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SAFETY DATA SHEET
(according to Regulation (EC) No 1907/2006 (REACH), ANNEX II)

AMMONIA



| SECTION 1: IDENTIFICATION OF THE SUBSTANCE AND OF THE COMPANY | | | | | |
|--|--|--|--|--|--|
| 1.1 Product identifier | | | | | |
| Trade name: | Ammonia | | | | |
| Other names: | Ammonia anhydrous | | | | |
| Name IUPAC/ international chemical name | Ammonia | | | | |
| INDEX No. and name as listed in Annex VI of CLP: | 007-001-00-5 ammonia, anhydrous | | | | |
| CAS No.: | 7664-41-7 | | | | |
| EINECS No.: | 231-635-3 | | | | |
| REACH registration No.: | 01-2119488876-14-0048 | | | | |
| Molecular formula | H3N | | | | |
| 1.2 Relevant identified uses of the subs | tance or mixture and uses advised against | | | | |
| Relevant identified uses: | Use of the substance in the manufacture of nitric acid, alkalis, dyes, pharmaceuticals, cosmetics, vitamins, synthetic textile fibres and plastics. (see ES 1) Use in photochemical processes, refrigerant systems, insulation products, inks & toners, coatings, thinners & paint removers, processing aid in chemical industry. Use as an extraction agent, in NOx/SOx reduction, processing aid in nutrition, neutralising agent, textile dye, washing & cleaning products, textile treatment. Use in pulp/paper treatment, leather treatment, wood treatment, metal surface treatment, the treatment of rubber/latex, the manufacture of semiconductors/electronics. Use in adhesives & sealants, polymer preparations, aircare products & preservatives. (see ES 2) Use as a laboratory chemical, refrigerant in cooling systems, water treatment chemical, fertiliser, coating, paint thinner or paint remover, photochemical. Use as a cleaning product, leather or other surface treatment product, pH regulatory or neutralisation agent, process aid for nutrition (see ES 3) Use in coatings, paints, thinners and removers; use in fillers, putties and plasters, use of washing and cleaning products, use in cosmetic & personal care products (see ES 4) | | | | |
| Uses advised against: | None | | | | |
| 1.3 Details of the supplier of the safety of | data sheet | | | | |
| Only Representative: | Zangas Hoch-und Tiefbau GmbH Schwindgasse 5/1/4 1040 Vienna Austria Phone: +43 1 274 16 366 www.zangasgroup.com E-mail: info@zangasgroup.com | | | | |
| Manufacturer: | PrJSC "AZOT" 72, Heroiv Kholodnoho Yaru Str., Cherkasy, Ukraine Phone: +38 0472 39-63-03 | | | | |
| E-mail address of the competent person responsible for the Safety Data Sheet | PrJSC "AZOT" REACH Department onr@azot.ck.ua | | | | |
| National contact: | Not available | | | | |
| 1.4 Emergency telephone number | | | | | |
| Emergency phone number: | Phone: +43 1 274 16 366 Opening hours: 9-18 (CET) Languages of the phone service: German, English Phone: + 38 (0472) 39 61 17 Opening hours: 0-24 Languages of the phone service: Ukrainian | | | | |



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| AZOI | CECTION 2. HAZADDO IDENTIFICATION | |
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| 2.4 Classification of the sub-to- | SECTION 2: HAZARDS IDENTIFICATION | |
| 2.1 Classification of the substance | | |
| | Classification in accordance with Regulation 127 | |
| Hazard statement(s): | H221 Flammable Gas [not applicable for solutions] H280 Contains gas under pressure; may explode if heated. H314 Causes severe skin burns and eye damage H331 Toxic if inhaled H400 Very toxic to aquatic life M-Factor (self-classification) = 1 | Flam. Gas 2 Liquefied gas Skin Corr. 1B Acute Tox. 3 Aquatic Acute 1 |
| 2.2 Label elements | , | |
| 2.2.1 | Labelling in accordance with Regulation 1272/2 | 2008 (CLP) |
| Hazard pictogram(s): | GHS 04 GHS 05 GHS 06 GHS 09 | |
| Signal word | Danger | |
| Hazard statement(s): | H221 Flammable gas H280 Contains gas under pressure; may e. H331 Toxic if inhaled H314 Causes severe skin burns and eye d H400 Very toxic to aquatic life | |
| Supplemental Hazard information (EU): | EUH071: Corrosive to the respiratory tract | |
| Precautionary statement(s): | | |
| Precautionary Statement Prevention Precautionary Statement Response | sources. No smoking P260 Do not breathe dust, fume, gas, mist, P280 Wear protective gloves, protective clo P303+P361+P353 IF ON SKIN (or hair): To Rinse skin with water or shower. | othing, eye protection and face protection ake off immediately all contaminated clothing. |
| Precautionary Statement Storage | P304+P340 IF INHALED: Remove person breathing P305+P351+P338 IF IN EYES: Rinse caut Remove contact lenses, if present and eas P310 Immediately call a POISON CENTER P403 Store in a well-ventilated place P405 Store locked up | iously with water for several minutes. y to do. Continue rinsing |
| B :: 01 1 B: 1 | P410+P403 Protect from sunlight. Store in | |
| Precautionary Statement Disposal 2.3 Other hazards: | Evaluation, Authorisation and Restriction o Not fulfilling PBT (persistent/bioaccumulation) Not fulfilling vPvB (very persistent/very bioaccumulation) | No.1907/2006 concerning the Registration, f Chemicals (REACH): ve/toxic) criteria; accummulative) criteria. |
| SECTI | ON 3: COMPOSITION/INFORMATION ON INC | GREDIENTS |
| 3.1 Substances | | |
| Name | INDEX No. as listed in Annex VI of CLP | Weight % content (or range) |
| Ammonia, anhydrous | 007-001-00-5 | Not less than 99,6 % (w/w) |
| 4.1 Description of first aid measures | SECTION 4: FIRST-AID MEASURES | |
| General notes: | masks with cartridges for ammonia must be lt may be dangerous to the person provid | ersonal risk or without suitable training. Gas be used for evacuation from the hazard zone. ing aid to give mouth-to-mouth resuscitation. wed before being removed. Urgent hospital |





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AZOT

| Following eye contact: | Immediately flush eyes with copious amounts of tepid water for at least 15 minutes. If irritation, pain, swelling, excessive tearing, or light sensitivity persists, the patient should be seen in a health care facility and referral to an ophthalmologist considered. |
|--|---|
| Following skin contact: | Immediately flush exposed area with copious amounts of tepid water for at least 15 minutes followed by washing area thoroughly with soap and water. The patient should be seen in a health care facility if irritation or pain persists. |
| Following ingestion: | Call a physician. If conscious, give the patient milk or water to drink immediately. Do not induce vomiting. |
| Following inhalation: | Move patient to fresh air. Monitor for respiratory distress. If cough or difficulty in breathing develops, evaluate for respiratory tract irritation, bronchitis, or pneumonitis. If trained to do so, administer supplemental oxygen with assisted ventilation as required. Administer artificial respiration if patient is not breathing. |
| Self-protection for the first aider: | None |
| 4.2 Most important symptoms and effect | ts, both acute and delayed |
| | Eye contact: Exposure to the eyes may cause temporary or permanent blindness. Eye exposure may result in conjunctivitis, lacrimation and/or corneal irritation. Total corneal epithelial loss may occur. |
| Acute effects/ | Skin contact: Ammonia is a severe irritant of the skin. Skin exposure to high concentrations of the gas may cause burning and blistering. Contact with liquid may cause severe skin burns. Concentrated ammonia may produce liquefaction necrosis and deep penetrating burns. |
| Delayed effects 4.3 Indication of any immediate medical | Inhalation: Ammonia is toxic and a severe irritant of the respiratory tract. It may cause a running nose, coughing, chest pain, cessation of respiration and death. It may cause severe breathing difficulties, which may be delayed in onset. Bronchospasm, laryngitis, tracheitis, wheezing, dyspnea, and laryngeal stridor may be noted. Mucosal burns to the tracheobronchial tree, pulmonary edema and associated hypoxemia frequently occur following exposure to concentrated ammonia. |

There is no specific antidote. Treat symptoms. Pulmonary edema prophylaxis.

| | SECTION 5: FIRE-FIGHTING MEASURES |
|---------------------------------|--|
| 5.1 Extinguishing media | |
| Suitable extinguishing media: | Fine water spray, normal foam, dry agent (carbon dioxide, dry chemical powder). Stopping the flow of gas rather than extinguishing the fire is usually the best procedure to follow when escaping gas is burning. Water spray can be used to bring down the vapour but should not be sprayed on pools of liquid ammonia. |
| Unsuitable extinguishing media: | None |

5.2 Special hazards arising from the substance or mixture

Formation of explosive gas mixtures in air. Explosion hazard. Heat of fire can build pressure in container and cause it to rupture. Nitrogen oxides, hydrogen, nitrogen can be emitted in case of fire.

5.3 Advice for fire fighters

Do not get water inside container. Move container from fire area if you can do it without risk. Apply cooling water to sides of containers which are exposed to flames until well after fire is out. Stay away from ends of tanks due to exploding potential when tanks are exposed to fire. Isolate area until gas has dispersed. Use water spray or foam to control vapour. Positive pressure selfcontained breathing apparatus (SCBA) should be used when there is a potential for inhalation of vapours and/or fumes. Chemical protective clothing that is safe for use with ammonia involved in a fire should be worn.

SECTION 6: ACCIDENTAL RELEASE MEASURES

6.1 Personal precautions, protective equipment and emergency procedures

6.1.1 For non-emergency personnel

Protective equipment: Gas masks with cartridges for ammonia.

Emergency procedures: Stop leak if you can do so without risk. Keep unnecessary people away, isolate hazard area and deny entry. Stay upwind, out of low areas, and ventilate closed spaces before entering. Evaluate the affected area to determine whether to evacuate or shelter-in-place by taping windows and doors, shutting off outside air intakes (attic fans, etc.), and placing a wet towel or cloth over the face (if needed).

6.1.2 For emergency responders:

With proper training, self-contained breathing apparatus (SCBA) and structural firefighter's protective clothing used in conjunction with water spray will provide limited protection in outdoor releases for short-term exposure. Fully encapsulating, vapor-protective clothing should be worn for spills and leaks with no fire. Use water spray or foam to control vapors. Mixing of water and liquid ammonia will increase vaporization rate. Do not put water on liquid ammonia unless more than 100 volumes of water are available for each volume of liquid ammonia.

6.2 Environmental precautions

Discard any product, residue, disposable container, or liner in an environmentally acceptable manner, in full compliance with federal, state, and local regulations.

6.3 Methods and material for containment and cleaning up



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6.3.1 For containment.

GAS: Do NOT spray water directly on the leak or ammonia container. For a small gas leak, increase ventilation and allow gas to vent to a safe area.

LIQUID: Use water spray to control vapours. Avoid any contact with liquid product. Stop the leakage. The excess of the product should be drained into suitable container. Allow the vapours to disperse. For small liquid spills, increase ventilation and allow the liquid to volatilise to safe area. For large spills, keep away from sources of ignition and sparks. Increase ventilation of spilled area. Self-contained breathing apparatus (SCBA) and appropriate protective clothing should be worn.

6.3.2 For cleaning up:

GAS: For larger gas leaks, use fire hoses equipped with fog nozzles to disperse gas down-wind.

LIQUID: Wash the affected area with great amount of water.

6.3.3 Other information: None

6.4 Reference to other sections

See section 8 for personal protective equipment and section 13 for waste disposal.

| | SECTION 7: HANDLING AND STORAGE | | | | | |
|--|--|--|--|--|--|--|
| 7.1 Precautions for safe handling | | | | | | |
| Protective measures: | Use proper personal protective equipment when working with or around ammonia. Avoid contact with eyes, skin and clothing. Safety shower and eyewash fountain should be provided in the ammonia handling area. | | | | | |
| Measures to prevent fire: | Keep away from sources of ignition. | | | | | |
| Measures to prevent aerosol and dust generation: | Local exhaust ventilation should be provided. | | | | | |
| Measures to protect the environment: | Prevent waste from contaminating the surrounding environment. | | | | | |
| Advice on general occupational hygiene: | Do not eat, drink or smoke in work areas. Remove contaminated clothing and protective equipment before entering eating areas. Wash hands, forearms and face thoroughly after handling chemical products, before eating, smoking and using the lavatory and at the end of the working period. Appropriate techniques should be used to remove potentially contaminated clothing. | | | | | |
| | Wash contaminated clothing before reusing | | | | | |
| 7.2 Conditions for safe storage, including | | | | | | |
| Technical measures/ Storage conditions: | Use transportable pressure equipment. Store in a well-ventilated area. Avoid heat, flames, sparks and other sources of ignition. | | | | | |
| Packing materials: | Use dedicated containers - do not rinse. | | | | | |
| Requirements for storage rooms and vessels: | | | | | | |
| Storage class: | 2A | | | | | |
| Further information on storage conditions: | None | | | | | |
| Incompatible products: | All strong acids and alkali, halogens, perchlorate, chloric acid, chlorine monoxide, chlorites, nitrogen tetroxide, compounds of chrome, selenium, silver, gold compounds, zinc, platinum, tin, sulphur, acetaldehyde, acrolein, boron, hydrogen, potassium and its salts, picric acid, nitrilchloride, nitrogen oxides, nitric acid, hydrazine. | | | | | |
| 7.3 Specific end use(s): | None | | | | | |
| SECTION | 8: EXPOSURE CONTROLS / PERSONAL PROTECTION | | | | | |
| 8.1 Control parameters | | | | | | |
| 8.1.1 National occupational exposure limit | values: Not available | | | | | |
| 8.1.2 National biological limit values: Not a | vailable | | | | | |
| 8.1.3 PNEC (Predicted No Effect Concentre | ation): | | | | | |
| Environmental protection target | PNEC | | | | | |
| Aqua – freshwater | 0.0011 mg/L | | | | | |
| Aqua - marine water | 0.0011 mg/L | | | | | |
| Aqua – intermittent releases | 0.089 mg/L | | | | | |
| Sediment | No exposure expected | | | | | |
| Soil | No exposure expected | | | | | |
| Sewage treatment plant | No exposure expected | | | | | |
| Food chain: oral (secondary poisoning) | No exposure expected | | | | | |
| Air | No exposure expected | | | | | |



