SAFETY DATA SHEET



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

1.1. Product identifier

Trade name or designation

of the mixture

VOLIBRIS TABLETS

Registration number

Synonyms VOLIBRIS 5 MG TABLETS * VOLIBRIS 10 MG TABLETS * AMBRISENTAN, FORMULATED

PRODUCT

Issue date 05-November-2014

Version number 02

Revision date 05-November-2014

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

Identified uses Medicinal Product.

> This safety data sheet is written to provide health, safety and environmental information for people handling this formulated product in the workplace. It is not intended to provide information relevant

to medicinal use of the product. In this instance patients should consult prescribing

information/package insert/product label or consult their pharmacist or physician. For health and safety information for individual ingredients used during manufacturing, refer to the appropriate

safety data sheet for each ingredient.

Uses advised against No other uses are advised.

1.3. Details of the supplier of the safety data sheet

GlaxoSmithKline UK 980 Great West Road

Brentford, Middlesex TW8 9GS UK

UK General Information (normal business hours): +44-20-8047-5000

Fmail Address: msds@gsk.com Website: www.qsk.com

1.4. Emergency telephone

number

TRANSPORT EMERGENCIES::

UK In-country toll call: +(44)-870-8200418 International toll call: +1 703 527 3887

available 24 hrs/7 days; multi-language response

SECTION 2: Hazards identification

2.1. Classification of the substance or mixture

Classification according to Directive 67/548/EEC or 1999/45/EC as amended

Exempt from requirements - product regulated as a medicinal product, cosmetic product or medical device.

Classification according to Regulation (EC) No 1272/2008 as amended

Exempt from requirements - product regulated as a medicinal product, cosmetic product or medical device.

2.2. Label elements

Label according to Regulation (EC) No. 1272/2008 as amended

Exempt from requirements - product regulated as a medicinal product, cosmetic product or medical device.

Caution - Pharmaceutical agent. See section 11 for additional information on health hazards. 2.3. Other hazards

SECTION 3: Composition/information on ingredients

3.2. Mixtures

Material name: VOLIBRIS TABLETS SDS MALTA

General information

Chemical name		%	CAS-No. / EC No.	REACH Registration No.	INDEX No.	Notes
MICROCRYSTALLINE CELLULOSE		OSE < 2	5 9004-34-6 232-674-9	-	-	
Classification:	DSD:	-				
	CLP:	-				
AMBRISENTAN		3,4 -	< 7 177036-94-1	-	-	
Classification:	DSD:	- Repr. Cat. 2;R60-61				
	CLP:	Repr. 1B;H	360FD			
MAGNESIUM STEARATE		< ′	1 557-04-0 209-150-3	-	-	
Classification:	DSD:	-				
	CLP:	-				
Talc		< ′	1 14807-96-6 238-877-9	-	-	
Classification:	DSD:	-				
	CLP:	-				
Titanium dioxide		< ′	1 13463-67-7 236-675-5	-	-	
Classification:	DSD:	-				
	CLP:	-				

Other components below reportable levels 60 - < 70

List of abbreviations and symbols that may be used above

CLP: Regulation No. 1272/2008. DSD: Directive 67/548/EEC.

M: M-factor

vPvB: very persistent and very bioaccumulative substance.

PBT: persistent, bioaccumulative and toxic substance.

#: This substance has been assigned Community workplace exposure limit(s).

SECTION 4: First aid measures

General information In the case of accident or if you feel unwell, seek medical advice immediately (show the label

where possible). Ensure that medical personnel are aware of the material(s) involved, and take

precautions to protect themselves.

4.1. Description of first aid measures

Inhalation Move to fresh air. If breathing is difficult, trained personnel should give oxygen. Call a physician if

symptoms develop or persist. Under normal conditions of intended use, this material is not

expected to be an inhalation hazard.

Immediately flush skin with plenty of water. Take off contaminated clothing and wash before reuse. Skin contact

Get medical attention if symptoms occur.

Eve contact Rinse thoroughly with plenty of water for at least 15 minutes and consult a physician.

