Directive 98/8/EC concerning the placing biocidal products on the market

Inclusion of active substances in Annex I or I A to Directive 98/8/EC

Assessment Report



Hydrogen cyanide Product-type 08 (wood preservatives)

25 May 2012

Annex I – the Czech Republic

| I | SIA | TEMENT OF SUBJECT MATTER AND PURPOSE | 4 |
|---|-------|---|----|
| | 1.1 | Procedure Followed | 4 |
| | 1.2 | Purpose of the assesment report | 6 |
| | 1.3 | Overall conclusion in the context of Directive 98/8/EC | 6 |
| 2 | OVE | ERALL SUMMARY AND CONCLUSIONS | 7 |
| | 2.1 | Presentation of the Active Substance | 7 |
| | 2.1.1 | Identity, Physico-Chemical Properties & Methods of Analyssis | 7 |
| | 2.1.2 | Intended Uses and Effficacy | 8 |
| | 2.1.3 | Classification and labelling | 8 |
| | 2.2 | Summary of the Risk Assessment | 10 |
| | 2.2.1 | Human Health Risk Assessment | 10 |
| | 2.3 | 2.1.1 Hazard Identification | 11 |
| | 2.3 | 2.1.2 Exposure assessment and risk characterisation | 13 |
| | 2.2.2 | Physical-chemical hazard | 16 |
| | 2.3 | 2.2.1 Risk characterisation for the physico-chemical properties | 17 |
| | 2.2.3 | Environmental risk assessment. | 17 |
| | 2 | 2.3.1 Fate and distribution in the environment | 17 |
| | 2 | 2.3.2 Effects assessment | 19 |

| Hydrogen cyanide | Product –type 08 | 25 May 2012 |
|--------------------------|---|---------------------------|
| 2.2.3.3 | PBT assessment | 22 |
| 2.2.3.4 | Risk characterization | 22 |
| 2.2.4 List | of endpoints | 26 |
| 3 DECISION. | | 26 |
| 3.1 Backgr | round to the proposed decision | 26 |
| 3.2 Propos | ed decision regarding the inclusion in Annex IErr | or! Bookmark not defined. |
| 3.3 Elemen Bookmark not | nts to be taken into account by Member States when authorise defined. | sing products Error! |
| 3.4 Require | ement for further informationErr | or! Bookmark not defined. |
| 3.5 Updati | ng this Assessment Report Err | or! Bookmark not defined. |
| Appendix I: LIST | OF ENDPOINTS | 29 |
| - | lentity, Physical and Chemical Properties, Details of Uses, I | |
| Chapter 2: M | 1ethods of Analysis | 38 |
| Chapter 3: Impa | act on human health | 41 |
| Chapter 4 :Fate | and Behaviour in the Environment | 49 |
| 3.6 Chapte | r 5: Effects on Non-target Species | 53 |
| Appendix II: LIST | Γ OF INTENDED USES | 55 |
| Annendix III: LIS | T OF STUDIES | 56 |

1 STATEMENT OF SUBJECT MATTER AND PURPOSE

1.1 Procedure Followed

This assessment report has been established as a result of the evaluation of Hydrogen Cyanide as product-type 08 (wood preservatives), carried out in the context of the work programme for the review of existing active substances provided for in Article 16(2) of Directive 98/8/EC concerning the placing of biocidal products on the market, with a view to the possible inclusion of this substance into Annex I or IA to the Directive.

Hydrogen Cyanide (CAS no. 74-90-8) was notified as an existing active substance, by Lučební závody Draslovka a.s. Kolín, hereafter referred to as the applicant, in product-type 08. Commission Regulation (EC) No 1451/2007 of 4 December 2007 lays down the detailed rules for the evaluation of dossiers and for the decision-making process in order to include or not an existing active substance into Annex I or IA to the Directive. In accordance with the provisions of Article 7(1) of that Regulation, the Czech Republic was designated as Rapporteur Member State to carry out the assessment on the basis of the dossier submitted by the applicant. The deadline for submission of a complete dossier for Hydrogen Cyanide as an active substance in Product Type 08 was 1 March 2006, in accordance with Annex V of Regulation (EC) No 2032/2003.

In accordance with provision of Article 4a of Regulation (EC) No. 2032/2003 as amended by Regulation (EC) No. 1048/2005 the Czech Republic applied for essential use of the active substance Hydrogen Cyanide on 18.11.2005.

On 16.2.2006, the competent authority of the Czech Republic received a dossier from the applicant. The Rapporteur Member State accepted the dossier as complete for the purpose of the evaluation on 28.2.2006.

On 24.1.2008, the Rapporteur Member State submitted, in accordance with the provisions of Article 14(4) and (6) of Regulation (EC) No 1451/2007, to the Commission and the applicant a copy of the evaluation report, hereafter referred to as the competent authority report. The Commission made the report available to all Member States by electronic means on 19.2.2008. The competent authority report

included a recommendation for the inclusion of Hydrogen Cyanide in Annex I to the Directive for product-type 08.

In accordance with Article 16 of Regulation (EC) No 1451/2007, the Commission made the competent authority report publicly available by electronic means on 25.2.2008. This report did not include any information that was to be treated as confidential in accordance with Article 19 of Directive 98/8/EC. In order to review the competent authority report and the comments received on it, consultations of technical experts from all Member States (peer review) were organised by the Commission. Revisions agreed upon were presented at technical and competent authority meetings and the competent authority report was amended accordingly.

On the basis of the final competent authority report, the Commission proposed the inclusion of Hydrogen Cyanide in Annex I to Directive 98/8/EC and consulted the Standing Committee on Biocidal Product on 25 May 2012.

In accordance with Article 15(4) of Regulation (EC) No 1451/2007, the present assessment report contains the conclusions of the Standing Committee on Biocidal Products, as finalised during its meeting held on 25 May 2012.

1.2 Purpose of the assesment report

This assessment report has been developed and finalised in support of the decision to include Hydrogen Cyanide in Annex I to Directive 98/8/EC for product-type 08. The aim of the assessment report is to facilitate the authorisation in Member States of individual biocidal products in product-type 08 that contain Hydrogen Cyanide. In their evaluation, Member States shall apply the provisions of Directive 98/8/EC, in particular the provisions of Article 5 as well as the common principles laid down in Annex VI.

For the implementation of the common principles of Annex VI, the content and conclusions of this assessment report, which is available at the Commission website, shall be taken into account.

However, where conclusions of this assessment report are based on data protected under the provisions of Directive 98/8/EC, such conclusions may not be used to the benefit of another applicant, unless access to these data has been granted.

1.3 Overall conclusion in the context of Directive 98/8/EC

It can be concluded from the evaluation that the proposed use of biocidal products based on hydrogen cyanide under the specified conditions fulfil the safety requirements laid down in Article 5 of Directive 98/8/EC. This conclusion is, thus, subject to

- i. compliance with the particular requirements in the following sections of this assessment report,
- ii. the implementation of the provisions of Article 5(1) of Directive 98/8/EC, and
- iii. the common principles laid down in Annex VI to Directive 98/8/EC.

Furthermore, these conclusions were reached within the framework of the uses that were proposed and supported by the applicant (see Appendix II). Extension of the use pattern betone those described will require an evaluation at product authorisation level in order to establish whether the proposed extensions of use will satisfy the requirements of Article 5(1) and of the common principles laid down in Annex VI to Directive 98/8/EC.

2 OVERALL SUMMARY AND CONCLUSIONS

2.1 Presentation of the Active Substance

2.1.1 Identity, Physico-Chemical Properties & Methods of Analyssis

| CAS number | 74-90-8 |
|--|---|
| Einecs number | 200-821-6 |
| Other No. | CIPAC NO. 126 |
| Chemical name, synonyms | Hydrogen cyanide, Hydrocyanic acid (water solution) |
| Molecular formula | HCN |
| Structural formula | H-C≡N |
| Molecular mass (g/mol) | 27.03 |
| Purity of the active substance as manufactured | Min. 97.6 % wt |
| Impurities | Water (1.18 -1.42 % wt) |
| Additives | Phosphoric acid (0.08-0.12 % wt), |
| | sulphur dioxide (0.9 – 1.1 % wt) |

Hydrogen cyanide is colourless liquid between -13.4 and +25.7°C (acid), and colourless gas with almond-like odour for higher temperatures. It is miscible with water and soluble in ethanol and ether. Octanol/water partition coefficient of 5 (log Kow = 0.66) indicates slight preference of the hydrophobic compartments. High values of vapour pressure (84 kPa at 20°C, 35 kPa at 0°C) and of Henry's law constant signalize rapid evaporation and rapid leakage from water solution. Specific density of vapours is

slightly below 1 (0.937 at 31°C) supports the assumption of an even distribution. The vapours are flammable and explosive in the range of concentrations in air of 5.6 to 40 v/v%.

The representative biocidal product named Uragan D2 (stabilized liquid hydrogen cyanide) is mixture of approx. 98 % of hydrogen cyanide (CAS No 74-90-8) with stabilizing additives. Uragan D2 is supplied completely soaked into a porous material in 1.5 kg gas-tight cans made of 0.45 mm steel. During fumigation it evaporates and brings about its effect as a gas.

Methods for analysis of the active substance as manufactured as well as methods for the determination of the additives and impurities have been described in sufficient detail. Methods for residue determinations in soil, water, air and blood have been validated and shown to be sufficiently specific, accurate, sensitive and to provide for appropriate LOQ with respect the toxicological and environmental endpoints of hydrogen cyanide.

Summary information on the identity and physico-chemical properties and analytical methods can be found in Appendix I to this document (List of Endpoints).

2.1.2 Intended Uses and Effficacy

Hydrogen cyanide is used as fumigant for professional use only to control wood destroying pests (insects, arthropods, nematodes, PT 08 –wood preservative) in empty storehouses, depositories, transport facilities, containers, libraries, other buildings or without any materials which are able to absorb hydrogen cyanide and which can not be made strict gastight.

Target organisms are all stages of wood destroying or disfiguring pests.

Universal efficacy against pests follows from the well-known mechanism of toxic action. This is confirmed by long term experience and data provided in support of the efficacy.

2.1.3 Classification and labelling

Proposal of the classification and labelling of the active substance

| Classification and labelling in compliance with Annex VI Regulation (EC) |
|--|
| No. 1272/2008 |

| Hazard classification | Flam. Liq. 1; |
|-----------------------|--|
| and Category Code(s) | Acute Tox.1; |
| | Aquatic Acute 1; Aquatic Chronic 1 |
| Hazard statement | H224; |
| Code(s) | H330; |
| | H400; H410 |
| Labelling | |
| Pictogram and | |
| Signal word Code(s) | Danger |
| Hazard statement | H224: Extremely flammable liquid and vapour |
| Code(s) | H330: Fatal if inhaled |
| | H410: Very toxic to aquatic life with long lasting effects |

| Precautionary statement | P210 Keep away from heat/sparks/open flames/hot surfaces. — No | |
|-------------------------|---|--|
| Code(s) | smoking. | |
| | P260 Do not breathe dust/fume/gas/mist/vapours/spray. | |
| | P262 Do not get in eyes, on skin, or on clothing. | |
| | P280/284 Wear protective gloves/protective clothing/eye protection/face | |
| | protection/respiratory protection. | |
| | P303+P361+P353 IF ON SKIN (or hair): Remove/Take off immediately | |
| | all contaminated clothing. Rinse skin with water/shower. | |
| | P304+P340 IF INHALED: Remove victim to fresh air and keep at rest in | |
| | a position comfortable for breathing. | |
| | P310 Immediately call a POISON CENTER or doctor/physician. | |
| | P273 Avoid release to the environment. | |

Proposal for classification of biocidal product Uragan D 2 is the same as that for the active substance.

2.2 Summary of the Risk Assessment

2.2.1 Human Health Risk Assessment

Human health risk assessment is based on data submitted by the applicant. Toxicology of hydrogen cyanide and generally of various sources of cyanide ion has long tradition: rich material has been accumulated on all relevant effects, and repeatedly analysed and discussed in peer-reviewed surveys. No new studies were therefore planned and performed by the applicant.

2.2.1.1 Hazard Identification

Dangerous properties as well as sub-cellular mechanisms of cyanide ion toxicity are thoroughly explored. Common mechanism of toxicity ,i.e. the toxic agent common to the below surrogates is CN⁻, and known toxicokinetics, e.g. slow releases of CN obviate occurrence of acutely cyanide dangerous peaks, justifies the use of toxicological data on inorganic cyanides and nitriles (aceton cyanhydrin, acetonitrile) as surrogates for missing or unreliable components of the toxicological profile of hydrogen cyanide. In addition to ample epidemiological and clinical evidence, literature provides a large quantity of experimental data; on the other hand most experimental studies collected did not meet requirements for a key study. The necessary validity and reliability is ensured by cross-comparison of results of many studies widely differing in the source of cyanide, routes of administration, endpoints, methods, species and interpretation approaches.

Toxicokinetics

Hydrogen cyanide is readily absorbed from orally administered water solutions or from fumigated food and oral absorption is 100 %. For respiratory route 100 % pulmonary retention is assumed. The rate of absorption of gaseous HCN by dry skin is by more than two orders of magnitude lower than absorption by inhalation.

Cyanides are readily distributed within the body by blood and up to 80 % of absorbed dose is metabolised to thiocyanate at a rate of 1 μ g/kg body weight per minute. At absorption rate exceeding 1.2 μ g/kg bw per minute the blood concentration of CN is expected to grow with duration of acute exposure in most subjects. Low affinity of HCN to lipids and relative rate of its metabolic transformation to thiocyanates indicate that cyanides do not accumulate in the organism.

Acute toxicity

Hydrogen cyanide is highly toxic on inhalation, its inhalation LC 50 ranging from 3778 mg/m³ for exposure time of 10 seconds to 158 mg/m³ for a 60 minute exposure. It is classified as very toxic (T+) with risk phrase R 26 (very toxic by inhalation) (CLP: Acute tox. 1; H330).

Due to low dermal uptake of gaseous hydrogen cyanide the acute toxicity via this route is low and no corresponding classification is required.

Hydrogen cyanide toxicity is due to the impairment of the tissue utilization of oxygen making the cells critically dependent on oxidative metabolism most vulnerable. Hence the effects on nervous and cardio vascular systems are the most critical ones.

None of the human or animal data meet requirements for labelling of hydrogen cyanide as a skin irritating substance, and hydrogen cyanide is not classified as irritant for eyes. Human data on respiratory irritation are mostly negative and do not justify classification either. Hydrogen cyanide does not present any structural alert for skin sensitization and sensitization properties of cyanides or nitriles have not been suggested by the experience in humans over a period of many years of production and use.

Repeated toxicity

The toxic effects found in studies using repeated oral dosing of cyanides are interpreted as being due to cumulated injury from repeated acute poisonings resulting from acutely dangerous peaks of readily absorbed cyanides. Such peaks and hence the acute effects avoided, the inhibition of thyroid function is the only critical long term effect. This effect is ascribed to goitrogenic potency of thiocyanate, the main metabolite of cyanides. The NOAEL for this effect from which long term AEL was derived is 10 mg/kg.bw per day. This NOAEL primarily draws on two chronic (2 year) studies, inhalatory (acetonitrile in rats and mice) and oral (HCN in diet, rats), NOAEL in both being \geq 10 mg CN/kg bw per day (top dose). This is further supported by several studies reporting daily doses of 4.7 to 26 mg cyanide/kg.bw (top doses used) being without effect in 13-week to 26 week studies.

Genotoxicity

Genotoxicity was observed only in cells with seriously lowered viability. HCN has been shown to posses no intrinsic genotoxic potential. This is based on negative outcome of various mutagenicity studies on bacteria, a relevant in vitro mutagenicity on mamalian cells, in vivo bone marrow chromosomal aberrations test in rats and test of inhibition of mouse testicular DNA synthesis.

Carcinogenicity

Carcinogenicity was explored in combined chronic toxicity – carcinogenicity study of acetonitrile in rats and mice and an extensive two-year inhalation studies with acetonitrile in rats and mice. Based on the data from these studies no carcinogenicity is expected at doses substantially below acutely toxic level. This is further confirmed by epidemiological studies in workers exposed for many years to hydrogen cyanide in concentrations exceeding 10 mg/m³ where no data leading to suspicion of hydrogen cyanide carcinogenicity were reported.

Reproductive toxicity

Various studies on reproductive toxicity were evaluated. These include teratology study with aceton cyanohydrin with rats, 13 week study via oral route (NaCN in drinking water) including reproduction toxicity in rats and mice, in vivo DNA synthesis inhibition in mouse, 10 week male fertility study (inhalation route to acetone cyanohydrin) in rats, female fertility study (inhalation route to acetone cyanohydrin) in rats. The NOAELs for reproductive toxicity end points range from 1 to 26 mg CN/kg bw, all the values being the top, or single, doses. All experimental studies permitting precise estimates of cyanide doses administered concur in a conclusion that decreased fertility, teratogenity, embryotoxicity or developmental toxicity is limited to doses severely toxic for the adults. This is further confirmed by epidemiological studies in workers exposed for many years to hydrogen cyanide in concentrations exceeding 10 mg/m³ where no data leading to suspicion of hydrogen cyanide being toxic for reproduction. Hence, NOAEL of 10 mg/kg bw determined for repeated toxicity covers also reproductive toxicity endpoints.

