SAFETY DATA SHEET

1. IDENTIFICATION OF THE SUBSTANCE/PREPARATION AND OF THE COMPANY/UNDERTAKING

Material: TIMENTIN INJECTABLE

Synonyms:
- TIMENTIN 1.6 GRAM INJECTABLE
- TIMENTIN 3 GRAM INJECTABLE
- TIMENTIN 3.1 GRAM INJECTABLE
- TIMENTIN 3.2 GRAM INJECTABLE
- TIMENTEN LIQUID
- TIMENTEN INJECTION
- CLAVENTIN INJECTION
- AUGPENIN INJECTION
- NDC NO. 0029-6571-26
- NDC NO. 0029-6571-40
- NDC NO. 0029-6579-21
- NDC NO. 0029-6571-31
- NDC NO. 0029-6571-56
- NDC NO. 0029-6575-57
- NDC NO. 0029-6579-76
- POTASSIUM CLAVULANATE AND TICARCILLIN DISODIUM, FORMULATED PRODUCT

Company Name:
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- Transport Emergency (EU): +44-1865-407333
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- Information and Advice: US number, available 24 hours
  Multi-language response

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- US General Information: +1-888-825-5249
- Transport Emergency (non EU): +1-703-527-3887
  US number, available 24 hours
  Multi-language response

2. COMPOSITION / INFORMATION ON INGREDIENTS

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>CAS RN</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>TICARCILLIN DISODIUM</td>
<td>4697-14-7</td>
<td>94 to 97</td>
</tr>
<tr>
<td>POTASSIUM CLAVULANATE</td>
<td>61177-45-5</td>
<td>3 to 6</td>
</tr>
</tbody>
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3. HAZARDS IDENTIFICATION

Fire and Explosion:
Assume that this product is capable of sustaining combustion.

Health:
- Exposure might occur via skin; eyes; ingestion; inhalation.
- May produce allergic skin reactions.
- Respiratory allergen.
- Possible effects of overexposure in the workplace include: symptoms of hypersensitivity (such as skin rash, hives, itching, and difficulty breathing); nausea; vomiting; diarrhoea.

Health effects information is based on hazards of components.
4. FIRST-AID MEASURES

Ingestion
Never attempt to induce vomiting. Do not attempt to give any solid or liquid by mouth if the exposed subject is unconscious or semi-conscious. Wash out the mouth with water. If the exposed subject is fully conscious, give plenty of water to drink. Obtain medical attention.

Inhalation
Using appropriate personal protective equipment, move exposed subject to fresh air. If breathing is difficult or ceases, ensure and maintain ventilation. Give oxygen as appropriate. The exposed subject should be kept warm and at rest. Obtain medical attention in cases of known or possible over exposure, or with symptoms including chest pain, difficulty breathing, loss of consciousness or other adverse effects, which may be delayed.

Skin Contact
Using appropriate personal protective equipment, remove contaminated clothing and flush exposed area with large amounts of water. Obtain medical attention if skin reaction occurs, which may be immediate or delayed.

Eye Contact
Wash immediately with clean and gently flowing water. Continue for at least 15 minutes. Obtain medical attention.

NOTES TO HEALTH PROFESSIONALS

Medical Treatment
Treat according to locally accepted protocols. For additional guidance, refer to the current prescribing information or to the local poison control information centre. Medical treatment in cases of overexposure should be treated as an overdose of penicillin antibiotic. In allergic individuals, exposure to this material may require treatment for initial or delayed allergic symptoms and signs. This may include immediate and/or delayed treatment of anaphylactic reactions.

Medical Conditions Caused or Aggravated by Exposure
Refer to prescribing information for detailed description of medical conditions caused by or aggravated by overexposure to this product. Ocular symptoms may be indicative of allergic reaction. Pulmonary symptoms may indicate allergic reaction or asthma. This material may cause or aggravate allergy to penicillin antibiotics.

Antidotes
No specific antidotes are recommended.

5. FIRE-FIGHTING MEASURES

Fire and Explosion Hazards
The combustibility of the product is not known, however the packaging is combustible.

Extinguishing Media
Water, dry powder or foam extinguishers are recommended. Carbon dioxide extinguishers may be ineffective.

Special Firefighting Procedures
For single units (packages): No special requirements needed. For larger amounts (multiple packages/pallets) of product: Since toxic, corrosive or flammable vapours might be evolved from fires involving this product and associated packaging, self contained breathing apparatus and full protective equipment are recommended for firefighters. If possible, contain and collect firefighting water for later disposal.

