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# Chapter 31

## LCA of Chemicals and Chemical Products

Peter Fantke and Alexi Ernstoff

**Abstract** This chapter focuses on the application of Life Cycle Assessment (LCA) to evaluate the environmental performance of chemicals as well as of products and processes where chemicals play a key role. The life cycle stages of chemical products, such as pharmaceuticals or pesticide active ingredients, are discussed and differentiated into extraction of abiotic and biotic raw materials, chemical synthesis and processing, material processing, product manufacturing, professional or consumer product use, and finally end-of-life treatment. LCA is put into perspective of other chemicals management frameworks and concepts including risk assessment, green and sustainable chemistry, and chemical alternatives assessment. A large number of LCA studies focuses on contrasting different feedstocks or chemical synthesis processes, thereby often conducting a cradle to (factory) gate assessment. While typically a large share of potential environmental impacts occurs during the early product life cycle stages, potential impacts related to chemicals that are found as ingredients or residues in products are dominated by the product use stage. Finally, methodological challenges in LCA studies in relation to chemicals are discussed from the choice of functional unit, over defining the system boundaries, quantifying emissions for many thousand marketed chemicals, to characterising these emissions in terms of toxicity and other impacts, and finally interpreting chemical-related LCA results. The chapter is relevant for LCA students and practitioners who wish to gain basic understanding of LCA studies of products or processes with chemicals as a key aspect.

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## 31.1 LCA and Chemicals: Introduction and Context

### 31.1.1 Chemicals and Their Relevance in Society

Chemicals are everywhere. Almost every second a new entry is added to the list of more than 100 million unique chemicals and substances registered in the Chemical Abstracts Service (CAS; [www.cas.org](http://www.cas.org)), the world's authority on chemical information. Since industrialisation, the welfare of modern society largely builds on extensively mining minerals and fossil fuels including coal, petroleum and natural gas to produce large quantities of synthetic chemicals ('synthetic' simply means man-made and should not be confused with 'artificial', which implies that a chemical does not occur naturally). Consequently, the enormity and diversity of the chemical industry is astounding and poses various challenges for the management of environmental and human health impacts related to chemicals production and use. In this chapter, we outline important aspects to know about chemicals in the context of LCA.

Fundamentally, chemicals are substances composed of one or more atoms, and make up every material thing on earth—including our bodies. The atomic composition of chemicals classifies them essentially as 'organic' (chemicals with molecules built on a skeleton of interlinked carbon atoms and primarily consisting of carbon, oxygen, and hydrogen) and 'inorganic' (chemicals with molecules generally lacking carbon-to-carbon bonds, but instead based on the rest of the elements, including metals). In this sense, 'organic' has nothing to do with 'organic food' or 'organic farming' or 'organic lifestyle' as these terms generally refer to promoting sustainability. The atomic composition, molecular structure and ionisation (positive/negative charge) all influence chemical reactivity and behaviour in the environment as well as in living organisms. Because of this, chemical behaviours can be predicted and tested, and chemicals can be designed by industries to fulfil biological (e.g. medical) and physical (e.g. solvent) functions.

Chemicals may also be classified according to functional groups (e.g. alcohols, amines, acids and bases), structural groups (e.g. polycyclic aromatic hydrocarbons), physical structure (e.g. nanotubes), feedstock sources (e.g. petrochemicals derived from fossil fuels, biochemicals derived from starch- and sugar-based feedstocks), physicochemical properties (e.g. volatile, lipophilic), use function (e.g. surfactants, warfare agents), means of creation (e.g. reaction intermediates, metabolites), main economic sector (e.g. cosmetics, agrochemicals), toxicity endpoints (e.g. carcinogens, neurotoxins, endocrine disruptors), and other aspects.

Established nomenclatures or patent names can be used to name chemicals. Most chemicals have an assigned CAS Registry Number except some metabolites of natural processes or grouped chemicals such as polychlorinated dibenzofurans. CAS numbers are the most discriminant method for chemical reference. Of the chemicals registered by CAS, more than ten thousand are currently in commercial use, some with annual production volume of millions of tonnes, while most chemicals are produced at less than thousand tonnes per year. Worldwide, the



68 production of chemicals has risen to several hundred million tonnes per year and  
69 sales were valued in 2013 at 3156 billion Euro with an average annual growth of  
70 10.3% between 2003 and 2012 (CEFIC 2014). China dominates world chemical  
71 sales with a share of 33.2% followed by the European Union (16.7%), USA  
72 (14.8%), and Japan (4.8%) in 2013.

