Options for an environmental risk assessment of intentional and unintentional chemical mixtures under REACH: the status and ways forward

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Abstract

It is acknowledged that a variety of chemicals enter the environment and may cause joint effects. Chemicals regulated under the European Chemicals Regulation REACH are often part of formulated mixtures and during their processing and use in various products they can be jointly released via sewage treatment plants or diffuse sources, and may combine in the environment. One can differentiate between intentional mixtures, and unintentional mixtures. In contrast to other substance-oriented legislations, REACH contains no explicit requirements for an assessment of combined effects, exposures and risks of several components. Still, it requires ensuring the safe use of substances on their own, in mixtures, and in articles. The available options to address intentional as well as unintentional mixtures are presented and discussed with respect to their feasibility under REACH, considering the responsibilities, communication tasks and information availability of the different actors (registrants, downstream-user and authorities). Specific mixture assessments via component-based approaches require a comprehensive knowledge on substances properties, uses, fate and behaviour, and the composition of the mixture under consideration. This information is often not available to the responsible actor. In principle, intentional mixtures of known composition can be assessed by the downstream-user. But approaches have to be improved to ensure a transparent communication and sound mixture assessment. In contrast, unintentional mixtures appear to be better addressable via generic approaches such as a mixture allocation factor during the chemical safety assessment, although questions on the magnitude, implementation and legal mandates remain. Authorities can conduct specific mixture risk assessments in well-defined and prioritized cases, followed by subsequent regulatory measures. In order to address intentional and unintentional mixtures within the current REACH framework, legal mandates together with guidance for the different actors are needed. Furthermore, further data on mixture compositions, uses and co-exposures need to be made accessible via shared databases.

Keywords: REACH, Mixtures, Environmental risk assessment, Mixture allocation factor, Co-exposure

Introduction: relevance of mixtures and their assessment

Currently, around 23 000 single substances with about 100 000 dossiers are registered under the European Chemicals Regulation REACH (1907/2006/EC) on “Registration, Evaluation, Authorisation and Restriction of Chemicals” [15]. A large proportion of these substances end up in a variety of mixtures, intentionally manufactured as such or containing by-products of other processes. Such intentional mixtures that contain a number of different components include wet formulations like varnishes, inks, paints (more than 20 ingredients), lacquers (10–50 ingredients), mortar

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(1–5 ingredients), or cleaning agents (5–10 ingredients) and dry formulations like pre-formulated granules [40]. During their processing and downstream use substances are blended together in formulations, are subsequently reformulated for different uses and end up in a variety of products. During their life cycle (from production, application and service life to waste) chemicals may be introduced into the environment in various ways. They are emitted from point sources such as production plants, enter compartments from different sources, such as industrial wastewater or municipal sewage treatment plant effluents, via diffuse pathways like urban run-off from streets or buildings, as well as via waste. One can hence differentiate between formulated intentional mixtures, and unintentional mixtures in the environment (Fig. 1).

Scientific evidence on environmental exposures to multiple hazardous chemicals is increasing [44]. Data from monitoring studies of European surface waters demonstrate the co-occurrence of multiple chemicals in time and space [11, 45]. Substance concentrations in European surface waters may exceed regulatory thresholds for single substances and potential risks have been shown to increase with the number of chemicals found [12, 46].

The scientific basis for an assessment of chemical mixtures is well established. It has been shown experimentally that chemicals may act jointly and that the combined toxicity may be higher than the toxicity of each of the single components on its own as reviewed by Kortenkamp et al. [43]. This may also be the case if substances are present at or below their regulatory thresholds, such as the EQS or NOEC [12, 57]. In most cases chemicals have been shown to act additively following the established concepts of Concentration Addition (CA) or Independent Action (IA). Synergisms or antagonisms, exceeding or reducing the additive ecotoxicities due to specific interactions between the chemicals are comparatively rare and restricted to specific cases [13, 47]. While CA was primary used to predict (eco)toxicities of substances with similar modes of action and IA for dissimilar modes of action, CA has been recognized as pragmatic standard in the environmental context.

Methodologies for assessing the combination effects of chemicals have been developed and experiences were gained in the academic as well as the regulatory context for mixtures of known composition. In particular, the State of the Art report by the Commission [43], the framework of the WHO International Programme on Chemical Safety (IPCS) [49] and several reviews of the Joint Research Centre and a report on behalf of the Swedish Government [4, 5, 40, 41, 56] provided analyses on the state of play. Recently, basic procedures have found

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**Fig. 1** Simplified overview on different levels of complexity to be considered for an environmental assessment for intentionally formulated as well as unintentional mixtures of chemicals reaching environmental compartments (surface/marine water, soil, air) via point sources (e.g. sewage treatment plants) or diffuse sources (e.g. urban run-off, depositions, or indirect exposures via man)
their way into overarching guidance of the Organisation for Economic Co-operation and Development (OECD) [51] and the European Food Safety Authority EFSA [50].

The issue of a risk assessment of chemical mixtures has been intensively discussed across the EU and OECD member states. Roughly 10 years ago, the Conclusions of the Environment Council [24], the Communication of the European Commission [18] together with the Opinions of the Scientific Committees SCHER/SCENHIR/SCCS [19], already expressed the need to address combined effects. In 2012, the European Commission proposed to develop technical guidelines and science-based approaches for the assessment of chemical mixtures, and to review the progress made. Most recently, the Conclusions of the Environment Council [25] on Chemicals together with the Chemicals Strategy for Sustainability in the context of the European Green Deal [20] again highlight the need for a consideration of multiple exposures and combined effects of chemicals. The accompanying comprehensive Staff Working Document of the European Commission [22] reports the progress made on the topic and future needs.

Although current risk assessments of chemicals most often focus on individual chemicals, environmental hazards and risks arising from combination effects and exposures are already addressed in several European substance-oriented regulations with respect to intentional mixtures. These are the legislations for plant protection products (Regulation (EC) No 1107/2009, PPPR), biocidal products (Regulation (EC) No 528/2012, BPR) as well as veterinary pharmaceuticals (Directive 2001/82/EC), which all require a product authorization by the authorities. These contain explicit legal requirements for the consideration of combined effects. Methods are specified in detailed guidance documents for plant protection products, e.g. [34], biocidal products [31], and veterinary pharmaceuticals [36]. These requirements are restricted to formulations, and do not consider re-formulations, sequential applications or simultaneous entries of several components. In the context of the media-oriented water framework directive (WFD) (2000/60/EC) [59], the method to derive Environmental Quality Standards (EQS) for mixtures is laid down in the common implementation plan Commission [17], without explicit obligations.