Neurotoxicity

The central nervous system is the primary target of acute cyanide toxicity due to its mechanism of toxic action which impairs the tissue utilization of oxygen. Studies exploring neurotoxicity include 13 week study via oral route (NaCN in drinking water) in rats and mice, 13 - 14 week inhalation study with acetone cyanohydrin in rats, 2 year inhalation study with acetonitrile in rats and mice, 180 day inhalation of cyanogens in rhesus monkeys. The NOAELs of these studies ranged from 4.7 (monkeys) to 26 mg/kg .bw (mice). As all these NOAELs are top doses, it is concluded that the neurotoxic endpoints are covered by NOAEL of 10 mg/kg bw.

Toxicological reference doses

Two AOELs have been defined for hydrogen cyanide covering the relevant exposure scenarios. Another condition that must be always fulfilled is that air concentrations of hydrogen cyanide never exceed AEC of 3 mg/m³ so as to avoid acutely dangerous peaks of cyanide in blood.

The AEC and acute AEL has been derived from human toxicokinetic data showing that the rate of spontaneous detoxication of cyanides in humans is 1 ug/kg body weight per minute. This rate of elimination balances, even under the very conservative assumption of 100% pulmonary retention, inhalation of air concentrations up to 3 mg/m³ with the concentration of CN in erythrocytes remaining on 24 hours exposure, safely below the concentration at which first subjective symptoms were reported. Thus 3mg/m³ define AEC the purpose of which is to prevent occurrence of acutely dangerous peaks in

blood. As accumulation of CN in blood depends also on the total amount of HCN absorbed applying AEC together with the relevant AEL, acute or chronic, prevents acutely dangerous CN peaks in blood from occurring in all the possible exposure scenarios. 24 hour exposure to 3mg/m³ corresponds to systemic dose of 1.44 mg/kg. bw assuming 100% pulmonary retention, inhalation rate of 1.25 m³/hour and body weight of 60 kg. The dose of 0.48 mg/kg bw is used as acute AEL which corresponds to 8 hour exposure under the above assumptions. The conservative assumptions used in the derivation of acute AEL of 0.48 mg/kg bw and the fact that the toxicokinetic data on which it is based come from studies on hospital patients treated for high blood pressure ensures protection of vulnerable groups as well as general population. This value is further supported by the absence of acute complaints in workers exposed for 8 hours to airborne HCN concentration below 20 mg/m³.

The derivation of long term AEL has been primarily based on two chronic studies, one for inhalatory and one for oral route of administration. In both the NOAEL was ≥ 10 mg CN/kg bw per day. Applying the standard assessment factor of 100 the long term AEL is 0.01 mg CN/kg bw per day. As the assessment factor accounts for both interspecies differences in toxicokinetics and toxicodynamics (coefficient < 10), and interindividual variability in heterogenous human population (coefficient >10) due to differences in thiocyanate elimination rate, thiosuphate and CN intake from diet, smoking etc. this AEL protects also the vulnerable groups as well as general population.

Exposure assessment and risk characterisation.

Operator exposure during fumigation:

During fumigation the personnel is required to use the prescribed personal protective equipment which rules out exposure to HCN during fumigation and ventilation. Organizational measures ensuring that operators will not come to contact with high concetrations of HCN vapours must be followed throughout the whole fumigation procedure including the ventilation phase and post ventilation phase until handing over the properly ventilated and cleared structures to the client/owner. Operators not wearing adequate PPE can only be exposed to concentration of HCN not exceeding 3mg/m³ while their internal exposure must not exceed long term AEL of 0.1 mg/kg bw. (i.e., in a case when operator should not wear adequate

PPE for the whole shift of 8 hours they can only be exposed to concentrations not exceeding 0.6 mg/m³

This then results in respiratory intake of less than 0.1 mg/kg bw (8 hours x 1.25 m³/hour x 0.6 mg/m³/60kg).

Post fumigation exposure:

Re- entry of operators into treated structures/ areas for inspection without use of the prescribed PPE including self-contained breathing apparatus is allowed only when gas concentrations dropped below AEC of 3mg/m³. The structure is entered then for the purpose of being handed over to the client. The time required for this hand-over should not exceed 1 hour resulting in an intake via inhalation of cyanide not exceeding 0.0625mg/kg.bw , which is safely below long term AEL.

Exposure of professionals during ventilation phase

During ventilation phase exclusion zone is determined so that the airborne HCN concentration at its border is 3 mg/m³ (AEC). An operator wearing prescribed PPE (i.e. a face mask with appropriate filter) is responsible for shifting the border if need be e.g. due to a change in weather conditions This operator is not exposed to HCN while wearing the PPE but exposure can take place during the breaks when the operator takes off the face mask. As a worst case such breaks are assumed to take up to 4 hours/day and the operator is required to find a place for these breaks, where the concentration of HCN in the air does not exceed 1 mg/m³. This then results in respiratory intake of HCN of 0.08 mg/kg bw when assuming body weight of 60 kg and inhalation rate 1.25 m³/h. This is 80% of long tem AEL of 0.1 mg/kg bw per day. If the operator is not to wear PPE for a major part of the 8 hour shift they must seek and stay in a place where the concentration of HCN does not exceed 0.6 mg/m³. This then results in respiratory intake of less than 0.1 mg/kg bw (8 hours x 1.25 m³/hour x 0.6 mg/m³/ 60kg).

Exposure of other users:

Hydrogen cyanide is intended for use by adequately trained professionals. Before delivery, the customer should declare the intended type of use and provide proof of his ability to handle the product safely. The manufacturer is, on the basis of delivery terms, entitled to carry out audits of the customer's premises.

Exposures of bystanders:

To avoid unacceptable exposure of by- standers and by- passers an exclusion zone is set around the fumigated structure which cannot be entered by any person except by adequately trained professionals

from the beginning of the fumigation till the handing over of the structure to the client. This zone is determined so as beyond its boundaries the concentration of airborne HCN never exceeds AEC of 3mg/m³. As by–passers are assumed to be exposed to HCN only infrequently acute AEL is relevant to assess the risk they undergo. 8 hour exposure to 3mg/m³ is needed before acute AEL is exceeded in adults which is more than passers-by can be reasonably assumed to spend near the frontier of the exclusion zone. Rather, reasonable assumption is that by-passers spent 30 minutes at the border of the exclusion zone. This corresponds to 0.03 mg /kg bw for adults when applying inhalation rate of 1.25 m³/hour and body weight of 60 kg (0.5 hours * 1.25 m³ *3 mg/m³ / 60 kg) and to 0.15 mg/kg bw for infants when applying inhalation rate 1 m³/hour and body weight of 10 kg. Thus, the systemic dose due to exposure of an adult passers- by corresponds to 6.3 % and that of an infant corresponds to 31% of acute AEL.

Exposure on the day following the hand over

On the day following its hand-over the fumigated structure is put to normal use. Eight hour exposure of persons entering it is assumed. HCN concentration in the air is bound to drop by several orders of magnitude by the time of the beginning of exposure if the first order kinetics with the rate constant derived from decrease during ventilation phase is assumed (i.e., during ventilation the drop was from 10 g/m³ to 3 mg/m³ in 24 hours thus giving rate constant of 0.34 hour¹ for first order kinetics). In reality, the post ventilation drop will be even more drastic as all the seals will be removed from windows, doors etc. and thus more air will be exchanged per unit of time. In addition, during the first part of the normal use its advisable to continue good ventilation of the object. Then during the ventilation the 8 hour exposure on the day following the hand over is calculated to be 0.008 mg /kg bw assuming inhalation rate of 1.25 m³/hour, body weight of 60 kg, 12 hours between the hand over and the beginning of the exposure, no drop of HCN concentration during the 8 hour exposure. This dose is 8% of chronic AEL thus posing no risk to human health.

2.2.2 Physical-chemical hazard

The relevant physical and chemical properties of biocidal product Uragan D2 are the same as that of hydrogen cyanide. Hydrogen cyanide is at normal pressure an extremely flammable gas/liquid. HCN vapours form explosive mixtures with air with upper explosive limit 40 % vol. and lower explosive limit 5.6 % vol.: the maximum concentration used in fumigation is below 5 %, nevertheless the danger of fire and explosion of vapours is high with regard to local concentration inhomogeneity.

2.2.2.1 Risk characterisation for the physico-chemical properties

When used conformably to special "Manual for Organization of hydrogen cyanide sanitation procedures", physical and chemical properties of hydrogen cyanide do not present risk to users.

2.2.3 Environmental risk assessment

2.2.3.1 Fate and distribution in the environment

Environmental fate and behaviour of HCN, due to its low boiling point, high vapour pressure at temperatures over 10 °C and lower relative density compared to density of air, is different from the fate and behaviour of other cyanide compounds. The main compartment where the most significant part of HCN liberated into the environment is transferred is the atmosphere. The persistence half-time of HCN in the atmosphere is 1-3 years. The most important mechanism of its degradation in the atmosphere is a reaction with hydroxyl radicals brought to the atmosphere by air humidity

Hydrogen cyanide is completely miscible with water. However, its ability to cross from the atmosphere into aqueous media, characterized by the value of Henry's law constant 5.2 kPa. m³. mol⁻¹, is low. Therefore, the part of hydrogen cyanide which is washed out from the atmosphere by precipitation is low as well. If hydrogen cyanide or cyanides enter aqueous media, equilibrium between the concentration of cyanide ions and undissociated hydrogen cyanide is established.

Biodegradation contributes to the elimination of cyanides from natural water. Cyanides occur in water most commonly in the form of hydrogen cyanide, cyanide ions and other cyanide compounds in a wide range.

In water, HCN and cyanide ion exist in equilibrium, their relative concentrations depend on pH and temperature. With pH lower than 8, more than 93 % free cyanides in water is in the form of undissociated hydrogen cyanide. HCN consequently hydrolyses to formamide which is further hydrolysed to ammonia and formate ion. However, the hydrolysis rate is slow and in the elimination of cyanide ion, it does not compete with evaporation and biodegradation.

Biodegradation of cyanides in surface water also depends on pH, cyanide concentration, temperature, availability of nutrients, and microbe adaptation. Cyanide ion is toxic for microorganisms at concentration 5-10 mg/l, but adaptation of microorganisms to this compound increases tolerance and microorganisms are able to decompose low cyanide concentrations.

In wastewater treatment plant conditions, adapted sludge is capable of decomposing cyanide concentrations lower than or equal to 100 mg/l.

Non-toxic concentrations of cyanides can be readily biodegraded, both aerobically and anaerobically. Aerobic degradation yields CO₂ and ammonia (that may be further converted to nitrate or nitrite); anaerobic biodegradation yields ammonia and methane.

In nature, degradation of free cyanide ions from aquatic environment occurs also due to these chemical processes: oxidation, hydrolysis, and photolysis, of which the last one plays only a negligible or very little role.

Hydrogen cyanide is very resistant to photolysis. The most important reaction of hydrogen cyanide in air is the reaction with photochemically generated hydroxyl radicals and subsequent rapid oxidation to carbon monoxide (CO) and nitric oxide (NO); photolysis and reaction with ozone are not important transformation processes, and reaction with singlet oxygen (O1D) is not a significant transformation process except at stratospheric altitudes where singlet oxygen is present in significant concentrations. The rate of hydroxyl radical reaction with hydrogen cyanide in the atmosphere depends on the altitude, and the rate of the reaction is at least one order of magnitude faster at lower tropospheric altitudes (0–8 km) than at upper tropospheric altitudes (10–12 km). Based on a reaction rate constant of 3x10⁻¹⁴ cm³/(molecule.sec) at 25 °C.

Photolysis in surface waters occurs, but is very low and its part in the degradation of cyanide ions from aquatic environment is insignificant.

Hydrogen cyanide hardly enters soil; its sorption ability to solid substances – sediment – is due to its high water solubility considered negligible.

Evaporation plays the biggest part in the dissipation of cyanides from water. In surface waters, this is a predominant fate of HCN.

Evaporation is influenced by several parameters, e.g.: temperature, pH, wind speed (in natural surface waters), and Henry's law constant.

At pH lower than 9.2, most of free cyanide in a solution exists in the form of HCN and volatile cyanides, and degradation (evaporation) proceeds faster. Evaporation is for HCN degradation from water more important than decomposition due to chemical reactions and biodegradation. This presumption applies to surface waters; elimination in ground waters shall take longer.

Most hydrogen cyanide from both natural and industrial sources reaches the atmosphere. HCN remains in the troposphere, only 2 % reaches the stratosphere.

In the atmosphere, HCN may be transported to long distances from the emission source.

HCN slowly degrades in air; its half-time is 1-3 years. In the atmosphere, it reacts with hydroxyl radicals brought there by air humidity, and through this reaction it decomposes. Although HCN is readily soluble in water, its elimination from the atmosphere through rain water is negligible.

HCN bioaccumulation in aquatic organisms is not expected. Bioconcentration factor for HCN was calculated - BCF 0.73. Neither HCN bioaccumulation in the food chain is expected.

Due to its usage as fumigant, using hydrogen cyanide for direct fumigation of food and feed is not expected. Since significant penetration of HCN into water or soil after treatment is not expected either, the risk of compartment-non-specific intoxication of people by the food chain may be considered negligible.

2.2.3.2 Effects assessment

Aquatic Compartment

The results of many experiments are published in the literature in which the toxicity of cyanides for fish, invertebrates and algae was investigated.

Acute toxicity for fish

Regarding fish toxicity, in some species of juvenile fish, the sensitivity is higher or the same at lower temperature, in other species the sensitivity to HCN is higher at higher temperatures. Generally, all measured values are within the classification highly toxic for aquatic organisms.

Observations from summary materials used are based on the article by Kovacs T. G., and G. Leduc. 1982. Acute toxicity of cyanide to rainbow trout (*Salmo gairdneri*) acclimated at different temperatures. Can . J. Fish. Aquat. Sci. 39: 1426-1429, in which dependency of temperature and HCN concentration effects on acute toxicity is documented. 96-hour mean LC50 values from the study conclusions:

$$LC50 = 0.028 \pm 0.004 \text{ mg.l}^{-1} \text{ at } 6 \text{ }^{\circ}\text{C}$$

$$LC50 = 0.042 \pm 0.004 \text{ mg.l}^{-1} \text{ at } 12 \text{ }^{\circ}\text{C}$$

$$LC50 = 0.068 \pm 0.004 \text{ mg.l}^{-1} \text{ at } 18 \text{ }^{\circ}\text{C}$$

Rainbow trout acclimated for the test temperature survived longer in lethal concentrations of cyanide. Toxicity curves clearly showed the temperature effect on the acute toxicity of cyanide is concentration dependent.

The LC50 = 0.042 mg.L⁻¹ value was selected for the risk assessment, with regard to temperatures at which acute toxicity test are performed according to current methods (Regulation (EC) 440/2008, EU method no. 203, temperature during the test 12-18 °C). Regarding the way of the substance use and the fact that the fumigation process is performed only at favourable climatic conditions, and regarding effects of other factors in the environment, no significant HCN concentration able to affect adversely aquatic organisms is expected to enter water. From this point of view the effect of temperature and concentration on the LC50 value is not important for the risk assessment.

Acute toxicity for invertebrates

A value from the test performed by the applicant was chosen as the key value, since this test was performed in the GLP system and according to the valid OECD methodology:

EC50 (Daphnia magna, 48 hours) = 1.07 mg.l⁻¹

Growth inhibition on algae

A value from the test performed by the applicant in the GLP system and according to the valid OECD methodology was chosen as the key value:

EC50 (Scenedesmus subspicatus, 72 hours) = 0.040 mg.l^{-1}

The calculation was performed with EUSES program, using a scenario for fumigation, with the following results:

PNEC Aqua: 4 x 10⁻⁵ mg/l.

Although hydrogen cyanide is highly toxic for aquatic organisms, exposure of the aquatic environment during fumigation is negligible.

Significant exposure of aqueous environment is not expected.

Sediment

No sediment tests are available.

The calculation was performed with EUSES program, using a scenario for fumigation, with the following results:

PNEC for fresh-water sediment-dwelling organisms 3.81 x 10⁻⁵ mg/kg wwt

Direct sediment exposure is not expected due to use pattern and physico-chemical properties of hydrogen cyanide.

Inhibition of microbial activity

A value for inhibition of microbial activity 25 mg/l was found in literature sources.

Hydrogen cyanide is a gas and its use pattern as a fumigant with direct release to the environment, there is no likelihood that the active ingredient will enter aerobic microbial treatment plants/sewage plants/water treatments plants. Consequently, there is no likelihood of exposure for STP micro-organisms.

The calculation was performed with EUSES program, using a scenario for fumigation, with the following results:

$$PNEC_{STP} = 2.5 \times 10^{-1} \text{ mg/L}$$

Terrestrial Compartment

The use is limited to closed spaces, hydrogen cyanide is used in the form of a gas for fumigation; the main environmental compartment it enters is air. Hydrogen cyanide tends to ascend to higher levels of the atmosphere. Direct release to the terrestrial compartment is not expected.

The calculation was performed with EUSES program, using a scenario for fumigation, with the following results:

$$PNEC_{soil} = 1.02 \times 10^{-5} \text{ mg/kg wwt}$$

Significant exposure of terrestrial environment is not expected.

Atmosphere

For the application of gaseous substances for fumigation, the general exposure scenario for the use of gaseous fumigants for the treatment of wood attacked by ligniperdous insects was proposed by working group of Environment Directorate OECD (OECD Series on Emission scenario Documents, Number 2, Emission Scenario Document for Wood Preservatives, Part 2, p. 93–96).