73 Over the last decades, there has been a shift in global chemicals production. As  
74 an example, polychlorinated biphenyls (PCBs) have been replaced by chlorinated  
75 paraffins in various applications. While PCBs have been primarily produced in  
76 USA and Europe with a total historical production volume of 1.3 million tonnes  
77 between 1930 and 1995, chlorinated paraffins are almost exclusively produced in  
78 China and reach production volumes of more than one million tonnes per year  
79 (Fantke et al. 2015). Databases, such as the European Chemicals Agency (ECHA)  
80 Registered Substances database ([echa.europa.eu/information-on-chemicals](http://echa.europa.eu/information-on-chemicals)), the  
81 Household Product Database ([householdproducts.nlm.nih.gov](http://householdproducts.nlm.nih.gov)), Hazardous  
82 Substances Data Bank through ToxNet ([toxnet.nlm.nih.gov](http://toxnet.nlm.nih.gov)), and the Chemical and  
83 Product Categories Database ([actor.epa.gov/cpcat/](http://actor.epa.gov/cpcat/)) attempt to keep track of  
84 chemicals, their uses, properties and/or toxicity, but large data gaps still remain.

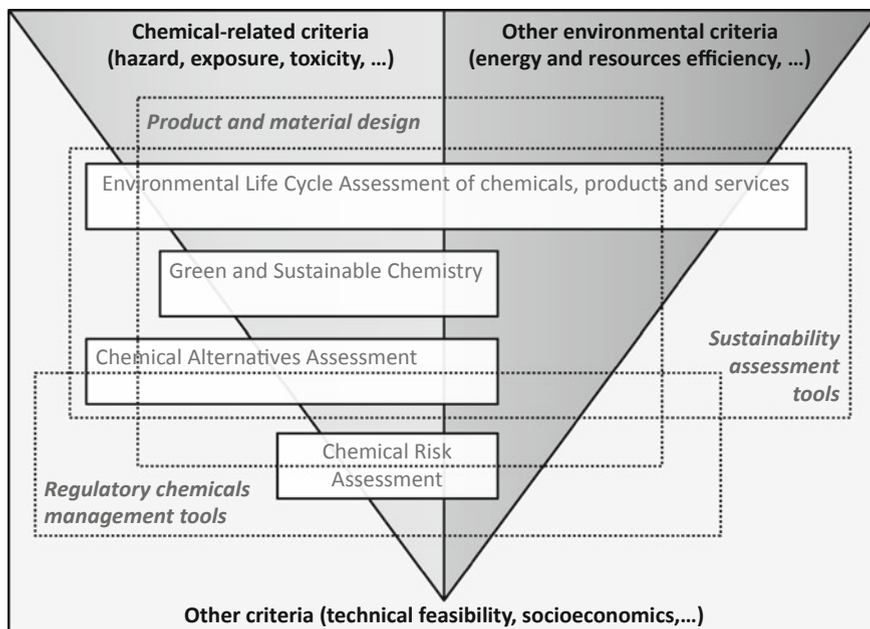
85 The chemical and pharmaceutical industries are a major driver of the welfare of  
86 modern society and scientific progress. These industries rely on the extraction,  
87 purification and synthesis of both naturally occurring and synthetic chemicals and  
88 are among the largest and most influential economic sectors at the global scale.  
89 Main production segments are petrochemicals (e.g. benzene, styrene), consumer  
90 chemicals (e.g. detergents, fragrances and flavours), speciality chemicals (chemicals  
91 used for providing a special performance or effect, e.g. paints, dyes, adhesives),  
92 basic inorganics (fertilisers, industrial gases like nitrogen and oxygen), and poly-  
93 mers (e.g. plastics, synthetic rubber and fibres). One of the largest segments is the  
94 production of organic chemicals with, e.g. formaldehyde, aromatics, acids, alcohols  
95 and esters providing the building blocks for drugs, agrochemicals, cosmetics and  
96 many other applications.

