

Safety Data Sheet

according to the REACH Regulation (EC) 1907/2006 amended by Regulation (EU) 2020/878 Revision date: 10/6/2022 Supersedes version of: 10/16/2018 Version: 3.31

SECTION 1: Identification of the substance/mixture and of the company/undertaking

1.1. Product identifier

Product form : Mixture

Product name : Rhizopon AA Powder 0.5% UFI : CADC-E38Q-287H-DCD7

Product group : Trade product

1.2. Relevant identified uses of the substance or mixture and uses advised against

1.2.1. Relevant identified uses

Main use category : Professional use

Use of the substance/mixture : Plant growth regulator that promotes root formation.

| Title | Life cycle stage | Use descriptors |
|-------------------------|------------------|-----------------|
| Rhizopon AA Powder 0.5% | Professional | PC27 |

Full text of use descriptors: see section 16

1.2.2. Uses advised against

No additional information available

1.3. Details of the supplier of the safety data sheet

Rhizopon B.V.

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The Netherlands

T + 31(0) 71 3415146 - F + 31(0) 71 3415829 <u>info@rhizopon.com</u> - <u>www.rhizopon.com</u>

1.4. Emergency telephone number

No additional information available.

SECTION 2: Hazards identification

2.1. Classification of the substance or mixture

Classification according to Regulation (EC) No. 1272/2008 [CLP]

Not classified

Adverse physicochemical, human health and environmental effects

To our knowledge, this product does not present any particular risk, provided it is handled in accordance with good occupational hygiene and safety practice.

2.2. Label elements

Labelling according to Regulation (EC) No. 1272/2008 [CLP]

EUH-statements : EUH210 - Safety data sheet available on request.

EUH401 - To avoid risks to human health and the environment, comply with the instructions

for use.

Extra phrases : SP1 Do not contaminate water with the product or its container.

2.3. Other hazards

This substance/mixture does not meet the PBT criteria of REACH regulation, annex XIII This substance/mixture does not meet the vPvB criteria of REACH regulation, annex XIII Contains no PBT/vPvB substances ≥ 0.1% assessed in accordance with REACH Annex XIII

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The mixture does not contain substance(s) included in the list established in accordance with Article 59(1) of REACH for having endocrine disrupting properties, or is not identified as having endocrine disrupting properties in accordance with the criteria set out in Commission Delegated Regulation (EU) 2017/2100 or Commission Regulation (EU) 2018/605 at a concentration equal to or greater than 0,1 %

SECTION 3: Composition/information on ingredients

3.1. Substances

Not applicable

3.2. Mixtures

Comments : Mentioned percentages are in (w/w %)

| Product name | Product identifier | % w/w (% w/w) | Classification according to Regulation (EC) No. 1272/2008 [CLP] |
|----------------------------|--|------------------|---|
| Talc (Mg3H2(SiO3)4) | CAS-No.: 14807-96-6 EC-No.: 238-877-9 REACH-no: 01-2120140278- 58 | > 50 | Not classified |
| 4-(indol-3-yl)butyric acid | CAS-No.: 133-32-4 EC-No.: 205-101-5 REACH-no: - | 0.1 – 1 | Acute Tox. 4 (Oral), H302 Repr. 2, H361fd |

Full text of H- and EUH-statements: see section 16

SECTION 4: First aid measures

4.1. Description of first aid measures

First-aid measures general : In all cases of doubt, or when symptoms persist, seek medical attention.

First-aid measures after inhalation : Remove person to fresh air and keep comfortable for breathing. In all cases of doubt, or

when symptoms persist, seek medical attention.

First-aid measures after skin contact : Wash skin with plenty of water and soap. If skin irritation or rash occurs: Get medical

advice/attention.

First-aid measures after eye contact : Immediately rinse with water for a prolonged period while holding the eyelids wide open.

 $\label{lem:contact} \mbox{Remove contact lenses, if present and easy to do. Continue rinsing. If eye irritation persists:}$

Get medical advice/attention.