According to the above mentioned general scenario it is assumed that at most 2 % w/w of the total amount of the fumigant released into a closed object is retained in treated objects or materials and 0.1 % of the fumigant is decomposed. The extent of fumigant emissions to air is then expressed as an amount of the fumigant released into the treated object (decreased by the part retained in the treated object and by the part which underwent decomposition) recalculated on days in dependence on the ventilation time. If these general principles are applied to an individual case of fumigation with hydrogen cyanide, for the determination of the total amount of hydrogen cyanide emissions to air the calculation may be based on

the volume of the treated object or working chamber in m^3 and on the hydrogen cyanide application concentration of 20 g/m^3 .

For a extremely large object with the volume around 100,000 m³, the consumption about 2,000 kg of hydrogen cyanide can be expected, for a large object with the volume around 10,000 m³, the consumption about 200 kg of hydrogen cyanide can be expected, for a smaller object around 1,000 m³ one tenth, i.e. 20 kg, can be expected, and for a small container around 100 m³ approx. 2 kg. For a smaller container with the volume of 300 m³, it is 6 kg.

Emission rates of active substance to atmosphere ($E_{atm, fumi}$) after fumigation acc. to OECD Series on Emission scenario Documents, Number 2, Emission Scenario Document for Wood Preservatives, Part 2, p. 93–96 for objects with volume of 100,000 m³, 10,000 m³, 1,000 m³, 300 m³ and 100 m³ are 1,960, 196, 19.6, 5.87 and 1.96 kg/d respectively during 24hr ventilation and 653, 65.3, 6.53, 1.19 and 0.653 kg/d for 72hr ventilation time.

If the decrease of the amount of ventilated hydrogen cyanide by its retention or decomposition is neglected, this amount should be ventilated in 24–72 hours. In the less favourable case, the whole applied amount of hydrogen cyanide should leave to air within 24 hours. The concentration of hydrogen cyanide in gas leaving the ventilated object will decrease from the initial value higher than 20 g/m³ practically to zero at the end of the ventilation phase.

2.2.3.3 PBT assessment

It can be reliably stated that hydrogen cyanide does not have properties of PBT or vPvB both because of its preferential detention in free atmosphere, its low ability to bioaccumulate, characterized by BCF =0.73, and low persistence from the point of view of definition values of those parameters.

Hydrogen cyanide does not fulfil the PBT or vPvB criteria.

2.2.3.4 Risk characterization

Risk for atmosphere

Hydrogen cyanide ventilated to air can cause damage by retaining in the air (and thus it could change the properties of atmosphere) and by indirect endangering human health and other parts of nature.

In air hydrogen cyanide behaves as small halogen-carbon compounds. It is capable of contributing to global warming, weakening the protective ozone layer, and increasing the ozone production in troposphere. However, the potential of those effects is small due to little penetration of hydrogen cyanide into stratosphere and due to a slow course of reactions by which ozone is formed in troposphere. The present conditions in atmosphere cannot be significantly changed by hydrogen cyanide entering the atmosphere after the end of fumigation, because the amount of hydrogen cyanide used for fumigation will

always be only a negligible part of the amount of this substance formed spontaneously by natural processes or released into atmosphere from other anthropogenic sources.

The amount of hydrogen cyanide released to air during individual applications in medium and large objects can be of the order of tens or hundreds of kilograms. From the regional point of view, such a small amount cannot cause any measurable change of hydrogen cyanide concentration in the atmosphere.

From the local point of view, it is necessary to know the distribution of concentrations in the vicinity of a treated object during its ventilation. According to the above mentioned typical emission scenario for fumigation, it was proposed to assume that the total applied amount of the fumigant is equal to the total flux of the fumigant emissions to air for the time of its ventilation. The above given reasoning gave us the flux of emissions 1–1000 kg HCN/day for 48-hours ventilation.

In 1970's, anthropogenic production of hydrogen cyanide into the atmosphere in the USA was estimated at approx. 20,000 t/y. Most of anthropogenic formed cyanides, around 90 %, were generated from motor vehicle exhaust fumes (7-9 mg/km for vehicles not equipped with a catalyst and approx. 0.6 mg/km for catalyst equipped vehicles). Further significant anthropogenic sources of hydrogen cyanide emissions to the atmosphere include its production and production of other organic as well as inorganic cyanide compounds. In 2000, the total world production of HCN reached 1.4 mil. tons. Large amount of hydrogen cyanide is released to the atmosphere from processing industries such as metallurgy, surface treatment of metals, gold and silver mining from low-grade ores. Significant sources of HCN anthropogenic emissions include also landfills and sludge setting lagoons to which wastes containing cyanides, emissions from municipal and industrial waste incinerators, emissions from incinerating organic substances with high content of nitrogen (polyurethane, acrylonitrile, polyamides etc.) are placed. A relatively small quantity comes from the usage of HCN for treatment of closed structures.

Overview on the calculated PEC in air (according to the EUSES calculation)

Concentration in air during emission episode

| Product Type | PEC _{air} [mg/m³] | |
|---|----------------------------|--|
| PT08 | | |
| PT08_1 - Fumigant applied in a container with volume 100 m ³ , amount of HCN used: 2 kg. | 5.44x 10 ⁻⁴ | |

| Product Type | PEC _{air} [mg/m³] |
|---|----------------------------|
| PT08_6 - Fumigant applied in a container with volume 300 m ³ , amount of HCN used: 6 kg. | 1.63x 10 ⁻³ |
| PT08_2 – Fumigant applied in a small standard structure with volume 1,000 m ³ , amount of HCN used: 20 kg. | 5.44 x 10 ⁻³ |
| PT08_3 – Fumigant applied in a large standard structure with volume 10,000 m³, amount of HCN used: 200 kg. | 5.44 x 10 ⁻² |
| PT08_4 – Fumigant applied in a large standard structure with volume 100,000 m³, amount of HCN used: 2,000 kg. | 5.44 x 10 ⁻¹ |

Concentration in air, 100 m from point source

| Product Type | PECair |
|---|-------------------------|
| •• | [mg/m³] |
| PT08 | |
| PT08_1 - Fumigant applied in a container with volume 100 m ³ , amount of HCN used: 2 kg. | 1.49x 10 ⁻⁶ |
| PT08_6 - Fumigant applied in a container with volume 300 m ³ , amount of HCN used: 6 kg. | 4.47x 10 ⁻⁶ |
| PT08_2 – Fumigant applied in a small standard structure with volume 1,000 m ³ , amount of HCN used: 20 kg. | 1.49 x 10 ⁻⁵ |
| PT08_3 – Fumigant applied in a large standard structure with volume 10,000 m ³ , amount of HCN used: 200 kg. | 1.49 x 10 ⁻⁴ |

| Product Type | PECair |
|---|-------------------------|
| 11oddet 1,pc | [mg/m³] |
| PT08_4 – Fumigant applied in a large standard structure with volume 100,000 m³, amount of HCN used: 2,000 kg. | 1.49 x 10 ⁻³ |

Real values of concentration will depend on dispersion conditions (direction and velocity of wind, vertical temperature gradient, terrain configuration, surrounding buildings, etc.). At climatic situations favourable for dissipation of emissions, under which the fumigation and ventilation should be carried out, the real ground concentration of hydrogen cyanide should be significantly lower due to the tendency of hydrogen cyanide molecules, which are lighter than air, to move up to higher layers of atmosphere.

The values of hydrogen cyanide concentrations 0.544 mg/m³, estimated as PEC_{local} for one-time application of 2,000 kg of HCN, are approximately 5 times lower than PEL or 20 times lower than the value of MAC for working atmosphere valid in a number of countries.

Risk for aquatic environment

The risk for water is not expected due to an insignificant potential exposure of aqueous environment during fumigation with hydrogen cyanide.

At fumigation, hydrogen cyanide is applied to hermetically closed spaces, which cannot in any way communicate with surface or underground waters. No water can be present in treated objects. Direct exposure of aquatic environment to hydrogen cyanide is thus completely excluded. Indirectly, aquatic environment could be exposed to hydrogen cyanide retained by precipitation or by descending fog. Fumigation and following ventilation should thus be carried out only under favourable temperature and dissipation conditions. Therefore, there is low probability of direct contact of ventilated hydrogen cyanide with rain or fog.

If hydrogen cyanide comes into contact with atmospheric precipitations, its ability to be adsorbed in aqueous phase is low, as indicated by a relatively high value of Henry's constant. The highest nominal concentration of hydrogen cyanide in ventilated air at the beginning of ventilation should be close to applied concentration 20 g/m³. In equilibrium with this concentration of hydrogen cyanide in air, the concentration of hydrogen cyanide dissolved in water should reach the theoretical value of approx. 400 µg HCN per litre. To reach this concentration, it would be necessary to keep constant initial concentration of hydrogen cyanide in air for sufficiently long time to establish equilibrium between aqueous and gaseous phases. In reality, even in the least favourable case, when an exposure of aquatic

system would occur, the concentration of hydrogen cyanide in the contaminated water would reach micrograms or even lower values. After contact with ground, this concentration would further decrease by dilution with non-contaminated water, by re-volatilization of hydrogen cyanide, and by neutralization of its toxic effects by conversion into less toxicologically important compounds, eventually by hydrolysis supported by bacterial enzymes.

Risk of secondary intoxication

The risk of food chain intoxication is negligible because of insignificant penetration of hydrogen cyanide into this chain

2.2.4 List of endpoints

In order to facilitate the work of Member States in granting or reviewing authorisations, and to apply adequately the provisions of Article 5(1) of Directive 98/8/EC and the common principles laid down in Annex VI of that Directive, the most important endpoints, as identified during the evaluation process, are listed in Appendix I.

3 DECISION

3.1 Background to the proposed decision

Hydrogen cyanide is intended for use by adequately trained professionals as fumigant for control of pests in buildings and other closed spaces. After sufficient exposure, hydrogen cyanide immediately kills all development stages of pests. No signs of resistance development were reported. Hydrogen cyanide is classified as extremely flammable and very toxic by inhalation. Inclusion of hydrogen cyanide in Annex I is feasible for the human health aspect because several safe uses are identified. Adverse health effects to operators during fumigation are ruled out by obligatory usage of adequate PPE and other safety measures. cute toxic effects to persons re-entering the fumigated area after ventilation are prevented by following the obligatory safety measures. The only effect of long-term operator exposures is inhibition of thyroid functions. This effect should be prevented by setting chronic AEL. However, it is recommended to check for a possible occurrence of this effect by appropriate functional testing. Adverse health effects of passers-by are prevented by setting an exclusion zone around the fumigated structure/area.

The environmental risk assessment has shown that the proposed usage of hydrogen cyanide presents no unacceptable risk to the environment and can thus be included in Annex I. Hydrogen cyanide entering the atmosphere after the end of fumigation forms only a negligible part of the amount of this substance formed spontaneously by natural processes or released into atmosphere from other anthropogenic sources. Due to its physico-chemical properties hydrogen cyanide used in fumigation does not contribute to increase in levels of local background HCN emission or its content in surface water, nor is it expected to bioaccumulate significantly in aquatic organisms. Accidental endangering of human and animal health by hydrogen cyanide being retained in the air on ventilation is minimized by following the strict measures proposed for fumigation procedure.

3.2 Proposed decision regarding the inclusion in Annex I

Hydrogen cyanide shall be included in Annex I to Directive 98/8/EC as an active substance for use in product-type 14 (rodenticides), subject to the following specific provisions:

The minimum purity of the active substance used for the evaluation was 976 g/kg.

Member states shall ensure that authorisations of products for use as a fumigant are subject to the following conditions:

- Product shall only be supplied to and used by professionals adequately trained to use them;
- Safe operational procedures during fumigation and venting shall be established for operators and bystanders;
- Products shall be used with adequate personal protective equipment including, where appropriate, self-contained breathing apparatus and gas-tight clothing;
- Re-entry into fumigated spaces shall be prohibited until the air concentration has reached safe levels for operators and bystanders by ventilation;
- Exposure during and after ventilation shall be prevented from exceeding safe levels for operators and bystanders by the establishment of a supervised exclusion zone;
- Prior to fumigation, any food and any porous material with a potential to absorb the active substance, except the wood intended to be preserved, shall either be removed from the space to be fumigated or protected from absorption by adequate means, and the space to be fumigated shall be protected against accidental ignition.

3.3 Elements to be taken into account by Member States when authorising products

Elements, which were not mentioned under the specific provisions of the decision but which need to be taken into account at product authorisation level:

- Studies proving efficacy including kinetics of HCN evaporation in a treated object shall be required at the product authorisation stage;
- Residential buildings fumigation is not recommended;
- Authorisation holders shall ensure that users of the product are provided with detailed instructions
 for use, specifying the safety measures to be observed to ensure a safe and efficient use of the
 product;
- An exclusion zone shall be determined at the border of which HCN concentration must not
 exceed 0.6 mg/m3 and shall be set according to assumed exposure duration so that long term
 AEL is not exceeded for operators. In the exclusion zone, the presence of bystanders shall be
 prohibited and operators shall wear appropriate personal protective equipment. The zone shall be
 supervised.
- After fumigation, fumigated spaces shall be ventilated until the air concentration is below the AEC of 0.6 mg/m³ in order to protect operators shall they have to re-enter the fumigated spaces, and must in any case be below 3mg/m³ for the re-entry of bystanders. Fumigated spaces shall be returned to their normal use no earlier than 24 hours after this concentration has been reached.

3.4 Requirement for further information

It is considered that the evaluation has shown that sufficient data have been provided to verify the outcome and conclusions, and permit the proposal for the inclusion of hydrogen cyanide for use in product-type PT 08 (wood preservatives) in Annex I to Directive 98/8/EC.

3.5 Updating this Assessment Report

This assessment report may need to be updated periodically in order to take account of scientific developments and results from the examination of any of the information referred to in Articles 7, 10.4 and 14 of Directive 98/8/EC. Such adaptations will be examined and finalised in connection with any amendment of the conditions for the inclusion of spinosad in Annex I to the Directive.

APPENDIX I: LIST OF ENDPOINTS

Chapter 1: Identity, Physical and Chemical Properties, Details of Uses, Further Information, and Proposed Classification and Labelling

| Active substance (ISO Common Name) | Hydrogen cyanide | |
|--|---|--|
| Function (e.g. fungicide) | Insecticide (Fumigant) | |
| | | |
| Rapporteur Member State | Czech Republic | |
| | | |
| Identity (Annex IIA, point II.) | | |
| Chemical name (IUPAC) | Hydrogen cyanide | |
| Chemical name (CA) | Hydrocyanic-acid | |
| CAS No | 74-90-8 | |
| EC No | 200-821-6 | |
| Other substance No. | Index no.: 006-006-00-X | |
| | | |
| Minimum purity of the active substance as | 976 g/kg | |
| manufactured (g/kg or g/l) | | |
| Identity of relevant impurities and additives | Sulphur dioxide $9-11$ | |
| (substances of concern) in the active substance as manufactured (g/kg) | stabilizing additive preventing spontaneous | |
| | polymerisation | |
| | Phosphoric acid 0.8-1.2 | |

Hydrogen cyanide Product –type 08 25 May 2012

stabilizing additive preventing spontaneous polymerisation

HCN

27.03 g/mol

Molecular formula

Molecular mass

Appearance (state purity)



Physical and chemical properties (Annex IIA, point III., unless otherwise indicated)

Melting point (state purity) -13.4°C (7.9°F)

Boiling point (state purity) 25.7°C (78.3°F) (acid)

Temperature of decomposition Not required – No decomposition or sublimation occur at the melting or boiling temperature. It is gas.

occur at the merting of coming temperature. It is gue

HCN is produced as liquid which is sorbed on surface of inert material. Boiling temperature of HCN in liquid state is 25.7 °C (78.3 °F). Due to the large surface of sorbed inert material, the evaporation is very fast. Therefore the active substance as used is gas only.

Gas/ colourless

Smells of bitter almonds.Olfactory threshold:

0.17ppm (wt/vol.) in water

0.58ppm (vol./vol.) in air

Relative density (state purity) Density 0.6884 g/cm3 (liquid at 20 °C/68 °F)

Relative density / Specific gravity 0.687 (liquid at

Surface tension

Specific density: vapours 0.937 at 31°C/ 87.8 °F

Not relevant. Active substance hydrogen cyanide is

gas. HCN is used in gas phase for fumigation as it evaporates from inert material to which it is sorbed.

Vapour pressure (in Pa, state temperature) 84 kPa (at 20°C / 68°F)

35 kPa (at 0°C / 32°F)

Henry's law constant (Pa.m3.mol -1) 5.1 kPa.m3.mol-1

Solubility in water (g/l or mg/l, state Substance is fully miscible with water.