97 Along with societal advantages, the rise of chemical industries has also caused  
98 various undesirable consequences. Health impacts of air pollution are increasing  
99 worldwide and there is currently insufficient information to fully assess the impacts  
100 of chemicals on humans and the environment. Rachel Carson's book *Silent Spring*  
101 published in 1962 documented the detrimental impacts of chemicals on wildlife and  
102 humans, especially related to using synthetic organic pesticides, and marked a  
103 major change in public awareness that eventually inspired regulation of industry  
104 and for example the creation of the United States Environmental Protection  
105 Agency. Since that time, a remarkable amount of research correlates and demon-  
106 strates impacts on human and ecosystem health as well as the environment (e.g. the  
107 ozone layer) caused by intentional and unintentional chemical releases both indoors  
108 and outdoors. Some reported impacts are directly related to the chemical industry,  
109 whereas other impacts are related to the use or disposal of chemicals by other  
110 industries. In the following sections, we overview strategies for chemical man-  
111 agement, focusing particularly on life-cycle assessments of chemicals production  
112 processes and chemical products.

### 31.1.2 Chemicals Management in Relation to LCA

Depletion of the ozone layer by chlorofluorocarbons used as refrigerants and solvents, soil and water pollution with heavy metals from ore mining and processing, pesticide emissions and residues in food, the formation of dioxins by incomplete combustion processes, and leaching of fertilisers into groundwater are just examples of the many problems associated with chemical releases to the environment. Hence, managing human and environmental risks posed by chemicals that are potentially toxic or may lead to other impacts is a major concern of regulators, industries, consumers and other stakeholders. As a consequence, the chemicals industry is one of the most regulated industries with main focus on regulating chemicals in consumer products and minimising chemical emissions to the indoor (workplace, public buildings and household) and outdoor environments along product life cycles. In the context of chemicals management, risk is defined as the probability of a chemical to cause an adverse effect (hazard) occurring as a result of a given contact between the chemical and humans or the environment (exposure). In reality, risks associated with chemical emissions from a given product or process can arise at specific points in space and time and depend on chemical background concentrations due to all release sources. In LCA, information on emission location and time as well as information on background concentrations, e.g. from sources outside the considered product system, is usually not available. Hence, modelled impacts in LCA are not interpreted in terms of actual risk, i.e. real environmental effects, but in terms of ‘potential impacts’ (Chap. 10) used as environmental performance indicators for comparing and optimising products or systems with respect to a defined functional unit (Hauschild 2005). However, models applied in LCA can also be advanced and adapted to consider background concentrations as well as spatiotemporal resolution (e.g. daily or seasonal changes), and in such cases estimated potential impacts can be interpreted as estimates of actual risk.

Chemicals management occurs from local to global scale, from specific product–chemical combinations to entire industries and from raw material acquisition to waste handling, depending on the intended scope and purpose. The Montreal Protocol on Substances that Deplete the Ozone Layer ([ozone.unep.org](http://ozone.unep.org)) and the Stockholm Convention on Persistent Organic Pollutants (POPs; [www.pops.int](http://www.pops.int)) are examples of global chemicals management treaties, whereas the Registration, Evaluation, Authorisation, and Restriction of Chemicals (REACH) is a recent example of an international legislative framework for managing industrial chemicals in the European Union. At all levels and scopes, effective chemicals management relies on assessment tools and guiding principles to ensure consistency and the achievability of defined goals. There are many examples of chemicals assessment tools and guidance, such as risk assessment, green and sustainable chemistry, chemical alternatives assessment, life cycle assessment, and a market for entrepreneurs to create industry-specific interfaces and applications. In the following sections, risk assessment (Sect. 31.1.3), green and sustainable chemistry



**Fig. 31.1** Conceptual relationships of main chemical management tools

155 (Sect. 31.1.4), and chemical alternatives assessment (Sect. 31.1.5) are discussed as  
 156 commonly used chemical management tools that have both complementary and  
 157 overlapping aspects with LCA as illustrated in Fig. 31.1.

