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Open Public Consultation on the Targeted Revision of the Regulation on Classification, Labelling and Packaging of Substances and Mixtures (CLP)

Fields marked with * are mandatory.

Introduction

The Regulation on the classification, labelling and packaging of substances and mixtures (http://publications.europa.eu/resource/cellar/e3f31046-b274-11eb-8aca-01aa75ed71a1.0013.02/DOC_1) (in short the CLP Regulation) covers almost all chemicals and products containing them, from industrial chemicals to house-hold ones, from fuels to pens, from solvents to detergents. For the purpose of this questionnaire, substances and mixtures are referred to as chemicals. The CLP Regulation aims to identify **hazards of chemicals**, such as causing cancer, disrupting aquatic life or causing allergy. Hazard identification relies on **scientific facts**. When hazards are identified for a chemical, products containing this chemical should be **labelled and/or packaged** before they are placed on the market. In addition to the hazard, labels also provide **advice on how to avoid and/or reduce exposure** to the hazardous chemical and how to deal with accidental exposure. Finally, the CLP regulation requires that **poison centres** receive information on the composition and hazards of chemicals to give the appropriate advice in case of poisoning accidents.

In other words, the first aim of the CLP Regulation is to **protect citizens and workers and the environment from dangerous substances and mixtures**. The second aim is to facilitate the **intra-EU exchange of chemicals** which can circulate freely within the European Internal Market when properly labelled and packaged according to the CLP criteria.

This public consultation will feed into the work of the European Commission in updating and improving the CLP Regulation, as pledged by the Commission in its 'Chemicals Strategy for Sustainability' (https://ec.europa.eu/environment/strategy/chemicals-strategy_en).

This questionnaire consists of **two sections**. This first section contains **general questions** to which all respondents are kindly invited to provide feedback. The second section focuses on **more technical points** of the CLP Regulation that requires prior knowledge and expertise.

About you

*Language of my contribution

English

*I am giving my contribution as

Business association

*First name

Laia

*Surname

Perez Simbor

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*Organisation name

255 character(s) maximum

ETRMA European Tyre and Rubber Manufacturers' Association

*Organisation size

Micro (1 to 9 employees)

Transparency register number

255 character(s) maximum

Check if your organisation is on the transparency register (<http://ec.europa.eu/transparencyregister/public/homePage.do?redir=false&locale=en>). It's a voluntary database for organisations seeking to influence EU decision-making.

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***Country of origin**

Please add your country of origin, or that of your organisation.

Belgium

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Only organisation details are published: The type of respondent that you responded to this consultation as, the name of the organisation on whose behalf you reply as well as its transparency number, its size, its country of origin and your contribution will be published as received. Your name will not be published. Please do not include any personal data in the contribution itself if you want to remain anonymous.

 Public

Organisation details and respondent details are published: The type of respondent that you responded to this consultation as, the name of the organisation on whose behalf you reply as well as its transparency number, its size, its country of origin and your contribution will be published. Your name will also be published.

I agree with the personal data protection provisions (<https://ec.europa.eu/info/law/better-regulation/specific-privacy-statement>)

Part I (general questions)

Question 0 - What is your level of knowledge of the following?

	Excellent knowledge	Good knowledge	Some knowledge	None
*The CLP regulation (legal text) and/or its implementation	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
*Chemical hazards	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>

Section 1 - New Hazard Classes

Following **new scientific evidence**, the Commission is considering introducing **new hazard classes** not currently covered by the CLP Regulation. This is expected to enhance the protection of human health and environment.

The European Commission has pledged to introduce an obligation for chemical producers and retailers to identify and explicitly label the following chemicals:

- **Endocrine disruptors.** Endocrine disruptors are chemicals that cause illness by interfering with the hormonal system of human beings or of wildlife (e.g. obesity of children, infertility, etc.);
- **Persistent, bio-accumulative and toxic chemicals.** These chemicals are not easily degraded in the environment, accumulate in wild plants and animals and are toxic to humans or plants or animals;
- **Persistent, mobile and toxic chemicals.** These chemicals are not easily degraded in the environment, pass from soil into water bodies and contaminate natural resources used to produce drinking water. They are also toxic to humans or plants or animals.