Solubility in organic solvents (in g/l or mg/l, state temperature) (Annex IIIA, point III.1)

temperature)

Soluble in ethanol, ether

Stability in organic solvents used in biocidal products including relevant breakdown products (IIIA, point III.2)

Not relevant. the active ingredient is actually the product. Hence, no organic solvents are used in the product

Partition coefficient (log POW) (state temperature)

Log Kow = +0.66 at 20 °C/68 °F

Hydrolytic stability (DT50) (state pH and temperature) (point VII.7.6.2.1)

Apparently at pH < 8.3 HCN is the dominant species, at pH < 7.99% will be as HCN molecule, and at pH > 10 CN

| Dissociation constant (not stated in Annex IIA or IIIA; additional data requirement from TNsG) | pKa of 9.2 |
|--|--|
| UV/VIS absorption (max.) (if absorption > 290 nm state ϵ at wavelength) | UV/VIS maximum ≤ 200 nm, no absorption above 290 nm |
| Photostability (DT50) (aqueous, sunlight, state pH) (point VII.7.6.2.2) | Airborne HCN undergoes slow photolysis. The overall atmospheric lifetime of HCN is 5 to 6 months. |
| Quantum yield of direct phototransformation in water at $\Sigma > 290$ nm (point VII.7.6.2.2) | None |
| Flammability | -17.8°C (flashpoint, closed cup) 538 °C / 1,000 °F (ignition point) |
| Explosive properties | Forms explosive gaseous mixtures with air with these explosive limits: upper: 40% vol. |
| | In alkali medium it may come under an autocatalytic polymerisation reaction running in an explosion speed. |

Hydrogen cyanide Product –type 08 25 May 2012

Summary of intended uses 1

| Object situation | and/or | Member State or | Produ ct name | Organisms controlled | Formulati Application | | | Applied amount per treatment | | | Remarks: | | |
|------------------|--------|--------------------|---------------------|----------------------|-----------------------|------------|---------------|------------------------------|----------------------|--------|----------|--|-----|
| | | Country | | | Тур | Con | metho | number | interval | g as/L | water | g as/m2 | |
| (a) | | | | (c) | e (d- | c. | d kind | min/ma x | between applications | _ | L/m2 | min / max | (m) |
| | | | | | f) | as | (f-h) | (k) | (min) | ax | ax | | |
| | | | | | | (i) | | | | | | | |
| Insect control - | pests | | URA - | All stages of target | Ga s | 97. 6 ± | Fumi gatio | Single use for | Single use. | | | Dosage: 10g/m3, i.e. in operating conditions | |

¹ Adapted from: EU (1998a): European Commission: Guidelines and criteria for the preparation of complete dossiers and of summary dossiers for the inclusion of active substances in Annex I of Directive 91/414/EC (Article 5.3 and 8,2). Document 1663/VI/94 Rev 8, 22 April 1998

| Hydrogen cyanide | Product –type 08 |
|------------------|------------------|
|------------------|------------------|

25 May 2012

| all stages | GAN | organisms - | 2.4 | n | killing | | 1kg/100m3. |
|------------------------|-----|-------------------------------|-----|---|---------|-------------|---|
| damaging | D2 | (e.g. | % | | pests | Further | Packing: Uragan D2 |
| goods stored in | | cockroach, cricket, mites, | | | | application | (stabilised liquid |
| storehouses, | | woodlouse, | | | | s only | hydrogen cyanide) is |
| depositories – | | larder beetle, | | | | upon new | supplied fully soaked |
| museums, temples, | | grain beetle, | | | | occurrence | into porous matter in |
| transport | | powder-post | | | | of pests. | closed gas-tight cans of 1.5kg Uragan D2. |
| vehicles – | | beetles, auger | | | | | 1.5kg Oragan D2. |
| railway | | beetle, wood- worm, spider | | | | | |
| wagons, sea | | beetle, | | | | | |
| and river boats, | | mealworm, | | | | | |
| airplanes, packages, | | sawyer beetle, | | | | | |
| containers, | | snout beetle, | | | | | |
| antiquities | | gelechid moth, | | | | | |
| (historical | | meal moth) | | | | | |
| wooden | | | | | | | |
| monuments), libraries, | | | | | | | |
| empty mills, | | | | | | | |
| etc. | | | | | | | |
| | | | | | | | |

| Hydrogen cyanide | | Product –type 08 | | | | | 25 May 20 | | | | |
|------------------|--|------------------|--|--|--|--|-----------|--|--|--|--|
| | | | | | | | | | | | |

Classification and proposed labelling (Annex IIA, point IX.)

| with rea | ard to ph | veical/ch | amical (| data |
|----------|------------|-----------|-----------|------|
| WILL IU2 | aiu io ini | vsicar/cm | ciiicai (| uata |

with regard to toxicological data

with regard to fate and behaviour data
with regard to ecotoxicological data

F+ - Extremely flammable.

R 12 Extremely flammable.

T+ Very toxic

R 26 Very toxic by inhalation.

No classification

N – Dangerous for the environment

R50/53 Very toxic to aquatic organisms, may cause long-term adverse effects in the aquatic environment.

(S 1/2) Keep locked up and out of reach of children.

S 7/9 Keep container tightly closed and in a well-ventilated place.

S 16 Keep away from sources of ignition – No smoking.

S 36/37 Wear suitable protective clothing and gloves.

S 38 In case of insufficient ventilation, wear suitable respiratory equipment.

S 45 In case of accident or if you feel unwell, seek medical advice immediately (show the label

| Hydrogen cyanide | Product –type 08 | 25 May 2012 |
|-----------------------|------------------|----------------|
| 11 valuedii e valiide | 1 Toduct type oo | 23 IVIA V 2012 |

where possible).

S60 This material and its container must be disposed of as hazardous waste.

S 61 Avoid release to the environment. Refer to special instructions / Safety data sheets.

Classification and labelling in compliance with Annex VI Regulation (EC) No. 1272/2008 (Annex IIA, point IX.)

| with regard to physical/chemical data | Flam. Liq. 1; |
|--|---|
| | H224: Extremely flammable liquid and vapour |
| with regard to toxicological data | Acute Tox.1; |
| | H330 Fatal if inhaled |
| with regard to fate and behaviour data | No classification |
| with regard to ecotoxicological data | Aquatic Acute 1; Aquatic Chronic |
| | H400 Very toxic to aquatic life |
| | H410 Very toxic to aquatic life with long lasting effects |
| | P210 Keep away from heat/sparks/open flames/hot surfaces. — No smoking. |
| | P260 Do not breathe dust / fume / gas / mist / vapours / spray. |
| | P262 Do not get in eyes, on skin, or on clothing. |
| | P280/284 Wear protective gloves/protective |
| | clothing/eye protection/face |

protection/respiratory protection.

P303+P361+P353: IF ON SKIN (or hair)
Remove/Take off immediately all contaminated clothing.Rinse skin with water/shower.

P304+P340 IF INHALED: Remove victim to fresh air and keep at rest in position comfortable for breathing.

P310 Immediately call a POISON CENTER or doctor/physician.

P273 Avoid release to the environment.

| RMS: Czech Republic | Hydrogen cyanide PT 08 | |
|---------------------|------------------------|--|
| | | |

Chapter 2: Methods of Analysis

Analytical methods for the active substance

Technical active substance (principle of method) (Annex IIA, point 4.1)

Assessment of the hydrogen cyanide content during its production is carried out by argentometric titration of cyanides by silver nitrate following chemisorption of hydrogen cyanide into sodium hydroxide solution.

Method principle

Titration of cyanide with nitrate in an alkaline medium leads first to dissolution of silver cyanide in NaCN excess. As soon as all cyanide ions are used for forming a complex anion, the first excessive drop of AgNO3 will make a silver cyanide precipitate.

Impurities in technical active substance

There are no impurities.

(principle of method) (Annex IIA, point 4.1)

Analytical methods for residues

Soil (principle of method and LOQ) (Annex IIA, point 4.2)

Modification of Standard Methods for the Examination of Water and Wastewater, American Public Health Association, Washington, Method No. 413:Cyanide

Method principle

All cyanides are isolated from acided sample by distillation with help of the inert gas, allowing for 5 to 10 fold enrichment, and after that are determined photometrically. Cyanides react with chloramine T to produce chlorcyan, which yields in combination with pyridine and barbiture acid at pH 4-5 in red-purple colouring. Its intensity is measured at a wavelength of 578nm.

The LOQ is 0.005 mg/l for enrichment factor 5.

Air (principle of method and LOQ) (Annex IIA, point 4.2)

1) The determination of cyanides content in workplace and storehouse atmospheres, and at combustion gases inlets from waste gas incinerators is done with COMPUR 4120 STATOX analyser operating with infrared detectors. Measuring range 0–50ppm (0–56mg.m-3). Manufacturer: Compur Monitors GmbH & Co. KG, Weissenseestrasse 101, D-81539 Munich, Germany.

And there is another possibility: Using detection tubes designed for hydrogen cyanide determination, type: hydrogen cyanide 2/a, No. CH 25701, Detection tubes manufacturer: Dräger Safety,

AG&Co.KGaA, Lubeck, Germany. measuring range for 5 pump strokes is: 2-30 ppm. The measuring range of the method depends on the number of strokes, e.g. for 40 strokes it is 0.25-3.75 ppm.

Water (principle of method and LOQ) (Annex IIA, point 4.2)

Modification of Standard Methods for the Examination of Water and Wastewater, American Public Health Association, Washington, Method No. 413:Cyanide

Method principle

All cyanides are isolated from acided sample by distillation with help of the inert gas allowing for 5 to 10 fold enrichment, and after that are determined photometrically. Cyanides react with chloramine T to produce chlorcyan, which yields in combination with pyridine and barbiture acid at pH 4-5 in red-purple colouring. Its intensity is measured at a wavelength of 578nm.

The LOQ is 0.005 mg/l for enrichment factor 5.

Body fluids and tissues (principle of method and LOQ) (Annex IIA, point 4.2)

Modification of Standard Methods for the Examination of Water and Wastewater, American Public Health Association, Washington, Method No. 413:Cyanide

Method principle

All cyanides are isolated from acided sample by distillation with help of the inert gas allowing for 5 to 10 fold enrichment and after that are determined photometrically. Cyanides react with chloramine T to produce chlorcyan, which yields in combination with pyridine and barbiture acid at pH 4-5 in red-

purple colouring. Its intensity is measured at a wavelength of 578nm.

The LOQ is 0.005 mg/l for enrichment factor 5.

Food/feed of plant origin (principle of method and LOQ for methods for monitoring purposes) (Annex IIIA, point IV.1)

In its use, HCN does not come in contact with food or feed.

Food/feed of animal origin (principle of method and LOQ for methods for monitoring purposes) (Annex IIIA, point IV.1)

In its use, HCN does not come in contact with food or feed.

Chapter 3 Impact on human health

Absorption, distribution, metabolism and excretion in mammals (Annex IIA, point 6.2)

Rate and extent of oral absorption:

HCN is a gas at body temperature. HCN and cyanates are readily absorbed from water solutions.

Rate of oral absorption is considered 100 %.

Rate and extent of dermal absorption:

Gaseous hydrogen cyanide may be absorbed by

skin; ratio of inhalatory/dermal absorption is

estimate to be 300/1.

Rate and extent of absorption on inhalation

HCN is readily absorbed on inhalation. The rate of

absorption is considered to be 100 %.

Distribution:

HCN is after absorption quickly, within seconds,

distributed by blood into all tissues.

Potential for accumulation:

Hydrogen cyanide does not accumulate in organism. Thiocyanate concentration in blood may increase as a result of repeated exposure to HCN

exposure.

Rate and extent of excretion:

CN is excreted as thiocyanate, renal clearance: half-

time of thiocyanate = 4 h - 2 d.

Acute toxicity (Annex IIA, point 6.1)

Rat LD50 oral 3.1 mg/kg bw (as cyanide, i.e. NaCN 5.7 mg/kg .bw

Rat LD50 dermal 6.7 mg/kg bw (rabbit, water solution of HCN)

Rat LC50 inhalation 493 mg/m3 (5 minutes)

173 mg/m3 (30 minutes)

158 /m3 (60 minutes)

Little change expected at longer exposures.

No primary data on skin irritation are available due to the inherent difficulty of performing such studies for gases in general. Apart from this, high toxicity of CN- makes it impossible to perform such studies using liquid HCN or solutions of cyanides as this would lead to immediate death of the animal following dermal absorption.

No primary data on eye irritation are available due to the inherent difficulty of performing such studies for gases in general. Apart from this, high toxicity of CN- makes it impossible to perform such studies using liquid HCN or solutions of cyanides as this would lead to immediate death of the animal following dermal absorption.

Eye irritation

Skin irritation

Skin sensitization (test method used and result)

Mild irritation reported in men.

No primary data on skin senstization are available due to the inherent difficulty of performing such studies for gases in general. Apart from this, high toxicity of CN- makes it impossible to perform such studies using liquid HCN or solutions of cyanides as this would lead to immediate death of the animal following dermal absorption.

Mild irritation is reported in men..

Repeated dose toxicity (Annex IIA, point 6.3)

Species/ target / critical effect

Lowest relevant oral NOAEL / LOAEL

Lowest relevant dermal NOAEL / LOAEL

Lowest relevant inhalation NOAEL / LOAEL

NOAEL: 10mg/kg/day, 2-year dietary study in rats

(summary in DOC IIIA 6.5b), (top dose)

Not available.

180 day ,rats and monkeys

LOAEL: 25 ppm cyanogens (corresponding to 25 ppm CN or 30mg HCN /m3), lower body weight, transient change in behaviour

NOAEL: 11 ppm cyanogens (corresponding to 11 ppm CN or 13.2 mg/ m3)

(Summary in DOC IIIA, section 6.4.3a).

Genotoxicity (Annex IIA, point 6.6)

No genotoxic risk

(Discussion in DOC IIA, section 3.6)

Carcinogenicity (Annex IIA, point 6.4) Non-carcinogenic

Species/type of tumour

No tumours have been observed at combined chronicity – carcinogenicity study in rats and mice

lowest dose with tumours

No tumours have been observed.

Reproductive toxicity (Annex IIA, point 6.8).

Species/ Reproduction target / critical effect

Lowest relevant reproductive NOAEL / LOAEL

Species/Developmental target / critical effect

Lowest relevant developmental NOAEL / LOAEL

No effects on reproduction were observed.

NOAELs ranged from 1-26 mg/kg, rats and mice, always top doses

No data available.

rat, NOAEL 3.3 mg CN/kg bw (top dose)

Neurotoxicity / Delayed neurotoxicity (Annex IIIA, point VI.1)

Species/ target/critical effect

Increased kill and serious neurological disorders (tremor, ataxia, cerebral cells kill) were observed in laboratory animals at concentrations 50mg/m3 HCN).

Lowest relevantl NOAEL / LOAEL.

NOAEL s, always top doses, in relevant studies ranged from 4.7 to 25 mg/kg.bw, rats monkeys and mice. Duration of studies ranged from 13 weeks to 2 years.

Other toxicological studies (Annex IIIA, VI/XI)

Goitrogenic effects found in exposed animals and humans.

Thyrotropic effects in rats at a dose in water 3mg/kg bw of KCN. (Summary in DOC IIIA.6.8.1b; discussion also in DOC IIA.3.9.2.)

Medical data (Annex IIA, point 6.9)

Inhalation of hydrogen cyanide in concentrations >120mg/m3 may be fatal.

Chronic **HCN** occupational exposure to concentrations approximately 17 mg/m3 revealed a high prevalence of neurological, cardiovascular and gastrointestinal symptoms at concentrations about 17 mg/m3, mild symptoms at concentrations in the rage 5 to 13 mg/m3. Thyroid enlargement has been observed in workers exposed still concentrations in air for two years, but no symptoms and toxic effects at concentrations <3.6 mg/m3.

Summary (Annex IIA, point 6.10)

ADI*

AOEC (Operator/Worker Exposure)

| Value | Study | Safety factor |
|---------------------|---|---------------|
| | | |
| 3 mg/m ³ | Toxicokinetic studies in human adults (Schulz et al., | |

| | | 1982,1984) | |
|---|----------------------------|---|-----|
| AEC (non professionals) | 3 mg/m ³ | Toxicokinetic studies in human adults (Schulz et al., 1982,1984) | |
| AOEL (Operator/Worker Exposure) (acute) | 0.48 mg/kg bw per day***. | Toxicokinetic 1 studies in human adults (Schulz et al., 1982,1984) | I |
| AEL (non professionals, by-standers) (acute) | 0.48 mg/kg bw per day***. | Toxicokinetic 1 studies in human adults (Schulz et al., 1982,1984) | |
| AOEL/AEL (Operator/Worker Exposure) (chronic) | 0.1 mg/kg bw per day*** | 2-year studies in rats (inhalation – NTP 1994, oral – Howard, Hanzal, 1955) | 100 |
| AOEL/AEL (Operator/Worker Exposure) (medium term) | 0.1 mg/kg bw per day*** | 2-year studies in 1 rats (inhalation – NTP 1994, oral – Howard, Hanzal, 1955) | .00 |
| Drinking water limit | 0.05mg/l ** | | |
| ARfD (acute reference dose) | 0.48 mg/kg bw* | ** | |

Hydrogen cyanide Product –type 08 25 May 2012

no residues in food or feed; AEL (chronic) may serve as estimate for ADI, DOC IIA 3.11

** Czech Republic

*** equal to AEL (acute), DOC IIA 3.11

Acceptable exposure scenarios (including method of calculation)

Production

Concentration of HCN in the production hall is continuously monitored and each surpassing of OEL is signalised. Workers are approx. 90% of working hours in the control room, isolated from the production hall.

Professional users

Recommended HCN occupational concentration in treated structures is 10,000mg/m3 (= 9,000 ppm).

Professional exposure of persons carrying out fumigation of closed spaces with hydrogen cyanide is for safety reasons reduced by using whole body gas-tight protective clothing (ČSN EN 464), special breathing apparatuses with filter-ventilation units (ČSN EN 132 and ČSN EN 133), rubber gloves (ČSN EN 374-1) and rubber boots (ČSN EN 346).