First-aid measures after ingestion : Never give anything by mouth to an unconscious person. Rinse mouth out with water. Do

NOT induce vomiting. Call a poison center or a doctor if you feel unwell.

4.2. Most important symptoms and effects, both acute and delayed

No additional information available

4.3. Indication of any immediate medical attention and special treatment needed

Treat symptomatically.

SECTION 5: Firefighting measures

5.1. Extinguishing media

Suitable extinguishing media : Water spray. Dry powder. Foam. Carbon dioxide (CO2).

5.2. Special hazards arising from the substance or mixture

Hazardous decomposition products in case of fire : Toxic fumes may be released.

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5.3. Advice for firefighters

Protection during firefighting

: Do not attempt to take action without suitable protective equipment. Self-contained breathing apparatus. Complete protective clothing.

SECTION 6: Accidental release measures

6.1. Personal precautions, protective equipment and emergency procedures

6.1.1. For non-emergency personnel

Emergency procedures : Ventilate spillage area.

6.1.2. For emergency responders

Protective equipment : Do not attempt to take action without suitable protective equipment. For further information

refer to section 8: "Exposure controls/personal protection".

6.2. Environmental precautions

Avoid release to the environment. For a large spillage, contain the spillage by bunding.

6.3. Methods and material for containment and cleaning up

Methods for cleaning up : Ventilate area. Dust deposited may be vacuum cleaned. Take up mechanically (sweeping,

shovelling) and collect in suitable container for disposal. Knock down dust with water spray

jet. After cleaning, flush traces away with water.

Other information : Dispose of materials or solid residues at an authorized site.

6.4. Reference to other sections

Concerning personal protective equipment to use, see section 8. Concerning disposal elimination after cleaning, see section 13.

SECTION 7: Handling and storage

7.1. Precautions for safe handling

Precautions for safe handling : Ensure good ventilation of the work station. Wear personal protective equipment. Use only

outdoors or in a well-ventilated area.

Hygiene measures : Do not eat, drink or smoke when using this product. Always wash hands after handling the

product.

7.2. Conditions for safe storage, including any incompatibilities

Storage conditions : Store in a well-ventilated place. Keep cool. Keep container tightly closed. Store under dry

conditions. Keep away from food, drink and animal feedingstuffs. Original packaging.

Storage temperature : 10 - 20 °C

Packaging materials : Polypropylene. Polyethylene.

7.3. Specific end use(s)

No supplementary information available.

SECTION 8: Exposure controls/personal protection

8.1. Control parameters

8.1.1 National occupational exposure and biological limit values

| Talc (Mg3H2(SiO3)4) (14807-96-6) | |
|---|-------------------------|
| United Kingdom - Occupational Exposure Limits | |
| Local name | Talc |
| WEL TWA (OEL TWA) [1] | 1 mg/m³ respirable dust |

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| Talc (Mg3H2(SiO3)4) (14807-96-6) | |
|----------------------------------|---------------------------------------|
| Regulatory reference | EH40/2005 (Fourth edition, 2020). HSE |