Hazardous Combustion Products
Toxic, corrosive or flammable thermal decomposition products are expected when the product is exposed to fire.

6. ACCIDENTAL RELEASE MEASURES

Personal Precautions
Wear protective clothing and equipment consistent with the degree of hazard.
Environmental Precautions
For large spills, take precautions to prevent entry into waterways, sewers, or surface drainage systems.

Clean-up Methods
Collect and place it in a suitable, properly labelled container for recovery or disposal.

Decontamination Procedures
No specific decontamination or detoxification procedures have been identified for this product. Water can be used for clean-up and decontamination operations.

7. HANDLING AND STORAGE

HANDLING
General Requirements
No special control measures required for the normal handling of this product. Normal room ventilation is expected to be adequate for routine handling of this product.

STORAGE
No storage requirements necessary for occupational hazards. Follow product information storage instructions to maintain efficacy.

8. EXPOSURE CONTROLS/PERSONAL PROTECTION

INGREDIENT	TICARCILIN DISODIUM
GSK Occupational Hazard Category 1
GSK Occupational Exposure Limit 1000 MCG/M3 (8 HR TWA) RESPIRATORY SENSITISER, SKIN SENSITISER

INGREDIENT	POTASSIUM CLAVULANATE
GSK Occupational Hazard Category 1
GSK Occupational Exposure Limit 5000 MCG/M3 (8 HR TWA)

Other Equipment or Procedures
Wear appropriate clothing to avoid skin contact. Wash hands and arms thoroughly after handling.

9. PHYSICAL AND CHEMICAL PROPERTIES

Appearance
Colour
Physical Form
Packaging
pH of Aqueous Solutions
White/light yellow.
Powder.
Vial.
5.5 to 7.5

10. STABILITY AND REACTIVITY

Stability
This product is expected to be stable.

Conditions to Avoid
None for normal handling of this product.

11. TOXICOLOGICAL INFORMATION

Oral Toxicity
Not expected to be toxic following ingestion.

Inhalation Toxicity
Adverse effects might occur following inhalation.

Skin Effects
Irritation is not expected following direct contact.

Eye Effects
Minor irritation might occur following direct contact with eyes.

Target Organ Effects
No specific target organ effects have been identified.
Sensitisation  Allergic skin reactions might occur following dermal exposure. Assessment based upon information from human exposure. Respiratory sensitisation (allergic) reactions might occur following exposure.

Genetic Toxicity  Not expected to be genotoxic under occupational exposure conditions. Assessment based upon effects of structurally similar substances.

Carcinogenicity  Not expected to produce cancer in humans under occupational exposure conditions. No components are listed as carcinogens by GSK, IARC, NTP or US OSHA.

Reproductive Effects  No adverse effects have been reported following extensive use or exposure in humans.

Pharmacological Effects  This material is a penicillin; an antibiotic.

12. ECOLOGICAL INFORMATION

Summary  No information is available about the potential of this product to produce adverse environmental effects. Local regulations and procedures should be consulted prior to environmental release.

PERSISTENCE/DEGRADATION

Biodegradation  Penicillins are generally susceptible to degradation by a number of micro-organisms found in wastewater treatment plants and the general environment. Resulting degradation products are readily mineralised by environmental micro-organisms.

13. DISPOSAL CONSIDERATIONS

Disposal Recommendations  Collect for recycling or recovery if possible. The disposal method for rejected products/returned goods must ensure that they cannot be re-sold or re-used.

Regulatory Requirements  Observe all local and national regulations when disposing of this product.

14. TRANSPORT INFORMATION

The SDS should accompany all shipments for reference in the event of spillage or accidental release. Only authorised persons trained and competent in accordance with appropriate national and international regulatory requirements may prepare dangerous goods for transport.

UN Classification and Labelling

Transport Information  Transportation and shipping of this product is not restricted. It has no known, significant hazards requiring special packaging or labelling for air, maritime, US or European ground transport purposes.

15. REGULATORY INFORMATION

The information included below is an overview of the major regulatory requirements. It should not be considered to be an exhaustive summary. Local regulations should be consulted for additional requirements.

EU Classification and Labelling  None.


Classification  This dosage form is exempt from the requirements of the OSHA Hazard Communication Standard.

Other US Regulations

TSCA Status  Exempt

16. OTHER INFORMATION
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.