### 158 31.1.3 Risk Assessment and Safety

159 Chemical risk assessment—also referred to as chemical safety assessment—is  
 160 implemented in various regulatory frameworks and is one of the most widely used  
 161 chemicals management tools. Risk assessment (*‘How risky is a situation?’*) as an  
 162 integral part of risk management (*‘What shall we do about it, if a situation is*  
 163 *risky?’*) essentially emerged at the start of the nineteenth century from studying  
 164 hazards and risks associated with different occupations. Risk assessment mainly  
 165 consists of hazard identification, dose-response assessment, exposure assessment  
 166 and risk characterisation. Depending on the context, ‘risk’ and ‘safety’ have dif-  
 167 ferent meanings with regulatory policy commonly seeking to minimise risk while  
 168 optimising safety. In this context, risk is generally defined as the probability of  
 169 harm, whereas safety is described as the absence of harm (Embry et al. 2014).

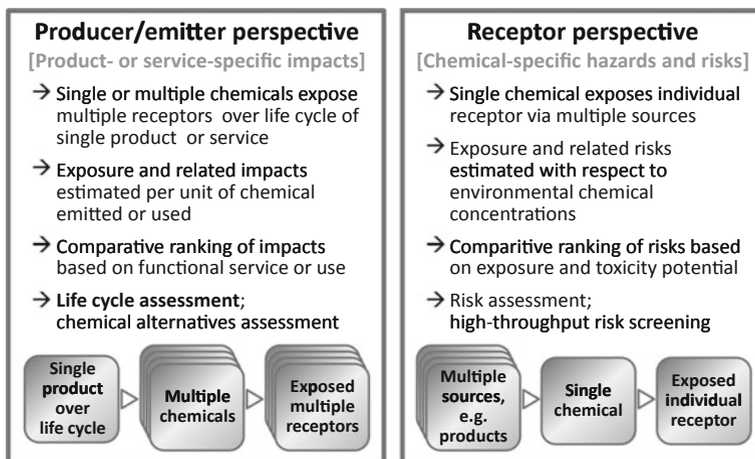
170 Chemical ‘safety’ is defined by legislators or regulators and can vary from  
 171 country to country and evolves over time as science progresses. In this sense, ‘safe’

172 is not synonymous with ‘natural’ as it is often perceived. In fact, using the word  
173 ‘natural’ is misleading in the context of chemical safety (and LCA) and there are  
174 many naturally occurring chemicals that have very harmful properties like arsenic,  
175 nicotine or radon. As a consequence, we need to acknowledge that it is not always  
176 the ‘natural’ chemicals or solutions that are most ‘environmentally friendly’—a  
177 common misconception in different science-policy fields and among consumers.  
178 Defining safety *thresholds*, e.g. chemical concentrations in different environmental  
179 media (e.g. ambient air, soil, water) or in food, is a common strategy in chemical  
180 risk assessment, and generally refers to levels below which a risk is considered  
181 ‘safe’ by a risk manager, meaning that any risk below threshold is regarded as  
182 ‘acceptable’. As an example, chemical exposure resulting in one additional cancer  
183 case or less over lifetime in a population of one million people is regarded as an  
184 acceptable risk, i.e. safe, in the U.S. (van Leeuwen and Vermeire 2007). Using units  
185 like ‘part per million’ (ppm) as in one cancer case in a million or ‘part per billion’ is  
186 common for describing (very small) amounts of chemicals in the environment. To  
187 get an impression of how much one ppm actually is, we can use 1 teaspoon of salt  
188 (5.5 g) in 5.5 tonnes of potato chips corresponding to one part of salt per one  
189 million parts of potato chips.

190 Thresholds are also applied when managing environmental systems and for  
191 developing chemical pollution control strategies, such as allowable nutrient releases  
192 from wastewater treatment plants or setting greenhouse gas emission targets, or in  
193 the context of ‘planetary boundaries’ in an attempt to assess if the pressure from  
194 chemical pollution (analogous to the amount of receiving environment required to  
195 dilute pollution to a threshold level) exceeds a planetary boundary (analogous to the  
196 amount of receiving environment available) for a ‘safe operating space’ for human  
197 activities (MacLeod et al. 2014). Chemical pollution levels have recently been  
198 expressed as ‘chemical footprints’ that can be compared with respective planetary  
199 or other boundaries for chemical pollution (Posthuma et al. 2014) to assess how  
200 companies or nations perform with respect to different chemicals management  
201 issues.