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| AZOT | | AQUITE (Q | |
|--|---|--|--|
| | | ACUTE (Systemic eff | |
| | Route | Workers | Effect Level (DNEL) |
| | Oral | Not quantifiable | General population 6.8 mg/kg bw/d |
| | Dermal | 68 mg/kg bw/d | 68 mg/kg bw/d |
| | Inhalation | 47.6 mg/m³ | 23.8 mg/m ³ |
| | | <u> </u> | <u> </u> |
| | | ACUTE (Local effect | |
| | Route | Workers | Effect Level (DNEL) |
| | Oral | Not quantifiable | General population Not quantifiable |
| | Dermal | Not quantifiable | Not quantifiable |
| | Inhalation | 36 mg/m³ | 7.2 mg/m³ |
| 8.1.4 DNEL: | | | <u> </u> |
| | <u></u> | ONG TERM (Systemic | |
| | Route | | Effect Level (DNEL) |
| | | Workers | General population |
| | Oral | Not quantifiable | 6.8 mg/kg bw/d |
| | Dermal Inhalation | 68 mg/kg bw/d 47.6 mg/m³ | 68 mg/kg bw/d 23.8 mg/m³ |
| | Ппаацоп | 47.0 mg/m² | 23.0 HIY/III |
| | | LONG TERM (Local ef | fects) |
| | Pouto | Derived No E | ffect Level (DNEL) |
| | Route | Workers | General population |
| | Oral | Not quantifiable | Not quantifiable |
| | Dermal | Not quantifiable | Not quantifiable |
| | Inhalation | 14 mg/m³ | 2.8 mg/m³ |
| 8.1.5 Monitoring procedures: Not availa | ble | | |
| 0.0 Evenouse controls | | | |
| 8.2 Exposure controls | | | |
| • | | | |
| 8.2.1 Appropriate engineering controls: Substance/mixture related measures tengineering controls (containment, LEV) | to prevent exposure during iden /) and protective equipment (glove | es, goggles/visor, protecti | ve clothing) as appropriate. |
| 8.2.1 Appropriate engineering controls: Substance/mixture related measures to engineering controls (containment, LEV Technical measures to prevent exposacceptable exposure levels, or respirate hoses, eye wash stations, or stock tank | to prevent exposure during iden /) and protective equipment (glove sure: Engineering controls should bry protection will be required to re | es, goggles/visor, protecti d be maintained to kee | ve clothing) as appropriate. ep ammonia concentrations wi |
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| Odour threshold: | 5 ppm |
|--|--|
| pH: | Not available |
| Melting point/Freezing point: | -77,7 °C |
| Initial boiling point and boiling range: | -33,33 °C |
| Flash-point: | The endpoint is not applicable as the substance is an inorganic gas; Aqueous solutions of ammonia did not show any flash point |
| Evaporation rate: | Not available |
| Flammability (solid, gas): | Flammable |
| Upper/lower flammability or explosive limits | 16 % / 25 % |
| Vapour pressure: | 8611 hPa at 20 °C |
| Vapour density: | Not available |
| Relative density: | Not applicable. The substance is a gas at room temperature: this endpoint is not relevant. The density of ammonia is calculated to be 0.717 kg/m³ at NTP (normal temperature and pressure); 0.769 kg/m³ at STP (standard temperature and pressure). |
| Solubility in water: | 482 g/L at 25 °C, 531 g/L at 20 °C. The substance is very soluble in water |
| Partition coefficient n-octanol/water: | Not applicable. As the substance is inorganic |
| Auto-ignition temperature | The auto flammability of anhydrous ammonia was measured to be 651°C |
| Decomposition temperature: | above 400 °C |
| Viscosity: | The substance is a gas at room temperature: this endpoint is not relevant. However the viscosity of anhydrous ammonia was measured to 0,475; 0,317; 0,276 & 0,255 centipoises (mPa.s) at -69, -50, -40 & -33.5°C respectively. |
| Explosive properties | Non explosive (anhydrous ammonia is not predicted to be explosive based on a theoretical assessment of its chemical structure) NB: Risk of explosion in case of contact or reaction with some substances (hypochlorites, mercury, silver). When ammonia is mixed with such gases as: oxygen, chlorine, bromine or iodine; its explosion may be initiated by direct sunlight or by UV radiation. |
| Oxidising properties | No (As the substance does not contain any oxygen or halogen atoms it is not expected to be an oxidising agent). |

9.2 Other information

SECTION 10: STABILITY AND REACTIVITY

10.1 Reactivity

Stable under recommended storage and handling conditions (see section 7, handling and storage).

10.2 Chemical stability

Stable under recommended storage and handling conditions (see section 7, handling and storage); hazardous polymerisation will not occur.

10.3 Possibility of hazardous reactions

The fire or explosion hazard results from its reactions with: acetaldehyde, acrolein, boron trifluoride, bromine, chlorine, chloric acid, chlorine trifluoride, chlorates, chlorosilane, ethylene oxide, fluorine, hydrogen bromide, hypochlorous acid, iodine, nitric acid, nitrogen dioxide, nitrogen trichloride, nitrosyl chloride, phosphorus pentoxide, picric acid, phosphorus and phosphorus hydride, arsenic hydride, antimony hydride, sodium and sulphur dichloride.

Reacts violently with copper, tin, zinc and their alloys, especially under humid conditions.

10.4 Conditions to avoid

Incompatible materials, open flame sources, heat sources / heat - risk of bursting.

10.5 Incompatible materials

Ammonia has potentially explosive or violent reactions with interhalogens, strong oxidisers, nitric acid, fluorine and nitrogen oxide. Ammonia forms sensitive explosive mixtures with air and hydrocarbons, ethanol and silver nitrate and chlorine. Explosive products are formed by the reaction of ammonia with silver chloride, silver oxide, bromine, iodine, gold, mercury and tellurium halides. See also Clause 10.3.

10.6 Hazardous decomposition products

Nitrogen oxides, hydrogen

Hydrogen is released upon heating above 454 °C. The decomposition temperature may be lowered to 300 °C by contact with certain metals such as nickel. At 690 °C or in the presence of an electric spark, ammonia decomposes into nitrogen and hydrogen gases, which may form a flammable mixture in the air.

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

SECTION 11: TOXICOLOGICAL INFORMATION

11.1 Information on toxicological effects

11.1.1 Acute toxicity





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| | 1,5 | | | | | | |
|---------------------|----------------------|--------|---|---------------------|-----------------|---|--|
| Route of exposure | Species | | Method | Effective dose | Exposure time | Results | |
| inhalation | rat male | Asse | ssment of acute inhalation toxicity in the rat | _ | 1 hour | LC50: 9850 mg/m³ air | |
| oral | rat (Wistar) male | | gavage valent or similar to OECD ne 401 (Acute Oral Toxicity) | _ | - | LD50: 350 mg/kg bw (male) (Probit analysis) | |
| dermal | | | No data s the substance is classified a by local effects at the site of co | | | | |
| 11.1.2 Serious ey | • | | Highly irritating | | - | , | |
| 11.1.3 Skin corro | sion/irritation | | Corrosive | | | | |
| 11.1.4 Respirator | ry or skin sensiti | zation | Not sensitising | | | | |
| 11.1.5 Germ cell | mutagenicity | | Negative | | | | |
| 11.1.6 Carcinoge | enicity: | | There is no evidence that the | e substance is c | arcinogenic | | |
| 11.1.7 Reproduct | | | The available data do not toxin | indicate that ar | mmonia is a re | productive or developments | |
| 11.1.8 STOT-sing | - | | Not available | | | | |
| 11.1.9 STOT-rep | · | | Not available | | | | |
| 11.1.10 Aspiratio | n hazard | | Not relevant | | | | |
| | | | SECTION 12: ECOLOGICAL | INFORMATIO | N | | |
| 12.1 Toxicity | | | T | | | | |
| Fish (freshwater, | short-term): | | LC ₅₀ : 0.068 mg/L | | | | |
| Fish (long-term): | | | LC50 for freshwater fish: 0.89 mg/L un-ionised ammonia | | | | |
| Freshwater invert | tebrates (short-te | erm): | EC50/LC50 for freshwater invertebrates: 110 mg/L LC50=101 mg/L (48 h; <i>Daphnia</i> magna) | | | | |
| Freshwater invert | tebrates (long-te | erm): | EC10/LC10 (NOEC): 0.79 mg/L | | | | |
| Freshwater algae | e: | | EC50/LC50: 2700 mg/L | | | | |
| Terrestrial plants: | : | | Not predicted Ammonia is used as a com not predicted | ponent of fertilis | sers; therefore | coxicity to terrestrial plants i | |
| Soil micro-organi | sms: | | Not predicted Toxicity to soil microorganis cycle | sms is unlikely: | ammonia is a | n intrinsic part of the nitrat | |
| Birds: | | | No data are available. A waiver is proposed on grounds of exposure | | | | |
| Mammals: | | | Not available | | | | |
| 12.2 Persistence | e and degradab | ility | | | | | |
| Abiotic degradati | on: | | | | | | |
| Hydrolysis: | | | Not predicted. In aqueous solution, ammon | ia is in equilibriu | ım with the amn | nonium ion. | |
| Excretion: | | | Ammonia is rapidly detoxified in mammals by conversion to urea by the urea cycle ir liver cells, and is subsequently excreted (as urea) in urine following glomerular filtration. | | | | |
| Phototransformat | tion/photolysis: | | | | | | |
| Phototransformat | tion in air: | | Photolytic degradation and reaction with photolytically produced hydroxyl radicals (Ohin the troposphere are major pathways for the removal of atmospheric ammonia | | | | |
| Phototransformat | tion in water: | | Not available | | | | |
| Phototransformat | tion in soil: | | Not available | | | | |
| Biodegradation: | | | T | | | | |
| Biodegradation in | | | Readily biodegradable | | | | |
| Biodegradation in | n sediments: | | No data are available and none are required | | | | |
| Biodegradation in | n soil: | | Ammonia is rapidly biode mineralisation | egraded in so | ii by the prod | cess of ammonification (| |
| 12.3 Bioaccumu | - | | | | | | |
| | | | product of normal metabolism | | | adapating of the control | |
| | | | monia levels are usually low a to nitrate, both processes which | | | ausorphon of the ammoniur | |

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15.2 Chemical safety assessment.

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|--|---|------------------------|---|---|
| 12.5 Results of PBT and vPvB assessm | ent | | | |
| According to Annex XIII of Regulation (Edinorganic. | C) No 1907/2006, no P | BT and vPvB assess | sment has been conduc | cted since ammonia is |
| 12.6 Other adverse effects: None | | | | |
| 12.7 Additional information: None | | | | |
| | SECTION 13: DISPOS | AL CONSIDERATIO | NS | |
| 13.1 Waste treatment methods: | | | | |
| 13.1.1 Product / Packaging disposal: | Dispose of according | to local authority reg | ulations. | |
| Waste codes / waste designations according to LoW (Commission Decision 2001/118/EC): | 06 10 99 Wastes not | otherwise specified | | |
| 13.1.2 Waste treatment-relevant information: | Waste must be dispose surface water without | | al regulations and shoul | d not be discharged to |
| 13.1.3 Sewage disposal-relevant information: | | - | dance with local authorit | ty regulations. |
| 13.1.4 Other disposal recommendations: | None | | | |
| | SECTION 14: TRANS | PORT INFORMATION | ON | |
| | ADR/RID | ADN/ADNR | IMDG | IATA |
| 14.1 UN number | UN1005 | Not regulated | UN 1005 | Not regulated |
| 14.2 UN proper shipping name | Ammonia, anhydrous | - | Ammonia, anhydrous | - |
| 14.3 Transport hazard class(es) | 2.3, 8 | - | 2.3, 8 | - |
| 14.4 Packing group | None | - | - | - |
| 14.5 Environmental hazards | No | No | No | No |
| 14.6 Special precautions for user | Not available | Not available | Not available | Not available |
| Additional information | Classification code: 2TC Label 2.3: toxic substance. Label 8: corrosive substance. | - | EmS number: 2-08 Label FS: 2-03 Marine pollutant: No | Transport forbidden on passenger aircraft - cargo aircraft only |
| 14.7 Transport in bulk according to Ann | ex II of MARPOL 73/78 | and the IBC Code | | |
| Not available | SECTION 15: REGUL | ATORY INFORMATI | ON | |
| 15.1 Safety, health and environmental re | | | | |
| EU Regulations | egulation/registation s | pecine for the subs | tunec or mixture | |
| Authorisations and\or restrictions on use: Authorisation: EU Regulation (EC) No. 1907/2006 (REACH) Annex XIV - List of substances subject to authorisation Substances of very high concern | None of the compone | nts are listed | | |
| Restrictions on use: Annex XVII - Restrictions on the manufacture, placing on the market and use of certain dangerous substances, mixtures and articles Other EU Regulations: | Not applicable | | | |
| Annex I of Seveso II Directive 96/82/EC: | | | | |
| Dangerous substances | CAS number | Qualify | ving quantity (tonnes) for | r the application of |
| | | Lower tier | - Upp | per tier |
| Anhydrous ammonia | 7664-41-7 | 50 | | 200 |
| National regulations (country): Not available | le | | | |

out for this substance.

In accordance with REACH Article 14, a Chemical Safety Assessment has been carried

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SECTION 16: OTHER INFORMATION

The information provided in this safety data sheet is correct to the best of our knowledge, information, and belief at the date of its publication. The information given is designed only as guidance for safe handling, use, processing, storage, transportation, disposal, and release and is not to be considered a warranty or quality specification. The information relates only to the specific material designated and may not be valid for such material used in combination with any other materials or in any proceed, unless specified in the text.

v. 3.0: Changes were made to comply with the Guidance on the compilation of safety data sheets (version 1.1)
v. 3.1: Changes were made to comply with Article 61 (CLP)

v. 4.0: Changes were made taking into account 5th and 8th ATP to CLP

v. 3.0: Page header; 1.1; 1.2; 1.3; 1.4; 2.1; 2.2; 2.3; 3.1; 4.1; 4.3; 6.1; 6.3; 7.1; 7.2; 8.1; 8.2; 9.1; 11.1; 12.2; 12.4; 12.6; 12.7; 13.1; 15.1

v. 3.1: Page header; 2.1.2; 2.2.2; 16.2

v. 4.0: Page header; 1.3; 2.2.1

v. 4.1: Page header; 1.3

v. 4.2: Page header; 1.3

v. 4.3: Page header; 1.3, 1.4

v. 4.4: Page header; 1.4

16.2 Abbreviations and acronyms:

- ADN European Agreement concerning the International Carriage of Dangerous Goods on Inland Waterway
- ADNR ADN Rhine
- · ADR Agreement on Dangerous Goods by Road
- CAS Chemical Abstracts Service
- CLP Classification, Labelling and Packaging of chemicals
- EC European Commission
- EC50 half maximal effective concentration
- EINECS European Inventory of Existing Commercial Chemical Substances
- EmS number Emergency schedule number
- ES Exposure Scenario
- GHS Globally Harmonized System of Classification and Labelling of Chemicals
- IATA International Air Transport Association
- IBC Code International Code for the Construction and Equipment of Ships Carrying Dangerous Chemicals in Bulk
- IMDG International Maritime Dangerous Goods
- IUPAC International Union of Pure and Applied Chemistry
- LC50 Lethal Concentration
- LD50 Lethal Dose
- LEV local exhaust ventilation
- LoW List of Wastes
- MARPOL International Convention for the Prevention of Pollution From Ships
- OECD Organization for Economic Co-operation and Development
- · PBT Persistent, bioaccumulative, toxic chemical
- PJSC Public Joint-Stock Company
- REACH Registration, Evaluation, Authorisation and Restriction of Chemicals
- RID International Rule for Transport of Dangerous Substances by Railway
- STOT Specific Target Organ Toxicity
- STP sewage treatment plant
- UN United Nations
- vPvB very persistent, very bioaccumulative

16.3 Key literature references and sources for data: CSR (Chemical Safety Report), Guidance on safe use etc.

| 16.4 Training advice: | In accordance with the local regulations |
|---------------------------|--|
| 16.5 Further information: | None |





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Exposure Scenario No. 1: Use of ammonia as an intermediate in the chemicals industry

1.1 Exposure Scenario

Ammonia is used by the chemicals industry to manufacture a range of other substances including: nitric acid, alkalis, dyes, pharmaceuticals, cosmetics, vitamins, synthetic textile fibres and plastics.