In view of the achievements of other substance-oriented legislations with respect to intentional mixtures, REACH seems to lag behind as there are no explicit legal obligations—except those referring to the CLP regulation (No 1272/2008/EC)—to assess and regulate the combined effects and exposures.

A number of authors have proposed ways forward for the regulatory assessment of combined exposures [6, 40, 44, 51, 55, 56]. Explicitly related to REACH, only few proposals for a mixture risk assessment are available [2, 8]. For the environment, Bunke et al. [10] provided an in-depth analysis of options for different actors under REACH to address intentional as well as unintentional mixtures. Recently, Rudén et al. [56] came up with more specific proposals for REACH to improve the consideration of combination effects and grouping of chemicals.

**Aims and methodology**

This regulatory review compiles the necessary background information on the functioning of REACH and central methods for a mixture assessment that is needed for a critical analysis of the available options and challenges to address intentional and unintentional mixtures within REACH. The present paper has a clear focus on the environment. The aim is to provide a comprehensive basis for the on-going discussions at EU level around the possible options and the European Commission’s proposal for an introduction of a so-called “mixture assessment factor” within REACH. Although the topic is relevant for other legislations and their interplay, the work focuses on REACH and assessments and regulatory measures for mixtures in the environment. For the analysis of possible policy options, the current legal obligations, different responsibilities, communication tasks, and information availability for the different actors (registrants, downstream-user and authorities) are considered.

For this purpose, recent reports on the issue were evaluated and referred to after a non-systematic and non-exhaustive literature research. This included the legal text of the REACH regulation, its accompanying guidance documents as well as experiences from regulatory practise. An early starting point was a development and research project on behalf of the German Environment Agency [10] complemented by own analyses.

**Background: how does REACH work? Data availability, tasks and communication of the different actors**

REACH is a typical single substance-oriented regulation and hence addresses single substances. In contrast to other substance regulations, REACH includes neither an authorization of single substances per se nor a product authorization of intentional mixtures. It also shifts the burden of proof: not authorities, but manufacturer and importer of chemicals (registrants) together with the downstream users (e.g. formulator) are responsible for the data generation, safety assessment and derivation of safe use conditions for single chemicals.

Although REACH aims at creating a comprehensive data basis for its 22,873 registered chemicals (ECHA dissemination website 07/2020 [62]), data availability and
Table 1  Information availability for an environmental assessment (requirements are laid down in Annex VII-IX of REACH and the respective chapters of ECHA GD) and tasks of the different actors in the supply chain (implying numerous subsequent downstream user and end user)

| Knowledge                        | Registrant                                                                 | Downstream user                                                                 | Authorities                                                                 |
|----------------------------------|---------------------------------------------------------------------------|--------------------------------------------------------------------------------|----------------------------------------------------------------------------|
| Obligations and tasks            | Assessment of hazards/exposures/risks of single substances, supporting (or excluding) certain uses | Adaptation or own derivation of $M_{safef}$ safe use conditions for own uses | Specific evaluations and regulatory risk management measures if unacceptable risks or hazards are identified |
| Number of substances addressed$^a$| In total: 22,873  
1–10 t/a: 3,249  
10–100 t/a: 2,923  
100–1000 t/a: 2,272  
> 1000 t/a: 2,455 (without intermediates) | All substances and indefinite number of formulations | SVHC identified in total: 197 (ED ENV: 12, PBT: 13) Restrictions (Annex XVII): 70 |
| Communication and assessment obligations | 1 t/a: Registration dossier and CSA, SDS (including PNEC, $M_{safef}$, RMM, OC for intended DU uses), CSR for NM  
> 10 t/a: CSR, with ES and eSDS for hazardous chemicals  
Self-classification of substances and PBT assessment | (e)SDS/mixture SDS (with ES for all substances) Optional DU CSR for substance(s) in a mixture Classification of a mixture | In case of identified concerns: Dossiers for substance evaluation (SEV) and/or proposals for regulatory measures (SVHC-identification, restrictions) CLH proposals for substances |
| Information available (details see below) | Study results from own tests or publicly available literature  
Generic information from DU associations | Receives (e)SDS from REG for substances used | MS CA/ECHA: Registration dossier (IUCLID, CSR). Evaluated on random basis (CCH) or concern (SEV) Additional information on request, during consultations or by Surveillance authorities ((e)SDS from DU, information on sites) |
| Effect data (see Annex VII–IX REACH) | Basic requirements depending on tonnage and properties (also non-test information or waiving):  
1–10 t/a Phys.-Chem  
10–100 t/a: short-term Algae, Daphnia  
100–1000 t/a: long-term fish or daphnia (ITS), short-term soil (invertebrates, plants, MO) if soil exposed  
> 1000 t/a: long-term soil (invertebrates, plants), sediment, birds Calculation of PNEC for each compartment | (e)SDS: Most sensitive PNEC for classified substances, PBT status | ECx and/or NOEC, PNECs and robust study summaries (RSS) available in technical IUCLID dossier and CSR Results from publicly available scientific studies |
| Exposure data | Calculation of PECs and RCRs for each use category and env. compartment Monitoring data usually not considered | Emission values, release factor, PEC and $M_{safef}$ received via eSDS from REG if indicated Concentrations used, OC and RMM for each use category | Access to PECs, RCRs, yearly processed tonnage, OC, RMM in CSR and IUCLID for identified use categories and compartments. Publicly available literature on monitoring data |
| Use of substance | Receives generic sector-specific information on uses, processed tonnages, OC and RMM from DU associations (own surveys, communication) | Information on ingredients (substances or preformulations) and own formulation. Information on subsequent uses only on demand | Use categories in CSR from REG No detailed information on specific uses (only via surveys or on request or very generic information in SPIN database) |

$^a$ ECHA dissemination Website (07/2020)
quality varies greatly for substances. Data availability is depending on production volumes and substance properties according to the data requirements defined in Annex VII–X. In particular for substances marketed at low tonnages < 10 t/a, ecotoxicity data are very limited. Exposure scenarios are only available for certain hazardous substances and at higher tonnages. Environmental data availability is often restricted to certain actors (Table 1).