202 Risk assessment approaches take a receptor perspective (Fig. 31.2, right-side  
203 box), where thresholds are set in order to protect specific receptors, i.e. exposed  
204 humans or ecosystem species. In a receptor perspective, all relevant sources of a  
205 chemical or target chemicals are typically considered. In contrast, impact assess-  
206 ment tools in LCA are generally not receptor-oriented or threshold-based. This is  
207 because LCA takes a ‘producer’ (or ‘emitter’) perspective (Fig. 31.2, left-side box)  
208 by comparing potential impacts relative to each other across compared products and  
209 life cycle stages, aiming at minimising impacts considering various receptors (entire  
210 human populations, freshwater ecosystems, marine ecosystems, etc.). Differences  
211 and commonalities of risk assessment and LCA have been contrasted elsewhere  
212 (e.g. Bare 2006; Pennington et al. 2006), and there are several attempts combining,  
213 blending or integrating both concepts (Harder et al. 2015).

214 An increasing number of chemicals is approved for use in commerce, e.g. in  
215 food contact materials, but often lacks adequate information to characterise risks  
216 (Neltner et al. 2013). In response, high-throughput screening (‘first tier’



**Fig. 31.2** Examples and underlying characteristics of dichotomous perspectives followed in different chemicals management approaches

217 assessments) of chemical risks has emerged as a strategy for prioritising and  
 218 ranking chemicals for more in-depth study ('higher-tier' assessments). First-tier  
 219 screening usually relies on ranking chemicals with respect to hazard (e.g. chemical  
 220 toxicity) combined with estimates of exposure. 'High-throughput' refers to pro-  
 221 cessing dozens to thousands of chemicals via resource efficient methodologies, such  
 222 as robotic in vitro bioassays (instead of animal in vivo experiments) and low-tier  
 223 computational models relying on databases (instead of data-intensive complex and  
 224 time-consuming modelling). LCA impact assessment models have been used in  
 225 high-throughput risk screening offering dual purpose and a promising area of  
 226 interdisciplinary overlap to manage chemical risks (e.g. Shin et al. 2015).

### 227 31.1.4 Green and Sustainable Chemistry

228 'Green chemistry' is a concept that was coined by the U.S. Environmental  
 229 Protection Agency in the early 1990s in response to the Pollution Prevention Act  
 230 and increasing attention to chemical pollution. This concept builds upon a set of 12  
 231 Principles of Green Chemistry defined by Anastas and Warner (1998) aiming at  
 232 reducing or eliminating hazardous substances in the design, manufacture and ap-  
 233 plication of chemical products. Thereby, 'green' refers to more environmentally  
 234 benign (less hazardous) chemicals. The concept of 'sustainable chemistry' is  
 235 broader than the scope of green chemistry and strives towards 'eco-efficiency'. In  
 236 addition to chemical hazards, sustainable chemistry centrally focuses on optimising  
 237 the use of finite resources, while reducing environmental impacts of chemical  
 238 production (OECD 2012). Sustainable chemistry—sometimes also referred to as

239 sustainable chemistry and engineering—is rooted in the concept of Sustainable  
240 Development established in Rio de Janeiro in 1992 at the United Nations  
241 Conference on Environment and Development and are guided by 9 Principles of  
242 Green Engineering postulated at the Sandestin conference (Abraham and Nguyen  
243 2003).

244 Green and sustainable chemistry are concepts focusing on the technological  
245 approaches aiming at the reduction of resource consumption and pollution pre-  
246 vention in chemical production processes rather than focussing on the assessment of  
247 chemicals in the environment. Hence, green and sustainable chemistry—often  
248 relying on comparing qualitative or semi-quantitative indicator results—are pri-  
249 marily applicable in the design phase of products to guide innovation and to support  
250 sustainable production goals.