Ammonia is used as an intermediate in the synthesis of a number of chemicals. It is used in the manufacture of nitric acid (HNO $_3$) which is used in making explosives such as TNT (2,4,6-trinitrotoluene); nitro-glycerine (which is also used as a vasodilator) and PETN (pentaerythritol nitrate). Ammonia is also used in the synthesis of alkalis: sodium hydrogen carbonate (sodium bicarbonate; NaHCO $_3$), soda ash (sodium carbonate, Na $_2$ CO $_3$), hydrogen cyanide (hydrocyanic acid; HCN) and hydrazine (N $_2$ H $_4$) used in rocket propulsion systems.

Ammonia is used to manufacture explosives such as ammonium nitrate (NH₄NO₃). It is also used as an intermediate in the synthesis of dyes, and synthetic 'man-made' fibres such as nylon, rayon and acrylics. It is also used in the manufacture of plastics such as phenolics and polyurethanes.

Ammonia is used in the manufacture of drugs such as sulphonamide which inhibit the growth and multiplication of bacteria that require *p*-aminobenzoic acid (PABA) and for the biosynthesis of folic acids, antimalarials and vitamins (e.g. B vitamins: nicotinamide and thiamine).

Ammonia is also used in the production of ammonium and nitrate salts used in fertilisers.

1.1.1 Description of Activities and Processes Covered in the Exposure Scenario

Processes using ammonia as an intermediate are carried out at large chemical manufacturing facilities. Due to the large size of these facilities, vessels and reactors for chemical synthesis and processing are housed outdoors. Some processes are carried out indoors. Processes are continuous or batch and are carried out in closed systems.

Most chemical manufacturing processes and units are operated automatically by a small number of operators located in separate control rooms. Operators may also carry out routine 'field' inspections around the facility to check that equipment is operating correctly. Other manual operations in the field may also be undertaken such as: preparation of equipment for mechanical or other work (e.g. maintenance), or taking samples or measurements. Workers unload ammonia stored in spheres onto tankers. Tanker unloading generally takes place in the open air and involves connecting or disconnecting pipes or hoses and opening or closing valves.

1.1.2 Operational Conditions Related to Frequency and Duration of Use

Chemical processes using ammonia as an intermediate are closed, continuous or batch processes which can run for long periods without interruption indoors or outdoors, for up to 24 hours/day, 330-360 days per year. Operational control and some field operations such as inspection tours are therefore also carried out continuously (e.g. in shifts covering 24 hour periods daily, without interruption of the processes). Although operators generally work standard shifts of 8 hours/day and a normal working week, with production continuing at weekends, longer shifts up to 12 hours /day can also be carried out. Operators will typically work for 220 days/year. During a typical shift, operators may spend 80 % of their time in a control room and 20 % of their time in the field. Field operation tours can be up to 6 hours/shift, every day. Sampling (10 minutes/sample) for quality control is routinely carried out. Other activities such as maintenance work, are carried out intermittently. Workers also unload ammonia from transportation vehicles into containers. All processes are supervised.

1.1.3 Risk Management Measures

Chemical processes using ammonia as an intermediate involve special equipment and high integrity contained systems with little or no potential for worker exposure. These facilities are usually housed outdoors, with workers being segregated in separate control rooms with no direct contact with chemical processing units The potential for industrial workers to be exposed to ammonia during these processes is therefore negligible since they are located in separate control rooms.

Workers may potentially be exposed to ammonia when conducting field activities (e.g. when operating valves, pumps or tanks etc). All operations are performed in a closed system. Pipelines and vessels are sealed and insulated and sampling is carried out with a closed sample loop. Extract ventilation is provided at openings and points where emission may occur. Anhydrous ammonia is stored in closed containers and tanks. Ammonia is transferred under containment. A good standard of general or controlled ventilation is applied when maintenance activities are carried out. Personal protective clothing (e.g. face/eye protection, helmet, gloves, boots and protective overalls) is worn when any potential contact may arise.

All technological devices have a proper quality certification, and are regularly controlled and maintained to avoid the uncontrolled discharge of ammonia.

Good occupational hygiene and exposure control measures are implemented to minimise the potential for worker exposure. Workers involved in the manufacture, sampling and transfer of anhydrous ammonia to road tankers are well-trained in these procedures and use appropriate protective equipment.

1.2 Exposure Estimation

1.2.1 Workers Exposure

The assessment of worker exposure to anhydrous and aqueous forms of ammonia used as an intermediate in chemical synthesis was carried out for processes relevant to this scenario as identified by PROC codes reflecting: use and storage of ammonia in closed systems with no likelihood of exposure (PROC 1), use in closed, continuous processes with occasional controlled exposure (PROC 2), formulation using closed batch processes (PROC 3), use in batch or other processes (PROC 4), mixing or blending in a batch process (PROC 5), maintenance and clean-down (PROC 8a), transfer (PROC 8b), transfer of ammonia into containers (PROC 9) and analysis of samples (PROC 15). A screening-level (Tier 1) assessment of worker exposure was carried out using the ECETOC Targeted Risk Assessment (TRA) model. The ECETOC TRA was used to predict dermal exposures (expressed as a daily systemic dose in mg/kg bw) and inhalation exposure concentrations (expressed as an airborne concentration in mg/m³) associated with each process defined by PROC codes.

Exposure to workers was assessed taking into account different operational conditions that may be associated with the use of

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ammonia as an intermediate in chemical synthesis and the impact of different exposure control measures. Exposures were determined for task durations of 1- 4 hours or >4 hours and assuming that process are carried out either outdoors, indoors without use of local exhaust ventilation (LEV) or indoors with the use of LEV. To reflect the use of personal protective equipment (PPE), dermal exposures were determined assuming either no gloves or gloves affording 90% protection of the hands are worn. To reflect the use of respiratory protective equipment (RPE), inhalation concentrations were determined assuming either no RPE or RPE affording 95% protection is worn.

The EČETOC TRA model uses a simple algorithm to determine dermal exposures that does not take the physical-chemical properties of a substance into account. The same dermal exposure where therefore predicted for anhydrous and aqueous forms of ammonia. Parameters used in the ECETOC TRA model to assess inhalation exposures were: molecular weight (35g.mol⁻¹ and 17 g.mol⁻¹ for aqueous and anhydrous forms respectively and vapour pressure (the vapour pressure of anhydrous forms of ammonia is 8.6 x 10⁵ Pa at 20°C, whereas the vapour pressure of aqueous ammonia solution between 5 and 25% w/w ranges from 5 x 10³ Pa to 4x10⁴ Pa at 20°C. Systemic dermal exposures have been determined for a worker with bodyweight 70 kg.

1.2.2 Acute/Short-term and Long-term Exposure

Potential systemic dermal exposures and inhalation exposure concentrations predicted by the ECETOC TRA model for processes associated with the use of ammonia in chemical synthesis are shown in Tables 87 and 88 respectively. ECETOC predicts a daily systemic dose following dermal exposure and a typical daily inhalation exposure concentration and does not specifically predict acute (short-term) and chronic (long-term) exposures.

Table 1.1 Dermal exposures to anhydrous or aqueous (in preparations of 5-25 % w/w) ammonia predicted using the ECETOC TRA model for industrial workers during chemical synthesis

Estimated Exposure PROC Description of activity Exposure assumptions Concentration mg/kg bw/d No gloves Gloves worn (90% Duration Use of ventilation worn reduction) Used in a closed process, no likelihood of exposure: Outdoors /Indoors without 0.34 PROC 1 1-4 hrs or >4 hrs 0.03 Storage (closed bulk or I FV container) Outdoors /Indoors without Use in a closed, 1-4 hrs or >4 hrs 1.37 0.14 continuous process with LEV PROC 2 occasional controlled Indoors with LEV 1-4 hrs or >4 hrs 0.14 0.01 exposure (e.g. sampling) Outdoors / Indoors without Use in closed batch 0.34 0.03 1-4 hrs or >4 hrs LEV process (synthesis or PROC 3 formulation) 1-4 hrs or >4 hrs Indoors with LEV 0.03 < 0.01 Use in batch process Outdoors / Indoors without 6.86 0.69 1-4 hrs or >4 hrs (synthesis) where **LEV** PROC 4 opportunity for exposure 0.69 1-4 hrs or >4 hrs Indoors with LEV 0.07 arises Outdoors / Indoors without 1-4 hrs or >4 hrs 13.71 1.37 Mixing or blending in LEV PROC 5 batch process 1-4 hrs or >4 hrs Indoors with LEV 0.07 0.01 Outdoors / Indoors without 13.71 1-4 hrs or >4 hrs 1.37 LEV PROC 8a Maintenance, clean down Indoors with LEV 0.14 1-4 hrs or >4 hrs 0.01 Outdoors / Indoors without Transfer 1-4 hrs or >4 hrs 6.86 0.69 (charging/discharging) LEV from/to vessels or large PROC 8b containers at dedicated Indoors with LEV 1-4 hrs or >4 hrs 0.69 0.07 facilities Outdoors / Indoors without 1-4 hrs or >4 hrs 6.86 0.69 Transfer into small LEV PROC 9 containers Indoors with LEV 1-4 hrs or >4 hrs 0.69 0.07 Outdoors / Indoors without 0.34 0.03 1-4 hrs or >4 hrs Quality control in a PROC 15 laboratory 0.03 1-4 hrs or >4 hrs Indoors with LEV < 0.01

Table 1.2 Inhalation exposure concentrations for anhydrous and aqueous (in preparations of 5-25 % w/w) ammonia predicted using the ECETOC TRA model for industrial workers during chemical synthesis

| | | | | Anhydrou | s ammonia | Aqueous w/w) | ammonia (5-25% |
|--|--------|-----------------------------|----------------------|----------|---|--------------|---------------------|
| Description of activity | PROC | Exposure a | Exposure assumptions | | Estimated Exposure Concentrations mg/m3 | | ns mg/m3 |
| | | Duration Use of ventilation | | No RPE | RPE (95% reduction) | No RPE | RPE (95% reduction) |
| Used in a closed process, no likelihood of | PROC 1 | 1-4 hrs or >4 hrs | Outdoors | 0.00 | NA | 0.01 | NA |