Exposure of wood in special hermetised chambers reduces substantially the potential exposure of operators.

Non-professional users

Non-professional usage is not permitted. .

Indirect exposure as a result of use

Structures (or subjects) treated by fumigation may be opened and used only after being thoroughly ventilated to 3mg/m3.

Exposure of bystanders and re-entering persons is discussed in DOC IIB 8.2.3.

Chapter 4 Fate and Behaviour in the Environment

Route and rate of degradation in water (Annex IIA, point 7.6, IIIA, point XII.2.1, 2.2)

| Hydrolysis of active substance and relevant metabolites (DT50) (state pH and temperature) | pH: - |
|---|--|
| | pH: - |
| | pH: - |
| Photolytic / photo-oxidative degradation of active substance and resulting relevant metabolites | Direct photolysis of HCN does not practically occur. |
| Readily biodegradable (yes/no) | No |
| Biodegradation in seawater | Hydrogen cyanide does not spread into sea water. |
| Non-extractable residues | - |
| Distribution in water / sediment systems (active substance) | Hydrogen cyanide does not spread into surface waters, groundwater and sediments. |
| Distribution in water / sediment systems (metabolites) | Hydrogen cyanide does not spread into surface waters, groundwater and sediments. |
| | |

Route and rate of degradation in soil (Annex IIIA, point VII.4, XII.1.1, XII.1.4; Annex VI, para. 85)

| Mineralization (aerobic) | Not applicable |
|---|----------------|
| Laboratory studies (range or median, with | Not applicable |
| number of measurements, with regression | |
| coefficient) | |
| | |

| Hydrogen cyanide | Product –type 08 | 25 May 2012 |
|--|-----------------------|-------------|
| Field studies (state location, range or m with number of measurements) | nedian Not applicable | |
| | Not applicable | |
| Anaerobic degradation | Not applicable | |
| Soil photolysis | Not applicable | |
| Non-extractable residues | Not applicable | |
| Relevant metabolites - name and/or code applied a.i. (range and maximum) | Not applicable | |
| Soil accumulation and plateau concentrat | ion Not applicable | |

| RMS: Czech Republic | Hydrogen cyanide PT 08 | |
|---------------------|------------------------|--|
|---------------------|------------------------|--|

Adsorption/desorption (Annex IIA, point XII.7.7; Annex IIIA, point XII.1.2)

Ka , Kd

Not applicable

Kaoc , Kdoc

pH dependence (yes / no) (if yes type of dependence)

Fate and behaviour in air (Annex IIIA, point VII.3, VII.5)

Direct photolysis in air

Direct photolysis of HCN does not practically occur.

Quantum yield of direct photolysis

Not applicable

Not applicable

Volatilization

Not applicable

Monitoring data, if available (Annex VI, para. 44)

study)

| Hydrogen cyanide | Produc | et –type 08 | 25 May 2012 |
|---|--------|-------------|-------------|
| Air (indicate location and type of study) | | No | |

3.1 Chapter 5: Effects on Non-target Species

Toxicity data for aquatic species (most sensitive species of each group) (Annex IIA, point 8.2, Annex IIIA, point 10.2)

| Species | Time-scale | Endpoint | Toxicity |
|-------------------------|------------|----------|------------|
| Fish | | | |
| Fish | 96 hrs. | LC50 | 0.042 mg/l |
| Salmo gairdnei | | | |
| Invertebrates | | | |
| Daphnia | 48 hrs. | EC50 | 1.07 mg/l |
| Daphnia magna | | | |
| Algae | | | |
| Scenedesmus subspicatus | 72 hrs. | EC50 | 0.04mg/l |
| Microorganisms | , | | |
| Data not found. | | | |

Effects on earthworms or other soil non-target organisms

| Acute | toxicity | to | Not applicable for intended usage of the substance. |
|----------------------|----------|----|---|
| (Annex IIIA, point X | | | |
| 71 | , | | |
| Reproductive | toxicity | to | Not applicable for intended usage of the substance. |
| | | | |

Hydrogen cyanide Product –type 08 25 May 2012 (Annex IIIA, point XIII.3.2)

Effects on soil micro-organisms (Annex IIA, point 7.4)

Nitrogen mineralization Not applicable for intended usage of the substance.

Carbon mineralization Not applicable for intended usage of the substance.

Effects on terrestrial vertebrates

Acute toxicity mammals Not applicable for intended usage of the substance. to (Annex IIIA, point XIII.3.3) Acute Not applicable for intended usage of the substance. toxicity birds to (Annex IIIA, point XIII.1.1) Dietary toxicity birds Not applicable for intended usage of the substance. to (Annex IIIA, point XIII.1.2) Reproductive toxicity birds Not applicable for intended usage of the substance. to (Annex IIIA, point XIII.1.3)

Effects on honeybees (Annex IIIA, point XIII.3.1)

Acute oral toxicity Not applicable for intended usage of the substance.

Acute contact toxicity Not applicable for intended usage of the substance.

Effects on other beneficial arthropods (Annex IIIA, point XIII.3.1)

| Hydrogen cyanide | Produc | t –type 08 25 May 2012 |
|--|--------|---|
| Acute oral toxicity | | Not applicable for intended usage of the substance. |
| Acute contact toxicity | | Not applicable for intended usage of the substance. |
| Acute toxicity | to | Not applicable for intended usage of the substance. |
| | | |
| Bioconcentration (Annex IIA, point 7.5) | | |
| Bioconcentration factor (BCF) | | BCF = 0.73 |
| | | Hydrogen cyanide has low bioaccumulation potential. |
| Depration time (DT50) | | Not applicable |
| (DT90) | | |
| Level of metabolites (%) in organic accounting for > 10% of residues | anisms | Not applicable |

APPENDIX II: LIST OF INTENDED USES

Hydrogen cyanide has been evaluated for its use in fumigation to kill wood destroying and disfiguring pests (Product Type 08 of the Biocidal Products Directive). It is applied as gas gradually evaporating from an inert sorbent and can be used only by authorised professional users.

The product URAGAN D 2 was submitted by the applicant for evaluation. It is the active substance as manufactured sorbed onto an inert sorbent. The prescribed concentration of hydrogen cyanide vapors in fumigated structures is 20g/m3.

Protection of wood is performed either by fumigating the whole buildings or by placing wooden items which are to be treated (furniture, pallets etc.) inside a structure or a specially modified container.

APPENDIX III: LIST OF STUDIES

Data protection is claimed by the applicant in accordance with Article 12.1(c) (i) and (ii) of Council directive 98/8/EC for all study reports marked "Y" in the "Data Protection Claimed" column of the table below. For studies marked Yes(i) data protection is claimed under Article 12.1(c) (i), for studies marked Yes(ii) data protection is claimed under Article 12.1(c) (ii). These claims are based on information from the applicant. It is assumed that the relevant studies are not already protected in any other Member State of the European Union under existing national rules relating to biocidal products. It was however not possible to confirm the accuracy of this information.

References listed by reference number in DOC IV A and IVB:

Supplementary literature listed by DOC III A or B section number:

| Reference No DOC IV "A" Section No DOC III A Author(s) Year Company, GLP (Un)Unput | e different from company) Report No. here relevant) / hed Data Protection Claimed (Yes/No) |
|---|---|
|---|---|