The registrants: chemicals safety assessment
Substances with volumes > 1 t/a have to be registered by the manufacturer and importer of single chemicals. For substance in volumes greater than 10 t/a registrants have to conduct a chemical safety assessment (CSA). An exposure and risk assessment need to be conducted for classified substances, substances assessed as persistent, bioaccumulative and toxic (PBT), or substances with hazardous effects (Article 14.4). Information on intrinsic properties, fate and behaviour, as well as hazards and risks are documented in a chemical safety report (CSR) as part of the registration dossier (REACH article 14.1).

For the risk assessment, risk characterization ratios (RCR = Predicted Environmental Concentration (PEC)/Predicted No Effect Concentration (PNEC)) are derived for each compartment (air, aquatic water and sediment, marine water and sediment, terrestrial environment) and for each intended use (described in a standardized way by means of the use descriptor system). An RCR < 1 indicates that the identified risks are acceptable. The PNEC is derived using effect concentration(s) from ecotoxicity tests with representative species (e.g. algae, daphnids, and/or fish) together with defined assessment factors taking into account the data availability and uncertainties related to the complexity of ecosystems as laid down in guidance R.7b of the European Chemicals Agency (ECHA). PECs are calculated on the basis of generic emission factors according to ECHA Guidance on environmental exposure assessment R.16 [29]. The PEC estimate represents local concentrations, considering daily release rates, dilution in the sewage treatment plant and receiving waters as well as regional background concentrations. The background concentrations are based on averaged, aggregated, annual concentrations of a substance across all life cycle steps for a standard region. The PEC and emission factors may be refined using so-called specific environmental release categories (spERCs), developed by industry associations, or monitoring data.

In order to estimate the maximum amounts of a substance that may be used during production, and hence "safely" emitted to environmental compartments, the so-called maximum safe use amount or “M_safe” (kg/day or kg/year) is calculated using the PEC and the emission days. In case unacceptable risks (RCR > 1) are identified, additional operational conditions (OCs) or risk mitigation measures (RMM), such as adapted filter systems or wastewater cleaning steps, are applied to achieve a safe use.

The registrant is obliged to communicate only the basic information for a substance via safety data sheets (SDS) or, in case of a required risk assessment, additionally the exposure scenarios in the so-called extended safety data sheets (eSDS) to the downstream user (REACH articles 14.4 and 31.7). In case of an eSDS, the basic substance information shall include the (most sensitive) PNECs, estimated safe use amounts (M_safe), safe use conditions, exposure scenarios and risk reduction measures.

The downstream users: check and adaptations for own use
Being responsible for the safe use of the substances, mixtures and articles they process, downstream users (DUs) are obliged to check via the (e)SDS whether the classification of substances and intentional mixtures is correct, whether their application is covered, and whether the described conditions ensure that risks are controlled (i.e. RCR < 1) for their intended use. In case of deviations, the DU has to conduct an own safety assessment, adapt the safe use conditions and/or amounts. If a use is not covered by the registrants’ exposure scenario, the DU has to notify ECHA, ask the registrant to integrate his use into the registration dossier, or provide an own DU CSR (REACH articles 37.4 and 31.2).

The DU is further obliged to communicate the result of his DU assessment, the information on single components, the classification and labelling of his formulated mixture, a corresponding exposure scenario, and summary of risk mitigation measures via the eSDS to subsequent DUs and end user (REACH Annex II and title IV, Article 31 and 38). The DU also informs the registrant on his uses via the so-called upstream communication in a very generalized form as use categories, respecting the confidentiality of recipes for formulations.

The regulatory measures of authorities
The regulatory authorities (ECHA, member state authorities) evaluate the standard data provided by the registrants in the technical dossiers and the chemical safety reports on a random basis during compliance checks of dossiers (CCH) and during testing proposal evaluations (TPE). Member state authorities may evaluate substances in more detail using all available information during the formal substance evaluation (SEV), if concerns and potential hazards/risks for the environment are indicated and justified. These substances are published on ECHA’s Community Rolling Action Plan (CoRAP) list [61]. Following these evaluations further test data or information...
on exposures or certain hazardous properties (e.g. endocrine disruption) may be requested from registrants. If certain hazards or a risk is concluded, further regulatory measures may be proposed by member state authorities. The identification as substance of very high concern (SVHC) leads to an inclusion in the candidate list [63] and certain information obligations to DU and consumers, a possible prioritization for inclusion in Annex XIV [67] and the authorization procedure. A second regulatory measure is the restriction of certain uses by inclusion of substances in the Annex XVII of REACH [64].
The challenge: communication of information between the actors

Within REACH, there is a discrepancy with respect to the degree of knowledge, and the different responsibilities between the registrants and DU (Fig. 2). The registrant has detailed information on the substance properties, but is not obliged to cover every single use condition for his substance during its lifecycle and only receives very generic information on the intended uses from the DUs (due to confidentiality reasons) via upstream communication. The DUs have more detailed information on their own specific uses such as the composition of a specific mixture they prepare and the obligation to forward basic information on the intentional mixtures they sell, formulate or process in the supply chain according to REACH titles IV and V via safety data sheets during downstream communication. However, they only receive limited information on their ingredients via the (e)SDS from the registrants and eventually previous suppliers. The regulatory authorities only get insight into chemical safety assessment reports of registrants, but usually will not receive information from DU.

Legal obligations enabling an environmental risk assessment of mixtures under REACH

Definitions for substances and mixtures

As stated above, REACH focuses on single substances. Besides substances as such, REACH refers to single substances in mixtures and in articles, where several substances are intentionally blended. So-called multi-constituent substances (MCS) and substances of unknown or variable composition and biological materials (UVCB) are defined as single substances under the REACH regulation. REACH defines a mixture as a combination of two or more substances, blended intentionally in a formulation. In light of around 23 000 registered single chemicals, the number of formulated intentional mixtures under REACH is high—concrete numbers remain unknown. Emissions of one single substance via different sources is often referred to as combined exposure under REACH. We use the term aggregated exposure. REACH does not refer to unintentional mixtures.