8.1.2. Recommended monitoring procedures

No additional information available

8.1.3. Air contaminants formed

No additional information available

8.1.4. DNEL and PNEC

| 8.1.4. DNEL and PNEC | | | |
|--|--------------------------------|--|--|
| Talc (Mg3H2(SiO3)4) (14807-96-6) | | | |
| DNEL/DMEL (Workers) | | | |
| Acute - systemic effects, inhalation | 2.16 mg/m³ | | |
| Acute - local effects, inhalation | 3.6 mg/m³ | | |
| Long-term - systemic effects, dermal | 43.2 mg/kg bodyweight/day | | |
| Long-term - local effects, dermal | 4.54 mg/m³ | | |
| Long-term - systemic effects, inhalation | 2.16 mg/m³ | | |
| Long-term - local effects, inhalation | 3.6 mg/m³ | | |
| DNEL/DMEL (General population) | DNEL/DMEL (General population) | | |
| Acute - systemic effects, inhalation | 1.08 mg/m³ | | |
| Acute - systemic effects, oral | 160 mg/kg bodyweight | | |
| Acute - local effects, inhalation | 1.8 mg/m³ | | |
| Long-term - systemic effects,oral | 160 mg/kg bodyweight/day | | |
| Long-term - systemic effects, inhalation | 1.08 mg/m³ | | |
| Long-term - systemic effects, dermal | 21.6 mg/kg bodyweight/day | | |
| Long-term - local effects, dermal | 2.27 mg/cm ² | | |
| Long-term - local effects, inhalation | 1.8 mg/m³ | | |
| PNEC (Water) | | | |
| PNEC aqua (freshwater) | 597.97 mg/l | | |
| PNEC aqua (marine water) | 141.26 mg/l | | |
| PNEC aqua (intermittent, freshwater) | 597.97 mg/l | | |
| PNEC aqua (intermittent, marine water) | 141.26 mg/l | | |
| PNEC (Sediment) | PNEC (Sediment) | | |
| PNEC sediment (freshwater) | 31.33 mg/kg dwt | | |
| PNEC sediment (marine water) | 3.13 mg/kg dwt | | |

8.1.5. Control banding

No additional information available

8.2. Exposure controls

8.2.1. Appropriate engineering controls

Appropriate engineering controls:

Ensure good ventilation of the work station.

8.2.2. Personal protection equipment

Personal protective equipment:

Gloves. Safety glasses. Protective clothing.

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Personal protective equipment symbol(s):







8.2.2.1. Eye and face protection

Eye protection:

Use eye protection according to EN 166, designed to protect powders and dusts

8.2.2.2. Skin protection

Skin and body protection:

Wear suitable protective clothing. Overall.

Hand protection:

In case of repeated or prolonged contact wear gloves. Recommendation: Wear suitable gloves tested to EN374. Suitable material: Nitrile rubber (NBR), Neoprene. Layer thickness: No data available. Breakthrough time: refer to the recommendations of the supplier. Choosing the proper glove is a decision that depends not only on the type of material, but also on other quality features, which differ for each manufacturer. The exact break through time has to be found out by the manufacturer of the protective gloves and has to be observed.

8.2.2.3. Respiratory protection

Respiratory protection:

Provide sufficient air exchange and/or exhaust. In case of inadequate ventilation wear respiratory protection.

8.2.2.4. Thermal hazards

No additional information available

8.2.3. Environmental exposure controls

Other information:

Do not eat, drink or smoke when using this product. Always wash your hands immediately after handling this product, and once again before leaving the workplace.

SECTION 9: Physical and chemical properties

9.1. Information on basic physical and chemical properties

Physical state : Solid
Colour : Light grey.
Appearance : Powder.
Odour : odourless.
Odour threshold : Not available

Melting point : 124 °C (4-(indol-3-yl)butyric acid) (OECD 102 method)

Freezing point : Not applicable

Boiling point : > 250 °C (4-(indol-3-yl)butyric acid) (OECD 102 method)

Flammability : Non flammable.

Explosive properties : Product is not explosive.

Explosive properties : Product is not explosive properties : Not oxidizing.

Explosive limits : Not applicable : Not applicable

Explosive limits : Not applicable
Lower explosion limit : Not applicable
Upper explosion limit : Not applicable
Flash point : Not applicable
Auto-ignition temperature : Not applicable

 $\label{eq:composition} Decomposition temperature \qquad \qquad : \ > 250 \ ^{\circ}\text{C} \ (4\text{-(indol-}3\text{-yl})butyric acid) \ (OECD \ 102 \ method)$

pH : 9 (10% solution in water) (20.5°C)

pH solution : Not available
Viscosity, kinematic : Not applicable
Viscosity, dynamic : Not applicable

Solubility : Water: insoluble in water (14,7 g/l @ pH7 20°C (4-(indol-3yl)butyric acid))