Those new obligations will complement existing requirements to identify hazards in chemicals.

Question 1 - Please indicate how important it is for you to know a chemical is ...?

(One single answer per row)

	Very important	Important	Not important	No opinion
*An endocrine disruptor with adverse effects on human health	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
*An endocrine disruptor with adverse effects on the environment (e.g. wild life)	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
*Persistent, bio-accumulative and toxic	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
*Persistent, mobile and toxic	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>

Question 2 - Imagine you want to buy or use a product which bears a label with one of the following hazards. Would you be ready to pay more for alternative products that have the same performance, but which do not have that hazard?

(One single answer per row)

	Yes	Probably	No	No opinion
	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>

Question 5a - Considering the example above, which pieces of the label would you like to keep?

(Select as many options as needed)

- Pictogram showing the risk (e.g., flame symbol for flammable chemical)
- Hazard statement and signal word (e.g., Danger It can cause cancer)
- Instructions of use
- Precautionary statements on how to store, dispose, prevent accidents etc.
- The name of the chemicals causing the hazard
- Additional specific labelling information (e.g. in case of chemicals containing lead, "Warning! contains lead")
- Identification code for poison centres (so called UFI code and allows poison centres to know the composition of a chemical)
- Other piece(s) of the label
- None of the provided options

Question 6 - Would you like to be able to consult labels of chemicals digitally in the future (e.g. on your computer or smartphone)?

It might be a digital consultation of the whole label or just part of it.

(Only one answer possible)

- Useful
- Not very useful
- Useless
- No opinion

Question 7 - Imagine you buy a detergent in bulk in a grocery. You have brought your own bottle which does not bear a label for this detergent. What would be the best option to inform you on the hazards and safety instructions?

(Only one answer possible)

- You do not need any information
- Information is displayed at the point of sale only
- Information is provided in the form of a document provided by the seller (leaflet or on the counter ticket)
- You can access the information digitally (scanning of a QR code for example)
- Other option(s)
- No opinion

Question 8 - Individual pens are very small items, with little room for a label and information about hazards. What would be the best option for you to inform on the hazardous substances they may contain and the safety instructions?

- You don't need any information
- Information displayed in the shop
- Information in the form of a document provided by the seller (leaflet or on the receipt)
- Information on the outer packaging, overwrapping a set of 10 pens
- Access the information digitally (scanning of a QR code for example)
- Other option(s)
- No opinion

Section 4 - Online sales**Question 9 - Online shopping of chemicals is becoming more and more common. Do you think it is important to receive the same safety information when you buy chemicals in a shop or online?**

- Yes
- No
- No opinion

Question 9a - When should you receive such information on hazards?

- Before ordering the chemical online
- When the chemical is delivered to you
- In both cases
- No opinion

Question 9a.i - Which information would you like to receive before ordering?

- Most important information (type of hazards, presence of hazardous components)
- All pieces of information which are on the label
- No opinion

Section 5 - Scope of the CLP regulation

Currently the product categories listed below are exempted from the CLP Regulation on classification and labelling.

- Medicines
- Veterinary medicines

- Cosmetics
- Medical devices (e.g. lens cleaning solutions)
- Food such as food additives, flavouring foodstuffs, or feed such as animal nutrition complement.

This is because hazards to human health are generally identified and dealt with by specific pieces of legislation. However, information on environmental hazards (such as "substance toxic to aquatic life") are not identified and information is not provided to the users of the above products.

Question 10 - When buying or using the product categories listed below, you might not be informed that they could be hazardous to the environment. What is your opinion?