| 4 DOC | | | Company, Report No. GLP (where relevant) / (Un)Unpublished | Protection Claimed (Yes/No) | Owner |
|---|------------|------|---|-----------------------------|-------|
| IV A1 | 2 | 2006 | Hazardous Substance Data Bank (HSDB), National Library of Medicine's TOXNET system (state in February 2006): Hydrogen cyanide *Peer reviewed* | N | n/a |
| A6.2, A6.7, A6.8.1, A6.9, A6.10,A 7.1.4 | 2 | 2004 | ATSDR 1997 Toxicological Profile for Cyanide, U.S. Department of Health and Human Services, September 2004. | N | n/a |
| DOC IV A3 Ram | mbeau M. 2 | 2001 | Delphine Benitez, S. Dupuis* and P. Ducom HYDROGEN CYANIDE AS AN IMMEDIATE ALTERNATIVE TO METHYL BROMIDE FOR STRUCTURAL FUMIGATIONS Ministry of Agriculture, Fisheries and Food. National Laboratory of Plant Protection, Research Unit on Fumigation and Stored Products Protection, Chemin d'Artigues, 33150 Bordeaux-Cenon, France [*e-mail: lnds@easynet.fr] | N | n/a |
| DOC IV A4 A3.5, A.6 | | | Data From SRC PhysProp Database | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|--------------------|-----------|--|----------------------------------|-------|
| DOC IV A5a, A5b A7.4.1.11 | | 1980 | US EPA (1980) Ambient Water Quality Criteria for Cyanides. 440/5-80-037 (published). | N | n/a |
| DOC IV A6 A6.1.1, A6.1.2 | Smyth H.F. | 1969 | Carpenter CP, Weil CS, et. Al. 1969. Range-finding toxicity data: List VII. Am Ind Hyg Assoc J 30: 470-476 | N | n/a |
| DOC IV A7 A6.1.1 | Ferguson H.C. | 1962 | Dilution of dose and acute oral toxicity. Toxicol Appl Pharmacol 4: 759-762. | N | n/a |
| DOC IV A8 A6.1.1, A6.1.1.1a, A6.1.2, A6.1.2a, A6.1.2d, A6.1.4.2a, A6.3.2 A6.9, A6.12 | Balantyne Bryan | 1988 | Toxicology and Hazard Evaluation of Cyanide Fumigation Powders, Applied Toxicology Department, Union Carbide Corporation, Danbury, Connecticut 06817, Clinical Toxicology, 26 (5&6), 325-335 | N | n/a |
| DOC IV A9 A6.3.2 | B. Ballantyne | 1983 b | Acute systemic toxicity of cyanides by topical application to the eye. J Toxicol, Cutan, Ocular Toxicol 2: 119-129 (DOC IVA/) | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|--|-----------|---|----------------------------------|-------|
| A6.1.4.2, A6.2 A6.1.2, A6.1.2c A.1.3, A6.3.2 | Ballantyne B. | 1983 a | . The influence of exposure route and species on the acute lethal toxicity and tissue concentrations of cyanide. In: Hayes AW, Schnell RC, Miya TS, eds. Developments in the science and practice of toxicology. New York, NY: Elsevier Science Publishers, 583-586 | N | n/a |
| DOC IV A11 A6.1.3 | Matijak- Schaper M Alarie Y. | 1982 | Toxicity of carbon monoxide, hydrogen cyanide and low oxygen. J Combust Toxicol 9:21-61. | N | n/a |
| DOC IV A12 A6.1.3a, A6.2, A6.4, A6.4a | J.M.McNerney, M.P.H., H.H.Schrenk, PhD., | 1960 | The Acute Toxicity of Cyanogen, Industrial Hygiene Foundation, 4400 Fifth Avenue, Pittsburg 13, Pennsylvania, Industrial Hygiene Journal, 121 – 124 | N | n/a |
| DOC IV A13 A6.1.4.2 | Blac P, Hoan M, Mallin K | 1985 | Cyanide intoxication among silver- reclaiming workers. J Am Med Assoc 253: 367-371 | N | n/a |
| DOC IV A14 A6.1.4.1, A6.1.4.2, A6.10 | El Ghawabi SH, Gaafar MA, El-Saharti AA, et al. | 1975 | Chronic cyanide exposure: A clinical, radioisotope, and laboratory study. Br J Ind Med 32:215-219. | N | n/a |
| DOC IV A15 A6.1.4.1, A6.2 A6.4, A.4c, A6.9 | Fairley A, Linton EC, Wild FE. | 1934 | The absorption of hydrocyanic acid vapour through the skin with notes on other matters relating to acute cyanide poisoning. J Hyg 34: 283-294 | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|--|-------|---|----------------------------------|-------|
| DOC IV A16 A6.1.4.2, A6.12 | Bonsall JL. | 1984 | Survival without sequelae following exposure to 500 mg/m³ hydrogen cyanide. Hum Toxicol 3:57-60 | N | n/a |
| DOC IV A17 A6.1.4.2, A6.2, A6.12 | Chandra H, Gupta BN, Bhargava SK, Clerk SH, Mahendre PN | 1980 | Chronic cyanide exposure: a biochemical and industrial hygiene study. Journal of Analytical Toxicology, 3:161–165. | N | n/a |
| DOC IV A18 A6.2 | Yamamoto K, Yamamoto Y, Hattori H, et al. | 1982. | Effects of routes of administration on the cyanide concentration distribution in the various organs of cyanide-intoxicated rats. Tohoku J Exp Med 137: 73-78 | N | n/a |
| DOC IV A19 A6.2 | Walton D.C., Witherspoon MG | 1926 | . Skin absorption of certain gases. J Pharmacol Exp Ther 26: 315-324 | N | n/a |
| A6.2, A6.7, A6.10, A6.12 | | 2004 | IPCS (WHO, CICAD 61: Hydrogen cyanide and cyanides: human health aspects). CICAD 61 | N | n/a |
| DOC IV A21 A6.2, A6.12 | Schultz V | 1984 | Clinical pharmacokinetics of nitroprusside, cyanide, thiosulfate and thiocyanate. Clinical Pharmacokinetics, 9:239–251. | N | n/a |
| DOC IV A22 A6.3.1 | Sousa A.B., Soto-Blanco B, Guerra JL, Kimura ET, Gorniak S | 2002 | Does prolonged oral exposure to cyanide promote hepatotoxicity and nephrotoxicity? Toxicology, 174:87–95. | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|---|------|--|----------------------------------|-------|
| DOC IV A23 A6.3.3 | Valade M.P. | 1952 | Central nervous systém lesions in chronic experimental poisoning with gaseous hydrocyanic acid. Bull Acad Natl Med (Paris) 136: 280-285. (in French) (DOC IVA/) | N | n/a |
| DOC IV A24 6.4.1 | Tewe O.O., Maner JH | 1981 | Performance and pathophysiological changes in pregnant pigs fed cassava diets containing different levels of cyanide. Research in Veterinary Science,30:147–151 | N | n/a |
| DOC IV A25 A6.4.1, A6.7, A6.7a | Howard J. W., R. F. Hanzal | 1955 | Chronic Toxicity for Rats of Food Treated with Hydrogen Gyanide, Hazleton Laboratories, Falls Church, Va., Agricultural and Food Chemistry, Volume 3 No.4 | N | n/a |
| DOC IV A26 A6.4.1, A6.9, A6.10 | Philbrick D.J., Hopkins JB, Hill DC, et al. | 1979 | Effects of prolonged cyanide and thiocyanate feeding in rats. J Toxicol Environ Health 5:579-592. | N | n/a |
| DOC IV A27 A6.4.1a, A6.6.1, A6.6.1a, A6.8.2 | NTP. | 1993 | Technical Report on toxicity studies of sodium cyanide (CAS No. 143-33-9) administered in drinking water to F344/N rats and B6C3Fl mice. Research Triangle Park, NC: National Toxicology Program, U.S. Department of Health and Human Services, Public Health Service, National Institutes of Health. NIH Publication 94-3386. NTP TOX 37. | N | n/a |
| DOC IV A28 A6.4.1 | | 1993 | US EPA ydrogen cyanide (CASRN 74-90-8). US Environmental Protection Agency, Integrated Risk Information System. | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|--|-------|---|----------------------------------|-------|
| DOC IV A29 A6.4.3, A6.4.3a | Lewis T.R., Anger WK, Te Vault RK | 1984. | Toxicity evaluation of sub-chronic exposures to cyanogen in monkeys and rats. J Environ Pathol Toxicol Oncol 5:151-163. | N | n/a |
| DOC IV A30 A6.2, A6.12 | Ansell & Lewis | 1970 | Ansell M, Lewis FAS, A review of cyanide concentrations found in human organs: A survey of literature concerning cyanide metabolism, "normal", non-fatal and fatal bydy cyanide levels. Journal of Forensic Medicine, 17: 148-155 | N | n/a |
| DOC IV A31 A6.6.1, A6.6.1b | Kushi A., Matsumoto T, Yoshida D. | 1983 | Mutagen from the gaseous phase of protein pyrolyzate. Agric Biol Chem 47: 1979-1982 | N | n/a |
| DOC IV A32 A6.6.1 | De Flora S., Camoirano A, Zanacchi P, et al | 1984 | Mutagenicity testing with TA97 and TA102 of 30 DNA-damaging compouds, negative with other Salmonella strains. Mutat Res 134:159-165. | N | n/a |
| DOC IV A33 A6.6.1 | Friedman M.A., Staub J. | 1976. | Inhibition of mouse testicular DNA synthesis by mutagens and carcinogens as a potential simple mammalian assay for mutagenesis. Mutat Res 37: 67-76 | N | n/a |
| DOC IV A34 A6.6.1 | Kubo T, Urano K, Utsumi H | 2002 | Mutagenicity characteristics of 255 environmental chemicals. J Health Sci 48(6):545-554. | N | n/a |
| DOC IV A35 A6.6.1 | Bhattacharya R., Laskshmana Rao PV. | 1997 | Cyanide induced DNA fragmentation in mammalian cell cultures. Toxicology 123:207-215 | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|---|-------|--|----------------------------------|-------|
| DOC IV A36 A6.6.1 | Henderson L., Wolfreys A, Fedyk J, et al. | 1998 | The ability of the Comet assay to discriminate between genotoxins and cytotoxins. Mutagenesis 13:89-94 | N | n/a |
| DOC IV A37 A6.6.1, A6.6.4 | Yamamoto H., Mohanan PV | 2002. | Melatonin attenuates brain mitochondria DNA damage induced by potassium cyanide in vivo and in vitro. Toxicology 179:29-36. | N | n/a |
| DOC IV A38 A6.6.4 | Friedman M.A., Staub J. | 1976 | Inhibition of mouse testicular DNA synthesis by mutagens and carcinogens as a potential simple mammalian assay for mutagenesis. Mutat Res 37: 67-76 | N | n/a |
| DOC IV A39 A6.9, A7.1.1.2.1 | Fechter L.D., Chen G, Johnson DL. | 2002 | Potentiation of noise-induced hearing loss by low concentrations of hydrogen cyanide in rats. Toxicol Sci 66(1):131-138. | N | n/a |
| DOC IV A40 A6.12 | Vladimír Pitschmann | 2004 | Vojenská chemie kyanovodíku HCN, , Brno 2004, str. 28,Borowitz J. L., Isom G.E. Baskin S.I. v knize Somani S.M. Romano J.A. (Eds.): Chemical Warfare Agents: Toxicity at Low Levels. CRC Press, Boca Raton 2001 | N | n/a |
| DOC IV A41 | Manyonda, I.T. | 1986 | Shaw, D.E, Foulkes, A., Osborn, D.E Industrial exposure to hydrogen cyanide: implications for treatment British Medical Journal, Volume 293, 1986 | N | n/a |
| DOC IV A42 A6.12 | Gettler A.O., Baine JO | 1938 | The toxicity of cyanide. American Journal of Medical Science, 195:182–198. | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|---|------|--|----------------------------------|-------|
| DOC IV A43 A7.1.1.1.1 | Krieble V. E | 1930 | McNally, J. G.: The Hydrolysis of Hydrogen Cyanide by Acids II, <i>J. Am. Chem, Soc.</i> , 1929, 51, 3368. | No | n/a |
| DOC IV A44 A7.1.1.1.1 | Krieble V. E | 1929 | McNally, J. G.: The Hydrolysis of Hydrogen Cyanide by Acids I, <i>J. Am. Chem, Soc.</i> , 1929, 51, 3368. | No | n/a |
| DOC IV A45 | | | Kirk-Othmer Encyclopedia of Chemical Technology (4 th Edition) | No | n/a |
| DOC IV A46 A7.1.1.2.1 | Klecka G.M., Landi LP, Bodner KM. | 1985 | Evaluation of the OECD activated sludge, respiration inhibition test. Chemosphere 14:1239-1251. | N | n/a |
| A7.1.1.11, A7.1.1.1.2, A7.1.1.2.1, A7.1.3, A7.1.4, A7.2, A7.3.1 | | | JACC No 53, Cyanides of Hydrogen, Sodium and Potasium, and acetone Cyanohydrin (CAS No. 74-90-8, 143-33-9, 151-50-8 and 75-86-5), ECETOC JACC REPORT No. 53 European Centre for Ecotoxicology and Toxicology of Chemicals Volume I | N | n/a |
| A7.1.1.11, A7.1.1.1.2, A7.1.1.2.1, A7.1.3, A7.1.4, A7.2, A7.3.1 | | | JACC No 53, Cyanides of Hydrogen, Sodium and Potasium, and acetone Cyanohydrin (CAS No. 74-90-8, 143-33-9, 151-50-8 and 75-86-5), ECETOC JACC REPORT No. 53 European Centre for Ecotoxicology and Toxicology of Chemicals, Volume II | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|---|------|--|----------------------------------|-------|
| DOC IV A50 A7.4.1.1 | Smith L.L., Broderius S.J., Osied D.M., Kimbal G.L., Koenst W.M., | | Acute Toxicity of Hydrogen Cyanide to Freshwater Fishes, Paper No. 9954, under Grant No. R802914 | N | n/a |
| DOC IV A51 | | 2007 | Crop Research Institute (CRI) Evaluation of URAGAN (HCN) Field Efficacy – CRI - 2007 | Y | |
| DOC IV A52 | Rambeau M. | 1999 | HYDROGEN CYANIDE AS AN IMMEDIATE ALTERNATIVE TO METHYL BROMIDE FOR STRUCTURAL FUMIGATIONS D. BENITEZ, S. DUPUIS, P. DUCOM | Y | |
| DOC IV A53 A4.2 | | 2002 | Modification of Standard Methods for the Examination of Water and Wastewater, American Public Health Association, Washington, Method No. 413:Cyanide | Y | |
| DOC IV A54 | Walton D.C | 1925 | Witherspoon MG. 1926. Skin absorption of certain gases. J Pharmacol Exp Ther 26: 315-324 | N | n/a |
| DOC IV A55 | | | Compur Statox 4120 | N | n/a |
| DOC IV A56 A6.8.1a | Benito Soto- Blanco, Silvana L. Go'rniak | 2004 | Prenatal toxicity of cyanide in goats—a model for teratological studies in ruminants. Theriogenology 62: 1012–1026 | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|---|------|--|----------------------------------|-------|
| DOC IV A57 A6.8.1b | Altamir Benedito de Sousa, Paulo C'esar Maiorka, Ivair Donizete Goncalves, L'ilian Rose Marques de S'a, Silvana Lima G'orniak | 2007 | Evaluation of effects of prenatal exposure to the cyanide and thiocyanate in Wistar rats. Reproductive Toxicology 23: 568–577 | N | n/a |
| DOC IV A58 | | 2002 | European Union Risk Assessment Report Acetonitrile European Commission Joint Research Centre Priority List Volume 18 | N | n/a |
| DOC IV A59 A6.2 | | 2005 | Acetone Cyanohydrin. Acute Exposure Guideline Levels August | N | n/a |
| DOC IV A60 A6.2 | Schultz V., Gross R, Pasch T, Busse J, Loescheke G | 1982 | Cyanide toxicity of sodium nitroprusside in therapeutic use with and without sodium thiosulfate. Klinische Wochenschrift,60:1393–1400. | N | n/a |
| DOC IV A61 A6.10 | Way J.L. | 1984 | Cyanide intoxication and its mechanism of antagonism. Annual Review of Pharmacology and Toxicology, 24:451–481. | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|---|------|---|----------------------------------|-------|
| DOC IV A62 | Banerjee et al | 1997 | Kishore K. Banerjee, PhD, A. Bishayee, PhD, P. Marimuthu, MSc Evaluation of Cyanide Exposure and Its Effect on Thyroid Function of Wokers in a Cable Industry JOEM, Volume 39, Number 3, March 1997 | N | n/a |
| DOC IV A63 A6.4.1 | Tewe O.O., Maner JH | 1985 | Cyanide, protein and iodine interactions in the performance and metabolism of rats. Journal of Environmental Pathology and Toxicology, 6:69–77. | N | n/a |
| DOC IV A64 A6.12 | Jackson L.C., Bloch EF, Jackson RT, Chandler JP, Kim YL, Malveaux F | 1985 | Influence of dietary cyanide on immunoglobulin and thiocyanate levels in the serum of Liberianadults. Journal of the National Medical Association, 77:777–782. | N | n/a |
| DOC IV A65 A6.2 | Schultz et al. | 1982 | Detoxification of cyanide in a newborn child Klinische Wochenschrift,60,527-528 | N | n/a |
| DOC IV A66 A6.2 | Schultz V., Bonn R, Kindler J | 1979 | [Kinetics of elimination of thiocyanate in 7 healthy subjects and 8 subjects with renal failure.] Klinische Wochenschrift, 57:243–247 (in German). | N | n/a |
| DOC IV A67 A6.2 | Schultz et al. | 1983 | Resorption of hydrocyanic acid from linseed, Leber Magen Darm 13: 10-14. | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|--------------------------------------|------|--|----------------------------------|-------|
| DOC IV A68 A6.2 | Schultz et al. | 1979 | Counteraction of cyanide poisoning by thiosulphate when administering sodium nitroprusside as hypotensive treatment, Klinische Wochenschrift 57, 905-907. | N | n/a |
| DOC IV A69 | Schulz V., | 1978 | Thiozyanat-Vergiftung bei der antihypertensiven Therapie mit Natriumnitroprussid Medizinische Universitätsklinik Köln und Department für Innere Medizin der Medizinischen Hochschule Hannover | N | n/a |
| DOC IV A70 | Olumide O.,Tewe and Jerome H. Manert | 1980 | Long-Term and Carry-Over Effect of Dietary Inorganic Cyanide (KCN) in the Life Cycle Performance and Metabolism of Rats Department of Animal Science, University of Ibadan, Ibadan, Nigeria. Centro International De Agricultura Tropical, Colombia, South America | N | n/a |
| DOC IV A71 A6.9 | Jackson L.C | 1988 | Behavioural effects of chronic sublethal dietary cyanide in an animal model: Implications for humans consuming cassava (Manihot esculenta). Human Biology, 60:597–614. | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|-------------------------------|------|--|----------------------------------|-------|
| DOC IV A72 A6.4.3, A6.5a, A6.6.1a, A6.7 | | | National Toxicology Program (NTP)(1994). Toxicology and carcinogenesis of acetonitrile (CAS N° 75-05-8) in F344/N rats and B6C3F1 mice (Inhalation studies). TR 447. NIH Publication N° 94-3363. US Department of Health and Human Services, Public Health Service, National Institutes of Health. | N | n/a |
| DOC IV A73 | R.C. Brandys, G.M. Brandys | 2006 | Global occupational exposure limits for over 5,000 specific chemicals Occupational & Environmental Health Consulting Services, Hinsdale, Ill. | N | n/a |
| DOC IV A74 | | 2002 | Technical Guidance Document on Risk Assessment Part II, BCF | N | n/a |
| DOC IV A75 | | 2006 | EN 335-1 Durability of wood and wood- based product, Definition of use classes- Part 1: General | N | n/a |
| DOC IV A76 A6.6.4 | Monsanto Co. | 1984 | CHO/HGPRT mammalian cell forward mutation assay, acetone cyanohydrin. St. Louis, MO, Monsanto Co. (Report PR-82-204). | N | n/a |
| DOC IV A77 A6.4.3, A6.8.2 | Monsanto Co. | 1985 | Male fertility study of Sprague-Dawley rats exposed by the inhalation route to acetone cyanohydrin. St. Louis, MO, Monsanto Co. (Report ML-82-144; US EPA/OPTS Public Files No. 878216404). | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|---------------------------------------|------|---|----------------------------------|-------|
| DOC IV A78 A6.8.2 | Monsanto Co. | 1985 | Female fertility study of Sprague Dawley rats exposed by the inhalation route to acetone cyanohydrin. St. Louis, MO, Monsanto Co. (Report ML-82 145; US EPA/OPTS Public Files No. 878216396). | N | n/a |
| DOC IV A79 A6.4.3 | Monsanto Co. | 1984 | Three-month inhalation toxicity of acetone cyanohydrin in male and female Sprague-Dawley rats. St. Louis, MO, Monsanto Co. (Report ML-82-143; US EPA/OPTS Public Files No. 878216397). | N | n/a |
| DOC IV A80 A7.4.2 | By E. A PARKIN , M.Sc., PH.D., D.I.C. | 1937 | THE TOXICITY OF HYDROGEN CYANIDE TO CERTAIN WOOD- BORING INSECTS Entomology Section, Forest products Research Laboratory, Princes Risborough, Bucks | N | n/a |
| DOC IV A81 A7.1.1.1.2 | D. J. Lary, | 2004 | Atmospheric pseudohalogen chemistry, Atmos. Chem. Phys. Discuss., 4, 5381– 5405, 2004 <u>www.atmos-chem-phys.org/acpd/4/5381/</u> SRef-ID: 1680- 7375/acpd/2004-4-5381 © European Geosciences Union | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|-------------------|------|--|----------------------------------|-------|
| DOC IV A82 A7.1.1.2.1 | Dumestre Alain | 1997 | , THERESE CHONE, JEAN-MARIE PORTAL, MYLENE GERARD, AND JACQUES BERTHELIN Cyanide Degradation under Alkaline Conditions by a Strain of Fusarium solani Isolated from Contaminated Soils, APPLIED AND ENVIRONMENTAL MICROBIOLOGY, 0099-2240/97/\$04.0010 July 1997, p. 2729–2734 | N | n/a |
| DOC IV A83 A7.1.1.2.1 | | | Cyanide Degradation under Alkaline Conditions by a Strain of Fusarium solani Isolated from Contaminated Soils,APPLIED AND ENVIRONMENTAL MICROBIOLOGY, 1997, p. 2729–2734 ALAIN DUMESTRE,† THERESE CHONE, JEAN-MARIE PORTAL, MYLENE GERARD, AND JACQUES BERTHELIN* | N | n/a |
| DOC IV A84 A7.1.4.2 | | | Hydrogen cyanide: An acute toxicity study with the daphnia Daphnia magna Strauss, Research Institute of Organic syntheses, Centre for ekotoxicology, toxicology an analytics, Pardubice – Rybitví, Czech Republic,Report No. 1514/L (unpublished), 2002-02-18 | N | n/a |

| Reference No DOC IV "A" Section No DOC III A | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|---|-----------|------|--|----------------------------------|-------|
| DOC IV A85 A7.1.4.3 | | | Growth Inhibition of Green Algae (Scenedesmus subspicatus Brinkmann 1953/SAG 86.81) by hydrogen cyanide liquid stabilized, Research Institute of Organic syntheses, Centre for ekotoxicology, toxicology an analytics, Pardubice – Rybitví, Czech Republic,Report No. 1522/L (unpublished), 2002-03-20 | N | n/a |

