*Chemical Safety Report (CSR)*

**USE:** The chemical safety report documents the chemical safety assessment undertaken as part of the REACH registration process, and is the key source from which the registrant provides information to all users of chemicals through the exposure scenarios. It also forms a basis for other REACH processes including substance evaluation, authorisation and restriction.

*EU Biocidal Products Regulation (BPR) Summary of Product Characteristics (SPC)*

**USE:** A Summary of Product Characteristics (SPC) is mandatory for all companies wishing to apply for product authorisations. In order to support companies in preparing their applications for the authorisation of their biocidal products, either on a national or European level, the report extracts SPC-related information for single products from IUCLID, which can then be imported directly into the [SPC Editor](https://echa.europa.eu/support/dossier-submission-tools/spc-editor).

*EU Biocidal Products Regulation (BPR) Confidentiality Report (BPR Article 66)*

**USE:** ECHA and the MSCA’s work together to evaluate the confidentiality claims made on biocidal products and active substances as part of their work for the Biocidal Product Regulation. The report supports ECHA and MSCAs during the evaluation of these confidentiality claims by listing all the relevant confidentiality claims which are to be assessed. The completed assessment is sent to ECHA via the R4BP messaging system <https://idp-authority.echa.europa.eu/idp/>.

*Literature References report for substance and mixture/product datasets and dossiers [RTF]*

**USE:** **Industry registrants** applying for a new product authorisation under the BPR regulation are required to attach a Product Authorisation Report (PAR) to their submission. This report allows registrants to extract all their literature references from a Mixture/Product dataset or dossier, and re-use these as part of their PAR as well as to quality check the study information they have included in the Mixture/Product dataset. **MSCAs** also use the report to check if all studies that should be included, are included in the registrant’s dossier. Note: the report can also be generated from a Substance dataset or dossier if needed.

*Literature References report for substance and mixture/product datasets and dossiers [CSV]*

**USE:** Under Article 10(a) of REACH, a registrant is required to have legitimate access to the full study they use or refer to, in their registration dossier. To support this process, the ‘Study report’ provides a table of all Literature references contained within a dataset or dossier, and provides additional study information on top of this, such as the Test Material Information and the Adequacy of the Study. The report has been specifically used to support cost sharing in registration consortia and to exchange study information between registrants. Note: the report can also be generated from a Mixture/Product.

*EU Biocidal Products Regulation (BPR) List of attachments for mixture/product datasets and dossiers [RTF]*

**USE:** During the evaluation and assessment of dossiers submitted under the Biocidal Products Regulation, registrants attach supporting information (in standalone documents) to individual IUCLID endpoints. To support this work, the report produces a table of all attachments inside the main Mixture/Product dataset or dossier, including all Substance and Mixture/Product components which are included in the composition of the main Mixture/Product.

*Table of attachments for Substances datasets and dossiers [RTF]*

**USE:** During the evaluation and assessment of dossiers submitted under REACH, registrants attach supporting information (in standalone documents) to individual IUCLID endpoints. To support this work, the report produces a table of all attachments inside a Substance dataset or dossier.

*REACH Classification and Labelling report (IUCLID section 2.1 GHS)*

**USE:** Under REACH Article 29(2)(b), one purpose of the Substance Information Exchange Forum (SIEF) is to agree on classification and labelling inside the joint submission. This report contains the classification and labelling information a lead registrant sends to the members of a joint submission in order to receive comments and approval on the classification and labelling.

*EU Biocidal Products Regulation (BPR) table of annotations*

**USE:** Under the Biocidal Products Regulation evaluation and assessment process, MSCAs often create IUCLID annotations on top of specific endpoints inside dossiers. These annotations are shared internally amongst the MSCAs. This report supports this process by listing the annotations which are found in a Mixture/Product dataset or dossier, and the endpoints they are attached to.

*List of use information (created for and by DOW Chemicals)*

**USE:** Managing the use of a chemical is complex and is often done by particular IT systems, such as SAP. This report outputs a CSV (comma separated file) of use information from IUCLID, which can then be injected into an external system to help align the use information contained in IUCLID and other systems.

*Classification, Labelling and Packaging (CLP) Regulation PCN Preview report [PDF] & PCN dossier viewer [HTML]*

***USE:*** *Based on Article 45 and Annex VIII to the CLP Regulation, companies placing hazardous mixtures on the market have an obligation to provide information on these mixtures to appointed bodies. As part of this obligation, there are now a PCN Preview Report [PDF] and a PCN dossier viewer [HTML] which summarises that information. The reports are in-built inside IUCLID and can be launched from the web interface. The PDF report displays the information in one single file; the HTML report organises the information in tabs, making the navigation more user friendly. Both reports are printable.*

*EU Biocidal Products Regulation (BPR) Cross references report*

**USE:** Under the Biocidal Products Regulation, the report provides an overview of the Biocidal Product Family with regards to which product compositions (family or single) have been referenced by which endpoint study. The report includes, if available, data waiving information.

*SCIP Notification Preview report*

**USE:** Based on Article 9(1)(i) of the  Waste Framework Directive (WFD), any supplier of an article containing a substance of very high concern (SVHC) on the Candidate List in a concentration above 0.1% weight by weight (w/w) on the EU market is required to submit information on that article to ECHA .The SCIP Notification preview report summarises that information.

*SCIP Notification Preview without sensitive information*

**USE:** Based on Article 9(1)(i) of the  Waste Framework Directive (WFD), any supplier of an article containing a substance of very high concern (SVHC) on the Candidate List in a concentration above 0.1% weight by weight (w/w) on the EU market is required to submit information on that article to ECHA . ECHA will publish the information, as received, on its website. In order to ensure the protection of confidential business information, ECHA will not make publicly available the required mandatory data that would allow the establishment of links between actors in the same supply chain.  The SCIP Notification preview report without sensitive information, summarises the information reported in IUCLID for the specific article  without including the information defined in SCIP as sensitive information to ensure the protection of confidential business information. This report will support companies to review what is the information that will be potentially disseminated in the SCIP database. The report can also be used to share information between companies. More detail information about Dissemination and confidentiality available at the document “[Dissemination and confidentiality in the SCIP Database](https://echa.europa.eu/documents/10162/28213971/dissemination_confidentiality_scip_en.pdf/e0efbea1-d8ec-b67c-de8f-1838b480db6d)".