However, during their production and life cycle, substances may be emitted from a production plant, a sewage treatment plant (point sources) or via a variety of pathways, and subsequently enter environmental compartments (diffuse sources). Hence, it is important to differentiate between intentional and unintentional mixtures and address different levels of complexity (Fig. 1).

In order to reach a common understanding terms such as technical mixtures, discharge mixtures, co- incidental mixtures, or environmental mixtures have been introduced by several authors [10, 41, 51], which are summarized in the Table 2.

Obligations under REACH enabling mixture assessments

Within REACH there are no clear legal obligations to address possible risks arising from joint effects and exposures of the components in a formulated mixture or later co-exposures in the environment. However, REACH explicitly addresses the safe use of single substances on their own, in (intentional) mixtures and in articles throughout their life cycle and obliges registrants and DU to assess and ensure safe use conditions (REACH Art. 17, Art. 10).

As laid down in the respective guidance documents [27, 28], DU have the obligation to check that their use of substances in mixtures is covered by the corresponding exposure scenario(s), but the simultaneous use of several hazardous substances is not explicitly addressed. For the registrant, there is no obligation for a mixture assessment—mixtures are not registered or listed. But potential emissions of a single substance via different sources and its uses during its life cycle and along the supply chain—so-called aggregated exposures—are taken in account for the calculation of the PEC by the registrants (see Annex I 6.2 and ECHA Guidance Documents R.12, R.16 and R.18) [9].

Classification and labelling of substances and mixtures

The regulation for classification, labelling and packaging (CLP) (Regulation (EC) No 1272/2008) [16] and its guidance on the application of the CLP criteria [30] are key instruments for REACH providing the basis for communication of hazard data on substances and intentional mixtures along the supply chain. Intentional mixtures have to be classified and labelled by companies considering additive effects of hazardous components by either testing the whole mixture as such or, by applying the additivity or summation method to determine the ecotoxicity. Self-classifications are conducted by each company on their own for substances and mixtures without further approval by authorities. These classifications may differ for the same substance between different companies that are coming to diverging conclusions and subsequently also vary for identical formulated mixtures. Lists on classified mixtures are not available. For single substances, authorities can propose harmonized classifications, which are included in Annex VI to the CLP Regulation. Both the self-notified as well as harmonized classified substances are listed in ECHAs inventories [65, 66].
Further obligations under REACH enabling future mixture assessments

In order to close data gaps and limit tests to only a few representatives, the assessment of multi-component substances (such as MCS or UVCB) makes use of the “summation method” based on the concept of CA to estimate overall ecotoxicities. This is described in REACH Annex I, section 0.4 and Annex XI section 1.5, as well as ECHA guidance R6 on grouping of chemicals [26] and the read across assessment framework (RAAF) [32]. For substances originating from petroleum or coal streams, the so-called PetCo substances, the Hydrocarbon block methodology (HCBM) [42] is used to assess the risks of complex substances based on the known properties of their similar constituents or blocks.

In the context of the authorities’ evaluations of substances that may give rise to concern, group-wise assessments of structural related substances are explicitly referred to in the legal text (REACH Title VI articles 44.1a, and 47). Also, REACH Annex XVII on restrictions contains entries for groups of related substances. In the context of granting authorizations for substances in Annex XIV, ‘all discharges, emissions and losses including risks arising from diffuse or dispersive use’ shall be taken into account (REACH Title VII, article 60.2).

Approaches for a specific mixture assessment

General principles

General procedures for specific mixture risk assessments are in principle the same for intended and unintentional mixtures and have been reviewed for example by [43, 50, 51]. Approaches for the environmental assessment of intentional mixtures have already been implemented in guidance documents in the context of the product authorization of plant protection products [37], biocidal products [31], and veterinary pharmaceuticals [36].

Procedures include the definition of the composition of the mixture under consideration, testing of whole mixtures when available or component-based approaches using the concepts CA or IA on the basis of ecotoxicity data (i.e. effect concentrations or reference values) to calculate mixture toxicities. Together with exposure concentrations of the components (i.e. predicted or measured), a summation of the risk characterization ratios (RCR-mix < 1), of the components often is used to characterize the risks. On this basis, the identification of possible drivers of ecotoxicity and/or risks is possible.

Composition of a mixture

While the composition and concentrations of ingredients of an intentional mixture are in the best case known to the formulator, the composition of unintentional mixtures is often difficult to define. Defining unintentional releases or mixtures in an environmental compartment rely on the availability of data on co-occurrences which often varies spatially and temporally. While effluents might be defined and assessed by production plant operators, an assessment of complex mixtures requires monitoring data on co-occurrences in environmental compartments and a prioritization of substance and mixtures of concern.

Criteria to include components

Both for intentional as well as unintentional mixtures, criteria are needed to define which components to consider for the mixture assessment as not all ingredients may be relevant. Besides substances that are classified as hazardous to the environment according to CLP/GHS, in particular substances of concern under REACH, such as identified SVHC (PBT, vPvB, PMT, vPvM, ED1) are important to be considered. The same would apply to priority (hazardous) substances (P(H)S) of the water framework directive and further substances classified as hazardous on a national level (e.g. German regulation on substances hazardous to water AWSV/22/2017 [71]).

Criteria can also be based on certain concentrations or percentages in a formulation. Here, it is to consider that possible additive effects may also occur in low individual concentrations of the components and that also substances with dissimilar modes of action may produce additive effects [43].

The “maximum cumulative ratio” (MCR) approach can be used as tool to decide on the need for a refined assessment of (intentional or unintentional) mixtures [52, 53]. It is based on the ratio between the toxicity of a mixture as predicted by CA and the toxicity of the most contributing chemical in the mixture. The calculation of the MCR involves a rough mixture risk assessment by summing up the risk quotients of all the components. Here, two basic scenarios would trigger a mixture assessment: either one or few constituents are driving the mixture toxicity (so-called “drivers”) or several substances contribute rather equally at lower individual concentrations to the overall toxicity [53]. Both scenarios would be relevant for an assessment: in some cases, a single-substance based assessment/regulation may be sufficient, while in other cases several components might need to be addressed.