Supplementary literature listed by section number

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|--------------|------|---|----------------------------------|-------|
| A3.1, A3.1.1, A3.2.1, A3.3.1, A3.3.2, A3.3.3, A3.5, A3.7, A3.9, A3.11, A3.12, A3.15 | ATSDR | 2004 | Agency for Toxic Substances and Disease Registry, U.S. Department of Health and Human Services: Draft Toxicological Profile for Cyanide, * Peer Reviewed*, str 141 (DOC IVA / A2) | N | n/a |
| A3.1, A3.3.1, A3.3.2, A3.3.3 | Budavari S., | 1989 | Merck index: An encyclopedia of chemicals, drugs, and biologicals. 11 th ed. Rahway, NJ: Merck &Co., Inc. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--------------------------------|---------------------------------------|------|--|----------------------------------|-------|
| A3.1.1, A3.11, A3.12, A3.15 | Jenks W.R. | 1979 | Cyanides. In: Grayson, M, ed. Kirk-Othmer encyclopedia of chemical technology. New York, NY: John Wiley and Sons, Inc., 307-334 | N | n/a |
| A3.11, A3.15 | Quincy | 1997 | Fire Protection Guide to Hazardous Materials. 12 ed, MA: National Fire Protection Association, 1997 | N | n/a |
| A3.2, A7.1.3 | Daubert T.E., Danner RP | 1989 | Physical and Thermodynamic Properties of Pure Chemicals Data Compilation Washington, DC: Taylor and Francis | N | n/a |
| A3.9, A7.4.2 | Hansch C., Leo, A., D. Hoekman. | 1995 | Exploring QSAR - Hydrophobic, Electronic, and Steric Constants. Washington, DC: American Chemical Society., p. 3 | N | n/a |
| A6.1.3 | | 1959 | Toxicology and Applied Pharmacology. (Academic Press, Inc., 1 E. First St., Duluth, MN 55802) V.1- 1959- v. 42, p. 417, (TXAPA9); | N | n/a |
| A6.1.3 | | 1987 | Fundamental and Applied Toxicology. (Academic Press, Inc., 1 E. First St., Duluth, MN 55802) V.1-40, 1981-97. For publisher information, see TOSCF2 v. 9, p. 236, (FAATDF) | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|---|----------------------------------|-------|
| A6.1.3 | AMRL | 1971 | The acute toxicity of brief exposures to hydrogen fluoride, hydrogen chloride, nitrogendioxide, and hydrogen cyanide singly and in combination with carbon monoxide. Wright-Patterson Air Force Base, OH: Aerospace Medical Research Laboratory. AD751442 | N | n/a |
| A6.1.3 | Hume A.S., Mozigo JR, McIntyre B, et al. | 1995 | Antidotal efficacy of alpha-ketoglutaric acid and sodium thiosulfate in cyanide poisoning. Clin Toxicol 33(6):721-724. | N | n/a |
| A6.1.3 | Monsanto Co.Report | 1985 | One-month inhalation toxicity of acetone cyanohydrin in male and female Sprague-Dawley rats. St Louis, Monsato Co. Report ML-81-178/810068 (US EPA/OPTS Public Files No. 878216393). | N | n/a |
| A6.1.3 A6.9 | Purser DA, Grimshaw P, Berrill KR. | 1984 | Intoxication by cyanide in fires: A study in monkeys using polyacrylonitrile. Arch Environ Health 39:394-400. IPCS, ATSDR | N | n/a |
| A6.10 | Maduh E.U., Johnson JD, Ardelt BK, Borowitz JL, Isom GE | 1988 | Cyanide-induced neurotoxicity: Mechanisms of attenuation by chlorpromazine. <i>Toxicology and Applied Pharmacology</i> , 96:60–67. | N | n/a |
| A6.10 | Pettersen JC, Cohen SD | 1993 | The effects of cyanide on brain mitochondrial cytochrome oxidase and respiratory activities. <i>Journal of Applied Toxicology</i> , 13:9–14. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|--|----------------------------------|-------|
| A6.10, A6.12 | Hardy H.L., Jeffries WM, Wasserman MM, Waddell WR | 1950 | Thiocyanate effect following industrial cyanide exposure. New England Journal of Medicine, 242:968–972 | N | n/a |
| A6.10, A6.12 | US EPA | 1990 | Summary review of health effects associated with hydrogen cyanide. Health issue assessment. Research Triangle Park, NC, US Environmental Protection Agency, Office of Research and Development (EPA/600/8-90/002F). | N | n/a |
| A6.12 | Anderson R.A., Harland WA | 1982 | Fire deaths in the Glasgow area. III. The role of hydrogen cyanide. <i>Medicine, Science and the Law</i> , 22:35–40 | N | n/a |
| A6.12 | | 1986 | American Conference of Governmental Industrial Hygienists. Documentation of the Threshold Limit Values and Biological Exposure Indices. 5th ed. Cincinnati, OH: American Conference of Governmental Industrial Hygienists, p. 314] | N | n/a |
| A6.12 | Abuye C., Kelbessa U, Wolde-Gebriel S | 1998 | Health effects of cassava consumption in south Ethiopia. <i>East African Medical Journal</i> , 75:166–170 | N | n/a |
| A6.12 | Alarie Y. | 2002 | Toxicity of fire smoke. Crit Rev Toxicol 32(4):259-289 | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|---|----------------------------------|-------|
| A6.12 | ATSDR | 1991 | Case studies in environmental medicine. Atlanta, GA, US Department of Health and Human Services, Public Health Service, Agency for Toxic Substances and Disease Registry. | N | n/a |
| A6.12 | Banea- Mayambu J.P., Tylleskar T, Gitebo N, Matadi N, Gebre- Medhin M, Rosling H | 1997 | Geographical and seasonal association between linamarin and cyanide exposure from cassava and the upper motor neurone disease konzo in former Zaire. <i>Tropical Medicine and International Health</i> , 2:1143–1151. | N | n/a |
| A6.12 | Birky M.M., Clarke FB | 1981 | Inhalation of toxic products from fires. Bulletin of the New York Academy of Medicine, 57:997–1013. | N | n/a |
| A6.12 | Boivin M.J. | 1997 | An ecological paradigm for a health behavior analysis of "Konzo," a paralytic disease of Zaire from toxic cassava. <i>Social Science and Medicine</i> , 45:1853–1862. | N | n/a |
| A6.12 | Chen K.K., Rose CL. | 1952 | 1952. Nitrite and thiosulfate therapy in cyanide poisoning. J Am Med Assoc 149:113-119. | N | n/a |
| A6.12 | Cherian M.A., Richmond I. | 2000 | Fatal methane and cyanide poisoning as a result of handling industrial fish: A case report and review of the literature. J Clin Pathol 53:794-795 | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|--|----------------------------------|-------|
| A6.12 | Cliff J., Coutinho J | 1995 | Acute intoxication from newlyintroduced cassava during drought in Mozambique. Tropical Doctor, 25:193. | N | n/a |
| A6.12 | Cliff J., Lundquist P, Rosling H, et al. | 1986 | Thyroid function in a cassava-eating population affected by epidemic spastic paraparesis. Acta Endocrinol (Copenh) 113:523-528. | N | n/a |
| A6.12 | Cooles P. | 1988 | Diabetes and cassava in Dominica. <i>Tropical</i> and Geographical Medicine, 40:272–273. | N | n/a |
| A6.12 | Delange F., Hershman JM, Ermans AM | 1971 | Relationship between the serum thyrotropin level, the prevalence of goiter and the pattern of iodine metabolism in Idjwi Island. <i>Journal of Clinical Endocrinology and Metabolism</i> , 33:261–268. | N | n/a |
| A6.12 | Delange F, Ermans AM | 1971 | Role of a dietary goitrogen in the etiology of endemic goiter on Idjwi Island. <i>American Journal of Clinical Nutrition</i> , 24:1354–1360. | N | n/a |
| A6.12 | DOA | 1976 | Estimates of the toxicity of hydrocyanic acid vapours in man. Aberdeen Proving Ground, MD: Department of the Army. EBTR76023. ADA02850 | N | n/a |
| A6.12 | Drinker P. | 1932 | Hydrocyanic acid gas poisoning by absorption through the skin. J Ind Hyg 14:1-2 | N | n/a |
| A6.12 | Dudley H.C., Sweeney TR, Miller JW. | 1942 | Toxicology of acrylonitrile (vinyl cyanide). II: Studies of effects of daily inhalation. J Ind Hyg Toxicol 24:255-258 | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|---|----------------------------------|-------|
| A6.12 | Ermans A.M., Delange F, Van Der Velden M, Kinthaert S | 1972 | Possible role of cyanide and thiocyanate in the etiology of endemic cretinism. <i>Advances in Experimental Medicine and Biology</i> , 30:455–486. | N | n/a |
| A6.12 | Ernesto M., Cardoso AP, Nicala D, Mirione E, Massaza F, Cliff J, Haque MR, Bradbury JH | 2002 | Persistent konzo and cyanogen toxicity from cassava in northern Mozambique. <i>Acta Tropica</i> , 82:357–362. | N | n/a |
| A6.12 | Gill G. | 1996 | Tropical diabetes? <i>Tropical Doctor</i> , 26:1–3. | N | n/a |
| A6.12 | Gosselin, R.E., R.P. Smith, H.C. Hodge | 1984 | Clinical Toxicology of Commercial Products. 5th ed. Baltimore: Williams and Wilkins,., p. III-126] | N | n/a |
| A6.12 | Gosselin, R.E., R.P. Smith, H.C. Hodge. | 1984 | Clinical Toxicology of Commercial Products. 5th ed. Baltimore: Williams and Wilkins,., p. III-127 | N | n/a |
| A6.12 | Howlett W.P., Brubaker GR, Mlingi N, et al. | 1990 | Konzo, an epidemic upper motor neuron disease studied in Tanzania. Brain 113:223-235 | N | n/a |
| A6.12 | Hugh-Jones P. | 1955 | Diabetes in Jamaica. Lancet, 2:891–897. Concise International Chemical Assessment Document 61 36 | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|---|----------------------------------|-------|
| A6.12 | JECFA | 1993 | Cyanogenic glycosides. In: Toxicological evaluation of certain food additives and naturally occurring toxicants. Geneva, World Health Organization, 39th Meeting of the Joint FAO/WHO Expert Committee on Food Additives (WHO Food Additives Series 30) | N | n/a |
| A6.12 | Kumar P., Das M, Kumar A. | 1992 | Health status of workers engaged in heat treatment (case hardening) plant and electroplating at cyanide bath. Indian J Environ Prot 12(3):179-183 | N | n/a |
| A6.12 | Lantum H. | 1998 | Spastic paraparesis–konzo in the Garoua Boulai Health District, East Province– Cameroon: A hidden Hydrogen cyanide and cyanides: Human health aspects 37 | N | n/a |
| A6.12 | Lasch E.E., El Shawa R. | 1981 | Multiple cases of cyanide poisoning by apricot kernels in children from Gaza. Pediatrics 68:5-7. | N | n/a |
| A6.12 | Leeser J.E., Tomenson JA, Bryson DD | 1990 | A cross-sectional study of the health of cyanide salt production workers. Macclesfield, ICI Central Toxicology Laboratory. | N | n/a |
| A6.12 | Liebowitz D., Schwartz H. | 1948 | Cyanide poisoning: Report of a case with recovery. Am J Clin Pathol 18: 965-970 | N | n/a |
| A6.12 | Lundquist P., Rammer L, Sorbo B | 1989 | The role of hydrogen cyanide and carbon monoxide in fire casualties: a prospective study. Forensic Science International, 43:9–14. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|---|----------------------------------|-------|
| A6.12 | Makene W.J., Wilson J. | 1972 | Biochemical studies in Tanzanian patients with ataxic tropical neuropathy. J Neurol Neurosurg Psychiatry 35:31-33 | N | n/a |
| A6.12 | McGlashan N.D. | 1967 | Geographical evidence on medical hypotheses. <i>Tropical and Geographical Medicine</i> , 19:333–344. | N | n/a |
| A6.12 | Ministry of Health, Mozambique. | 1984 | Mantakassa: An epidemic of spastic paraparesis associated with chronic cyanide intoxication in a cassava staple area of Mozambique. 1. Epidemiology and clinical and laboratory findings in patients. Bull WHO 62:477-484 | N | n/a |
| A6.12 | Mittal S.; Gupta, K. S. and Gupta, Y. K.: | 1982 | Kinetics & Mechanism of Acid Hydrolysis of Formamide, Acetamide, Propanamide & Butanamide over an Extended Concentration Range: Kinetic Evidence for Fast Protonation Pre-equilibrium, <i>Indian J. Chem.</i> , 21A, 357–360. | N | n/a |
| A6.12 | Mittal, S.; Gupta, K. S. and Gupta, Y. K | 1981 | Kinetics of Carboxylic Acid Catalysed Hydrolysis of Formamide: Evidence for Specific Hydronium Ion Catalysis, <i>Indian J. Chem</i> , 20A, 1220–1221. | N | n/a |
| A6.12 | Miyakawa, S.; Cleaves, H. J. | 2001 | : Implications Based on the Hydrolytic Stabilities of Hydrogen Cyanide and Formamide, <i>Journal of Applied Chemistry</i> , 196 | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|---|----------------------------------|-------|
| A6.12 | Nahrstedt A.F. | 1993 | Cyanogenesis and food plants. In: van Beek TA, Breteler H, eds. <i>Proceedings of the International Symposium on Phytochemistry and Agriculture, 22–24 April 1992, Wageningen</i> . Oxford, Oxford University Press, pp. 107–129. | N | n/a |
| A6.12 | NIOSH | 1976 | Health hazard evaluation report. Cincinnati, OH: U.S. Department of Health, Education, and Welfare, Center for Disease Control, National Institute for Occupational Safety and Health. No. 74-129-268 | N | n/a |
| A6.12 | Okafor P.N., Okorowko CO, Maduagwu EN | 2002 | Occupational and dietary exposures of humans to cyanide poisoning from large-scale cassava processing and ingestion of cassava foods. Food and Chemical Toxicology, 49:1001–1005. | N | n/a |
| A6.12 | Oluwole OSA, Onabolu AO, Cotgreave IA, Rosling H, Persson A, Link H | 2002 | Low prevalence of ataxic polyneuropathy in a community with high exposure to cyanide from cassava foods. <i>Journal of Neurology</i> , 249:1034–1040. | N | n/a |
| A6.12 | Osuntokun BO | 1981 | Cassava diet, chronic cyanide intoxication and neuropathy in the Nigerian Africans. <i>World Review of Nutrition and Dietetics</i> , 36:141–173 | N | n/a |
| A6.12 | Peden NR, Taha A, McSorley PD, et al. | 1986 | Industrial exposure to hydrogen cyanide: Implications for treatment. Br Med J 293:538. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|--|----------------------------------|-------|
| A6.12 | Pentore R, Venneri A, Nichelli P | 1996 | Accidental choke cherry poisoning: early symptoms and neurological sequelae of an unusual case of cyanide intoxication. <i>Italian Journal of Neurological Science</i> , 17:233–235. | N | n/a |
| A6.12 | Pijoan M | 1942 | Cyanide poisoning from choke cherry seed. American Journal of Medical Science, 204:550. | N | n/a |
| A6.12 | Rabinovitch, B. S. and Winkler, C. A.: | 1942 | The Hydrolysis of Aliphatic Nitriles in Concentrated Hydrochloric Acid Solutions, <i>Canad. J. Res.</i> , 20B, 221–230. | N | n/a |
| A6.12 | Rieders F | 1971 | Noxious gases and vapors. I: Carbon monoxide, cyanides, methemoglobin, and sulfhemoglobin. In: De | N | n/a |
| A6.12 | Sayre JW, Kaymakcalan S | 1964 | Cyanide poisoning from apricot seeds among children in Central Turkey. New England Journal of Medicine, 270:. | N | n/a |
| A6.12 | Singh BM, Coles N, Lewis P, et al. | 1989 | The metabolic effects of fatal cyanide poisoning. Postgrad Med J 65:923-925 | N | n/a |
| A6.12 | Suchard JR, Wallace KL, Gerkin RD | 1998 | Acute cyanide toxicity caused by apricot kernal ingestion. Ann Emerg Med 32(6):742-744. | N | n/a |
| A6.12 | Sullivan, J.B. Jr., G.R. Krieger | 1992 | (eds.). Hazardous Materials Toxicology- Clinical Principles of Environmental Health. Baltimore, MD: Williams and Wilkins, p. 704 | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|---|----------------------------------|-------|
| A6.12 | Swai AB, McLarty DG, Mtinangi BL, Tatala S, Kitange HM, Mlingi N, Rosling H, Howlett WP, Brubaker GR, Alberti KG | 1992 | Diabetes is not caused by cassava toxicity. A study in a Tanzanian community. <i>Diabetes Care</i> , 15:1378–1385 | N | n/a |
| A6.12 | Tylleskar T, Banea M, Bikangi N, et al. | 1992 | Cassava cyanogens and konzo, an upper motoneuron disease found in Africa [erratum in Lancet 1992 Feb 15;339(8790):440]. Lancet 339(8787):208-211. | N | n/a |
| A6.12 | Tylleskar T, Legue FD, Peterson S, et al. | 1994 | Konzo in the Central African Republic. Neurology 44(5):959-61 | N | n/a |
| A6.12 | VanderLaan WP, Bissell A. | 1946 | Effects of propylthiouracil and of potassium thiocyanate on the uptake of iodine by the thyroid gland of the rat. Endocrinology 39:157-160. | N | n/a |
| A6.12 | Wexler J, Whittenberger JL, Dumke PR. | 1947 | The effect of cyanide on the electrocardiogram of man. Am Heart J 34:163-173 | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|--|----------------------------------|-------|
| A6.12 | WHO | 1985 | (1985) Diabetes mellitus. Report of a WHO Study Group. Geneva, World Health Organization, 131 pp. Available at http://whqlibdoc.who.int/trs/WHO_TRS_727 . pdf (WHO Technical Report Series 727) | N | n/a |
| A6.12 | Zuidema PJ | 1959 | Cirrhosis and disseminated calcification of the pancreas in patients with malnutrition. Tropical and Geographical Medicine, 11:70–74. | N | n/a |
| A6.2 A6.12 | Aminlari et al., Aminlari M, Vaseghi T, Karpar MA | 1994 | The cyanide-metabolizing enzyme rhodanese in different parts of the raspiratory systems in sheep and dog. <i>Toxicology and Applied Pharmacology</i> , 124: 64-71 | N | n/a |
| A6.2 | Ansell & Lewis, | 1970 | A review of cyanide concentrations found in human organs: A survey of literature concerning cyanide metabolism, "normal", non-fatal and fatal bydy cyanide levels. Journal of Forensic Medicine, 17: 148-155 | N | n/a |
| A6.2, | Williams, 1959 – Williams RT | 1959 | Detoxification mechanisms, 2nd ed. London, Chapman and Hall, p. 393 | N | n/a |
| A6.2, A6.12 | Dahl | 1989 | The cyanide-metabolizing enzyme rhodanese in rat nasal respiratory and olfactory mucosa. <i>Toxicology Letters</i> , 45: 199-205 | N | n/a |
| A6.3.1 | Olusi SO, Oke OL, Odusote A | 1979 | Effects of cyanogenic agents on reproduction and neonatal development in rats. <i>Biology of the Neonate</i> , 36:233–234. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|--|----------------------------------|-------|
| A6.3.1 | Pritsos CA. | 1996 | Mitochondrial dysfunction and energy depletion from subchronic peroral exposure to cyanide using the Wistar rat as a mammalian model. Toxic Subst Mech 15(3):219-229. | N | n/a |
| A6.3.2 | Hugod C. | 1981 | Myocardial morphology in rabbits exposed to various gas-phase constituents of tobacco smoke: an ultrastructural study. Atherosclerosis 40: 181 - 190. | N | n/a |
| A6.7, A6.8.1 | Doherty P.A., Ferm VH, Smith RP. | 1982 | Congenital malformations induced by infusion of sodium cyanide in the Golden hamster. Toxicol Appl Pharmacol 64:456-464. | N | n/a |
| A6.8.1 | Frakes R.A., Sharma RP, Willhite CC, Gomez G | 1986 | Effect of cyanogenic glycosides and protein content in cassava diets on hamster prenatal development. Fundamental and Applied Toxicology, 7:191–198 | N | n/a |
| A6.9 | Hertting G.O., Kraupp E, Schnetz E, Wuketich ST | 1960 | Investigation about the consequences of a chronic administration of acutely toxic doses of sodium cyanide to dogs. <i>Acta Pharmacologica et Toxicologica</i> , 17:27–43 | N | n/a |
| A6.9 | Lessell S. | 1971 | Experimental cyanide optic neuropathy. Arch Ophthalmol 86:194-204 ATSDR | N | n/a |
| A7,4:2 | EPA | 1992 | U.S. Environmental Protection Agency. Fed Regist 57:26248. | N | n/a |
| A7.1.1.1 | Colt A. W.; Walton, J. H. | 1937 | The Reaction of Hydrogen Cyanide with Sulfuric and Phosphoric Acids, <i>J. Phys. Chem.</i> , , 41, 351. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|--|----------------------------------|-------|
| A7.1.1.1.1 | Hine J.; King, R. SM.; Midden, W. R. and Sinh, | 1981 | A.: Hydrolysis of Formamide at 80°C and pH 1–9, <i>J. Org. Chem.</i> , 46, 3186–3189. | N | n/a |
| A7.1.1.1.1 | Krieble V. E. and McNally, J. G | 1929 | The Hydrolysis of Hydrogen Cyanide by Acids I, J. Am. Chem, Soc., 51, 3368. | N | n/a |
| A7.1.1.1.1 | Krieble, V. E. and McNally, J. G | 1933 | The Hydrolysis of Hydrogen Cyanide by Acids II, J. Am. Chem, Soc., , 55, 2326. | N | n/a |
| A7.1.1.1.1 | Krieble, V. E. and McNally, J. G.: | 1943 | The Hydrolysis of Hydrogen Cyanide in Acetic Acid Solutions with Mineral Acids as Catalysts, <i>J. Am. Chem, Soc.</i> , 65, 1479. | N | n/a |
| A7.1.1.1.1 | Marsh J. D. F. and Martin, M. J | 1957 | The Hydrolysis and Polymerization of Hydrogen Cyanide in Alkaline Solutions, <i>J. Appl. Chem.</i> , 7, 205–209. | N | n/a |
| A7.1.1.1.1 | Salem, S. M. and Sidahmed, I. M | 1985 | Solvent Effect on the Kinetic Study of the Alkaline Hydrolysis of Formamide in Acetone–Water Mixtures, <i>J. Chin. Chem. Soc.</i> , 32, 451–456. | N | n/a |
| A7.1.1.1.1 | Salem, S. M. and Sidahmed, I. M | 1986 | .: The Acid Hydrolysis of Formamide in Water-Acetone Mixtures, <i>Egypt. J. Chem.</i> , 29, 521–528. | N | n/a |
| A7.1.1.1.1 | Sanchez, R. A.; Ferris, J. P. and Orgel, L. E | 1967 | .: Studies in Prebiotic Synthesis II. Synthesis of Purine Precursors and Amino Acids from Aqueous Hydrogen Cyanide, <i>J.Mol. Biol.</i> , 30, 223–253. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|---|----------------------------------|-------|
| A7.1.1.1.1 | Skundric, B. and Penavin, J | 1984 | .: Acid Catalysed Amide Hydrolysis in Water- Ethanol Mixtures. Medium Interactions Study, Zeitschrift Physikalische Chemie Neue Folge, 141, 29-31. | N | n/a |
| A7.1.1.1.1 | Tan, T. C. and Teo, W. K.: | 1987 | Destruction of Cyanides by Thermal Hydrolysis, <i>Plat. and Surf. Fin.</i> 74 (4), 70–73. | N | n/a |
| A7.1.1.1.1 | White, J.M., Jones, D.D., Huang, D., et al. | 1988 | Conversion of cyanide to formate and ammonia by a pseudomonad obtained from industrial wastewater. J Indust Microbiol 3:263-272. | | n/a |
| A7.1.1.1.1 | Wiegand, G. H. and Tremelling, M. | 1972 | The Kinetics and Mechanism of the Decomposition of KCN in Aqueous Alkaline Medium Hydrolysis of Simplest Nitrile, HCN, <i>J. Org. Chem.</i> , 37, 914. | N | n/a |
| A7.1.1.1.2 | Abbas, M., Guo, J., Carli, B., Mencaraglia, F., Carlotti, M., and Nolt, I | 1987 | Stratospheric distribution of HCN from far infrared observations, Geophys. Res. Lett., 14, 531–534, | N | n/a |
| A7.1.1.1.2 | Anderson, D | 1983 | The troposphere-stratosphere radiation-field at twilight – A spherical model, - Planet. Space Sci., 31, 1517–1523 | N | n/a |
| A7.1.1.1.2 A7.3.1 | Cicerone, R. and Zellner, R. | 1983 | The atmospheric chemistry of hydrogen-cyanide (HCN), J. Geo-phys. Res., 88, 689–696, | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|---|------|--|----------------------------------|-------|
| A7.1.1.2 | DeMore W. B., Howard, C. J., Sander, S. P., Ravishankara, A. R., Golden, D. M., Kolb, C. E., Hampson, R. F., Molina, M. J., and Kurylo, M. J. | 2000 | Chemical Kinetics and Photochemical Data for Use in Stratospheric Modeling, Supplement to Evaluation 12: Update of Key Reactions, JPL Publ. 00-3 | N | n/a |
| A7.1.1.2.1 | | | Reviews of the environmental effects of pollutants. V. Cyanide. Cincinnati, OH: U. S. Environmental Protection Agency Health Effects Research Laboratory, Office of Research and Development PB289920. | N | n/a |
| A7.1.1.2.1 | Akcil, A, Mudder, T. | 2003 | Microbial destruction of cyanide wastes in gold mining: Process review. Biotechnol Lett 25:445-450. | N | n/a |
| A7.1.1.2.1 | Chapatwala, K.D., Babu, G.R.V., Wolfram, J.H., | 1993 | Screening of encapsulated microbial cells for the degradation of inorganic cyanides. J Ind Microbiol 11(2):69-72. | N | n/a |
| A7.1.1.2.1 | Gaudy, A.F, Gaudy, E.T, Feng,Y.J, et al. | 1992 | U.S. Environmental Protection Agency. Fed Regist 57:26248.7 1982. Treatment of cyanide waste by the extended aeration process. J Water Pollut Control Fed 54:153-164. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|--|----------------------------------|-------|
| A7.1.1.2.1 | Kunz D.A., Nagappan, O., Silva-Avalos, J., et al. | 1992 | Utilization of cyanide as a nitrogenous substrate by <i>Pseudomonas fluorescens</i> NCIMB 11764: Evidence for multiple pathways of metabolic conversion. Appl Environ Microbiol 58(6):2022-2029. | N | n/a |
| A7.1.1.2.1 | Ludzack F.J., Moore, W.A., Krieger, H.L., et al. | 1951 | Effect of cyanide on biochemical oxidation in sewage and polluted water. Sewage Ind Wastes 23:1298-1307. | N | n/a |
| A7.1.1.2.1 | Malaney G.W., Sheets WD, Quillin R. | 1959 | Toxic effects of metallic ions on sewage microorganisms. Sewage Ind Wastes 31:1909-1915. | N | n/a |
| A7.1.1.2.1 | Meyers, P.R., Rawlings, D.E., Woods, D.R., et al. | 1993 | Isolation and characterization of a cyanide dihydratase from <i>Bacillus pumilus</i> C1. J Bacteriol 175(19):6105-6112. | N | n/a |
| A7.1.1.2.1 | Pettet, A.E.J, Mills, E.V. | 1954 | Biological treatment of cyanides with and without sewage. J Appl Chem 4:434-444. | N | n/a |
| A7.1.1.2.1 | Raef SF, Characklis WG, Kessick MA, et al | 1977 | Fate of cyanide and related compounds in aerobic microbial systemsII. Microbial degradation. Water Res 11:485-492. | N | n/a |
| A7.1.1.2.1 | Raybuck, S.A., | 1992 | Microbes and microbial enzymes for cyanide degradation. Biodegradation 3(1):3-18. | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|--|----------------------------------|-------|
| A7.1.1.2.1 | Richards, D.J., Shieh, W.K | 1989 | Anoxic-oxic activated-sludge treatment of cyanides and phenols. Biotechnol Bioeng 33:32-38. | N | n/a |
| A7.1.1.2.1 | Shivaraman, N., Kumaran, P., Pandey, R.A., et al. | 1985 | Microbial degradation of thiocyanate, phenol and cyanide in a completely mixed aeration system. Environ Pollut Ser A 39:141-150. | N | n/a |
| A7.1.1.2.1 | Silva-Avalos, J., Richmond, M.G., Nagappan, O., et al. | 1990 | Degradation of the metal-cyano complex tetracyanonickelate (II) by cyanide-utilizing bacterial isolates. Microbiol 56:3664-3670. | N | n/a |
| A7.1.1.2.1 | US EPA | 1978 | Reviews of the environmental effects of pollutants. V. Cyanide. Cincinnati, OH: U.S. Environmental Protection Agency Health Effects Research Laboratory, Office of Research and Development. PB289920. | N | n/a |
| A7.1.3 | Gaffney J.S. et al | 1987 | Environ Sci Technol 21: 519-23 (1987) | N | n/a |
| A7.1.3 | Roy WR | 1994 | ; Groundwater Contamination From Municipal Landfills in the USA. in Contam Groundwaters, Adriano DC et al eds. Sci Rev Northwood, UK | | n/a |
| A7.3.1 | Callahan, M.A., M.W. Slimak, N.W. Gabel, et al. | 1979 | Water-Related Environmental Fate of 129 Priority Pollutants. Volume I. EPA-440/4 79- 029a. Washington, DC: U.S. Environmental Protection Agency, | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|------------------------|------|---|----------------------------------|-------|
| A7.3.1 | EPA | 1984 | Health effects assessment for cyanide. Washington, DC: U.S. Environmental Protection Agency. EPA540186011. | N | n/a |
| A7.3.1, A7.4.2 | ЕРА | 1979 | Cyanides. In: Water-related environmental fate of 129 priority pollutants. Vol. 1. Washington, DC: U.S. Environmental Protection Agency, Office of Water Planning and Standards, Office of Water and Waste Management. EPA440479029a. PB80204373. 12-1-12-12. | N | n/a |
| A7.4.2 | EPA | 1978 | Reviews of the environmental effects of pollutants. V. Cyanide. Cincinnati, OH: U.S. Environmental Protection Agency Health Effects Research Laboratory, Office of Research and Development. PB289920. | N | n/a |
| A7.4.2 | ЕРА | 1980 | Water quality criteria documents: Availability. U.S. Environmental Protection Agency. Fed Regist 45:79318-79379. | N | n/a |
| A7.4.2 | EPA | 1985 | Ambient water quality for cyanide - 1984. Washington, DC: Office of Water Regulations and Standards, Criteria and Standards Division. EPA440584028. PB85227460. | N | n/a |
| A7.4.2 | Franke C. et al; | 1994 | Chemosphere 29: 1501-14 | N | n/a |
| A7.4.2 | v Meylan W.M. et al | 1999 | Environ. Toxicol. Chem. 18: 664-72 | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|--|----------------------------------|-------|
| IIA section 3.1 | E.H. Jeffery, M.A Walling, M.e. Tumbleson | 2002 | Nutritional toxicological pathology in Handbook of Toxicologic Pathology; Editors: W.A. Haschek and C.E. Rousseaux ; Volume 1, Second Edition ,Academic Press Second Edition | N | n/a |
| IIA section 3.1 | J. Van Sande, C. Massart, R. Beauwens, A. Schoutens, S. Costagliola , J.E. # Dumont, J. Wolff, | | Anion selectivity by the sodium iodide symporter, Endocrinology 144 (2003) 247–252. | N | n/a |
| IIA section 3.1 | Li, J. Zhang, Z. Li, | | Prevention of iodine deficiency in high fluoride areas in Tianjin City, China, Fluoride (1998) 18. | N | n/a |
| IIA section 3.1 | Lisandro Irizarry, Nadine A Youssef, Anton A Wray | | Toxicity, Thyroid Hormone: Treatment & Medication in eMedicine Specialties #(http://emedicine.medscape.com/article/8196 92-overview) | N | n/a |