Ingestion If swallowed, rinse mouth with water (only if the person is conscious). If ingestion of a large

amount does occur, call a poison control centre immediately. Do not induce vomiting without

advice from poison control center.

4.2. Most important symptoms

and effects, both acute and delayed

The following adverse effects have been noted with therapeutic use of this material: headache; flushing; dizziness.

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

Material name: VOLIBRIS TABLETS

No specific antidotes are recommended. Treat according to locally accepted protocols. For additional guidance, refer to the current prescribing information or to the local poison control

information centre.

SECTION 5: Firefighting measures

General fire hazards No unusual fire or explosion hazards noted.

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5.1. Extinguishing media

Suitable extinguishing

media

Water. Foam. Dry chemical powder. Carbon dioxide (CO2).

Unsuitable extinguishing

media

None known.

5.2. Special hazards arising from the substance or mixture

During fire, gases hazardous to health may be formed.

5.3. Advice for firefighters

Special protective equipment for firefighters

Self-contained breathing apparatus and full protective clothing must be worn in case of fire.

Special fire fighting

procedures

Move containers from fire area if you can do so without risk.

Specific methodsUse standard firefighting procedures and consider the hazards of other involved materials.

SECTION 6: Accidental release measures

6.1. Personal precautions, protective equipment and emergency procedures

For non-emergency

personnel

Keep unnecessary personnel away. Keep people away from and upwind of spill/leak. Keep out of low areas. Do not touch damaged containers or spilled material unless wearing appropriate protective clothing. For personal protection, see section 8.

For emergency responders

Keep unnecessary personnel away. Use personal protection recommended in Section 8 of the

SDS.

6.2. Environmental precautions

Avoid discharge into drains, water courses or onto the ground.

6.3. Methods and material for containment and cleaning up

Stop the flow of material, if this is without risk. Prevent entry into waterways, sewer, basements or

confined areas. Following product recovery, flush area with water.

6.4. Reference to other

sections

For personal protection, see section 8. For waste disposal, see section 13 of the SDS.

SECTION 7: Handling and storage

7.1. Precautions for safe

handling

Avoid prolonged exposure. Observe good industrial hygiene practices.

7.2. Conditions for safe storage, including any incompatibilities

Store in original tightly closed container. Store away from incompatible materials (see Section 10

of the SDS).

7.3. Specific end use(s) Medicinal Product.

SECTION 8: Exposure controls/personal protection

8.1. Control parameters

Occupational exposure limits

001
(3.5 M

Components	Туре	Value	Note			
AMBRISENTAN (CAS 177036-94-1)	8 HR TWA	20 mcg/m3				
•	OHC	3	Reproductive hazard			
MICROCRYSTALLINE CELLULOSE (CAS 9004-34-6)	OHC	1				
alaniaal limitualusa	No histograph synaptic limits noted for the ingradient(s)					

Biological limit values No biological exposure limits noted for the ingredient(s).

Recommended monitoring

procedures

Follow standard monitoring procedures.

Derived no-effect level (DNEL) Not available.

Predicted no effect concentrations (PNECs)

Not available.

Exposure guidelines

8.2. Exposure controls

Appropriate engineering

General ventilation normally adequate.

controls

Individual protection measures, such as personal protective equipment

General information

Personal protection equipment should be chosen according to the CEN standards and in discussion with the supplier of the personal protective equipment. Follow all local regulations if

personal protective equipment (PPE) is used in the workplace.

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Not normally needed. If contact is likely, safety glasses with side shields are recommended. (e.g. Eye/face protection

EN 166).

Skin protection

- Hand protection Not normally needed. For prolonged or repeated skin contact use suitable protective gloves. Select

suitable chemical resistant protective gloves (EN 374) with a protective index 6 (>480min

permeation time).

Not normally needed. Wear suitable protective clothing as protection against splashing or - Other

contamination. (EN 14605 for splashes, EN ISO 13982 for dust).