\footnote{Persistent, Bioaccumulative and Toxic, very Persistent and very Bioaccumulative, Persistent, Mobile and Toxic, very Persistent and very Mobile, Endocrine Disrupter.}
Assessment of combined effects
In cases where a real mixture is available on hand and testable, a whole mixture approach is one way to assess the overall toxicity. However, this is only applicable for certain formulations (for example, during classification and labelling), or samples from emissions or environmental compartments and only valid for this specific situation.

Component-based approaches, i.e. CA and IA are well established to predict mixture toxicities based on available ecotoxicity data, usually effect concentrations. In particular, CA is widely accepted as feasible and sound default approach in the environmental context [43]: it relies on effect concentrations for the individual components (i.e. EC10 or EC50, NOECs or LOECs), what makes it less demanding than IA. CA also works considerably well in case of different modes of actions, where IA typically is used. However, CA often predicts higher toxicities than IA and therefore is known to be more conservative, i.e. precautious. For rare cases, specific analyses for the consideration of potential synergisms might be meaningful. In the REACH context, one would most often have to deal with the reported PNEC value(s) based on differing underlying data (QSAR estimates, ecotoxicological standard data for few species from different trophic levels). These would be used when calculating mixture risks. Here, using CA may be challenging as PNECs may rely on data for species from different trophic levels.

Assessment of combined exposures
The concentrations of the individual components of a “whole mixture” may be analysed by means of chemical analyses to gain information on the composition.

Theoretically, similar to other substance-oriented regulations, predicted environmental exposure concentrations for intentional mixtures could be determined by summation of the calculated PECs by the registrants and/or DU for the respective environmental compartments.

More sophisticated approaches such as co-exposure modelling are valuable to characterize exposures of unintentional mixtures in the environment on a European scale, but rely on information on uses, production volumes, and fate properties [39]. Knowledge on measured co-exposures of major components in sewage treatment plants [23] and surface waters may get better available via the European Data Platform IPCHEM [70]. Besides identifying relevant substances for a mixture assessment, monitoring and/or modelling data can be used to better define background concentrations for the PEC estimate that also might take aggregated and combined exposures to further substances into account.

Risk characterizations to estimate a safe use conditions for the environment
Under other substance-oriented frameworks, a risk characterization of intentional mixtures is in principle done via component-based approaches by a summation of PEC/PNEC ratios for the relevant ingredients by applicants and authorities during product authorization in order to estimate a safe use. This is unfeasible for REACH due to the aforementioned limited availability of information on the mixture compositions (see also Fig. 2).

For unintentional mixtures present in the environment, potential risks can be evaluated by means of modelling approaches or comparisons of predicted or measured exposure information together with effect data or threshold values (PNECs, EQS) for single substances. This relies on the availability of data on substance properties, fate, use amounts and exposures. For example, van Gils et al. [39] used production volumes, fate properties and publicly available ecotoxicity data (e.g. PNEC) to predict possible joint risks of chemicals on a European scale. Malaj et al. [46] predicted risks of joint occurrences of chemicals in European surface waters based on measured concentrations from monitoring programmes and environmental quality standards.

Approaches for a generic assessment: mixture allocation factor
Specific assessments are very laborious, need a sound data basis and conflict with different responsibilities and knowledge of the actors within REACH. Hence, generic assessment approaches may be a pragmatic default option that could be established during the environmental risk assessment and/or management of chemicals, in particular to take account for unintentional co-occurrences in the environment. The safety factors applied within REACH for PNEC derivation are only covering uncertainties due to biotic interactions and the complexity of ecosystems but are not covering combined effects or co-occurrences of substances [48].

The use of a so-called mixture assessment/allocation factor (MAF), has been proposed and is currently discussed at EU level [2, 3, 8, 10, 21, 22] to address risks of unintentional mixtures during the single substances “chemicals safety assessment”. This would go beyond the single substance paradigm of the current legislations aiming at a reduction of the toxic pressure due to joint occurrences, effects and risks. The current proposals envisage a reduction of the current size of the RCR [2, 8, 10, 21]. Hence, the RCR would be interpreted as contribution of a substance to the overall risk of unintentional multiple substances in the environment instead of the single
substances’ risks [2]. The appropriate magnitude of such a measure should be determined based on scientific evidence that is available from monitoring, modelling and/or experimental studies [23, 56]. Recently, [3] provided a possible approach to derive an appropriate order of magnitude of a MAF. Still, central questions and challenges remain with respect to an implementation (Table 3).

**Options within the different REACH processes**

In the following paragraphs, the different options to address intentional and unintentional mixtures within the different REACH processes are discussed. This is illustrated in Fig. 3.

**Chemical safety assessment to ensure safe use for intentional mixtures**

Following the setup of REACH, a consideration of intentional mixtures during the CSA of the registrant to ensure safe uses would be appropriate. But in order to do so, the registrant would require information on the further use of his substance in intentional mixtures and detailed sector-specific information on uses and compositions of formulated mixtures as well as regularly updated use amounts. In theory, this could be communicated upstream via reporting templates, such as the templates for the “safe use of mixtures information” (SUMI), developed by CEFIC [14]. However, this is challenging in light of the presumably enormous number of intentional mixtures and confidential information on compositions. Backhaus et al. [2] proposed to use the product categories (PC) as specified in ECHA Guidance Document R.12 on the use descriptor system to generate exposure scenarios for groups of chemicals. However, a specification for somehow standardized mixtures would either be very generic or lead to a tremendous number of possible PCs. A compilation of typical compositions and/or the self-classifications of intentional mixtures in databases accessible for authorities and registrants would, however, be valuable in order to gain knowledge on the substances sources during the life cycle and possible later co-occurrences of substances. The improved use of the SPIN database as product register was recently proposed by Rudén et al. [56].

**Downstream assessment of safe use for intentional mixtures**

As the downstream formulators know compositions and are obliged to classify hazards and ensure the safe use of their own intentional mixtures, they seem to be the appropriate actors to address the potential combined effects, exposures and risks of substances and mixtures they use. However, in particular small and medium-sized enterprises (SMEs) are extremely challenged by the limited information in the (e)SDS together with limited capacities to generate data and conduct own assessments.