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| (closed bulk or container) | 1-4 hrs or >4 hrs | Indoors without LEV | 0.01 | NA | 0.01 | NA |
|--|--|---|--|---|--|--|
| | >4hrs | Outdoors | 24.79 | 1.24 | 30.63 | 1.53 |
| Use in a closed, | >4hrs | >4hrs Indoors without LEV | | 1.77 | 43.75 | 2.19 |
| continuous process with | >4hrs Indoors with LEV | | 3.53 | 0.18 | 4.38 | 0.22 |
| occasional controlled PROC exposure (e.g. sampling) | 1-4 hrs | Outdoors | 14.88 | 0.74 | 18.38 | 0.92 |
| exposure (e.g. sumpling) | 1-4 hrs | Indoors without LEV | 22.25 | 1.06 | 26.25 | 1.31 |
| | 1-4 hrs | Indoors with LEV | 2.13 | 0.11 | 2.63 | 0.13 |
| | >4hrs | Outdoors | 49.58 | 2.48 | 61.25 | 3.06 |
| | >4hrs | Indoors without LEV | 70.83 | 3.54 | 87.5 | 4.38 |
| Use in closed batch | >4hrs | Indoors with LEV | 7.08 | 0.35 | 8.75 | 0.44 |
| process (synthesis or PROC formulation) | 1-4 hrs | Outdoors | 29.75 | 1.49 | 36.75 | 1.84 |
| , | 1-4 hrs | Indoors without LEV | 42.5 | 2.13 | 52.50 | 2.63 |
| | 1-4 hrs | Indoors with LEV | 4.25 | 0.21 | 5.25 | 0.26 |
| | >4hrs | Outdoors | 49.58 | 2.48 | 61.25 | 3.06 |
| Lies in botch process | >4hrs | Indoors without LEV | 70.83 | 3.54 | 87.5 | 4.38 |
| Use in batch process (synthesis) where | >4hrs | Indoors with LEV | 7.08 | 0.35 | 8.75 | 0.44 |
| opportunity for exposure | 1-4 hrs | Outdoors | 29.75 | 1.49 | 36.75 | 1.84 |
| arises | 1-4 hrs | Indoors without LEV | 42.5 | 2.13 | 52.5 | 2.63 |
| | 1-4 hrs | Indoors with LEV | 4.25 | 0.21 | 5.25 | 0.26 |
| | >4hrs | >4hrs Outdoors | | 6.20 | 153.13 | 7.66 |
| | >4hrs | >4hrs Indoors without LEV | | 8.85 | 218.75 | 10.94 |
| Mixing or blending in | >4hrs | s Indoors with LEV | | 0.89 | 21.88 | 1.09 |
| batch process PROC | 1-4 hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| | 1-4 hrs | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| | 1-4 hrs | Indoors with LEV | 10.63 | 0.53 | 13.13 | 0.66 |
| | >4hrs | >4hrs Outdoors | | 6.20 | 153.13 | 7.66 |
| | >4hrs | >4hrs Indoors without LEV | | 8.85 | 218.75 | 10.94 |
| Maintana de la PROC | >4hrs | >4hrs Indoors with LEV | | 0.89 | 21.88 | 1.09 |
| Maintenance, clean down 8a | 1-4 hrs | 1-4 hrs Outdoors | | 3.72 | 91.88 | 4.59 |
| | 1_// hre | 1-4 hrs Indoors without LEV | | | | |
| | 1-4 1113 | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| | 1-4 hrs | Indoors without LEV Indoors with LEV | 106.25 10.63 | 5.31 0.53 | 131.25 13.13 | |
| | | | | | | 6.56 |
| Transfer of ammonia | 1-4 hrs | Indoors with LEV | 10.63 | 0.53 | 13.13 | 6.56 0.66 |
| (charging/discharging) | 1-4 hrs >4hrs | Indoors with LEV Outdoors | 10.63 74.38 | 0.53 3.72 | 13.13 91.88 | 6.56 0.66 4.59 |
| (charging/discharging) PROC 8h | 1-4 hrs >4hrs >4hrs | Indoors with LEV Outdoors Indoors without LEV | 10.63 74.38 106.25 | 0.53 3.72 5.31 | 13.13 91.88 131.25 | 6.56 0.66 4.59 6.56 |
| (charging/discharging) from/to vessels or large | 1-4 hrs >4hrs >4hrs >4hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV | 10.63 74.38 106.25 3.19 | 0.53 3.72 5.31 0.16 | 13.13 91.88 131.25 3.94 | 6.56 0.66 4.59 6.56 0.20 |
| (charging/discharging) from/to vessels or large containers at dedicated PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors | 10.63 74.38 106.25 3.19 44.63 | 0.53 3.72 5.31 0.16 2.23 | 13.13 91.88 131.25 3.94 55.13 | 6.56 0.66 4.59 6.56 0.20 2.76 |
| (charging/discharging) from/to vessels or large containers at dedicated PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV | 10.63 74.38 106.25 3.19 44.63 63.75 | 0.53 3.72 5.31 0.16 2.23 3.19 | 13.13 91.88 131.25 3.94 55.13 78.75 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 |
| (charging/discharging) from/to vessels or large containers at dedicated PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs 1-4 hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors without LEV | 10.63 74.38 106.25 3.19 44.63 63.75 1.91 | 0.53 3.72 5.31 0.16 2.23 3.19 0.1 | 13.13 91.88 131.25 3.94 55.13 78.75 2.36 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 0.12 |
| (charging/discharging) from/to vessels or large containers at dedicated facilities PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs 24hrs >4hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors without LEV Outdoors Outdoors | 10.63 74.38 106.25 3.19 44.63 63.75 1.91 99.17 | 0.53 3.72 5.31 0.16 2.23 3.19 0.1 4.96 | 13.13 91.88 131.25 3.94 55.13 78.75 2.36 122.50 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 0.12 6.13 |
| (charging/discharging) from/to vessels or large containers at dedicated facilities PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs 24hrs >4hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors with LEV Outdoors Indoors without LEV | 10.63 74.38 106.25 3.19 44.63 63.75 1.91 99.17 141.67 | 0.53 3.72 5.31 0.16 2.23 3.19 0.1 4.96 7.08 | 13.13 91.88 131.25 3.94 55.13 78.75 2.36 122.50 175.00 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 0.12 6.13 8.75 |
| (charging/discharging) from/to vessels or large containers at dedicated facilities PROC 8b PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs >4hrs >4hrs >4hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors with LEV Outdoors Indoors without LEV Indoors without LEV | 10.63 74.38 106.25 3.19 44.63 63.75 1.91 99.17 141.67 | 0.53 3.72 5.31 0.16 2.23 3.19 0.1 4.96 7.08 0.71 | 13.13 91.88 131.25 3.94 55.13 78.75 2.36 122.50 175.00 17.50 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 0.12 6.13 8.75 0.88 |
| (charging/discharging) from/to vessels or large containers at dedicated facilities PROC 8b PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs 24hrs >4hrs >4hrs 1-4 hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors without LEV Outdoors Indoors without LEV Indoors with LEV Outdoors | 10.63 74.38 106.25 3.19 44.63 63.75 1.91 99.17 141.67 14.17 59.50 | 0.53 3.72 5.31 0.16 2.23 3.19 0.1 4.96 7.08 0.71 2.98 | 13.13 91.88 131.25 3.94 55.13 78.75 2.36 122.50 175.00 17.50 73.50 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 0.12 6.13 8.75 0.88 3.68 |
| (charging/discharging) from/to vessels or large containers at dedicated facilities PROC 8b PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs >4hrs >4hrs 1-4 hrs 1-4 hrs >4hrs >4hrs >4hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors without LEV Indoors without LEV Indoors with LEV Outdoors Indoors with LEV | 10.63 74.38 106.25 3.19 44.63 63.75 1.91 99.17 141.67 14.17 59.50 85.00 | 0.53 3.72 5.31 0.16 2.23 3.19 0.1 4.96 7.08 0.71 2.98 4.25 | 13.13 91.88 131.25 3.94 55.13 78.75 2.36 122.50 175.00 17.50 73.50 105.00 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 0.12 6.13 8.75 0.88 3.68 5.25 |
| (charging/discharging) from/to vessels or large containers at dedicated facilities PROC 8b PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs >4hrs >4hrs >1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors without LEV Indoors with LEV Outdoors Indoors with LEV Outdoors Indoors without LEV Indoors without LEV | 10.63 74.38 106.25 3.19 44.63 63.75 1.91 99.17 141.67 14.17 59.50 85.00 8.5 | 0.53 3.72 5.31 0.16 2.23 3.19 0.1 4.96 7.08 0.71 2.98 4.25 0.43 | 13.13 91.88 131.25 3.94 55.13 78.75 2.36 122.50 175.00 17.50 73.50 105.00 10.50 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 0.12 6.13 8.75 0.88 3.68 5.25 0.53 |
| (charging/discharging) from/to vessels or large containers at dedicated facilities PROC 8b PROC 8b | 1-4 hrs >4hrs >4hrs >4hrs 1-4 hrs 1-4 hrs 1-4 hrs 1-4 hrs >4hrs >4hrs 1-4 hrs >4hrs >4hrs >4hrs >4hrs >4hrs >4hrs >4hrs >4hrs -4hrs -4 hrs -4 hrs -4 hrs -4 hrs -4 hrs | Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors with LEV Outdoors Indoors without LEV Indoors without LEV Indoors with LEV Outdoors Indoors with LEV Outdoors Indoors without LEV Indoors without LEV Indoors without LEV | 10.63 74.38 106.25 3.19 44.63 63.75 1.91 99.17 141.67 14.17 59.50 85.00 8.5 35.42 | 0.53 3.72 5.31 0.16 2.23 3.19 0.1 4.96 7.08 0.71 2.98 4.25 0.43 1.77 | 13.13 91.88 131.25 3.94 55.13 78.75 2.36 122.50 175.00 17.50 73.50 105.00 10.50 43.75 | 6.56 0.66 4.59 6.56 0.20 2.76 3.94 0.12 6.13 8.75 0.88 3.68 5.25 0.53 2.19 |





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1.3 General Public / Consumer Exposure

Industrial uses of anhydrous and aqueous ammonia are carried out at industrial sites from which members of the public are excluded. Members of the public will not be exposed to anhydrous or aqueous ammonia during industrial end-use.

1.3.1 Indirect Exposure of Humans via the Environment (oral)

Ammonia is ubiquitous in the environment with <30% of emissions resulting from fertiliser uses and from non-agricultural sources (ref. 'Ammonia in the UK' - DEFRA).

In addition, there is no evidence that ammonia bioaccumulates as the log Kow value is 0.23. Since the trigger of BCF >100 (log Kow>3) is not met, the derivation of PNECs to protect against secondary poisoning is not required.

The risk of indirect exposure of humans via the environment is therefore not considered.

1.4 Environmental Exposure

First tier conservative environmental exposure estimations were carried out using EUSES 2.1 and with the specified defaults. Second tier worst case environmental exposure estimations were carried out using EUSES 2.1 to take into account more realistic factors that affect the environmental concentrations.

1.4.1 Environmental Releases

The environmental releases are determined primarily by tonnage and the ERC in the first tier with conservative estimations and defaults being implemented in EUSES. For the second tier assessments in EUSES, more realistic inputs were chosen to best suit the description of the production and uses of anhydrous ammonia. Emission defaults are those specified by the ECHA "Guidance on information requirements and chemical safety assessment: Chapter R.16: Environmental Exposure Estimation". Regional data and emission fractions were calculated using EUSES. Full EUSES inputs are shown below.

Table 1.3 Predicted Releases to the Environment Tier 1

| ERC | Compartments | Predicted releases | Measured release | Explanation / source of measured data | |
|-----|--------------------------------------|----------------------------------|--|---|--|
| | Release to air | 1.21 x 10 ⁵ kg/day | Predicted values are those calculated by EU the tonnage data and defaults for ERC6A. | | |
| 6A | Release to wastewater | 4.85 x 10 ⁴ kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC6A. | |
| | Soil (direct only) Agricultural soil | NA | - | No directly loss to soil is expected for this ERC. | |

^{*}The predicted releases were estimated using the EUSES 2.1 program.

In reality removal of ammonia in sewage treatment plants is highly efficient being removed first by nitrification to nitrate followed by denitrification resulting in the release of nitrogen gas. Complete consumption within the STP can be assumed and this has been used in the tier 2 assessment within EUSES.

Table 1.4 Predicted Releases to the Environment Tier 2

| ERC | Compartments | Predicted releases | Measured release | Explanation / source of measured data |
|-----|--------------------------------------|-------------------------------|------------------|---|
| | Release to air | 1.21 x 10 ⁵ kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC6A. |
| 6A | Release to wastewater | 4.85 x 10 ⁴ kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC6A. |
| | Soil (direct only) Agricultural soil | NA | - | No direct loss to soil is expected for this ERC |

1.4.2 Exposure concentration in sewage treatment plants (STP)

Table 1.5 Tier 1 Concentrations in sewage

| ERC for Compartment: | Estimated exposure concentrations | | Measured exposure concentrations | | Explanation / source of measured data |
|-------------------------------------|-----------------------------------|------|----------------------------------|------|---------------------------------------|
| | Value | unit | value | unit | |
| Waste water before treatment ERC 6A | 2.42 x 10 ⁻⁴ | mg/L | NA | mg/L | |
| ERC 6A Sewage (STP effluent) | 3.02 x 10 ³ | mg/L | NA | mg/L | |
| ERC 6A Local freshwater | 302 | mg/L | NA | mg/L | 10-fold dilution by receiving waters |

Table 1.6 Tier 2 Concentrations in sewage

| ERC for Compartment: | Estimated exposure concentrations | | Measured exposure concentrations | | Explanation / source of measured data |
|-------------------------------------|-----------------------------------|------|----------------------------------|------|---------------------------------------|
| | Value | unit | value | unit | |
| Waste water before treatment ERC 6A | 2.42 x 10 ⁻⁴ | mg/L | NA | mg/L | |
| ERC 6A Sewage (STP effluent) | 0 | mg/L | NA | mg/L | Based on efficient removal by STP |
| ERC 6A Local freshwater | 0 | mg/L | NA | mg/L | 10-fold dilution by receiving waters |

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| 1.4.3 Exposure concentration in aquatic pelagic compartment | | | | | | |
|---|-------------------------|-------------------------------|--------------------------------------|--|--|--|
| Table 1.7 Tier 1 Predicted Environmental Concentrations (PEC) in aquatic compartment | | | | | | |
| Compartments | PEC aquatic (local | mg/L) | Justification | | | |
| ERC6A Freshwater (in mg/L) | 302 | | | | | |
| ERC 6A Marine water (in mg/L) | 30.2 | | 10-fold dilution by receiving waters | | | |
| Table 1.8 Tier 2 Predicted Environm | ental Concentrations | (PEC) in aquatic compartment | | | | |
| Compartments | PEC aquatic (local | mg/L) | Justification | | | |
| ERC6A Freshwater (in mg/L) | 2.19 x 10 ⁻³ | | | | | |
| ERC6A Marine water (in mg/L) | 5.37 x 10 ⁻⁴ | | | | | |
| 1.4.4 Exposure concentration in | n sediments | | | | | |
| Table 1.9 Tier 1 Predicted Enviror | nmental Concentration | s (PEC) in aquatic sediment c | ompartment | | | |
| Compartments | | PEC aquatic (local) | | | | |
| ERC6A Freshwater sediment (in mg/ | kg) | 327 | | | | |
| ERC6A Marine sediment (in mg/kg) | | 32.7 | | | | |
| Table 1.10 Tier 2 Predicted Environ | mental Concentrations | (PEC) in aquatic sediment co | pmpartment | | | |
| Compartments | | PEC aquatic (local) | | | | |
| ERC6A Freshwater sediment (in mg/ | kg) | 2.37 x 10 ⁻³ | | | | |
| ERC6A Marine sediment (in mg/kg) | | 5.82 x 10 ⁻⁴ | | | | |
| 1.4.5 Exposure concentrations | in soil and groundwa | ater | | | | |
| Upon contact with soil, ammonia will be rapidly converted by a variety of bacteria, actinomycetes and fungi to ammonium (NH ₄ ⁺) by the process of ammonification or mineralization. Ammonium is then rapidly converted to nitrate. Nitrate is subsequently taken up | | | | | | |

Upon contact with soil, ammonia will be rapidly converted by a variety of bacteria, actinomycetes and fungi to ammonium (NH₄[†]) by the process of ammonification or mineralization. Ammonium is then rapidly converted to nitrate. Nitrate is subsequently taken up and utilised by plants or returned to the atmosphere following denitrification; the metabolic reduction of nitrate into nitrogen or nitrous oxide (N2O) gas. The most likely fate of ammonium ions in soils is conversion to nitrates by nitrification. Therefore accumulation of concentrations of ammonia in soil and groundwater will not be expected.

1.4.6 Atmospheric compartment

Table 1.11 Tier 1 local concentrations in air

| ERC | | Estimated local exposure concentrations | Explanation / source of data | | |
|-----|-----------------------------|---|------------------------------|--|--|
| | During emission (mg/m3) | 33.7 | Estimated using EUSES 2.1 | | |
| 6A | Annual average (mg/m3) | 30.5 | Estimated using EUSES 2.1 | | |
| | Annual deposition (mg/m²/d) | 43.9 | Estimated using EUSES 2.1 | | |
| | | | | | |

Table 1.12 Tier 1 Predicted Exposure Concentration (PEC) in air

| ERC | | Local concentration | PEC air (local+regional) | Justification |
|-----|--|---------------------|--------------------------|----------------------------|
| 6A | Annual average PEC in air, total (mg/m3) | 30.5 | 30.5 | Estimated using EUSES 2.1. |

Table 1.13 Tier 2 local concentrations in air

| ERC | | Estimated local exposure concentrations | Explanation / source of data |
|-----|-----------------------------|---|------------------------------|
| | During emission (mg/m3) | 33.7 | Estimated using EUSES 2.1 |
| 6A | Annual average (mg/m3) | 30.5 | Estimated using EUSES 2.1 |
| - | Annual deposition (mg/m²/d) | 43.8 | Estimated using EUSES 2.1 |

Table 1.14 Tier 2 Predicted Exposure Concentration (PEC) in air

| ERC | | Local concentration | PEC air (local+regional) | Justification |
|-----|--|---------------------|--------------------------|----------------------------|
| 6A | Annual average PEC in air, total (mg/m3) | 30.5 | 30.5 | Estimated using EUSES 2.1. |

1.4.7 Exposure concentration relevant for the food chain (Secondary poisoning)

In terms of secondary poisoning, there is no evidence that ammonia bioaccumulates as the log Kow value is 0.23. Since the trigger of BCF >100 (log Kow>3) is not met, the derivation of PNECs to protect against secondary poisoning is not required. Risk characterisation ratios cannot therefore be derived.

1.4.8 Regional exposure levels and environmental concentrations.

Anhydrous ammonia is produced and used at many sites throughout a region and this may lead to a degree of regional exposure. Regional exposure has been modelled for this exposure scenario using the regional module of EUSES 2.1.

