- Dossier preparation manuals
- Create support request
- <u>IUCLID user community</u>
- Additional information
- Video tutorials

SuperUser EPA/ORD/CCTE/SCDCD

- User Settings
- Logout
- Dashboard
- Cesium formate

Filtered aggr_1

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View Dossiers Validate

- Export to i6z
- Extract to dataset
- Create component PDF/RTF
- Create document PDF/RTF
- Compare
- Generate report
- Dissemination preview

Type at least 3 characters

REACH Complete

Cesium formate

- 1 General information
- 2 Classification & Labelling and PBT assessment
- 3 Manufacture, use and exposure
- 4 Physical and chemical properties
- 5 Environmental fate and pathways 10
- 6 Ecotoxicological information 24
 - 7 Toxicological information
- 32 o 166f589c-b5c3-4839-86b0-58598d07c238
 - o 7.1 Toxicokinetics, metabolism and distribution
 - 7.2 Acute Toxicity

```
7.3 Irritation / corrosion
            4

    7.4 Sensitisation

            2
         • 7.5 Repeated dose toxicity
               • 7.5.1 Repeated dose toxicity: oral
                     ■ 3557fc51-303f-4021-af2d-
                        2dd83aaedf5e
                     4e1a78b6-ba19-440b-8027-
                        3377f04dcb28
                     ■ 00a1aef3-4337-4d0d-9261-
                        713b4a7297e3
                     • e48c889e-667e-47f8-9a83-
                        6b7f80d2b71e
                     ■ 07582398-b407-4cc2-8729-
                        def9c53ac76b
                     ■ 31c3911e-0a44-4d3c-8a13-
                        de5a3f0acaa5
               • 7.5.2 Repeated dose toxicity: inhalation
               • 7.5.3 Repeated dose toxicity: dermal
               • 7.5.4 Repeated dose toxicity: other
                 routes

    7.6 Genetic toxicity

    7.7 Carcinogenicity

    7.8 Toxicity to reproduction

    7.9 Specific investigations

         • 7.10 Exposure related observations in humans
         • 7.11 Toxic effects on livestock and pets
         • 7.12 Additional toxicological information
   • 8 Analytical methods
     9 Residues in food and feedingstuffs
   • 10 Effectiveness against target organisms
   • 11 Guidance on safe use
     12 Literature search
     13 Assessment reports
   • 14 Information requirements
   • Inherited templates
UUID 00a1aef3-4337-4d0d-9261-713b4a7297e3 Hide empty fields
       Compare Document
            0
       1
 Administrative data
                          Data source
                                          Materials and methods
                                                                      Results and discussion
                                                                                                  Overall remarks, attachments
 Applicant's summary and conclusion
```

4

| Administrative data |
|---|
| |
| Endpoint short-term repeated dose toxicity: oral |
| Type of information experimental study |
| Adequacy of study supporting study |
| Robust study summary |
| Used for classification |
| Used for SDS |
| Study period |
| Reliability 1 (reliable without restriction) |
| Rationale for reliability incl. deficiencies |
| Data waiving |
| Justification for data waiving |
| Justification for type of information |
| Attached justification |
| # Attached justification Reason / purpose Actions |
| Cross-reference |
| # Reason / purpose for cross-reference Related information Remarks Actions |
| Data source |
| Reference |
| • study report Unnamed 2008 2008-09-03 |
| Data access |
| Data protection claimed |
| Materials and methods |
| Test guideline |
| # Qualifier Guideline Version / remarks Deviations Actions 1 Qualifier according to guideline |

OECD Guideline 407 (Repeated Dose 28-Day Oral Toxicity Study in Rodents)

Version / remarks

Deviations

| Frequency of treatment |
|---|
| Doses / concentrations |
| # Dose / conc. Remarks Actions |
| No. of animals per sex per dose |
| Control animals |
| Details on study design |
| Positive control |
| Examinations |
| Observations and examinations performed and frequency |
| Sacrifice and pathology |
| Optional endpoint(s) |
| Other examinations |
| Statistics |
| Any other information on materials and methods incl. tables |
| Results and discussion |
| Results of examinations |
| Clinical signs |
| Description (incidence and severity) |
| Mortality |
| Description (incidence) |
| Body weight and weight changes |
| Description (incidence and severity) |
| Food consumption and compound intake (if feeding study) |
| Description (incidence and severity) |
| Food efficiency |
| Description (incidence and severity) |
| Water consumption and compound intake (if drinking water study) |
| Description (incidence and severity) |
| Ophthalmological findings |
| Description (incidence and severity) |
| Haematological findings |

| Description (incidence and severity) |
|---|
| Clinical biochemistry findings |
| Description (incidence and severity) |
| Endocrine findings |
| Description (incidence and severity) |
| Urinalysis findings |
| Description (incidence and severity) |
| Behaviour (functional findings) |
| Description (incidence and severity) |
| Immunological findings |
| Description (incidence and severity) |
| Organ weight findings including organ / body weight ratios |
| Description (incidence and severity) |
| Gross pathological findings |
| Description (incidence and severity) |
| Neuropathological findings |
| Description (incidence and severity) |
| Histopathological findings: non-neoplastic |
| Description (incidence and severity) |
| Histopathological findings: neoplastic |
| Description (incidence and severity) |
| Other effects |
| Description (incidence and severity) |
| Details on results |
| Effect levels |
| # Key result Dose descriptor Effect level Based on Sex Basis for effect level Remarks on result Actions 1 Key result Dose descriptor NOAEL Effect level ca. 15 mg/kg bw/day (nominal) Based on Sex male/female Basis for effect level • other: Remarks on result |

| # Key result Critical effects observed Lowest effective dose / conc. System Organ Treatment related Dose response relationship Relevant for humans Actions 1 | |
|--|---|
| Key result | |
| Critical effects observed | |
| yes | |
| Lowest effective dose / conc. | |
| 15 mg/kg bw/day (nominal) | |
| System | |
| haematopoietic | |
| Organ | |
| • not specified | |
| Treatment related | |
| yes | |
| Dose response relationship | |
| yes | |
| Relevant for humans | |
| not specified | |
| Any other information on results incl. tables | |
| Overall remarks, attachments | |
| Overall remarks | _ |
| Attachments | |
| # Type Attached (confidential) document Attached (sanitised) documents for publication Remarks Actions | |
| Illustration (picture/graph) | |
| Applicant's summary and conclusion | |
| Conclusions | _ |
| Executive summary | |
| TOP | |
| | |
| | |
| | |
| Dashboard | |
| Cubatanaca | |
| Substances | |
| Mixtures / Products | |
| | |
| Articles | |
| • | |
| Categories | |
| Toolbox | |
| • Template | |
| Manage Reports | |

Inventory manager

- ContactLegal entity
- Sites

- Reference substance
- Test material
- Literature reference

User management

- User Settings Users
- Roles

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- About
- Help