	An issue which should be immediately solved	An issue where future improvement would be welcomed	Not an issue	No opinion
Medicines	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Veterinary medicines	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Medical devices (e.g. lens cleaning solutions)	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Cosmetics	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Food or feed, such as additives	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>

Question 11 - in case you you would like to share anything else in addition to the previous questions and in the view of the targeted revision of the CLP regulation (optional):

Question 12 - in case you would like to share a document in the view of the targeted revision of the CLP regulation, please upload it below (optional):

Part II - Questions for experts

This section should be answered by people having an excellent or good understanding of the CLP, from a legal or implementation perspective, or of chemical hazards.

Section 1 - New hazard classes

Endocrine disruptors

The World Health Organisation (WHO) has defined criteria (<https://www.who.int/ipcs/publications/en/ch1.pdf>) for endocrine disruptors which are the basis for the existing criteria for endocrine disruptors in plant protection and biocide products.

Question 13 - For known endocrine disruptors, do you think...?

- The WHO's definition and criteria should be taken over, word for word, in the foreseen EU CLP criteria.
- The foreseen CLP criteria should be the criteria in place for plant protection products (<http://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX:32018R0605>) or for biocide products (http://eur-lex.europa.eu/eli/reg_del/2017/2100/oj), which are based on the WHO definition and criteria.
- It is necessary to further refine WHO's definition and criteria and/or existing criteria for plant protection and biocide products to develop the foreseen CLP criteria.

Question 14 - Are you in favour of a sub-categorisation for chemicals with a high level of certainty on their endocrine disrupting properties, as for mutagenic chemicals (e.g. Categories 1A and 1B)?

- Yes
- No
- No opinion

Question 14a - Please detail why and how a subcategorisation should be provided. Please indicate whether there should differences between human health and the environment.

document attached

Question 15 - What would you suggest as criteria for a second category for chemicals with a lower level of certainty on their endocrine disrupting properties (human health and environment), as for mutagenic chemicals?

document attached

Question 16 - According to you, what would be the best statement on a label for chemicals identified as toxic to reproduction and as an ED according to the foreseen ED criteria?

- May cause infertility or damage to the unborn child
- May cause infertility or damage to the unborn child via an endocrine mode of action
- May cause infertility or damage to the unborn child
- May cause endocrine-related adverse effects on human health
- Other option(s)
- No opinion

(Very) persistent, (very) bio-accumulative and toxic substances

The introduction of criteria for persistent, bio-accumulative and toxic (PBT) or very persistent and very bi-accumulative (vPvB) substance in the CLP Regulation is expected, based on the criteria laid down in Annex XIII of the REACH regulation (<https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX%3A02006R1907-20210705>).

Question 17 - Do such criteria as provided in Annex XIII of REACH need to be updated before their foreseen introduction into the CLP Regulation?

- Yes
- No
- No opinion

Question 18 - Do you think a category for suspected PBT (and one for suspected vPvB) would be needed?

- Yes
- No
- No opinion

Question 19 - According to you, what is the best statement on a label for chemicals on the foreseen PBT, vPvB hazard classes?

If a chemical is identified as PBT and carcinogen category 1, its label should display:

(Only one answer possible)

- May cause cancer
- Persistent, bio-accumulative and toxic (PBT)
- May cause cancer
- Persistent (P)
- Bio-accumulative (B)
- Other option(s)
- No opinion

(Very) persistent, (very) mobile and toxic substances

The foreseen introduction of criteria for **persistent, mobile and toxic (PMT) or very persistent and very mobile (vPvM) substances** aims at improving protection, from chemical contamination, of water bodies when **used for drinking water purposes** (to protect human health).

Question 20 - Do you think environmental toxicity should be part of the toxicity criterion?

- Yes
- No
- No opinion

Question 21 - do you think a category for suspected PMT (and one for vPvM) would be needed?

- Yes
- No
- No opinion

Question 22 - According to you, what is the best statement on a label for chemicals on the foreseen PMT, vPvM hazard classes?