Respiratory protection No personal respiratory protective equipment normally required. When workers are facing

concentrations above the exposure limit they must use appropriate certified respirators.

Thermal hazards Wear appropriate thermal protective clothing, when necessary.

Always observe good personal hygiene measures, such as washing after handling the material Hygiene measures

> and before eating, drinking, and/or smoking. Routinely wash work clothing and protective equipment to remove contaminants. For advice on suitable monitoring methods, seek guidance from a qualified environment, health and safety professional. New or expectant mothers might be at greater risk from overexposure. Risk assessments must take this into consideration. Female employees anticipating pregnancy or with a confirmed pregnancy must be encouraged to notify an occupational health professional or their line manager. This will act as the trigger for individual

re-assessment of the employee's work practices.

Environmental exposure controls

Hazard guidance and control recommendations Environmental manager must be informed of all major releases.

SECTION 9: Physical and chemical properties

9.1. Information on basic physical and chemical properties

Appearance

Physical state Solid. Tablet **Form**

Not available. Colour Not available Odour **Odour threshold** Not available. Not available. Not available. Melting point/freezing point Initial boiling point and boiling Not available.

range

Not available. Flash point **Evaporation rate** Not available. Flammability (solid, gas) Not available.

Upper/lower flammability or explosive limits

Flammability limit - lower

Not available.

Flammability limit - upper

Not available.

(%)

Not available. Vapour pressure Vapour density Not available. Not available. Relative density

Solubility(ies)

Not available. Solubility (water) Solubility (other) Not available. Not available. Partition coefficient

(n-octanol/water)

Not available. Auto-ignition temperature **Decomposition temperature** Not available. Not available. **Viscosity** Not available. **Explosive properties** Not available. Oxidizing properties

9.2. Other information No relevant additional information available.

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SECTION 10: Stability and reactivity

10.1. ReactivityThe product is stable and non-reactive under normal conditions of use, storage and transport.

10.2. Chemical stability Material is stable under normal conditions.

10.3. Possibility of hazardous

reactions

No dangerous reaction known under conditions of normal use.

10.4. Conditions to avoid Contact with incompatible materials.10.5. Incompatible materials Strong oxidising agents. Fluorine.

10.6. Hazardous None known. Irritating and/or toxic fumes and gases may be emitted upon the product's

decomposition products decomposition.

SECTION 11: Toxicological information

General information Occupational exposure to the substance or mixture may cause adverse effects.

Information on likely routes of exposure

InhalationUnder normal conditions of intended use, this material is not expected to be an inhalation hazard.Skin contactHealth injuries are not known or expected under normal use. Dust or powder may irritate the skin.Eye contactHealth injuries are not known or expected under normal use. Dust or powder may irritate eye

tissue

Ingestion Health injuries are not known or expected under normal use. May be harmful if swallowed.

However, ingestion is not likely to be a primary route of occupational exposure.

Symptoms None known.

The following adverse effects have been noted with therapeutic use of this material: headache;

flushing; dizziness.

11.1. Information on toxicological effects

Acute toxicity Health injuries are not known or expected under normal use. May be harmful if swallowed.

Components Species Test results

AMBRISENTAN (CAS 177036-94-1)

Acute

Oral

LD50 Rat

Rat >= 3160 mg/kg

MAGNESIUM STEARATE (CAS 557-04-0)

Acute

Oral

LD50 Rat > 2000 mg/kg

MICROCRYSTALLINE CELLULOSE (CAS 9004-34-6)

Acute

Dermal

LD50 Rabbit > 2000 mg/kg

Oral

LD50 Rat > 2000 mg/kg

Titanium dioxide (CAS 13463-67-7)

Acute

Inhalation

LC50 Rat 6820 mcg/m3

Oral

LD50 Rat > 24 g/kg

Chronic

Inhalation

LOEC Rat 8,6 mg/m3, 1 years TiO2 accumulated in

interstitial macrophages, aggregated interstitial cells and particle laden macrophrages in lymphoid tissue.

