# FUNGICIDE

# **Isofetamid** New chemical class of SDHI

**Physico-Chemical Properties** 

Chemical structure

Molecular weight : 359.48 Molecular formula : C<sub>20</sub>H<sub>25</sub>NO<sub>3</sub>S

Vapour pressure : 4.2 x 10<sup>-7</sup> Pa (25°C) Water solubility : 5.33 mg/L (20°C) Form : White Solid (powder) Development code : IKF-5411

Isofetamid is a novel SDHI (Succinate Dehydrogenase Inhibitor, FRAC code 7) fungicide discovered and under development by ISK.

Isofetamid is a new chemical group (phenyl-oxo-ethyl thiophene amide) based on its thiophene carboxamide moiety. Due to this unique chemical structure, Isofetamid remains highly effective against the majority of fungal isolates that have developed resistance to other SDHI fungicides.

As a broad-spectrum fungicide, Isofetamid exhibits excellent activity against a broad range of fungi, but is especially effective on the Ascomycota (such as *Botrytis* spp., *Sclerotinia* spp., *Monilinia* spp., *Venturia* spp.) at low dose rates.

In addition to its outstanding efficacy, Isofetamid has no negative impacts on beneficial insects and mites, making it an excellent choice for integrated pest management programs.

H<sub>3</sub>C

сн.

tolyl)-2-oxoethyl]-3-methylthiophene-

Class : Phenyl-oxo-ethyl thiophene amide

IUPAC name : N-[1,1-dimethyl-2-(4-isopropoxy-o-

2-carboxamide



## **Toxicology & Ecotoxicology**

Rat LD <sup>50</sup> (oral) : > 2,000 mg/kg (f) Rat LD <sup>50</sup> (dermal) : > 2,000 mg/kg (m/f) Rat LC <sup>50</sup> (inhalation) : > 4.82 mg/L (m/f)	
Skin irritation : non irritant (rabbit) Eye irritation : slightly irritating to eyes (rabbit) Skin sensitization : not a sensitizer (mouse, guinea )	pig)
Avian LD <sub>50</sub> (acute oral) : $> 2,000$ mg/kg (quail, m/f) Avian LD <sub>50</sub> (subacute oral) : $> 5,000$ ppm in feed (quail)	
Fish LC <sub>50</sub> : $> 7.12$ mg/L (carp, 96 h)	
Bees $LD_{50}$ (acute oral) : > 30 µg a.i./bee (48 h) Bees $LD_{50}$ (acute contact) : > 100 µg a.i./bee (48 h)	
Daphnia magna EC₅₀ : 4.7 mg/L (48 h)	

## Product

Trade Names	KENJA, ZENBY, KRYOR, HAREGI, KABUTO, ASTUN, etc.			
Formulations	40%SC			
<b>Registered Countries</b>	Asia	China, Japan, Korea		
Europe Belgium, Bulgaria, Czech Republic, France, Germany, Greece, Hungary, Italy, Luxe Poland, Portugal, Romania, Spain, Slovenia, UK, etc.		Belgium, Bulgaria, Czech Republic, France, Germany, Greece, Hungary, Italy, Luxembourg, Poland, Portugal, Romania, Spain, Slovenia, UK, etc.		
	Oceania	Australia		
	Americas	Brazil, Canada, Chile, Colombia, Ecuador, Mexico, Peru, USA, etc.		

Always read and follow the product label instructions in your country.

## **Characteristics**

- SDHI class (FRAC code 7) with broad-spectrum fungicidal activity
- Good persistence and rainfastness
- Flexible molecular structure makes it effective on major SDHI resistant isolates
  Extension of shelf life by pre-harvest application
- · Inhibits all growth stages of fungal life cycle

• High safety for crop and beneficial organisms



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## **Isofetamid** New chemical class of SDHI

## Mode of Action

Isofetamid acts specifically on the succinate dehydrogenase (SDH) of Complex II, a key enzyme of the mitochondrial respiratory chain at the crossroads of two metabolic pathways essential to fungal cell life. By inhibiting SDH, Isofetamid impairs energy (ATP) production by the respiratory chain and the synthesis of amino acids, lipids and fatty acids (metabolites essential to cell function) at the Krebs cycle stage.

### Advantages of Isofetamid for resistance management

Isofetamid can control numerous isolates with confirmed resistance to other SDHI fungicides, including SdhB H272R and H272Y, which are the two most common field-collected isolates.

Research has confirmed that Isofetamid fits the mutated binding pocket of SDHI-resistant fungal isolates (SdhB H272R and H272Y). It is hypothesized that the unique molecular structure of Isofetamid gives the molecule flexibility at the binding site, allowing Isofetamid to retain efficacy on these mutants. Other SDHI fungicides have a rigid structure, are unable to bind at sites where mutations have occurred, and are therefore ineffective as control options.



Isofetamid SDHI A SDHI B SDHI C SDHI D Standard S S S S S SdhB H272Y s R S MR R SdhB H272R S R S S S SdhB N230I R R s MR MR SdhB P225F R R R R R SdhB H272L R R R MR R S:Sensitive, MR:Moderately Resistant, R: Resistant

**Fungicidal spectrum** 

<i>Botrytis</i> spp.	Cercospora spp.	
<i>Monilinia</i> spp.	Mycosphaerella spp.	
<i>Sclerotinia</i> spp.	Passalora spp.	
<i>Venturia</i> spp.	Pseudocercospora spp.	
Corynespora spp.	<i>Elsinoë</i> spp.	
<i>Didymella</i> spp.	Ramularia spp.	
Wilsonomyces spp.	Alternaria spp.	
Cladosporium spp.	Rhizopus spp.	
<i>Diaporthe</i> spp.	and Powdery mildew	etc.

#### Control of fruit rots on strawberries during storage









### Multiple disease control (Cucumber field trials in Japan)

Disease name	Commorpial	Disease Incidence or Severity (%)		
	standard	lsofetamid 266 ppm	Standard	Untreated control
Sclerotinia rot Sclerotinia sclerotiorum	Dicarboximides 250 ppm	1.4	0.5	44.5
Gray mold Botrytis cinerea	Anilino-pyrimidines 200 ppm	0.2	1.6	34.4
Gummy stem blight Didymella bryoniae	Dicarboximides 500 ppm	0.3	0.4	16.8
Corynespora leaf spot Corynespora cassiicola	TPN 400 ppm	0.9	2.4	11.7
Powdery mildew Podosphaera xanthii	Quinoxaline 83 ppm	0.5	6.7	64.5



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Simulation modeling of enzyme 3D structure