Recently, the European Chemical Industry Council CEFIC [14] proposed a “lead component identification methodology” (LCID) together with reporting templates for the “safe use of mixtures information” (SUMI) for downstream-user to improve the communication along the supply chain for hazardous substances in intentional mixtures.

LCID is a working tool aiming at an improved assessment of the safe use conditions of classified substances in intentional mixtures [14], but it aims not per se at the assessment of joint toxicities, exposures or risks. The gaps of the methodology with respect to the environment were analysed by Reihlen et al. [54], as well as Galert and Hassold [38]. Although based on Concentration Addition, a prioritization approach (MCR) is used to determine a lead component for which the safe use is to be ensured by adapting the initial Msafe and/or RMM provided by the registrants. Here, only substances that are classified with respect to the protection goal (for the environment: hazardous to the aquatic environment) and those for which information is available in the safety data sheets provided by the registrant (using the most sensitive PNEC leading to classification) are considered. Also, SVHCs are currently not included as these are separately assessed. Sub-lead substances are only considered when it makes up at least 10% of the lead substance. The method has its origin in human health assessment, where derived risk mitigation measures aim at the appropriate protective equipment for workers (e.g. gloves) and it is assumed, that “if the risk of most hazardous substance(s) is/are controlled, then risks for other substance in the mixture are controlled”. However, this is different for the environmental assessment, because here, specific safe use amounts and environmental release concentrations are to be derived and also further substances and smaller amounts may be released to the environment and contribute to effects. With LCID, further potentially relevant components in a mixture, in particular those that are not yet classified under CLP, or certain substances of concern seem not to be considered sufficiently. One attempt could be the introduction of “Mixture Assessment Triggering Substances” (MATS), as proposed by Bunke et al. [10] which could be defined by authorities on the basis of certain hazardous properties, high concentrations in the environment or potential risks and would trigger the need for an assessment of mixtures which contain such substances.
Table 3 Central considerations to be addressed for deriving a defined "mixture allocation factor" to take account for unintentional environmental co-occurrences of chemicals

| Open question                              | Options                                                                 | Needs and challenges                                                                 |
|--------------------------------------------|-------------------------------------------------------------------------|---------------------------------------------------------------------------------------|
| Process and responsible actor?             | MAF set by authorities to be implemented in legal text, supported by ECHA GD. Applicable during Chemical safety assessment of registrants and/or Downstream user assessments | The premise is that chemicals are not reaching environmental compartments alone but co-occur and may produce joint effects and risks—which is currently not addressed during the single substance’s CSA nor DU assessments. Takes account for limited information and resources of REG and DU and follows the legal setup of REACH that safe use ensured during CSA. |
| Location for implementation in legal text? | In Annex I on CSA factor could be introduced on: PNEC (reflects intrinsic properties and includes AFs for 1 substance) PEC (reflects exposure for compartment/use) RCR (reflects predicted risk for use and compartment) Msafe (reflects safe use amounts) | Communication along supply chain to be ensured. Confusion with existing uncertainty factors to be avoided. Clear wording, definitions and references in legal text and guidances. |
| Uniform or specific factors?              | Either the same or specific factors for: different environmental compartments different uses and product categories different tonnages | One uniform MAF more pragmatic, assumed to affect substances close to RCR of 1, easier to communicate. Implications, evidences and benefits would need to be justified for possible differentiated factors. |
| Definition of the appropriate order of magnitude? | The order of magnitude or "size" of a MAF can be based on: Evidence on averaged environmental co-occurrences and predicted risks Knowledge on predicted exposures or emissions from substances/mixtures Analysis of current chemical safety assessments and potential impact of a factor on the reduction of safe use amounts, predicted emissions or risks | Review and analyses of available scientific studies on environmental co-occurrence and possible risks of chemicals. Use of modelling approaches to estimate EU level co-exposures and current risk underestimations Analyses of current CSAs. |
| Refinements or exemptions?                | Refinements may justifiable in certain situations (no exposure due to close system) if sufficient and reliable data are available to conduct specific assessment | Conceivable that a generic default factor may be refined via specific assessments in cases where data available or exposures can be excluded. |
| Chemicals and types of mixtures covered?   | It needs to be defined whether a MAF should cover possible risks due to exposures of chemicals regulated under REACH co-occurring other substances, such as pesticides, biocides or pharmaceuticals | The coverage needs to be clearly defined. Covering substances not falling within REACH is legally challenging. |
| Possibilities for a later review?          | The magnitude or specification of a MAF could be readjusted After a fixed time span In case of new scientific evidence | A re-assessment could be in accordance with the REACH Review and could imply a surveillance of chemicals exposures and predicted risks. |
Due to the previously mentioned missing legal obligations, but most of all limited resources, and the limited information availability on co-exposures, specific assessments of unintentional mixtures seem unfeasible for registrants and downstream user. Hence, a generic approach or “mixture allocation factor” as proposed by [2, 8, 10, 56] could be a feasible option to address unintentional mixtures during the CSA. Legally, a MAF could be implemented in REACH Annex I describing the obligations for risk characterization during the CSA as proposed by [8, 21]. An allocation factor could be included during PNEC, PEC, RCR or $M_{\text{safe}}$ estimations (as these are communicated downstream) for each use of a substance to safeguard for co-exposures. It is to be assumed, that in particular those substances with RCRs close to 1 for the respective uses and compartments would be affected by such as factor. Defining the magnitude of such a factor, still requires analyses of, e.g. environmental monitoring data on concentrations, co-occurrences and effects. Also, the benefits and impacts on chemical safety assessments and resulting environmental emissions, and whether the most relevant substances would be addressed, have to be carefully analysed (see also Table 3).