NOAEC Rat 250 mg/m3, 2 years Highest dose

5 mg/m3, 24 months

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Components **Species Test results Subacute** Inhalation LOEL Rat 0,1 - 35 mg/m3, 4 weeks Mild macrophage hyperplasia, no change in bronchio-alveolar lavage fluid. **NOAEC** 26 mg/m3, 3 weeks No evidence of Guinea pig significant inflammation in respiratory tract. Oral **NOAEL** Rat 100000 ppm, 14 Day Dietary study, highest dose tested. **Subchronic** Inhalation

Rat

Health injuries are not known or expected under normal use. Skin corrosion/irritation

Irritation Corrosion - Skin

LOEC

TITANIUM DIOXIDE 0, Literature data Result: Non-irritant

Species: Guinea pig 0, Literature data Result: Non-irritant Species: Human

Acute dermal irritation; OECD 404, Literature data

Result: Non-irritant Species: Rabbit

AMBRISENTAN Reconstituted Human Epidermis (RHE)

Result: Negative - Not likely to be a significant irritant.

3,2 - 20 mg/m3, 8 min Accumulation of TiO2 in macrophages and evidence of

pulmonary inflammation.

Irritation Corrosion - Skin: P.I.I. value

MAGNESIUM STEARATE

Serious eye damage/eye

Health injuries are not known or expected under normal use. Dust or powder may irritate eye tissue.

irritation

TITANIUM DIOXIDE

OECD 405, Literature data

Result: Mild irritant Species: Rabbit

Reconstituted Human Corneal Epithelium (HCE) **AMBRISENTAN**

Result: Negative; not likely to be a severe irritant

Eye / Kay and Calandra class - Intact

MAGNESIUM STEARATE

Recovery Period: 2 days

Respiratory sensitisation Not available.

Skin sensitisation Health injuries are not known or expected under normal use.

Sensitisation

TITANIUM DIOXIDE 5 % Optimisation Test, Literature data - Vehicle: petrolatum

> Result: negative Species: Guinea pig

Test Duration: 48 hour exposure

AMBRISENTAN OECD 429 / Local Lymph Node Assay, Maximum

concentration = 50%; vehicle = DMF

Result: negative Species: Mouse

TITANIUM DIOXIDE Patch test, Literature data

Result: negative Species: Human

Germ cell mutagenicity Health injuries are not known or expected under normal use.

Mutagenicity

AMBRISENTAN Ames Assay, GLP assay

> Result: negative Ames, Literature data Result: negative

AMBRISENTAN Chromosomal Aberration Assay In Vitro

Result: positive

Material name: VOLIBRIS TABLETS

TITANIUM DIOXIDE

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^{*} Estimates for product may be based on additional component data not shown.

Mutagenicity

TITANIUM DIOXIDE Micronucleus Assay in vitro, CHO cells, Literature data

Result: negative

Micronucleus Assay in vitro, cultured human peripheral

lymphocytes, Literature data

Result: positive

AMBRISENTAN Micronucleus Assay, GLP assay

Result: negative Species: Rat

TITANIUM DIOXIDE Syrian Hamster Embryo (SHE) cell transformation assay

Result: negative

AMBRISENTAN Unscheduled DNA Synthesis in vivo

Result: negative

Species: Rat
TITANIUM DIOXIDE WIL2-NS HPR

WIL2-NS HPRT/ t-Thioguanidine - Human B-Cell

lymphoblastoid, Literature data

Result: positive

Carcinogenicity Health injuries are not known or expected under normal use. Contains a material (titanium

dioxide) classified as a carcinogen by external agencies. Carcinogenic activity was seen in inhalation studies using laboratory animals. High concentrations or doses administered over an

extended period of time were required to produce adverse effects.

TITANIUM DIOXIDE 0,5 mg/m3, Literature data

Result: negative Species: Rat

Test Duration: 24 months

0,72 - 14,8 mg/m3, Literature data

Result: negative Species: Mouse

10 - 250 mg/m3, Dietary study - Literature data.