Partition coefficient n-octanol/water (Log Kow) : Not available

Vapour pressure : 1.1 Pa @20°C (4-(indol-3-yl)butyric acid) (EEC A.4/OECD 104)

Vapour pressure at 50 °C : Not available

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Density : 0.72 g/ml Relative density : Not applicable Relative vapour density at 20 °C : Not applicable Not available Particle size Particle size distribution Not available Particle shape Not available Particle aspect ratio Not available Particle aggregation state Not available Particle agglomeration state : Not available Particle specific surface area : Not available Particle dustiness : Not available

| 4-(indol-3-v | d)butyric acid | (133-32-4) |
|---------------|----------------|------------|
| T (IIIGOI O) | INDULYIIO GOIG | 1100 02 7/ |

| Vapour pressure | 1.0X 10-5 Pa @ 20 °C |
|-----------------|----------------------|
| Vapour prosouro | 1.0% 10 0 1 4 @ 20 0 |

9.2. Other information

9.2.1. Information with regard to physical hazard classes

No additional information available

9.2.2. Other safety characteristics

No additional information available

SECTION 10: Stability and reactivity

10.1. Reactivity

The product is non-reactive under normal conditions of use, storage and transport.

10.2. Chemical stability

Stable under normal conditions.

10.3. Possibility of hazardous reactions

No dangerous reactions known under normal conditions of use.

10.4. Conditions to avoid

None under recommended storage and handling conditions (see section 7). Avoid creating or spreading dust.

10.5. Incompatible materials

Oxidizing agent. Strong acids.

10.6. Hazardous decomposition products

Under normal conditions of storage and use, hazardous decomposition products should not be produced.

SECTION 11: Toxicological information

11.1. Information on hazard classes as defined in Regulation (EC) No 1272/2008

Acute toxicity (oral) : Not classified
Acute toxicity (dermal) : Not classified
Acute toxicity (inhalation) : Not classified

| Rhizopon AA Powder 0.5% | |
|-------------------------|--------------|
| LD50 oral rat | > 5000 mg/kg |
| LD50 dermal rabbit | > 2000 mg/kg |
| LC50 Inhalation - Rat | 2.4 mg/l |

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| 4-(indol-3-yl)butyric acid (133-32-4) | |
|---------------------------------------|--|
| LD50 oral | 1925 g/kg mouse |
| LD50 dermal rat | > 750 mg/kg |
| ATE oral | 500 mg/kg bodyweight |
| Skin corrosion/irritation | : Not classified pH: 9 (10% solution in water) (20.5°C) |
| Serious eye damage/irritation | : Not classified pH: 9 (10% solution in water) (20.5°C) |
| Respiratory or skin sensitisation | : Not classified |
| Germ cell mutagenicity | : Not classified |
| Carcinogenicity | : Not classified |
| Reproductive toxicity | : Not classified |
| STOT-single exposure | : Not classified |
| STOT-repeated exposure | : Not classified |
| Talc (Mg3H2(SiO3)4) (14807-96-6) | |
| NOAEL (oral, rat, 90 days) | 100 mg/kg bodyweight Animal: rat, Guideline: OECD Guideline 452 (Chronic Toxicity Studies) |
| Aspiration hazard | : Not classified |
| Rhizopon AA Powder 0.5% | |
| Viscosity, kinematic | Not applicable |

11.2. Information on other hazards

No additional information available

SECTION 12: Ecological information

12.1. Toxicity

Ecology - general : The product is not considered harmful to aquatic organisms nor to cause long-term adverse

effects in the environment.