Table 1.15 Tier 1 regional concentrations in the environment



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| | | | Measured regional concentrations | exposure | Explanation / source of measured data |
|---------------------------------|---------------------------------|-------------------|----------------------------------|-------------------|---------------------------------------|
| | PEC value | unit | Measured value | unit | |
| ERC 6a Freshwater | 2.68 x 10 ⁻² | mg/L | NA | mg/L | |
| ERC 6a Marine water | 2.67 x 10 ⁻³ | mg/L | NA | mg/L | |
| ERC 6a Freshwater sediments | 2.56 x 10 ⁻² | mg/kg | NA | mg/kg | |
| ERC 6a Marine sediments | 2.56 x 10 ⁻³ | mg/kg | NA | mg/kg | |
| ERC 6a Agricultural soil | 1.00 x 10 ⁻³ | mg/kg | NA | mg/kg | |
| ERC 6a Grassland | 1.47 x 10 ⁻³ | mg/kg | NA | mg/kg | |
| ERC 6a Air | 2.24 x 10 ⁻³ | mg/m ³ | NA | mg/m ³ | |
| Table 1.16 Tier 2 regional cond | centrations in the en | vironment | | | |
| | Predicted region Concentrations | al Exposure | Measured regional concentrations | exposure | Explanation / source of measured data |
| | PEC value | unit | Measured value | unit | |
| ERC 6a Freshwater | 2.19 x 10 ⁻³ | mg/L | NA | mg/L | |
| ERC 6a Marine water | 5.37 x 10 ⁻⁴ | mg/L | NA | mg/L | |
| ERC 6a Freshwater sediments | 2.09 x 10 ⁻³ | mg/kg | NA | mg/kg | |
| ERC 6a Marine sediments | 5.15 x 10 ⁻⁴ | mg/kg | NA | mg/kg | |
| ERC 6a Agricultural soil | 9.88 x 10 ⁻⁴ | mg/kg | NA | mg/kg | |
| ERC 6a Grassland | 1.39 x 10 ⁻³ | mg/kg | NA | mg/kg | |
| ERC 6a Air | 2.12 x 10 ⁻³ | mg/m ³ | NA | mg/m ³ | |





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Exposure Scenario No. 2: Industrial end-use: Use of ammonia as a processing aid, non-processing aid and auxiliary agent

2.1 Exposure scenario

Anhydrous liquid and aqueous solutions of ammonia are used by a range of industry sectors in a broad number of applications. These include industrial use as a reactive or non-reactive processing aid in continuous or batch processes, as an auxiliary agent or as substance in a closed system. Common industrial end-uses of ammonia are shown below in table 2.1.

Table 2.1 Common industrial end-uses of ammonia

| Table 2.1 Common industrial end-uses of ammonia | | | | | | | | |
|--|-------------|-----|-----|-----|--------|---|--|--|
| Industrial end-use | Type of use | | | | | Description of use | | |
| industrial end-use | ₽ - | Z o | R a | ⋖ : | O S | Description of use | | |
| Use as developing agent in photochemical processes | х | | | | | Ammonia is used as a developing agent in photochemical processes such as white printing, blue printing and in the diazo duplication press. | | |
| Use of refrigerant systems | | х | | | х | Anhydrous liquid ammonia is used as a refrigerant in household, commercial and industrial systems due to its high heat of vaporisation and relative ease of liquefaction. | | |
| Insulation products | | Х | | | | | | |
| Inks and toners | Х | Х | | | | Ammonia vapours are used as a reagent in treating writing or ink marks | | |
| Coatings, thinners, paint removers | Х | Х | | | | | | |
| Processing aid in chemicals industry | | | Х | | | | | |
| Use as an extraction agent | | | Х | | | Ammonia is used as an extraction agent in the mining industry to extract metals like copper, nickel and molybdenum from their ores. | | |
| Treatment of gas (NOx and SOx reduction) | | | X | | x | Ammonia is used in stack emission control systems to neutralise sulphur oxides from combustion of sulphur-containing fuels, as a method of NOx control in both catalytic and non-catalytic applications and to enhance the efficiency of electrostatic precipitators for particulate control. | | |
| Processing aid in nutrition | | | Х | | Х | The food and beverage industry use ammonia as a source of nitrogen required for yeast and micro-organism | | |
| Use as neutralising agent | | | Х | | х | Ammonia is used by the petrochemical industry in neutralizing the acid constituents of crude oil and in the protection of equipment from corrosion | | |
| Textile dyes | | | Χ | | | | | |
| Treatment of water | х | | х | | | Aqueous ammonia is used in water and waste-water treatment areas to control pH, to regenerate weak anion exchange resins and as an oxygen scavenger in boiled water treatment. In water disinfection, aqueous ammonia is added to water containing free chlorine to produce a chloramines disinfectant. | | |
| Use as washing and cleaning products | х | | X | | | Weak ammonia solutions are used extensively within industry, by professionals and consumers as commercial and household cleaners and detergents cleaning products. Commercial ammonia cleaning products contain up to 30% ammonia whereas household products contain 5-10% ammonia | | |
| Treatment of textiles | | Χ | Χ | | | Liquid ammonia is used to increase the quality of textiles | | |
| Treatment of pulp and paper | | Х | Х | | | Ammonia is used in the pulp and paper industry to pulp wood and as a casein dispersant to coat paper. | | |
| Treatment of leather | | Х | Х | | | The leather industry utilises ammonia as a curing agent, as a slime and mould preservative in tanning liquors and as a protective agent for leather and furs in storage | | |
| Treatment of wood | Х | | Х | | | Anhydrous ammonia fumes are used to darken wood in a process called "ammonia fuming" | | |
| Treatment of metal surfaces | Х | | х | | | Ammonia is used in metal treatment processes such as nitriding, carbonitriding, bright annealing, furnace brazing, sintering, sodium hydride descaling, atomic hydrogen welding and other application where protective atmospheres are required. | | |
| Treatment of rubber/latex | | х | х | | | Concentrated aqueous ammonia is used in the rubber industry as a preservative for natural and synthetic latex due to its antibacterial and alkaline properties and as a stabiliser to prevent pre-mature coagulation (e.g. "ammoniation" of natural rubber latex. | | |
| Manufacture of semiconductors/electronic s | | | | х | | Ammonia is used in the electronics industry in the manufacturing of semiconductor chips. | | |



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| Adhesives, sealants | Х | | Х | | |
|----------------------|---|---|---|---|---|
| Polymer preparations | Χ | | Х | | |
| Aircare products | | | | Х | |
| Preservatives | | Х | | | Ammonia is uses as a preservative for the storage of high moisture corn |

2.1.2 Description of activities and processes covered in the exposure scenario

Operational conditions pertaining to the broad range of industrial end-use scenarios involving anhydrous and aqueous forms of ammonia vary considerably across applications and industrial sector of use. A full characterisation of the frequency and duration of tasks is beyond the scope of this report. For the purpose of worker exposure estimation, activities and processes associated with the industrial end-use of ammonia have been represented generically, based on the process categories (e.g. PROC codes) defined by REACH guidance. Processes and activities relevant to ES 2 are described as: use and storage of ammonia in closed systems with no likelihood of exposure (PROC 1), use in close, continuous process with occasional controlled exposure (PROC 2), formulation using closed batch processes (PROC 3), use in batch or other processes (PROC 4), mixing or blending in a batch process (PROC 5), industrial spraying (PROC 7), maintenance and clean down (PROC 8a), transfer (PROC 8b), transfer of ammonia into containers (PROC 9), brush and roller applications (PROC 10), treatment of articles by dipping and pouring (PROC 13), and analysis of samples (PROC 15) and hand-mixing (PROC 19).

2.1.3 Operational conditions related to frequency and duration of use

Operational conditions pertaining to the broad range of industrial end-use scenarios involving anhydrous and aqueous forms of ammonia vary considerably across applications and industrial sector of use. A full characterisation of the frequency and duration of tasks is beyond the scope of this report. For the purposes of worker exposure estimation, operational conditions have been represented generically based on the assumptions that tasks may be either 1-4 hours or >4 hours in duration and processes can be carried out either outdoors, indoors without LEV or indoors with LEV.

2.1.4 Risk Management Measures

Industrial end-uses of anhydrous and aqueous forms of ammonia involve special equipment and high integrity contained systems with little or no potential for worker exposure. Facilities may be housed outdoors, with workers being segregated in separate control rooms with no direct contact with chemical processing units The potential for industrial workers to be exposed to ammonia during these processes is therefore negligible since they are located in a separate control room.

Workers may potentially be exposed to ammonia when conducting field activities (e.g. when operating valves, pumps or tanks etc). All operations are performed in a closed system. Pipelines and vessels are sealed and insulated and sampling is carried out with a closed sample loop. Extract ventilation is provided at openings and points were emission may occur. Anhydrous ammonia is stored in closed containers and tanks. Ammonia is transferred under containment. A good standard of general or controlled ventilation is applied when maintenance activities are carried out. Personal protective clothing (e.g. face/eye protection, helmet, gloves, boots and protective overalls) is worn when any potential contact may arise.

All technological devices have a proper quality certification, and are regularly controlled and maintained to avoid the uncontrolled discharge of ammonia.

Good occupational hygiene and exposure control measures are implemented to minimise the potential for worker exposure. Workers involved in the manufacture, sampling and transfer of anhydrous ammonia to road tankers are well-trained in these procedures and use of appropriate protective equipment.

2.2 Exposure Estimation

The assessment of worker exposure to anhydrous and aqueous forms of ammonia in industrial end-use applications was carried out for processes relevant to this scenario as identified by PROC codes reflecting: use and storage of ammonia in closed systems with no likelihood of exposure (PROC 1), use in closed, continuous processes with occasional controlled exposure (PROC 2), formulation using closed batch processes (PROC 3), use in batch or other processes (PROC 4), mixing or blending in a batch process (PROC 5), industrial spraying (PROC 7), maintenance and clean down (PROC 8a), transfer (PROC 8b), transfer of ammonia into containers (PROC 9), brush and roller applications (PROC 10), treatment of articles by dipping and pouring (PROC 13), and analysis of samples (PROC 15) and hand-mixing (PROC 19).

A screening-level (Tier 1) assessment of worker exposure was carried out using the ECETOC Targeted Risk Assessment (TRA) model. The ECETOC TRA was used to predict dermal exposures (expressed as a daily systemic dose in mg/kg bw) and inhalation exposure concentrations (expressed as an airborne concentration in mg/m3) associated with each process defined by PROC codes.

Exposure to workers was assessed taking into account different operational conditions that may be associated with the industrial end-use of ammonia and the impact of different exposure control measures. Exposures were determined for task durations of 1- 4 hours or >4 hours and assuming that process are carried out either outdoors, indoors without use of local exhaust ventilation (LEV) or indoors with the use of LEV. To reflect the use of personal protective equipment (PPE), dermal exposures were determined assuming either no gloves or gloves affording 90% protection of the hands are worn. To reflect the use of respiratory protective equipment (RPE), inhalation exposure concentrations were determined assuming either no RPE or RPE affording 95% protection is worn.

The ECETOC TRA model uses a simple algorithm to determine dermal exposures that does not take the physical-chemical properties of a substance into account. The same dermal exposure where therefore predicted for anhydrous and aqueous forms of ammonia. Parameters used in the ECETOC TRA model to assess inhalation exposures were: molecular weight (35 g.mol⁻¹ and 17 g.mol⁻¹ for aqueous and anhydrous forms respectively) and vapour pressure (the vapour pressure of anhydrous forms of ammonia is 8.6 x 10⁵ Pa at 20°C, whereas the vapour pressure of aqueous ammonia solution between 5 and 25% w/w ranges from 5 x 10³ Pa to 4 x10⁴ Pa at 20°C. Systemic dermal exposures have been determined for a worker with bodyweight 70 kg.





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2.2.1 Acute/Short term and long-term exposure

Potential systemic dermal exposures and inhalation exposure concentrations predicted by the ECETOC TRA model for the industrial end-use of ammonia are shown in Tables 2.2 and 2.3 respectively. ECETOC predicts a daily systemic dose following dermal exposure and a typical daily inhalation exposure concentration and does not specifically predict acute (short-term) and chronic (long-term) exposures.

Table 2.2 Dermal exposures to anhydrous or aqueous (in preparations of 5-25 % w/w) ammonia predicted using the ECETOC

TRA model for industrial workers during industrial end-use processes

| Description of activity | PROC | Exposure assum | ptions | Estimated Exposure Concentration mg/kg bw/d | | | |
|---|----------|---|--------------------------------|---|-----------------------------|--|--|
| | | Duration | Use of ventilation | No gloves worn | Gloves worn (90% reduction) | | |
| Used in a closed process, no likelihood of exposure: Storage (closed bulk or container) | PROC 1 | 1-4 hrs or >4 hrs | Outdoors /Indoors without LEV | 0.34 | 0.03 | | |
| Use in a closed, | | 1-4 hrs or >4 hrs | Outdoors /Indoors without LEV | 1.37 | 0.14 | | |
| continuous process with occasional controlled exposure (e.g. sampling) | PROC 2 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.14 | 0.01 | | |
| Use in closed batch | DD00 2 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 0.34 | 0.03 | | |
| process (synthesis or formulation) | PROC 3 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.03 | <0.01 | | |
| Use in batch process | | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 6.86 | 0.69 | | |
| (synthesis) where opportunity for exposure arises | PROC 4 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.69 | 0.07 | | |
| Mixing or blending in | PROC 5 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 13.71 | 1.37 | | |
| batch process | | 1-4 hrs or >4 hrs | Indoors with LEV | 0.07 | 0.01 | | |
| Industrial agreesing | PROC 7 | 1-4 hrs or >4 hrs Outdoors / Indoors without LE | | 42.86 | 4.29 | | |
| Industrial spraying | | 1-4 hrs or >4 hrs | Indoors with LEV | 2.14 | 0.21 | | |
| Maintenance, clean | PROC 8a | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 13.71 | 1.37 | | |
| down | PROC 6a | 1-4 hrs or >4 hrs | Indoors with LEV | 0.14 | 0.01 | | |
| Transfer | | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 6.86 | 0.69 | | |
| (charging/discharging) from/to vessels or large containers at dedicated facilities | PROC 8b | 1-4 hrs or >4 hrs | Indoors with LEV | 0.69 | 0.07 | | |
| Transfer into small | PROC 9 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 6.86 | 0.69 | | |
| containers | 11003 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.69 | 0.07 | | |
| Roller application or | PROC 10 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 27.43 | 2.74 | | |
| brushing | PROC 10 | 1-4 hrs or >4 hrs | Indoors with LEV | 1.37 | 0.14 | | |
| Treatment of articles by | PROC 13 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 13.71 | 1.37 | | |
| dipping and pouring | 1 100 13 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.69 | 0.07 | | |
| Quality control in a | PROC 15 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 0.34 | 0.03 | | |
| laboratory | 1 100 13 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.03 | <0.01 | | |
| And-mixing with intimate contact and PPE only | PROC 19 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 141.73 | 14.13 | | |

Table 2.3 Inhalation exposure concentrations for anhydrous and aqueous (in preparations of 5-25 % w/w) ammonia predicted using the ECETOC TRA model for industrial workers during industrial end-use processes

| doing the Louise | THOUGH TOT ITTUGORITA | i montoro aaning inaaotinar | 31.14 400 | produced |
|------------------|-----------------------|-----------------------------|-----------|----------|
| | | | Anhy | /drous |

| Description of activity | DDOO | Exposure assumptions | | Anhydrous ammonia Estimated Exposure Concentration mg/m3 | | Aqueous ammonia (5-25% w/w) | |
|---|--------|----------------------|--------------------|--|---------------------|---|---------------------|
| Description of activity | PROC | | | | | Estimated Exposure Concentration mg/m3 | |
| | | Duration | Use of ventilation | No RPE | RPE (95% reduction) | No RPE | RPE (95% reduction) |
| Used in a closed process, no likelihood of exposure | PROC 1 | 1-4 hrs or >4 hrs | Outdoors | 0.00 | NA | 0.01 | 0.00 |