Result: Inflammation at all doses with alveolar/bronchiolar

adenoma at the highest concentration.

Species: Rat

Test Duration: 24 months

25000 - 50000 ppm, Dietary study

Result: negative Species: Mouse

25000 - 50000 ppm, Dietary study - Literature data.

Result: negative Species: Rat

7,2 - 14,8 mg/m3, Literature data

Result: Lung tumour Species: Rat

Test Duration: 24 months

IARC Monographs. Overall Evaluation of Carcinogenicity

Titanium dioxide (CAS 13463-67-7) 2B Possibly carcinogenic to humans.

Reproductive toxicity Suspected of damaging fertility or the unborn child.

Reproductivity

AMBRISENTAN Embryo-foetal development

Result: Foetal toxicity, malformations in multiple species;

endothelin receptor antagonist class effect

Fertility, Male

Result: Testicular toxicity, reduced fertility in mice and rats

Specific target organ toxicity -

single exposure

None known.

Specific target organ toxicity -

repeated exposure

None known.

Not available.

Aspiration hazard

Mixture versus substance information

No information available.

Other information Caution - Pharmaceutical agent. Occupational exposure to the substance or mixture may cause

adverse effects.

SECTION 12: Ecological information

12.1. Toxicity Not expected to be harmful to aquatic organisms.

Material name: VOLIBRIS TABLETS

Components **Species Test results**

AMBRISENTAN (CAS 177036-94-1)

Aquatic

Acute

EC50 Algae 10 - 100 mg/l, 96 hours QSAR Estimate Algae Crustacea EC50 Daphnia 10 - 100 mg/l, 48 hours QSAR Estimate Fish EC50 Fish 10 - 100 mg/l, 96 hours QSAR Estimate

MAGNESIUM STEARATE (CAS 557-04-0)

Aquatic

Acute

Fish EC50 Orange-red killfish (Adult Oryzias 130 mg/l, 96 hours

latipes)

Talc (CAS 14807-96-6)

Aquatic

Acute

Fish EC50 Zebra fish (Adult Brachydanio rerio) > 100 g/l, 24 hours Static renewal test

Titanium dioxide (CAS 13463-67-7)

Aquatic

Acute

Crustacea EC50 > 1000 mg/l, 48 hours Static test Water flea (Daphnia magna)

12.2. Persistence and

degradability

Photolysis

Half-life (Photolysis-atmospheric)

17 Hours Estimated MAGNESIUM STEARATE

UV/visible spectrum wavelength

MAGNESIUM STEARATE 210 nm

Biodegradability

Percent degradation (Aerobic biodegradation-inherent)

MAGNESIUM STEARATE 77 %, 28 days BOD

Percent degradation (Aerobic biodegradation-ready)

MAGNESIUM STEARATE 95 %, 22 days Sturm test

Percent degradation (Aerobic biodegradation-soil)

MAGNESIUM STEARATE 50 %, 13 days

12.3. Bioaccumulative potential

Partition coefficient

n-octanol/water (log Kow)

AMBRISENTAN 1,2 (Measured).

Bioconcentration factor (BCF)

MAGNESIUM STEARATE > 9999 Estimated

12.4. Mobility in soil

Adsorption

Soil/sediment sorption - log Koc

MAGNESIUM STEARATE 5,86 Estimated

Mobility in general Not available. Not available. 12.5. Results of PBT

and vPvB assessment

Not available. 12.6. Other adverse effects

SECTION 13: Disposal considerations

13.1. Waste treatment methods

Dispose of in accordance with local regulations. Empty containers or liners may retain some Residual waste

product residues. This material and its container must be disposed of in a safe manner (see:

Disposal instructions). Avoid discharge into water courses or onto the ground.

Empty containers should be taken to an approved waste handling site for recycling or disposal. Contaminated packaging

Since emptied containers may retain product residue, follow label warnings even after container is

emptied.

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^{*} Estimates for product may be based on additional component data not shown.

EU waste codeThe Waste code should be assigned in discussion between the user, the producer and the waste

disposal company.