If a chemical is identified as PMT and carcinogen category 1, its label should display:

(Only one answer possible)

- May cause cancer
- Persistent, mobile and toxic (PMT)
- May cause cancer
- Persistent (P)
- Mobile (M)
- Other option(s)
- No opinion

Other hazard classes

Question 23 - In the environmental classification of chemicals, do you consider it relevant to use toxicity data obtained on terrestrial organisms to complement the information on toxicity for aquatic organisms?

(Please rate from 0 - not relevant to 10 - very relevant)

0

Question 24 - Immunotoxicity effects are currently covered under the hazard classes 'Specific target organ toxicity' and 'Reproductive toxicity' (in case of developmental immunotoxicity). Do you consider relevant to develop a separate specific hazard class/criteria for Immunotoxicity?

(Please rate from 0 - not relevant to 10 - very relevant)

0

Question 25 - Neurotoxicity effects are currently covered under the hazard classes 'Specific target organ toxicity' and 'Reproductive toxicity' (in case of developmental neurotoxicity). Do you consider relevant to develop a separate specific hazard class/criteria for neurotoxicity ?

(Please rate from 0 - not relevant to 10 - very relevant)

0

Possible impacts of the new hazard classes

Question 26 - The CLP regulation requires to use all available data to identify hazards in chemicals. Data may come from REACH registration(s) or public scientific literature. To what extent do you think that the data currently available on chemicals are sufficient to perform an assessment for the foreseen hazard classes mentioned above?

	Totally sufficient (with specific data on all substances)	Sufficient (incl. read-across and bridging)	Only partially sufficient covered (incl. read-across and bridging)	Not sufficient at all	No opinion/Not relevant to me or my organisation
Endocrine disruptors (human health)	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Endocrine disruptors (environment)	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
PBT/vPvB	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
PMT/vPvM	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>

Question 27 - Considering the suggested new criteria for additional hazard classes, do you foresee a need to invest significant resources to get the expertise to assess the hazards of chemicals?

	Need to invest in significant additional resources	Need to invest in some additional resources	Need to invest in little additional resources	No investment needed at all	No opinion or not relevant to me or my organisation
Endocrine disruptors (human health)	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
Endocrine disruptors (environment)	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
PBT/vPvB	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
PMT/vPvM	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>

Question 28 - Do you or your organisation/company already have an estimate of the number of impacted chemicals due to the potential new hazard classes?

- Yes (it will unfold a series of more detailed questions)
 No information or no opinion

Question 28a - What percentage of products would need to be re-classified and re-labelled due to foreseen criteria for ED, PMT/vPvM, PBT/vPvB?

0

Question 28b - In view of the foreseen new criteria for EDs, PMTs/vPvMs, PBTs/vPvBs, what percentage of products would you expect to be re-formulated voluntarily to avoid they contain ED, PMT/vPvM, PBT /vPvB?

Are you or is your company or organisation related to plant protection products or biocidal products?

- Yes
 No

Section 2 - Classification

Question 29 - In order to increase the number of substances with harmonised classification, to what extent do you agree to the following statements?

The European Commission should also have the right to initiate European classification for some substances

The European Commission should help Member States to submit more dossiers.

Question 30 - Setting toxicological/ecotoxicological values such as DNEL/DMEL, PNEC is part of the hazard assessment. These values are currently derived in accordance with REACH or specific sectorial regulations (e.g. food contact materials, cosmetics, biocidal products, workers protection). As part of the 'One substance, one assessment' concept, the Commission intends to include a procedure to harmonise values for some toxicological/ecotoxicological parameters in CLP. Such harmonised values could be then used for risk assessment in the different EU chemicals legislations.

How important would you rate the harmonisation of toxicological/ecotoxicological values?

	Important	Neutral	Not important	No opinion
Harmonising DNELs (Derived No-Effect Limits) in CLP	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
Harmonising DMELs (Derived Minimum-Effect Limits) in CLP	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
Harmonising PNECs (Predicted No-Effect Concentrations) in CLP	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>

Question 31 - How would you assess the possible impact of the harmonisation of toxicological/ecotoxicological parameters (e.g. DNELs or PNECs)?

