C)	25
Solid Phase (if solid and aqueou	as phases were not reported separately, report the Combined Phase)
Half-life (days)	42.0
Soil Texture	Clay
Sand (%)	14
Silt (%)	28
Clay (%)	58
Organic Matter (%)	5
Organic Carbon (%)	3
рН	6.4
Soil Mapping Unit	-
Soil Taxonomy Classification	-

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Initial Concentration (mg/kg)	-					
Major Degradate(s)	Common Name	Chemical Structure	Chemical Name	CAS	CAS Number	
Degradate #1	HPA	See back		-	-	
Major Degradate #1 (HPA):	Days to Maxim	um Concent	ration	42		
	Maximum Concentration mg/kg				0.582	
	Maximum Con % of ap			17.0		
·	Half-life (days))				
Aqueous Phase						
Half-life (days)	87.7					
Initial Concentration (mg/L)	0.858					
Major Degradate(s)	Common Name	Chemical Structure	Chemical Name	CAS	CAS Number	
Degradate #1	THPA	See back	-	-		
Degradate #2	HPA	See back	-	-	-	
Major Degradate #1 (THPA):	Days to Maxim	um Concent	ration	1		
	Maximum Con mg/L	centration		0.360		
· .	Maximum Con % of ap	41.9	41.9			
	Half-life (days)			-		
Major Degradate #2 (HPA):	Days to Maxim	um Concent	ration	360		
	Maximum Con mg/L	centration		0.625		
	Maximum Con % of ap			72.8		
	Half-life (days)			-		
MRID	45914602					
Source	DER					

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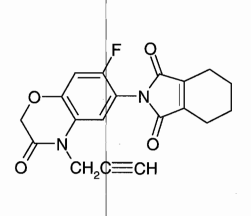
Date DER Reviewed	July 15, 2003 by Dynamac Corporation.
Comments	[Tetrahydrophthaloyl- ¹⁴ C] label. Study conducted for 360 days. Pond water and sediment collected from Greenville, Mississippi. In water, redox potentials were -237.3 to -184.0 mV and oxygen content was 0.00 ppm.

Contractor submittal date: July 15, 2003. Chemical structures of the parent and major degradates identified are included on the following pages.

Chemical Structures of Parent and Major Degradates Identified in the Study

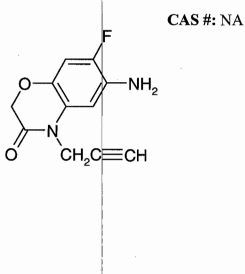
Parent compound: Flumioxazin

- **IUPAC name:** N-(7-fluoro-3,4-dihydro-3-oxo-4-prop-2-ynyl-2H-1,4-benzoxazin-6-yl)cyclohex-1-ene-1,2-dicarboxamide
- CAS name: 2-[7-Fluoro-3,4-dihydro-3-oxo-4-(2-propynyl)-2H-1,4-benzoxazin-6-yl]-4,5,6,7-tetrahydro-1Hisoindole-1,3(2H)-dione
- **CAS #:** 103361-09-7



Degradate 1: APF

IUPAC name: 6-Amino-7-fluoro-4-(2-propenyl)-2H-1,4-benzoxazin-3(4H)-one CAS name: NA



Degradate 2: DAPF (Dihydro-APF)

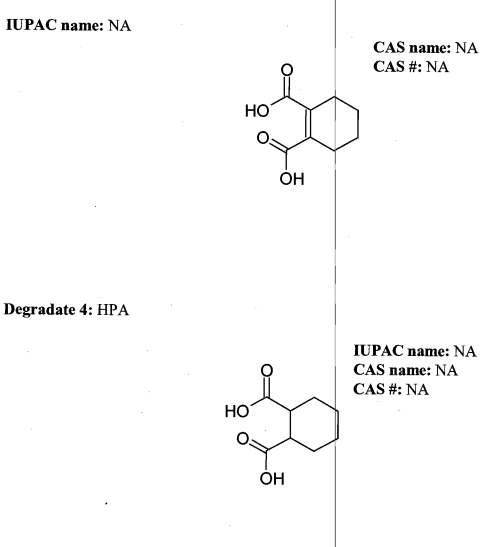
IUPAC name: NA CAS name: NA

۰ŅΗ O CH₂CH -CH2

CAS #: NA

Degradate 3: THPA

IUPAC name: NA



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