Disposal methods/information Collect and reclaim or dispose in sealed containers at licensed waste disposal site. Do not

discharge into drains, water courses or onto the ground. Dispose in accordance with all applicable

regulations.

Special precautionsDispose in accordance with all applicable regulations.

SECTION 14: Transport information

ADR

Not regulated as dangerous goods.

RID

Not regulated as dangerous goods.

ADN

Not regulated as dangerous goods.

IATA

Not regulated as dangerous goods.

IMDG

Not regulated as dangerous goods.

14.7. Transport in bulk according to Annex II of

MARPOL Annex II applies to liquids used in a ship's operation that pose a threat to the marine

environment. These materials may not be transported in bulk.

MARPOL73/78 and the IBC Code

SECTION 15: Regulatory information

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

EU regulations

Regulation (EC) No. 1005/2009 on substances that deplete the ozone layer, Annex I

Not listed

Regulation (EC) No. 1005/2009 on substances that deplete the ozone layer, Annex II

Not listed.

Regulation (EC) No. 850/2004 On persistent organic pollutants, Annex I as amended

Not listed

Regulation (EC) No. 689/2008 concerning the export and import of dangerous chemicals, Annex I, part 1 as amended Not listed.

Regulation (EC) No. 689/2008 concerning the export and import of dangerous chemicals, Annex I, part 2 as amended

Regulation (EC) No. 689/2008 concerning the export and import of dangerous chemicals, Annex I, part 3 as amended Not listed.

Regulation (EC) No. 689/2008 concerning the export and import of dangerous chemicals, Annex V as amended Not listed.

Regulation (EC) No. 166/2006 Annex II Pollutant Release and Transfer Registry

Not listed.

Regulation (EC) No. 1907/2006, REACH Article 59(10) Candidate List as currently published by ECHA Not listed.

Authorisations

Regulation (EC) No. 1907/2006, REACH Annex XIV Substances subject to authorization, as amended

Not listed.

Restrictions on use

Regulation (EC) No. 1907/2006, REACH Annex XVII Substances subject to restriction on marketing and use as amended

Not listed

Directive 2004/37/EC: on the protection of workers from the risks related to exposure to carcinogens and mutagens at work

Not listed.

Directive 92/85/EEC: on the safety and health of pregnant workers and workers who have recently given birth or are breastfeeding

Not listed.

Material name: VOLIBRIS TABLETS

Other EU regulations

Directive 96/82/EC (Seveso II) on the control of major-accident hazards involving dangerous substances

Directive 98/24/EC on the protection of the health and safety of workers from the risks related to chemical agents at work Not listed.

Directive 94/33/EC on the protection of young people at work

Not listed.

Other regulations The product is classified and labelled in accordance with EC directives or respective national laws.

This Safety Data Sheet complies with the requirements of Regulation (EC) No 1907/2006.

National regulations 15.2. Chemical safety

assessment

Follow national regulation for work with chemical agents. No Chemical Safety Assessment has been carried out.

SECTION 16: Other information

List of abbreviations Not available.

References GSK Hazard Determination

Information on evaluation method leading to the classification of mixture

The classification for health and environmental hazards is derived by a combination of calculation

methods and test data, if available.

Full text of any statements or R-phrases and H-statements under Sections 2 to 15

o 15 R60 May impair fertility.

R61 May cause harm to the unborn child.

H360FD May damage fertility. May damage the unborn child.

Revision information Product and Company Identification: Product and Company Identification

Composition / Information on Ingredients: Ingredients

Exposure Controls / Personal Protection: Physical & Chemical Properties: Regulatory Information: United States

GHS: Classification

Training information Follow training instructions when handling this material.

Disclaimer

The information and recommendations in this safety data sheet are, to the best of our knowledge,

accurate as of the date of issue. Nothing herein shall be deemed to create any warranty, express or implied. It is the responsibility of the user to determine the applicability of this information and

the suitability of the material or product for any particular purpose.

Material name: VOLIBRIS TABLETS