	Important	Neutral	Not important	No opinion
Increase the level of protection of human health and the environment	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Ensure level playing field across sectors	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
Increase workload of the Risk Assessment Committee	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Increase of burden and regulatory requirements	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>

Question 32 - Currently CLH dossiers can be submitted by national competent authorities and in some cases by companies. Once received, the dossiers are checked for accordance.

What is your opinion about the three following statements?

The system should allow prioritisation of substances for which serious concerns are raised (e.g. priority given to substances highly suspected of being an endocrine disruptor, once the criteria are adopted).

The system should allow low prioritisation of substances of lower concerns.

No need to modify the current approach as the system already contained a prioritisation mechanism (National Authorities' priorities, ECHA screening).

Question 33 - Currently economic operators (manufacturers, importers, downstream users, distributors) are not allowed to submit a proposal to ECHA to revise an existing harmonised classification for an Annex VI entry. Only Member states can submit such a proposal. Please select the preferred option amongst the following ones:

- The system should not change to avoid a proliferation of CLH revision requests by stakeholders
- The CLH revision request by a stakeholder should be addressed first at the EU Commission for decision on the need of an action at Community level. If accepted by Commission, the request will be provided to ECHA against the payment of a fee covering all expected costs.
- The revision request by a stakeholder should be allowed and be provided to ECHA against the payment of a fee covering all expected costs.

Question 34 - To derive the correct classification of certain chemicals, the use of animal testing is still necessary.

Would you be confident to classify (your) products on the basis of alternative methods only?

- In the case the result of a test performed with an alternative method is positive, to classify (your) chemicals accordingly:
 - Yes
 - No
- In the case the result of a test performed with an alternative method is negative, not to classify (your) chemicals for that hazard class:
 - Yes
 - No

Question 35 - Currently, where the notification to the classification and labelling inventory (C&L inventory) results in different entries for the same substance, manufacturers and importers shall make every effort to come to an agreed entry in the inventory. Despite this obligation, different entries for the same substances are very frequent and significantly reduce the usefulness of the inventory.

Please provide your views on the potential following options below.

	Agree	Disagree	No opinion
The system should not change.	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>
The obligation to come to an agreed entry should be strengthened.	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
ECHA should be able to remove/refuse notifications that seem incorrect after having informed the notifier.	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>

Section 3 - Labelling

Question 36 - Did you experience issues with double or contradicting labelling obligations (CLP v. other legislation)?

- Yes
- No

Question 37 - How do you rate the economic impact (cost savings) of the following five policy options?

	Significant savings	No significant savings	No opinion
Exempt small products (pens, lighters) from certain labelling requirements	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Exempt bulk chemicals (fuels) from certain labelling requirements	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Allow a wide use of multilanguage labels / fold-out labels	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
Provide certain obligatory labelling information digitally instead of on the label	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>
Provide additional information digitally	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>

Question 38 - How do you rate the health, safety and environmental impacts of the following policy options? Please justify your choice in box below

	Significant positive impacts	No significant impacts (neutral)	Significant negative impacts	No opinion
Exempt small products (pens, lighters) from certain labelling requirements	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Exempt bulk chemicals (fuels) from certain labelling requirements	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Allow a wide use of multilanguage labels / fold-out labels	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
Provide certain obligatory labelling information digitally instead of on the label	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>
Provide additional information digitally	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>

Section 4 - Online sales

Question 39 - Some chemicals purchased online from non-EU countries often do not comply with EU law (e.g. are not providing obligatory safety information). In those cases, it is very difficult to identify the responsible company and take corrective measures.

In such a case, do you think the online service providers, platforms should be considered responsible?

- Yes
 No
 No opinion

Question 40 - How would you rate the need to apply the same CLP obligations (e.g. labelling, classification and notifications to poison centres) also to hazardous chemicals purchased online (compared to traditional purchase)?

4

Question 41 - How would you rate the need to have a responsible actor for compliance with CLP located in the EU also for chemicals purchased online?

2

Question 42 - What in your view are the major problems with online sales to ensure a level-playing field between companies?