Addressing emissions from production or sewage treatment plants

While REACH provides the prospective assessment of chemicals and sets the limits for a “safe” use, production volumes and potential emissions, further associated regulations are in close interaction. In order to avoid and minimize industrial emissions, production plant and industrial wastewater treatment plant operators are obliged to follow the Industrial Emissions Directive (IED, Directive 2010/75/EU [60]) and its Best available technique Reference Documents (BREFs), which set provisions for surveillance. They are also obliged to achieve a defined clearance level according to Directive 91/271/EEC on urban waste water treatment. Emission thresholds only exist for very few chemicals or sum parameters for groups (e.g. for benzenes). Therefore, proposals were made by the EU-project “Hazardous Chemicals in IED BREFs” (HAZBREF) [69], to improve the data basis for single chemicals. A register for substances emitted by industrial sites was proposed [58]. The German Pollutant Release and Transfer Register (PRTR) [72] and its European counterpart (E-PRTR) [68] could be used and extended as data source for emissions. This could support documentation and assessments of groups of substances and co-occurrences closer to the emissions via point sources.

A whole effluent assessment of multiple co-occurring substances is not done on a regular basis neither in industrial nor in municipal treatment plants. Although the problem of so-called micropollutants in sewage treatment plants has been well recognized, the focus is still on single substances [1]. A fourth sewage treatment stage is discussed as a generic solution to reduce the toxic pressure [35]. However, many substances enter environmental compartments via various other diffuse pathways and are not captured. Here, the environmental monitoring for selected priority substances under the Water Framework Directive (2000/60/EC) and Annex I to Directive 2013/39/EU on environmental quality standards (EQSs) could be improved. The existing specifications to...
calculate an EQSmix certainly help to address unintentional mixtures [17].

**Specific assessments by authorities to clarify possible mixture risks**

Following the existing instruments of REACH, specific assessments of unintentional co-occurrence in the environment can be conducted by authorities (ECHA and member states) in certain cases, i.e. when additional mixture risks or hazards are indicated. This could be done as part of the screening process in preparation of regulatory assessments, risk management option analyses (RMOA), or during the formal substance evaluation (SEV) to clarify justified concerns. For this purpose, groups of substances, co-occurrences, or mixture scenarios would need to be defined together with the suspected risks to justify inclusion on the CoRAP and a formal SEV. During substance evaluations, the respective CSRs and all other available information could be analysed for defined groups of substances. Subsequently, further data on uses, combined releases or exposures could be requested to be able to conclude on possible mixture risks [10].

Currently, ECHA increases efforts on grouping approaches for a common assessment in order to prioritize substances with similar properties (e.g. bisphenols or PFAS) for further regulatory actions and to avoid regrettable substitutions. Rudén et al. [56] proposed to strengthen group-wise assessments. Grouping could be used to define a sort of cumulative assessment groups of substances with similar structures or hazard profiles that might pose joint risks. Also, groups or scenarios of dissimilar substances could be defined that enter the environment via similar sources and/or are typically co-occurring in the environment.

In order to decide which unintentional mixture or co-occurrence is relevant for further assessments and possible regulation, the definition of a typical “priority” mixture and its key components is needed. Bunke et al. [10] proposed to define priority mixtures that have a high likelihood for combined risks on the basis of their composition, critical uses or relevant co-exposures exceeding predicted risks. For this purpose, risk modelling approaches as provided for example by Gils et al. [39] seem to be promising providing knowledge on relevant exposures on a European or regional scale and evidences for possible mixture risks. As basis, knowledge on production volumes, uses, substance properties would be needed. An improved environmental monitoring that considers environmental co-occurrences of substances and ways to deal with high temporal and spatial variability is also crucial.

**Targeted regulatory measures to address drivers of unintentional mixture risks**

In cases, where potential risks of co-occurring substances can be demonstrated, regulatory measures might be taken in order to reduce substance concentrations and impact on ecosystems using the instruments established within REACH.

Restriction seem to be the most suitable approach to regulate mixtures of “definable” composition as some experiences were gained. A restriction was already achieved for a group of 4 phthalates in 2017 (Annex XVII, entry 51) after a proposal of DK proving their joint toxicity, exposure and risks for human health, although the justification and proof of co-exposure was challenging. This way a threshold concentration for the sum of a group of substances is set, as e.g. also for the existing restriction for polyaromatic hydrocarbons (PAH) in consumer products. A recent example with respect to environmental risks is the envisaged restriction proposal for PFAS [33]. Currently, the focus for these defined co-exposure scenarios is on well-known and partly already regulated similar substances. This is in line with the proposal of [56] to “flag” suspected substances with similar properties as already restricted ones in order to regulate further relevant substances. However, restrictions could also be possible for groups of dissimilar substances with similar use and exposure profiles.

For those substances identified as SVHCs according to article 57 (PBT or ED), the identification process might be sped up by flagging substances as “suspected” belonging to a structural similar group as proposed by Rudén et al. [56]. However, groups can easily contain hundreds of substances with different hazard profiles as for example currently seen for the group of bisphenols. Although assessments may be conducted group-wise, a possible identification as SVHC would be based on the single substances’ properties. However, evidences for co-exposures have already been used as an argument during the assessment of the “equivalent level of concern” (ELOC) during SVHC identification according to article 57f (for example for the PFAS “HFPO-DA” [33]). Subsequently it would be an option to consider evidence on co-exposures of similar substances during the prioritization process for inclusion of a substance into Annex XIV and the following authorization procedure. In addition to the obligation to clarify common uses in the application for authorization, the consideration of co-exposures in the environment could be assessed during decision-making for granting or denying an authorization.
Conclusions and further needs

Clear mandates and guidance to address combined effects and exposures

Currently, REACH is a single substance-oriented framework and does not explicitly consider the possible joint additive effects and exposures of chemicals neither with respect to intended nor unintentional mixtures. Under the premise that chemicals are neither used alone nor are entering environmental compartments alone but co-occur and may produce joint effects and risks, clear mandates and legal obligations would be needed in the legal text as already proposed by Bunke et al. [10] and Rudén et al. [56]. Experiences under PPPR and BPR show that such a mandate could facilitate and trigger the further development of tools and ensure an enforcement of a mixture assessments. The development of detailed guidance and tools for the different actors under REACH, in particular DUs, is essential and would need to be integrated in the respective ECHA guidance documents as proposed by Bunke et al. [10] in more detail.

