Intertek Health Environmental & Regulatory Services (HERS) supports its customers with the know-how to stay compliant with the ANNEX VIII of CLP-Regulation – ensuring the safe and regulatory compliant placement of mixtures on the European Market.

**Background**

In case of a poisoning caused by bites, medicines, plants or hazardous mixtures, the medical advice for citizens or healthcare professionals by Poison Centers is essential. Since 1988, Member States of the EU have to appoint a body (e.g., a Poison Centre), collecting relevant data about hazardous mixtures (and all other above-mentioned cases) to ensure a prompt reaction, supporting first aiders and medical staff in poisoning incidences. In the CLP-Regulation (EC No 1272/2008), Article 45 defines the notification obligations for the chemical industry and formulators of mixtures.

A review, carried out by the Commission, lead to the conclusion that there is considerable variation in the existing notification systems and the information requirements for each country in the EU. To align the Article 45 CLP-Regulation notification requirements ANNEX VIII of CLP Regulation was implemented in March 2017. ANNEX VIII harmonizes the information requested by the appointed bodies and the submission format.

Additionally, a Unique Formula Identifier (UFI) is introduced, to be affixed on the label, establishing a link between a product and the specific information submitted to the appointed body.

All these new requirements lead to new challenges for industry, importers and distributors – so, be prepared!

**Your Challenge**

Key challenges facing the industry, when hazardous mixtures are placed on the market:

- Analysis of your portfolio to define notification deadline for consumer/ professional and industrial use products
- Lack of information on Mixtures and Mixtures in Mixtures – data gathering
- Manage portfolio and assigned UFI Codes
- Missing time/staff to handle notification preparation & submission
- Keep notifications up to date or request new UFIs and submit new notifications

**Our Solutions**

HERS provides to clients the necessary regulatory and technical support to fulfill the obligations under Article 45 of CLP Regulation. Our regulatory experts analyze the portfolio and assist the clients in notifying their hazardous mixtures to the relevant timeline.

Our services include:

- Portfolio screening for uses & their notification timelines
- Portfolio clustering for group/single submissions
- Requesting & assigning of UFI Codes
- Screening for missing notification information, e.g., Mixture in Mixture information
- Data gathering by communication with supplier outside of the EU
- Prepare & submit notification
- Update notifications
- Consultancy on how to fulfill obligations best

**The Intertek Advantage**

Intertek’s scientific & regulatory consultants have been successfully delivering expert advice for over 30 years. We optimize success and minimize client risk through broad expertise and knowledge. We are ideally equipped to help clients achieve success in a fast-paced, changing global market.

Intertek is a leading Total Quality Assurance provider to industries worldwide. Our network of more than 1,000 laboratories and offices and over 44,000 people in more than 100 countries, delivers innovative and bespoke Assurance, Testing, Inspection and Certification solutions for our customers’ operations and supply chains. Intertek Total Quality Assurance expertise, delivered consistently with precision, pace and passion, enabling our customers to power ahead safely.

**FOR MORE INFORMATION**

Germany
+49 711 27311 196
info.hers@intertek.com

Italy
+39 051 0562930
chemicals.pharma.italy@intertek.com

France
+33 2 78 94 01 73
sante.beaute@intertek.com

United Kingdom
+44 (0) 116 263 9629
hersuk-regtox@intertek.com

intertek.com/reach/clp/