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| | | 1-4 hrs or >4 hrs | Indoors without LEV | 0.01 | NA | 0.01 | 0.00 |
|---|--------|----------------------|------------------------|--------|------|--------|-------|
| | | >4hrs | Outdoors | 24.79 | 1.24 | 30.63 | 1.53 |
| Use of ammonia in a | | >4hrs | Indoors without LEV | 35.42 | 1.77 | 43.75 | 2.19 |
| closed, continuous process with occasional | PROC 2 | >4hrs | Indoors with LEV | 3.53 | 0.18 | 4.38 | 0.22 |
| controlled exposure (e.g. | PROC 2 | 1-4 hrs | Outdoors | 14.88 | 0.74 | 18.38 | 0.92 |
| sampling) | | 1-4 hrs | Indoors without LEV | 22.25 | 1.06 | 26.25 | 1.31 |
| | | 1-4 hrs | Indoors with LEV | 2.13 | 0.11 | 2.63 | 0.13 |
| | | >4hrs | Outdoors | 49.58 | 2.48 | 61.25 | 3.06 |
| Here of a constant of the least | | >4hrs | Indoors without LEV | 70.83 | 3.54 | 87.5 | 4.38 |
| Use of ammonia in closed batch process (synthesis | PROC 3 | >4hrs | Indoors with LEV | 7.08 | 0.35 | 8.75 | 0.44 |
| or formulation) | | 1-4 hrs | Outdoors | 29.75 | 1.49 | 36.75 | 1.84 |
| | | 1-4 hrs | Indoors without LEV | 42.5 | 2.13 | 52.50 | 2.63 |
| | | 1-4 hrs | Indoors with LEV | 4.25 | 0.21 | 5.25 | 0.26 |
| | | >4hrs | Outdoors | 49.58 | 2.48 | 61.25 | 3.06 |
| Use of ammonia in batch | | >4hrs | Indoors without LEV | 70.83 | 3.54 | 87.5 | 4.38 |
| process (synthesis) | PROC 4 | >4hrs | Indoors with LEV | 7.08 | 0.35 | 8.75 | 0.44 |
| where opportunity for exposure arises | 111001 | 1-4 hrs | Outdoors | 29.75 | 1.49 | 36.75 | 1.84 |
| exposure anses | | 1-4 hrs | Indoors without LEV | 42.5 | 2.13 | 52.5 | 2.63 |
| | | 1-4 hrs | Indoors with LEV | 4.25 | 0.21 | 5.25 | 0.26 |
| | PROC 5 | >4hrs | Outdoors | 123.96 | 6.20 | 153.13 | 7.66 |
| | | >4hrs | Indoors without LEV | 177.08 | 8.85 | 218.75 | 10.94 |
| Mixing or blending in | | >4hrs | Indoors with LEV | 17.71 | 0.89 | 21.88 | 1.09 |
| batch process | FROC 5 | 1-4 hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| | | 1-4 hrs | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| | | 1-4 hrs | Indoors with LEV | 10.63 | 0.53 | 13.13 | 0.66 |
| | | >4hrs | Outdoors | NA | NA | 306.25 | 15.31 |
| | | >4hrs | Indoors without LEV | NA | NA | 437.5 | 21.88 |
| Industrial spraying | PROC 7 | >4hrs | Indoors with LEV | NA | NA | 21.88 | 1.09 |
| madatilai apraying | 111007 | 1-4 hrs | Outdoors | NA | NA | 183.75 | 9.19 |
| | | 1-4 hrs | Indoors without LEV | NA | NA | 262.5 | 13.13 |
| | | 1-4 hrs | Indoors with LEV | NA | NA | 13.13 | 0.66 |
| | | >4hrs | Outdoors | 123.96 | 6.20 | 153.13 | 7.66 |
| Transfer of ammonia | | >4hrs | Indoors without LEV | 177.08 | 8.85 | 218.75 | 10.94 |
| (charging/discharging) from/to vessels or large | PROC | >4hrs | Indoors with LEV | 17.71 | 0.89 | 21.88 | 1.09 |
| containers at non- | 8a | 1-4 hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| dedicated facilities | | 1-4 hrs | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| | | 1-4 hrs | Indoors with LEV | 10.63 | 0.53 | 13.13 | 0.66 |
| Transfer of access to | | >4hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| Transfer of ammonia (charging/discharging) | PROC | >4hrs | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| from/to vessels or large containers at dedicated | 8b | >4hrs | Indoors with LEV | 3.19 | 0.16 | 3.94 | 0.20 |
| facilities | | 1-4 hrs | Outdoors | 44.63 | 2.23 | 55.13 | 2.76 |
| | | 1-4 hrs | Indoors without | 63.75 | 3.19 | 78.75 | 3.94 |





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| | | | LEV | | | | |
|---------------------------|--------|---------|------------------------|--------|------|--------|-------|
| | | 1-4 hrs | Indoors with LEV | 1.91 | 0.1 | 2.36 | 0.12 |
| | | >4hrs | Outdoors | 99.17 | 4.96 | 122.50 | 6.13 |
| | | >4hrs | Indoors without LEV | 141.67 | 7.08 | 175.00 | 8.75 |
| Transfer of ammonia into | PROC 9 | >4hrs | Indoors with LEV | 14.17 | 0.71 | 17.50 | 0.88 |
| small containers | PROC 9 | 1-4 hrs | Outdoors | 59.50 | 2.98 | 73.50 | 3.68 |
| | | 1-4 hrs | Indoors without LEV | 85.00 | 4.25 | 105.00 | 5.25 |
| | | 1-4 hrs | Indoors with LEV | 8.5 | 0.43 | 10.50 | 0.53 |
| | | >4hrs | Outdoors | NA | NA | 153.13 | 7.66 |
| | | >4hrs | Indoors without LEV | NA | NA | 218.75 | 10.94 |
| Roller application or | PROC | >4hrs | Indoors with LEV | NA | NA | 21.88 | 1.09 |
| brushing | 10 | 1-4 hrs | Outdoors | NA | NA | 91.88 | 4.59 |
| | | 1-4 hrs | Indoors without LEV | NA | NA | 131.25 | 6.56 |
| | | >4hrs | Outdoors | NA | NA | 13.13 | 0.66 |
| | | >4hrs | Outdoors | 123.96 | 6.20 | 153.13 | 7.66 |
| | | >4hrs | Indoors without LEV | 177.08 | 8.85 | 218.75 | 10.94 |
| Treatment of articles by | PROC | >4hrs | Indoors with LEV | 17.71 | 0.89 | 21.88 | 1.09 |
| dipping and pouring | 13 | 1-4 hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| | | 1-4 hrs | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| | | 1-4 hrs | Indoors with LEV | 10.63 | 0.53 | 13.13 | 0.66 |
| | | >4hrs | Indoors without LEV | 35.42 | 1.77 | 43.75 | 2.19 |
| Use as a laboratory agent | PROC | >4hrs | Indoors with LEV | 3.54 | 0.18 | 4.38 | 0.22 |
| Ose as a laboratory agent | 15 | 1-4 hrs | Indoors without LEV | 21.25 | 1.06 | 26.25 | 1.31 |
| | | 1-4 hrs | Indoors with LEV | 2.13 | 0.11 | 2.63 | 0.13 |
| | | <4 hrs | Outdoors | NA | NA | 153.13 | 7.66 |
| Hand-mixing with intimate | PROC | <4 hrs | Indoors without LEV | NA | NA | 218.75 | 10.94 |
| contact and PPE only | 19 | 1-4 hrs | Outdoors | NA | NA | 91.88 | 4.59 |
| | | 1-4 hrs | Indoors without LEV | NA | NA | 131.25 | 6.56 |

2.3 General public / consumer exposure

Industrial uses of anhydrous and aqueous ammonia are carried out at industrial sites from which members of the public are excluded. Members of the public will not be exposed to anhydrous or aqueous ammonia during industrial end-use.

2.3.1 Indirect exposure of humans via the environment (oral)

Ammonia is ubiquitous in the environment with <30% of emissions resulting from fertiliser uses and from non-agricultural sources (ref. 'Ammonia in the UK' - DEFRA).

In addition, there is no evidence that ammonia bioaccumulates as the log Kow value is 0.23. Since the trigger of BCF >100 (log Kow>3) is not met, the derivation of PNECs to protect against secondary poisoning is not required.

The risk of indirect exposure of humans via the environment is therefore not considered.

2.4 Environmental exposure

First tier conservative environmental exposure estimations were carried out using EUSES 2.1 and with the specified defaults. Second tier worst case environmental exposure estimations were carried out using EUSES 2.1 to take into account more realistic factors that affect the environmental concentrations.

2.4.1 Environmental releases

The environmental releases are determined primarily by tonnage and the ERC in the first tier with conservative estimations and defaults being implemented in EUSES. For the second tier assessments in EUSES, more realistic inputs were chosen to best suit the description of the production and uses of anhydrous ammonia. Emission defaults are those specified by the ECHA "Guidance on information requirements and chemical safety assessment: Chapter R.16: Environmental Exposure Estimation". Regional data and emission fractions were calculated using EUSES. Full EUSES inputs are shown below.

Table 2.4 Predicted Releases to the Environment Tier 1



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| ERC | Compartments | Predicted releases | Measured release | Explanation / source of measured data |
|-----|--------------------------------------|----------------------------------|------------------|---|
| | Release to air | 7.15 x 10 ⁴ kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC4. |
| 4 | Release to wastewater | 7.52 X 10 ⁴ kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC4. |
| | Soil (direct only) Agricultural soil | NA | - | No direct loss to soil is expected for this ERC. |
| | Release to air | 3.76 X 10 ⁴ kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC5. |
| 5 | Release to wastewater | 3.76 X 10 ⁴ kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC5. |
| | Soil (direct only) Agricultural soil | NA | - | No direct loss to soil is expected for this ERC. |
| | Release to air | 75.2 kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC6B. |
| 6B | Release to wastewater | 3760 kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC6B. |
| | Soil (direct only) Agricultural soil | NA | - | No direct loss to soil is expected for this ERC. |
| | Release to air | 3760 kg/day | - | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC7. |
| 7 | Release to wastewater | 3760 kg/day - | | Predicted values are those calculated by EUSES using the tonnage data and defaults for ERC7. |
| | Soil (direct only) Agricultural soil | NA | - | No direct loss to soil is expected for this ERC. |

^{*}The predicted releases were estimated using the EUSES 2.1 program.

In reality removal of ammonia in sewage treatment plants is highly efficient being removed first by nitrification to nitrate followed by denitrification resulting in the release of nitrogen gas. Complete consumption within the STP can be assumed and this has been used in the tier 2 assessment within EUSES.

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| Tak | ole 2.5 RMMs and | l magazir | ad values for | tier 2 associ | emont | | | | | |
|--|---|------------|-------------------------------|---|----------------------------------|--------------|---|--|------------------------------|--|
| | ption of RMM | Details | | 1 | | accon | nt in EUSES | | Comments | |
| Efficient removal of 0 mg/L (Local) Lo | | | Lowering of Applied at | owering of calculated concentration in STP effluent. oplied at both a local and a regional level. All regional missions to STP. | | | | | | |
| Tak | ole 2.6 Predicted F | Releases | to the Enviro | nment Tier | 2 | | | | | |
| ERC | Compartments | | Predicted | releases | Meas relea | sured ise | - | / source of measur | | |
| | Release to air | | 7.15 x 10 ⁴ l | kg/day | - | | Predicted values are those calculated by EUSES usin the tonnage data and defaults for ERC4. | | | |
| 4 | Release to wast | | 7.52 X 10 ⁴ kg/day | | - | | | alues are those calcudate and defaults for | ulated by EUSES using ERC4. | |
| | Soil (direct only) Agricultural soil | | NA | | - | | | s to soil is expected f | | |
| | Release to air | | 3.76 X 10⁴ | kg/day | - | | the tonnage | data and defaults for | | |
| 5 | Release to wast | | 3.76 X 10 ⁴ kg/day | | - | | | alues are those calcudate and defaults for | ulated by EUSES using ERC5. | |
| | Soil (direct only) Agricultural soil | | NA | | - | | | s to soil is expected f | | |
| | Release to air | | 75.2 kg/day | / | - | | the tonnage | data and defaults for | | |
| 6B | Release to wast | | 3760 kg/da | У | - | | | alues are those calcudate and defaults for | ulated by EUSES using ERC6B. | |
| | Soil (direct only) Agricultural soil |) | NA | | - | | | s to soil is expected f | | |
| | Release to air | | 3760 kg/da | у | the to | | the tonnage | d values are those calculated by EUSES using age data and defaults for ERC7. | | |
| 7 | Release to wastewater 3760 kg/day | | у | | | | alues are those calcudate and defaults for | lated by EUSES using ERC7. | | |
| | Soil (direct only) Agricultural soil | 1 | NA | | - No direct loss to soil is expe | | | s to soil is expected f | or this ERC. | |
| 2.4.2 E | xposure concent | tration in | sewage tre | atment pla | nt (STP | ') | | | | |
| Tak | ole 2.7 General er | mission fr | actions from | the municip | al STP | , | | | | |
| Fractio | n description | | | | | Fract | ion amount | | unit | |
| Fraction | n of emission direc | cted to ai | r by STP | | 0.583 | | | | % | |
| | n of emission direc | | | | 12.4 | | | | % | |
| Fraction | n of emission direc | cted to sl | udge by STP | 1 | 0.13 | | | | % | |
| Fraction | n of emission degi | raded by | STP | | 86.8 | | | | % | |
| 2.4.3 E | xposure concent | tration in | aquatic pel | agic compa | artmen | t | | | | |
| Tak | ole 2.8 Tier 1 Pred | dicted En | vironmental (| Concentratio | ns (PE | C) in ac | quatic compart | ment | | |
| Compa | ırtments | | | PEC ac | quatic (| local m | ng/L) | Justification | | |
| | reshwater (in mg | • | | 468 | | | | | | |
| ERC4 N | Marine water (in m | ng/L) | | 46.8 | | | | | | |
| | reshwater (in mg | | | 234 | | | | | | |
| | Marine water (in m | • . | | 23.4 | | | | | | |
| ERC6B Freshwater (in mg/L) | | 23.4 | 23.4 | | | | | | | |
| | ERC6B Marine water (in mg/L) | | 2.34 | | | | | | | |
| | reshwater (in mg. | | | 23.4 | | | | | | |
| | ERC7 Marine water (in mg/L) | | 2.34 | | | | | | | |
| | ole 2.9 Tier 2 Pred | dicted En | vironmental (| | | | | 1 | | |
| | rtments | | | PEC ac | | local m | ng/L) | Justification | | |
| | reshwater (in mg. | • | | 2.82 x | | | | | | |
| | Marine water (in m | - | | 6.06 x | | | | | | |
| ERC5 F | ERC5 Freshwater (in mg/L) | | | 1.46 x | 1.46 x 10 ⁻³ | | | | | |