Hazardous to the aquatic environment, short–term

(acute)

: Not classified

Hazardous to the aquatic environment, long-term

: Not classified

(chronic)

| alc (Mg3H2(SiO3)4) (14807-96-6) | | |
|---------------------------------------|--|--|
| LC50 - Fish [1] | 89.581 – 110 g/l | |
| LC50 - Fish [2] | 110000 mg/l Test organisms (species): other: | |
| EC50 - Crustacea [1] | 36.812 g/l | |
| EC50 72h - Algae [1] | 7.203 g/l | |
| EC50 96h - Algae [1] | 7202.7 mg/l Test organisms (species): other: | |
| NOEC (chronic) | 1459798 mg/l Test organisms (species): other: Duration: '30 d' | |
| 4-(indol-3-yl)butyric acid (133-32-4) | | |
| LC50 - Fish [1] | 250 mg/l 96 h, Oncorhynchus mykiss (Rainbow trout) | |
| LC50 - Fish [2] | 210 mg/l 96 h, Leuciscus idus (golden orfe) | |
| EC50 - Crustacea [1] | 112 mg/l 48 h - Daphnia magna | |
| EC50 72h - Algae [1] | 101 mg/l EyC50 | |
| EC50 72h - Algae [2] | 118 mg/l EbC50, 48 h - P.subcapitata | |

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12.2. Persistence and degradability

| Rhizopon AA Powder 0.5% | |
|---------------------------------------|---|
| Persistence and degradability | Potentially biodegradable. |
| 4-(indol-3-yl)butyric acid (133-32-4) | |
| Persistence and degradability | Readily biodegradable (Modified Sturm test; OECD 301B). |

12.3. Bioaccumulative potential

| Rhizopon AA Powder 0.5% | |
|---|--|
| Bioaccumulative potential | Bioaccumulation unlikely. |
| Talc (Mg3H2(SiO3)4) (14807-96-6) | |
| Partition coefficient n-octanol/water (Log Pow) | -9.4 @ 25 °C / pH 7 |
| 4-(indol-3-yl)butyric acid (133-32-4) | |
| Bioaccumulative potential | Low potential for bioaccumulation (Log Kow < 4). |

12.4. Mobility in soil

| Rhizopon AA Powder 0.5% | |
|-------------------------|---|
| Ecology - soil | No supplementary information available. |

12.5. Results of PBT and vPvB assessment

Rhizopon AA Powder 0.5%

This substance/mixture does not meet the PBT criteria of REACH regulation, annex XIII

This substance/mixture does not meet the vPvB criteria of REACH regulation, annex XIII

12.6. Endocrine disrupting properties

No additional information available

12.7. Other adverse effects

Additional information : Avoid release to the environment.

SECTION 13: Disposal considerations

13.1. Waste treatment methods

Regional legislation (waste) : Disposal must be done according to official regulations.

Waste treatment methods : Dispose of contents/container in accordance with licensed collector's sorting instructions.

SECTION 14: Transport information

In accordance with ADR / IMDG / IATA / RID

| ADR | IMDG | IATA | RID |
|-------------------------------|----------------|----------------|----------------|
| 14.1. UN number or ID number | | | |
| Not applicable | Not applicable | Not applicable | Not applicable |
| 14.2. UN proper shipping name | | | |
| Not applicable | Not applicable | Not applicable | Not applicable |

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| ADR | IMDG | IATA | RID |
|--|----------------|----------------|----------------|
| 14.3. Transport hazard class(es) | | | |
| Not applicable | Not applicable | Not applicable | Not applicable |
| 14.4. Packing group | | | |
| Not applicable | Not applicable | Not applicable | Not applicable |
| 14.5. Environmental hazards | | | |
| Not applicable | Not applicable | Not applicable | Not applicable |
| No supplementary information available | | | |

14.6. Special precautions for user

Overland transport

Not applicable

Transport by sea

Not applicable

Air transport

Not applicable

Rail transport

Not applicable

14.7. Maritime transport in bulk according to IMO instruments

Not applicable

SECTION 15: Regulatory information

15.1. Safety, health and environmental regulations/legislation specific for the substance or mixture

15.1.1. EU-Regulations

REACH Annex XVII (Restriction List)

Contains no REACH substances with Annex XVII restrictions

REACH Annex XIV (Authorisation List)

Contains no REACH Annex XIV substances

REACH Candidate List (SVHC)

Contains no substance on the REACH candidate list

PIC Regulation (Prior Informed Consent)

Contains no substance subject to Regulation (EU) No 649/2012 of the European Parliament and of the Council of 4 July 2012 concerning the export and import of hazardous chemicals.