Sound data basis on co-occurrences via accessible common databases

As has become clear, extensive knowledge gaps have to be closed and data made available to the different actors along the supply chain (upstream and downstream) to enable single substance and mixture assessments for compliant safe use conditions. This concerns data on substance properties, uses, fate and behaviour, and amounts produced as well as environmental monitoring data. To reach this, the ECHA dissemination website [62], which currently compiles data on the publicly available substance properties, could be extended with available data on exposures, uses and typical formulations provided that these are made available by companies at least on a generic level. Here, it would be crucial that issues with confidential business information can be addressed in an appropriate way.

With respect to substance concentrations in environmental compartments more monitoring data are needed, as for example provided in a limited way via the retrospective media-oriented frameworks such as the WFD. The EU platform IPCHEM [70] is becoming central and besides data on single substances already includes data on co-occurrences. In order to better link exposures to effects on organisms in the environment and establish possible causal relationships, effect-based tools gain more and more consideration [7]. Furthermore, modelling approaches may allow for the identification of environmental co-occurrences and possible risks on regional or EU level on the basis of available data and again feed into a common data basis.

Safe use for intentional mixtures

From a prospective point of view, mixtures should be tackled at the source and emissions reduced before substances enter environmental compartments. Following the aims of REACH to ensure a high level of protection and the burden of proof for a safe use of chemicals during their life cycle, also the safe use of intentional mixtures should be ensured during the CSA by registrants and DUs. The proposed LCID methodology and SUMI reporting templates seem to be valuable communication tools for mixtures, but still need substantial improvements. They should be further followed up with respect to their acceptance and practicability to reach a transparent communication of information between the actors. Furthermore, case studies and analyses are warranted checking their soundness, benefits and impacts to ensure safe use amounts of formulated mixtures. Possible emissions, joint effects and possible risks for the environment arising from an intentional mixture need to be taken in account.

Generic mixture assessment factor to address unintentional co-exposures in CSA

Following the principal setup of REACH, also the unintentional co-occurrence of multiple chemicals in the environment should be considered during the CSA. This would account for the fact that single substances are not released to an unpolluted environment. Specific mixture assessments go beyond the information availability on mixture compositions, resources and the direct responsibilities of registrants or DU. Additionally, such assessments are challenging due to the high spatial-temporal variability of substances in the environment and the difficulty to trace substances back to their source and allocate responsible companies. A generic approach such as a mixture allocation factor (MAF) may be a feasible solution, although efforts are needed to define an appropriate size and assess its benefits, possible impacts and challenges for a possible implementation. Further considerations are needed whether such a factor would only address chemicals regulated under REACH and how to deal with co-occurrences with other substances such as pesticides or pharmaceuticals, which is outside the scope of REACH.

Specific assessment and targeted regulation of identified drivers of possible mixture risks

In certain cases when indications for remaining risks are available and drivers are identified, targeted mixture risk assessments could be done by the regulatory authorities using the established REACH instruments as for example substance evaluations and subsequent restrictions if
regulatory measures are needed. This implies comprehensive data on substance’s co-occurrences and properties and the definition and prioritization of relevant components of an unintentional environmental mixture. Assessments can be done on the basis of the current legal obligations. Some experiences are already available and current efforts with respect to grouping and prioritization approaches would need to be stepped up.

Increase cross-talk between frameworks

A (formalized) cross-talk between substance and media-oriented frameworks would be needed to link retrospective data on environmental occurrences and effects better to the prospective risk management measures for substances and mixtures of concern. Overall, a close interaction of the different substance and media-oriented regulations is needed and should be considered when analysing options to assess and regulate mixtures of concern.

Abbreviations

BP(R): Biocidal products (regulation); BREFS: Best available technique Reference Documents in the context of the Industrial Emissions Directive; CA: Concentration Addition; CCH: Compliance check of dossiers, conducted by ECHA for certain selected registration dossiers; CLP: Classification, Labelling and Packaging of substances and mixtures (EC 1272/2008) regulation; CoRAP: ECHA’s Community Rolling Action Plan; CSA: Chemical safety assessment; CSR: Chemical safety report; DU: Downstream user; EC: European Commission; ECHA: European Chemicals Agency; EC10: Concentration, where 10% effect is observed; ED: Endocrine disrupters; ELOC: Equivalent level of concern during SVHC identification under REACH; EQS: Environmental quality standard in the context of the water framework directive; eSOS: Extended safety data sheet; HCBM: Hydrocarbon block methodology; IA: Independent Action; IED: Industrial Emissions Directive; LCID: Lead component identification methodology; LOEC: Statistically determined and tested lowest observed effect concentration; MAF: Mixture assessment/allocation factor; MATS: Mixture Assessment Triggering Substances; MCR: Maximum cumulative ratio; Msafe: Maximum safe use amount (kg/day or kg/year); MSC: Multi constituent substances; NOEC: No observed effect concentration, lowest tested concentration with no significant effect; OCs: Operational conditions; P(H)S: Priority (hazardous) substances in the context of the water framework directive; PBT/vPvB: Substances, which are persistent, bioaccumulative and toxic/very persistent and very bioaccumulative according to REACH Annex XVII; PC: Product category; PEC: Predicted Environmental Concentration; PECV: Prediction of the Environment Concentration; PECvV: Substances, which are persistent, mobile and toxic/very persistent and very mobile; PNEC: Predicted No Effect Concentration; PPP(R): Plant protection products (regulation); PRTR: German Pollutant Release and Transfer Register; QSR: Quantitative structure activity relationships; RCR: Risk characterization ratio; REACH: Regulation EC 1907/2006; SME: Small and medium-sized enterprises; spERCs: Specific environmental release categories; SUMI: Reporting template for the safe use of mixtures information; SVHC: Substances of very high concern according to REACH; TPE: Testing proposal evaluation, conducted by ECHA if tests are proposed by registrant; UVCB: Substances of unknown or variable composition and biological materials.

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Disclaimer

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Authors’ contributions

All authors contributed with their expertise to the manuscript. EH mainly contributed with conceptualization of figures and text and writing the final version of the manuscript. WJ in particular contributed with details on exposure assessments and supply chain requirements and designed the figures. JS in particular reviewed, adapted and improved the entire manuscript and fine-tuned all the figures. All authors read and approved the final manuscript.

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Availability of data and materials

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