| Section No DOC IIIA | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|------------------------|--|------|---|----------------------------------|-------|
| IIA section 3.1 | Pablo Enrique Pedraza, Maria-Jesus Obregon, Hector Francisco Escobar- Morreale, Francisco Escobar del Rey, and Gabriella Morreale de Escobar | | Mechanisms of Adaptation to Iodine Deficiency in Rats: Thyroid Status Is Tissue Specific. Its Relevance for Man Endocrinology 175(5):2098–2108 | N | n/a |
| IIA section 3.1 | V F H Brauer, H Below1, A Kramer1, D Führer and R Paschke | 2006 | The role of thiocyanate in the etiology of goiter in an industrial metropolitan area; European Journal of Endocrinology, Vol 154, Issue 2, 229-235 Copyright © by European Society of Endocrinology | N | n/a |

| Reference No DOC IV Section No DOC III | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|--------------|------|---|----------------------------------|-------|
| B3.4 B3.6 | | | Hazardous Substance Data Bank (HSDB), National Library of Medicine's TOXNET system (state in February 2006): Hydrogen cyanide *Peer reviewed* | N | n/a |
| DOC IV A12 B6.4 B6.4.a | J.M.McNerney | | M.P.H., H.H.Schrenk, PhD., 1960, The Acute Toxicity of Cyanogen, Industrial Hygiene Foundation, 4400 Fifth Avenue, Pittsburg 13, Pennsylvania, Industrial Hygiene Journal, April 1960, 121 – 124 | N | n/a |
| DOC IV A15 B6.4 B6.4.c | A. Fairley | | E.C.Linton, F.E.Wild , The Absorption of Hydrocyanic Acid Vapour through the Skin (with notes on other matters relating to acute cyanide poisoning), Journal of Hyg., Volume 34, October 1934, No. 3: 283 - 294 | N | n/a |
| B3.1.1 B3.1.2 B3.2 B3.6 | | | ATSDR 2004 - Agency for Toxic Substances and Disease Registry, U.S. Department of Health and Human Services: Draft Toxicological Profile for Cyanide, Sept. 2004, * Peer Reviewed*, str 141 | N | n/a |

| Reference No DOC IV Section No DOC III | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|-------------|------|---|----------------------------------|-------|
| DOC IV A20 B3.6 | | | HYDROGEN CYANIDE AND CYANIDES: HUMAN HEALTH ASPECTS Published under the joint sponsorship of the United Nations Environment Programme, the International Labour Organization, and the World Health Organization, and produced within the framework of the Inter-Organization Programme for the Sound Management of Chemicals. | N | n/a |
| B6.4 B6.4.b | D.C. Walton | | M.G. Witherspoon 1926. Skin absorption of certain gases. J Pharmacol Exp Ther 26: 315-324 | N | n/a |
| DOC IV B1 5_10_2a_PT08 | E.A. Parkin | 1937 | The toxicity of hydrogen cyanide to certain wood-boring insects. E.A. Parkin, J.R. Busvine. Ann. Appl. Biol., 24:131-143 | N | n/a |
| DOC IV B2 5_10_2b_PT08 | | | Research Report, Study of Hydrogen cyanide penetration into wood implex 2007, Ministry of the Interior Central Office of Fire Rescue Service of the Czech Republic, Institute of Population Protection | N | n/a |
| DOC IV B3 5_10_2c_PT08 | | 2009 | Crop Research Institute (CRI) Evaluation of URAGAN (HCN) Field Efficacy - CRI – 2009 | Y | |

| Reference No DOC IV Section No DOC III | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|-----------|------|---|----------------------------------|-------|
| DOC IV B4 5-10-2a-PT18 5-10-2_PT18 | | 2007 | Crop Research Institute (CRI) Evaluation of URAGAN (HCN) Field Efficacy - CRI - 2007 | Y | |
| DOC IV B5 5-10-2b_PT18 5-10-2_PT18 | | | M. RAMBEAU, D. BENITEZ, S. DUPUIS, P. DUCOM HYDROGEN CYANIDE AS AN IMMEDIATE ALTERNATIVE TO METHYL BROMIDE FOR STRUCTURAL FUMIGATIONS Laboratoire National d'Etudes des Techniques de Fumigation et de Protection des Denrées Stockées | N | n/a |
| DOC IV B6 PT08 | | 2011 | Timber-wood research and development institute, Prague, state enterprise DETERMINATION OF THE ERADICANT ACTION OF URAGAN D2 AGAINST THE LARVAE OF DOMESTIC LONGHORN BEETLE (HYLOTRUPES BAJULUS), ACCORDING TO ČSN EN 1390 | Y | |
| DOC IV B7 PT08 | | 2011 | Timber-wood research and development institute, Prague, state enterprise DETERMINATION OF THE ERADICANT ACTION OF URAGAN D2 AGAINST THE EGGS AND GROWN DOMESTIC LONGHORN BEETLE (HYLOTRUPES BAJULUS) | Y | |

| Reference No DOC IV Section No DOC III | Author(s) | Year | Title. Source (where different from company) Company, Report No. GLP (where relevant) / (Un)Unpublished | Data Protection Claimed (Yes/No) | Owner |
|--|-------------------|------|---|----------------------------------|-------|
| B3.1.1 B3.1.2 | | | Budavari S, ed. 1989, Merck index: An encyclopedia of chemicals, drugs, and biologicals. 11 th ed. Rahway, NJ: Merck &Co., Inc. | N | n/a |
| B3.2 B3.4 B3.6 | Jenks W R. | | Cyanides. In: Grayson, M, ed. Kirk-Othmer encyclopedia of chemical technology. New York, NY: John Wiley and Sons, Inc., 307-334, 1979 | N | n/a |
| B3.2 B3.4 | | | Fire Protection Guide to Hazardous Materials. 12 ed. Quincy, MA: National Fire Protection Association, 1997 | N | n/a |
| B3.6 | Lide, D.R. (ed.). | | CRC Handbook of Chemistry and Physics. 79th ed. Boca Raton, FL: CRC Press Inc., 1998-1999., p. 3-197 | N | n/a |