(Please select as many answers as needed)

- Wrong or incomplete advertising
 Wrong or incomplete information on the webpage where the order is placed
 Wrong or incomplete labelling/packaging of chemicals
 Other problems than listed above
 No problem
 No opinion

Question 43 - What in your view are the major problems with online sales to ensure the same level of health, safety and environmental protection?

(Please select as many answers as needed)

- Wrong or incomplete advertising
 Wrong or incomplete information on the webpage where the order can be placed
 Wrong or incomplete labelling/packaging of products
 No poison centre notifications
 None of the options above

Question 44 - Do you think that the CLP regulation should address problematic issues arising from on-line sales of hazardous substances and mixtures?

- Yes
 No
 No opinion

Section 5 - Scope of the CLP regulation

Question 45 - Do you consider that there are gaps or overlaps between Article 1(5) of the CLP regulation and provisions in other legislations or that the wording is unclear?

	O v e r l a p s	G a p s	L a c k o f c l a r i t y	E v e r y t h i n g i s c l e a r	N o o p i n i o n
Medicines as defined in Directive 2001/83/EC (https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX%3A02001L0083-20190726)	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>
Veterinary medicines as defined in Directive 2001/82/EC (https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX%3A02001L0082-20090807)	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>
Medical devices as defined in Regulation (EU) 2017/745 (https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX%3A02017R0745-20200424) and Directive 98/79/EC (https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX%3A01998L0079-20120111)	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>
Cosmetics as defined in Regulation (EC) No 1223/2009 (https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX%3A02009R1223-20210617)	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>
Food and feeding stuffs as defined in Regulation (EC) No 178/2002 (https://eur-lex.europa.eu/legal-content/EN/TXT/?uri=CELEX%3A02002R0178-20210526), including flavouring of foodstuffs, animal nutrition and feed additives	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>

Question 46 - Currently neither the CLP nor the specific ('sectorial') legislation applying to the products listed in the table below require that information on classification and labelling of environmental hazards is provided to the users.

In your view, what would be the best option to make users aware of these environmental hazards?

	Add an obligation to classify and label according to CLP for environmental hazards.	Add an obligation to assess and label according to sectorial legislation	Promote voluntary use of CLP classification and labelling for environmental hazards	No opinion
Medicines	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Veterinary medicines	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Medical devices	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Cosmetics	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Food and feeding stuffs, including flavouring of foodstuffs, animal nutrition and feed additives	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>

Section 6 - Notifications to poison centres

Question 47 - CLP states that mixtures classified on the basis of their health and physical effects shall be submitted to appointed bodies (poison centres) in the Member States to provide emergency health response. CLP also provides that hazardous substances shall be notified to ECHA's classification and labelling inventory (C&L inventory) which is publicly accessible.

For poison centre purposes, is it useful to submit information also on substances?

- Yes
 No
 No opinion

Question 48 - What are in your view the most suitable transitional periods until the new rules become applicable for the different aspects amended under CLP?

	As soon as possible	18 months	24 months	36 months	48 months	No opinion
Introduction of new hazard classes	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>
Harmonised DNEL, PNEL, PNEC	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>
Improvements to CLH process (prioritisation mechanism, ECHA dossier submitter)	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>
Improve self-classifications	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>	<input type="radio"/>
Remove certain exemptions from CLP (medical devices, medicines, cosmetics etc.)	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Simplify labelling	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>
Tackle online sales lack of compliance	<input checked="" type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>
Improve notification to poison centres	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input checked="" type="radio"/>

Question 48a - Please provide the reasons for the above proposed timelines for the applicability period.

Section 7 - final (additional) feedback

Question 49 - in case you would like to share anything else in addition to the previous questions to experts and in the view of the targeted revision of the CLP regulation (optional):

Question 50 - in case you would like to share a document in the view of the targeted revision of the CLP regulation, please upload it below (optional):

20211108 OPC_ED_CLP_ETRMA_supporting-document_FINAL_Clean.pdf

Contact

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