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| ERC5 Marine water (in mg/L) | 3.17 x 10 ⁻⁴ | |
|------------------------------|-------------------------|--|
| ERC6B Freshwater (in mg/L) | 4.54 x 10 ⁻⁵ | |
| ERC6B Marine water (in mg/L) | 5.19 x 10 ⁻⁶ | |
| ERC7 Freshwater (in mg/L) | 1.46x 10 ⁻⁴ | |
| ERC7 Marine water (in mg/L) | 3.17 x 10 ⁻⁵ | |

2.4.4 Exposure concentration in sediments

Table 2.10 Tier 1 Predicted Environmental Concentrations (PEC) in aquatic sediment compartment

| Compartments | PEC aquatic (local) |
|--------------------------------------|---------------------|
| ERC4 Freshwater sediment (in mg/kg) | 507 |
| ERC4 Marine sediment (in mg/kg) | 50.7 |
| ERC5 Freshwater sediment (in mg/kg) | 253 |
| ERC5 Marine sediment (in mg/kg) | 25.3 |
| ERC6B Freshwater sediment (in mg/kg) | 25.3 |
| ERC6B Marine sediment (in mg/kg) | 2.53 |
| ERC6D Freshwater sediment (in mg/kg) | 0.026 |
| ERC6D Marine sediment (in mg/kg) | 0.00274 |
| ERC7 Freshwater sediment (in mg/kg) | 25.3 |
| ERC7 Marine sediment (in mg/kg) | 2.53 |

Table 2.11 Tier 2 Predicted Environmental Concentrations (PEC) in aquatic sediment compartment

| Compartments | PEC aquatic (local) |
|--------------------------------------|-------------------------|
| ERC4 Freshwater sediment (in mg/kg) | 3.05 x 10 ⁻³ |
| ERC4 Marine sediment (in mg/kg) | 6.56 x 10 ⁻⁴ |
| ERC5 Freshwater sediment (in mg/kg) | 1.58 x 10 ⁻³ |
| ERC5 Marine sediment (in mg/kg) | 3.43 x 10 ⁻⁴ |
| ERC6B Freshwater sediment (in mg/kg) | 4.91 x 10 ⁻⁵ |
| ERC6B Marine sediment (in mg/kg) | 5.62 x 10 ⁻⁶ |
| ERC7 Freshwater sediment (in mg/kg) | 1.58 x 10 ⁻⁴ |
| ERC7 Marine sediment (in mg/kg) | 3.43 x 10 ⁻⁵ |

2.4.5 Exposure concentrations in soil and groundwater

Upon contact with soil, ammonia will be rapidly converted by a variety of bacteria, actinomycetes and fungi to ammonium (NH₄⁺) by the process of ammonification or mineralization. Ammonium is then rapidly converted to nitrate. Nitrate is subsequently taken up and utilised by plants or returned to the atmosphere following denitrification; the metabolic reduction of nitrate into nitrogen or nitrous oxide (N2O) gas. The most likely fate of ammonium ions in soils is conversion to nitrates by nitrification. Therefore accumulation of concentrations of ammonia in soil and groundwater will not be expected.

Table 2.12 Tier 1 Predicted Exposure Concentration (PEC) in air

| ERC | | Local concentration | PEC air (local+regional) | Justification |
|-----|--|---------------------|--------------------------|----------------------------|
| 4 | Annual average PEC in air, total (mg/m3) | 18 | 18 | Estimated using EUSES 2.1. |
| 5 | Annual average PEC in air, total (mg/m3) | 9.45 | 9.45 | Estimated using EUSES 2.1. |
| 6B | Annual average PEC in air, total (mg/m3) | 0.0189 | 0.0189 | Estimated using EUSES 2.1. |
| 6D | Annual average PEC in air, total (mg/m3) | 6.62 | 6.62 | Estimated using EUSES 2.1. |
| 7 | Annual average PEC in air, total (mg/m3) | 0.945 | 0.945 | Estimated using EUSES 2.1. |

Table 2.13 Tier 2 Predicted Exposure Concentration (PEC) in air

| ERC | | Local concentration | PEC air (local+regional) | Justification |
|-----|--|---------------------|-----------------------------|----------------------------|
| 4 | Annual average PEC in air, total (mg/m3) | 18 | 18 | Estimated using EUSES 2.1. |
| 5 | Annual average PEC in air, total (mg/m3) | 9.45 | 9.45 | Estimated using EUSES 2.1. |
| 6B | Annual average PEC in air, total (mg/m3) | 0.0189 | 0.0189 | Estimated using EUSES 2.1. |
| 7 | Annual average PEC in air, total (mg/m3) | 0.945 | 0.945 | Estimated using EUSES 2.1. |



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2.4.6 Exposure concentration relevant for the food chain (Secondary poisoning)

In terms of secondary poisoning, there is no evidence that ammonia bioaccumulates as the log Kow value is 0.23. Since the trigger of BCF >100 (log Kow>3) is not met, the derivation of PNECs to protect against secondary poisoning is not required. Risk characterisation ratios cannot therefore be derived.

2.4.7 Regional exposure levels and environmental concentrations

Anhydrous ammonia is produced and used at many sites throughout a region and this may lead to a degree of regional exposure. Regional exposure has been modelled for this exposure scenario using the regional module of EUSES 2.1.

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Exposure Scenario No. 3: Wide dispersive end-use: Professional uses of anhydrous and aqueous ammonia

3.1 Exposure scenario

Anhydrous liquid ammonia (>99.5 % wt) and aqueous ammonia solution (5-25% wt) are used by professional workers in a broad number of applications. Common applications include: use as a laboratory chemical, a refrigerant in cooling systems, a water treatment chemical, a fertiliser, a coating, paint thinner or paint remover, a photochemical, a cleaning product, a leather or other surface treatment product, a pH regulator or neutralisation agent and a process aid for nutrition.

Typical activities associated with the professional uses of ammonia where exposures can arise include operating equipment containing ammonia (e.g. opening and closing valves), transferring ammonia from storage containers using pipe or hoses, maintaining equipment and applying ammonia-based products (e.g. fertiliser, cleaning or surface treatment products).

3.1.1 Operational conditions related to frequency and duration of use

Operational conditions pertaining to the broad range of professional end-use scenarios involving anhydrous and aqueous forms of ammonia vary considerably across applications. A full characterisation of the frequency and duration of tasks is therefore beyond the scope of this report. For the purposes of worker exposure estimation, operational conditions have been represented generically based on the assumption that tasks may be either 1-4 hours or >4 hours in duration and that processes may be carried out either outdoors, indoors without LEV or indoors with LEV.

3.1.2 Risk management measures

Activities involving the use of ammonia by professionals can be regarded as wide dispersive uses: e.g. activities which deliver uncontrolled exposures. Professional workers are expected to follow good occupational hygiene practices and apply appropriate exposure control measures to minimise the potential for exposure. Workers should be trained in procedures involving the handling, sampling and transfer of ammonia and in the use of appropriate protective equipment. A good standard of general or controlled ventilation should be applied. Personal protective clothing (e.g. face/eye protection, helmet, gloves, boots and protective overalls) should be worn when any potential contact may arise. Any professional working directly with anhydrous ammonia as required to wear eye, face and respiratory protection.

3.2 Exposure estimation

3.2.1 Workers exposure

The assessment of worker exposure to anhydrous and aqueous ammonia during professional uses was carried out for process categories relevant to this scenario as identified by PROC codes: use and storage of ammonia in closed systems with no likelihood of exposure (PROC 1), use in closed, continuous processes with occasional controlled exposure (PROC 2), formulation using closed batch processes (PROC 3), use in batch or other processes (PROC 4), mixing or blending in a batch process (PROC 5), maintenance and clean-down (PROC 8a), transfer (PROC 8b), transfer of ammonia into containers (PROC 9), brush and roller applications (PROC 10), spraying (PROC 11), treatment of articles by dipping and pouring (PROC 13), and analysis of samples (PROC 15), hand-mixing (PROC 19) and heat and pressure transfer in closed systems (PROC 20).

A screening-level (Tier 1) assessment of worker exposure was carried out using the ECETOC Targeted Risk Assessment (TRA) model. The ECETOC TRA was used to predict dermal exposures (expressed as a daily systemic dose in mg/kg bw) and inhalation exposure concentrations (expressed as an airborne concentration in mg/m³) associated with each process defined by PROC codes. Exposure to workers was assessed taking into account different operational conditions that may be associated with the professional use of ammonia and the impact of different exposure control measures. Exposures were determined for task durations of 1- 4 hours or >4 hours and assuming that process are carried out either outdoors, indoors without use of local exhaust ventilation (LEV) or indoors with the use of LEV. To reflect the use of personal protective equipment (PPE), dermal exposures were determined assuming either no gloves or gloves affording 90% protection of the hands are worn. To reflect the use of respiratory protective equipment (RPE), inhalation exposures concentrations were determined assuming either no RPE or RPE affording 95% protection is worn.

The ECETOC TRA model uses a simple algorithm to determine dermal exposures that does not take the physical-chemical properties of a substance into account. The same dermal exposures where therefore predicted for anhydrous and aqueous forms of ammonia. Parameters used in the ECETOC TRA model to assess inhalation exposures were: molecular weight (35 g.mol⁻¹ and 17 g.mol⁻¹ for aqueous and anhydrous forms respectively and vapour pressure (the vapour pressure of anhydrous forms of ammonia is 8.6 x 10⁵ Pa at 20°C, whereas the vapour pressure of aqueous ammonia solution between 5 and 25% w/w ranges from 5 x 10³ Pa to 4 x10⁴ Pa at 20°C. Systemic dermal exposures have been determined for a worker with bodyweight 70 kg.

3.2.2 Acute/Short Term and Long Term Exposure

Potential systemic dermal exposures and inhalation exposure concentrations predicted by the ECETOC TRA model for processes associated with the professional use of ammonia are shown in Tables 3.1 and 3.2 respectively. ECETOC predicts a daily systemic dose following dermal exposure and a typical daily inhalation exposure concentration and does not specifically predict acute (short-term) and chronic (long-term) exposures.

Table 3.1 Dermal exposures to anhydrous or aqueous (in preparations of 5-25 % w/w) ammonia predicted using the ECETOC TRA model for professional workers

| Description of activity | PROC | Exposure assun | nptions | Estimated Exposure Concentrati mg/kg bw/d | | |
|---|--------|---|-------------------------------|---|-----------------------------|--|
| | | Duration Use of Ventilation | | No gloves worn | Gloves worn (90% reduction) | |
| Used in a closed process, no likelihood of exposure | PROC 1 | 1-4 hrs or >4 hrs | Outdoors /Indoors without LEV | 0.34 | 0.03 | |
| Use of ammonia in a closed, continuous | PROC 2 | 1-4 hrs or >4 hrs Outdoors /Indoors without LEV | | 1.37 | 0.14 | |

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| process with occasional controlled exposure (e.g. sampling) | | 1-4 hrs or >4 hrs | Indoors with LEV | | 0.01 |
|---|------------|----------------------|--------------------------------|--------|-------|
| Use of ammonia in closed batch process (synthesis | PROC 3 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 0.34 | 0.03 |
| or formulation) | 1 100 3 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.03 | <0.01 |
| Use of ammonia in batch process (synthesis) where | PROC 4 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 6.86 | 0.69 |
| opportunity for exposure arises | PROC 4 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.69 | 0.07 |
| Mixing or blending in | PROC 5 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 13.71 | 1.37 |
| batch process | PROCS | 1-4 hrs or >4 hrs | Indoors with LEV | 0.07 | 0.01 |
| Transfer of ammonia (charging/discharging) | PROC | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 13.71 | 1.37 |
| from/to vessels or large containers at non-dedicated facilities | 8a | 1-4 hrs or >4 hrs | Indoors with LEV | 0.14 | 0.01 |
| Transfer of ammonia (charging/discharging) | PROC | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 6.86 | 0.69 |
| from/to vessels or large containers at dedicated facilities | 8b | 1-4 hrs or >4 hrs | Indoors with LEV | 0.69 | 0.07 |
| Transfer of ammonia into | PROC 9 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 6.86 | 0.69 |
| small containers | | 1-4 hrs or >4 hrs | Indoors with LEV | 0.69 | 0.07 |
| Roller application or | PROC 10 | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 27.43 | 0.14 |
| brushing | | 1-4 hrs or >4 hrs | Indoors with LEV | 1.37 | 10.71 |
| Non industrial spraying | PROC | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 107 | 10.71 |
| Non industrial spraying | 11 | 1-4 hrs or >4 hrs | Indoors with LEV | 2.14 | 0.21 |
| Treatment of articles by | PROC | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 13.71 | 1.37 |
| dipping and pouring | 13 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.69 | 0.07 |
| Use as a laboratory agent | PROC | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 0.34 | 0.03 |
| | 15 | 1-4 hrs or >4 hrs | Indoors with LEV | 0.03 | <0.01 |
| Hand-mixing with intimate contact and PPE only | PROC 19 | 1-4 hrs or >4 hrs | Indoors with LEV | 141.73 | 14.14 |
| Heat and pressure | PROC | 1-4 hrs or >4 hrs | Outdoors / Indoors without LEV | 1.71 | 0.17 |
| transfer fluids in dispersive use but closed | FROC | | | | |

Table 3.2 Inhalation exposure concentrations for anhydrous and aqueous ammonia (in preparations of 5-25 % w/w) predicted using the ECETOC TRA model for professional workers

| Description of activity | PROC | Evposuro o | ssumptions | Anhydro | ous ammonia | 25% w/w) Estimated Exposure Concentration mg/m3 | |
|--|--------|---------------------------------------|--------------------|-----------|-----------------------------|---|---------------------|
| Description of activity | PROC | Exposure a | ssumptions | | ed Exposure ration mg/m3 | | |
| | | Duration | Use of ventilation | No RPE | RPE (95% reduction) | No RPE | RPE (95% reduction) |
| Used in a closed process, | PROC 1 | 1-4 hrs or >4 hrs Outdoors | | 0.00 | NA | 0.01 | 0.00 |
| no likelihood of exposure | | 1-4 hrs or >4 hrs Indoors without LEV | | 0.01 | NA | 0.01 | 0.00 |
| Use of ammonia in a | PROC 2 | >4hrs Outdoors | | 24.79 | 1.24 | 30.63 | 1.53 |
| closed, continuous process with occasional | | >4hrs Indoors without LEV | | 35.42 | 1.77 | 43.75 | 2.19 |
| controlled exposure (e.g. | | >4hrs Indoors with LEV | | 3.53 | 0.18 | 4.38 | 0.22 |