POP Regulation (Persistent Organic Pollutants)

Contains no substance subject to Regulation (EU) No 2019/1021 of the European Parliament and of the Council of 20 June 2019 on persistent organic pollutants

Ozone Regulation (1005/2009)

Contains no substance subject to REGULATION (EU) No 1005/2009 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 16 September 2009 on substances that deplete the ozone layer.

Explosives Precursors Regulation (2019/1148)

Contains no substance subject to Regulation (EU) 2019/1148 of the European Parliament and of the Council of 20 June 2019 on the marketing and use of explosives precursors.

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Drug Precursors Regulation (273/2004)

Contains no substance(s) listed on the Drug Precursors list (Regulation EC 273/2004 on drug precursors)

15.1.2. National regulations

No additional information available

15.2. Chemical safety assessment

No chemical safety assessment has been carried out

For the following substances of this mixture a chemical safety assessment has been carried out:

4-(indol-3-yl)butyric acid

SECTION 16: Other information

Indication of changes:

according to Regulation (EC) No. 1907/2006 (REACH) with its amendment Regulation (EU) 2020/878.

| Indication of changes | | | |
|-----------------------|----------------------|----------|----------|
| Section | Changed item | Change | Comments |
| | Revision date | Modified | |
| | Supersedes | Modified | |
| | Version | Modified | |
| | Regulatory reference | Modified | |
| 1.1 | UFI | Added | |

| Abbreviations and acronyms: | |
|-----------------------------|---|
| ADR | European Agreement concerning the International Carriage of Dangerous Goods by Road |
| ATE | Acute Toxicity Estimate |
| CLP | Classification Labelling Packaging Regulation; Regulation (EC) No 1272/2008 |
| DNEL | Derived-No Effect Level |
| EC50 | Median effective concentration |
| IATA | International Air Transport Association |
| IMDG | International Maritime Dangerous Goods |
| LC50 | Median lethal concentration |
| LD50 | Median lethal dose |
| PBT | Persistent Bioaccumulative Toxic |
| PNEC | Predicted No-Effect Concentration |
| REACH | Registration, Evaluation, Authorisation and Restriction of Chemicals Regulation (EC) No 1907/2006 |
| RID | Regulations concerning the International Carriage of Dangerous Goods by Rail |
| STP | Sewage treatment plant |
| vPvB | Very Persistent and Very Bioaccumulative |

Data sources

: ECHA (European Chemicals Agency). zRMS Nederland (REGULATION (EC) No 1107/2009).

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Other information

: DISCLAIMER OF LIABILITY The information in this SDS was obtained from sources which we believe are reliable. However, the information is provided without any warranty, express or implied, regarding its correctness. The conditions or methods of handling, storage, use or disposal of the product are beyond our control and may be beyond our knowledge. For this and other reasons, we do not assume responsibility and expressly disclaim liability for loss, damage or expense arising out of or in any way connected with the handling, storage, use or disposal of the product. This SDS was prepared and is to be used only for this product. If the product is used as a component in another product, this SDS information may not be applicable.

| Full text of H- and EUH-statements: | |
|-------------------------------------|---|
| Acute Tox. 4 (Oral) | Acute toxicity (oral), Category 4 |
| EUH210 | Safety data sheet available on request. |
| EUH401 | To avoid risks to human health and the environment, comply with the instructions for use. |
| H302 | Harmful if swallowed. |
| H361fd | Suspected of damaging fertility. Suspected of damaging the unborn child. |
| Repr. 2 | Reproductive toxicity, Category 2 |

| Full text of use descriptors | |
|------------------------------|---------------------------|
| PC27 | Plant protection products |

Safety Data Sheet (SDS), EU

This information is based on our current knowledge and is intended to describe the product for the purposes of health, safety and environmental requirements only. It should not therefore be construed as guaranteeing any specific property of the product.