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|---|------------|--------------------|------------------------------|-----------------|--------------|-----------------|-------|
| sampling) | | 1-4 hrs | Outdoors | 14.88 | 0.74 | 18.38 | 0.92 |
| | | 1-4 hrs | Indoors without LEV | 22.25 | 1.06 | 26.25 | 1.31 |
| | | 1-4 hrs | Indoors with LEV | 2.13 | 0.11 | 2.63 | 0.13 |
| | | >4hrs | Outdoors | 49.58 | 2.48 | 61.25 | 3.06 |
| | | >4hrs | Indoors without LEV | 70.83 | 3.54 | 87.5 | 4.38 |
| Use of ammonia in closed | | >4hrs | Indoors with LEV | 7.08 | 0.35 | 8.75 | 0.44 |
| batch process (synthesis or formulation) | PROC 3 | 1-4 hrs | Outdoors | 29.75 | 1.49 | 36.75 | 1.84 |
| or formalation) | | 1-4 hrs | Indoors without LEV | 42.5 | 2.13 | 52.50 | 2.63 |
| | | 1-4 hrs | Indoors with LEV | 4.25 | 0.21 | 5.25 | 0.26 |
| | | >4hrs | Outdoors | 49.58 | 2.48 | 61.25 | 3.06 |
| | | >4hrs | Indoors without LEV | 70.83 | 3.54 | 87.5 | 4.38 |
| Use of ammonia in batch process (synthesis) where | | >4hrs | Indoors with LEV | 7.08 | 0.35 | 8.75 | 0.44 |
| opportunity for exposure | PROC 4 | 1-4 hrs | Outdoors | 29.75 | 1.49 | 36.75 | 1.84 |
| arises | | 1-4 hrs | Indoors without LEV | 42.5 | 2.13 | 52.5 | 2.63 |
| | | 1-4 hrs | Indoors with LEV | 4.25 | 0.21 | 5.25 | 0.26 |
| | | >4hrs | Outdoors | 123.96 | 6.20 | 153.13 | 7.66 |
| | | >4hrs | Indoors without LEV | 177.08 | 8.85 | 218.75 | 10.94 |
| Mixing or blending in | | >4hrs | Indoors with LEV | 17.71 | 0.89 | 21.88 | 1.09 |
| batch process | PROC 5 | 1-4 hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| • | | 1-4 hrs | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| | | 1-4 hrs | Indoors with LEV | 10.63 | 0.53 | 13.13 | 0.66 |
| | | >4hrs | Outdoors | 123.96 | 6.20 | 153.13 | 7.66 |
| Transfer of ammonia | PROC 8a | >4hrs | Indoors without LEV | 177.08 | 8.85 | 218.75 | 10.94 |
| (charging/discharging) | | >4hrs | Indoors with LEV | 17.71 | 0.89 | 21.88 | 1.09 |
| from/to vessels or large | | 1-4 hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| containers at non- dedicated facilities | | 1-4 hrs | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| | | 1-4 hrs | Indoors with LEV | 10.63 | 0.53 | 13.13 | 0.66 |
| | | >4hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| Transfer of ammonia | | >4hrs | Indoors without LEV | 106.25 | 5.72 | 131.25 | 6.56 |
| (charging/discharging) | DDOC | >4hrs | Indoors with LEV | 3.19 | 0.16 | 3.94 | 0.20 |
| from/to vessels or large | PROC 8b | 1-4 hrs | Outdoors | 44.63 | 2.23 | 55.13 | 2.76 |
| containers at dedicated facilities | | 1-4 hrs | Indoors without LEV | 63.75 | 3.19 | 78.75 | 3.94 |
| | | 1-4 hrs | Indoors with LEV | 1.91 | 0.1 | 2.36 | 0.12 |
| | | >4hrs | Outdoors | 99.17 | 4.96 | 122.50 | 6.13 |
| | | | Indoors without LEV | | | | 8.75 |
| T | | >4hrs >4hrs | Indoors with LEV | 141.67 14.17 | 7.08 0.71 | 175.00 17.50 | 0.88 |
| Transfer of ammonia into small containers | PROC 9 | | | + | 2.98 | | 3.68 |
| San contamore | | 1-4 hrs 1-4 hrs | Outdoors Indoors without LEV | 59.50 85.00 | 4.25 | 73.50 105.00 | 5.25 |
| | | | | | | | |
| | | 1-4 hrs | Indoors with LEV | 8.5 | 0.43 | 10.50 | 0.53 |
| | | >4hrs | Outdoors | NA | NA | 153.13 | 7.66 |
| | | >4hrs | Indoors without LEV | NA | NA | 218.75 | 10.94 |
| Roller application or | PROC | >4hrs | Indoors with LEV | NA | NA | 21.88 | 1.09 |
| brushing | 10 | 1-4 hrs | Outdoors | NA | NA | 91.88 | 4.59 |
| | | 1-4 hrs | Indoors without LEV | NA | NA | 131.25 | 6.56 |
| | | >4hrs | Outdoors | NA | NA | 13.13 | 0.66 |
| | PROC | >4hrs | Outdoors | NA | NA | 613.20 | 30.66 |
| Non-industrial spraying | 11 | >4hrs | Indoors without LEV | NA | NA | 876.00 | 43.80 |
| | | >4hrs | Indoors with LEV | NA | NA | 175.20 | 8.76 |



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| | | 1-4 hrs | Outdoors | NA | NA | 367.92 | 18.40 |
|--------------------------------------|------------|---------|---------------------|--------|------|--------|-------|
| | | 1-4 hrs | Indoors without LEV | NA | NA | 525.60 | 26.28 |
| | | >4hrs | Outdoors | NA | NA | 105.12 | 5.26 |
| | | >4hrs | Outdoors | 123.96 | 6.20 | 153.13 | 7.66 |
| | | >4hrs | Indoors without LEV | 177.08 | 8.85 | 218.75 | 10.94 |
| Treatment of articles by | PROC | >4hrs | Indoors with LEV | 17.71 | 0.89 | 21.88 | 1.09 |
| dipping and pouring | 13 | 1-4 hrs | Outdoors | 74.38 | 3.72 | 91.88 | 4.59 |
| | | 1-4 hrs | Indoors without LEV | 106.25 | 5.31 | 131.25 | 6.56 |
| | | 1-4 hrs | Indoors with LEV | 10.63 | 0.53 | 13.13 | 0.66 |
| | | >4hrs | Indoors without LEV | 35.42 | 1.77 | 43.75 | 2.19 |
| Llas as a laboratory agent | PROC 15 | >4hrs | Indoors with LEV | 3.54 | 0.18 | 4.38 | 0.22 |
| Use as a laboratory agent | | 1-4 hrs | Indoors without LEV | 21.25 | 1.06 | 26.25 | 1.31 |
| | | 1-4 hrs | Indoors with LEV | 2.13 | 0.11 | 2.63 | 0.13 |
| | | <4 hrs | Outdoors | NA | NA | 153.13 | 7.66 |
| Hand-mixing with intimate | PROC 19 | <4 hrs | Indoors without LEV | NA | NA | 218.75 | 10.94 |
| contact and PPE only | | 1-4 hrs | Outdoors | NA | NA | 91.88 | 4.59 |
| | | 1-4 hrs | Indoors without LEV | NA | NA | 131.25 | 6.56 |
| | | >4hrs | Outdoors | 24.79 | 1.24 | 30.63 | 1.53 |
| Heat and pressure transfer fluids in | PROC | >4hrs | Indoors without LEV | 35.42 | 1.77 | 43.75 | 2.19 |
| | | >4hrs | Indoors with LEV | 7.08 | 0.35 | 8.75 | 0.44 |
| dispersive use but closed | 20 | 1-4 hrs | Outdoors | 14.88 | 0.74 | 18.38 | 0.92 |
| systems | | 1-4 hrs | Indoors without LEV | 21.25 | 1.06 | 26.25 | 1.31 |
| | | 1-4 hrs | Indoors with LEV | 4.25 | 0.21 | 5.25 | 0.26 |

3.3 General public / Consumer exposure

Professional workers are expected to conduct risk assessment to ensure that members of the public are excluded from operational activities and are not inadvertently exposed to ammonia.

3.3.1 Indirect exposure of humans via the environment (oral)

Ammonia is ubiquitous in the environment with <30% of emissions resulting from fertiliser uses and from non-agricultural sources (ref. 'Ammonia in the UK' - DEFRA).

In addition, there is no evidence that ammonia bioaccumulates as the log Kow value is 0.23. Since the trigger of BCF >100 (log Kow>3) is not met, the derivation of PNECs to protect against secondary poisoning is not required.

The risk of indirect exposure of humans via the environment is therefore not considered.

3.4 Environmental exposure

The majority of ammonia in the environment originates from natural sources, predominantly decaying organic matter.

Wide dispersive professional uses of ammonia are diverse and widespread. The resulting environmental exposure is not expected to add significantly to already present background levels of ammonia in the environment. An additional assessment for environmental exposure for wide dispersive uses has therefore not been performed.





(according to Regulation (EC) No 1907/2006 (REACH), ANNEX II)

AMMONIA

Revision date: 01.10.2024 Version 4.4



Exposure Scenario No. 4: Wide-dispersive end-use: Consumer use of aqueous ammonia

4.1 Exposure scenario

Consumers may be exposed to aqueous solutions of ammonia (containing up to 25 % w/w ammonia) when using a variety of products. In this section, exposures have been assessed for the consumer use of representative common products for which default data and scenarios are available (e.g. in the ECETOC TRA model or the ConsExpo 4.1 model and RIVM Factsheets). Although data are not available for other uses (e.g. inks and toners, water treatment chemicals etc.) exposures arising from these uses is not expected to be any worse than for the representative products and is therefore considered to be addressed by this assessment.

Consumers may be exposed to aqueous ammonia when using a variety of common household products including Do-It-Yourself (DIY) products such as coatings, paints, thinners and removers (PC9a) and fillers, putties and plasters (PC 9b), washing and cleaning products (e.g. all-purpose liquid; PC 35), cosmetic, personal care products such as hair dyes (PC 39) and fertilisers (PC12). Aqueous ammonia (at concentrations up to 25 % w/w) is added to water-based paints and other DIY products as a stabiliser. The composition of these products typically contains 0.2 % ammonia solution (at 25 % w/w ammonia) giving a final concentration in the product of 0.05 % w/w ammonia. Cleaning products are typically aqueous solutions containing 5 -10 % ammonia and will be diluted with water prior to use. Cosmetic products such a hair dyes contain ammonia at a maximum concentration of 4 % w/w.

Primary routes of exposure for consumers using common household products containing ammonia are the dermal and inhalation routes. Consumers are not expected to ingest ammonia during the normal use of household products and oral exposures are unlikely.

Consumer exposure to ammonia will depend on a number of factors including the frequency and duration of use. DIY products are likely to be used intermittently during the year, for several consecutive days. Cleaning products are expected to be used more frequently e.g. several times per week. Consumers are likely to use hair dye products several times per year, possibly up to once/month. The durations of product use per day are likely to vary across the applications.

4.2 Exposure estimation

Consumer exposure to aqueous ammonia associated with the use of common household products (e.g. DIY, cleaning and cosmetic products) was assessed using ConsExpo version 4.1 and default assumptions presented in RIVM Factsheets relevant to the scenario being assessed. Dermal exposures were assessed using either the constant rate, dermal contact with product model or the instant application model, as appropriate. Inhalation exposure concentrations were assed using either the evaporation model or the spraying model as appropriate. To assess possible worst case scenarios, it was assumed that consumers will use DIY product at least once per month. Consumer exposure from the use of cleaning products has been assessed using the default scenario in ConsExpo 4.1 for the application of an all-purpose liquid cleaner: e.g. Household ammonia (10 % w/w ammonia) is diluted 1:80 times with water to give a final concentration of 0.125 % w/w. It is assumed that consumers will use cleaning products daily. In a worst case scenario, consumers are assumed to use hair dye once per month.

Consumer exposure arising from the use of fertilisers (containing up to 25% w/w ammonia) was assessed using the ECETOC TRA model and default parameters for the scenario PC12 fertilisers: lawn and garden preparations. It is assumed that amateur gardeners will apply fertilisers twice per year.

It is reasonable to assume that consumers will not always read product labels or follow advice recommended by the manufacturer. In a worst-case assessment of consumer exposure, no use of gloves or other PPE has been assumed.

Table 4.1 shows the dermal exposures predicted by ConsExpo for consumer uses of common household products containing aqueous ammonia. Dermal exposures are presented as: acute systemic dermal exposures reflecting the total exposure during one event and as chronic systemic exposures reflecting the exposure per event averaged over a year taking into account the use frequency. In a conservative assessment of dermal exposures, it was assumed that 100 % of the dose is absorbed dermally.

Table 4.1 also shows the inhalation exposure concentrations predicted by ConsExpo for consumer uses of common household products containing aqueous ammonia. Inhalation exposure concentrations are presented as acute airborne concentrations of ammonia associated with one event and as chronic exposure concentrations reflecting the concentration per event averaged over a year taking into account the use frequency.

Table 4.1 Dermal exposures to aqueous ammonia and inhalation exposure concentrations predicted by ConsExpo for consumers using common household products (e.g. DIY, cleaning and cosmetic products)

| Scenario | Ammonia % w/w | Use frequency | e ut | c zi o ţ C | e ut | : e n 5 at a n ii c n o h C |
|--|------------------|-------------------|----------------------|-----------------------|------|-----------------------------|
| PC9 Coatings, paints, thinne | rs, removers (| 0.05% w/w am | nmonia) | | | |
| Applying waterborne paint using brush and roller | 0.05 | 1 event /month | 0.03 | 8.2x 10 ⁻⁵ | 7 | 0.0018 |
| Spraying paint from a can (application) | 0.05 | 1 event /month | 0.013 | 6.8x 10 ⁻⁵ | 0.67 | 5.1x10 ⁻⁵ |
| Applying general coatings | 0.05 | 1 event /month | 0.0021 | 1.9x10 ⁻⁶ | 6.7 | 2.4x10 ⁻⁴ |
| Applying paint remover | 0.05 | 1 event /month | 0.0042 | 1.1x10 ⁻⁵ | 3.2 | 3.6x10 ⁻⁴ |
| PC9b Fillers, putties, plasters | s etc (0.05 % v | //w ammonia) | | | | |
| Applying filler | 0.05 | 1 event /month | 4.2x10 ⁻⁴ | 3.4x10 ⁻⁶ | 0.37 | 5.1x10 ⁻³ |
| PC35 Washing and cleaning | products (0.12 | 25 % w/w amn | nonia) | | | |
| Applying all-purpose liquid | 0.125 | 104 times/ | 0.41 | 0.12 | 3.3 | 0.16 |



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SAFETY DATA SHEET

(according to Regulation (EC) No 1907/2006 (REACH), ANNEX II)

AMMONIA

Revision date: 01.10.2024 Version 4.4

AZOT

| cleaner / detergent | | year | | | | | |
|---------------------------|----------------|-------------------|---------|-------|----|----|--|
| PC39 Cosmetics, person | nal care produ | cts (4% w/w amm | onia) | | | | |
| | | 1 event / | | | | | |
| Applying hair dye | 4 | month | 67 | 2.203 | NA | NA | |
| PC12 Fertilisers: lawn ar | nd garden pre | parations (25 % w | /w ammo | onia) | | | |
| | | 2 | | | | | |
| Applying fertilisers | 25 | events/year | 35.7 | | NA | NA | |

4.3 Environmental exposure

The majority of ammonia in the environment originates from natural sources, predominantly decaying organic matter. Wide dispersive consumer uses of ammonia are diverse and widespread. The resulting environmental exposure is not expected to add significantly to already present background levels of ammonia in the environment. An additional assessment for environmental exposure for wide dispersive uses has therefore not been performed.