251 Green chemistry in relation to LCA has been discussed in more detail elsewhere  
252 (e.g. Anastas and Lankey 2000). In summary, compared to green and sustainable  
253 chemistry, LCA aims at fully quantifying potential impacts associated with a  
254 chemical product or production system over its entire life cycle. Using LCA in early  
255 stages of chemical product and process design of various sectors including  
256 emerging technologies (e.g. bio- and nanotechnology) has provided insight into the  
257 relationship between chemical and process parameter selection and related impacts  
258 on humans and the environment (Kralisch et al. 2015). LCA results have moreover  
259 demonstrated that quantitative methods are needed to assess the environmental  
260 performance of ‘green’ chemicals (Tufvesson et al. 2013). This is especially rele-  
261 vant as green chemistry usually focuses on optimisation of (production) processes,  
262 including some specific end-of-life problems related to chemicals, which may still  
263 risk sub-optimisation when a full life cycle perspective is lacking.

264 Using LCA in early product development stages, for example before a product  
265 has been created and marketed, comes with methodological and practical chal-  
266 lenges, such as low data availability, uncertainty related to future product appli-  
267 cations, and unclear scale of production for a changing market. Therefore, LCA has  
268 mostly been applied to chemical products and processes that are already well  
269 established and operational at the market scale, which leads to LCA results often  
270 being reactive instead of proactive.

### 271 ***31.1.5 Chemical Alternatives Assessment***

272 Chemical alternatives assessment (CAA) aims to identify, compare and select safer  
273 alternatives to substitute (replace) harmful chemicals in materials, processes and  
274 products on the basis of their hazards, performance and economic viability (Hester  
275 and Harrison 2013). CAA emerged from the U.S. Environmental Protection  
276 Agency’s Design for Environment (DfE) program in the late 1990s to promote less  
277 hazardous chemicals in various products and applications, and to avoid unintended  
278 consequences of harmful alternatives resulting in incremental improvements or  
279 even ‘regrettable substitution’ situations (Fantke et al. 2015). Ideally, CAA tools

280 would evaluate hazard, exposure, life cycle and social impacts, economic feasibility  
 281 and technical performance of alternative solutions, and consider chemicals, materi-  
 282 als, products or technologies, and behavioural changes as viable solutions options.  
 283 In reality, however, most CAA tools focus only on comparisons of hazard scores  
 284 and exclusively consider chemicals as potential solutions. Several existing CAA  
 285 tools have been compiled into the OECD Substitution and Alternatives Assessment  
 286 Toolbox ([www.oecdsaatoolbox.org](http://www.oecdsaatoolbox.org)).

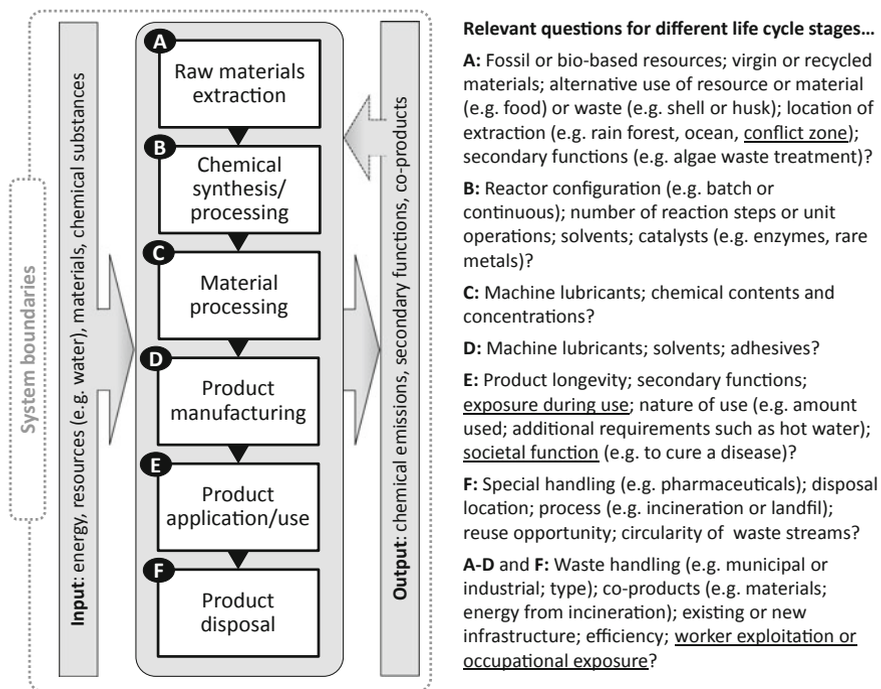
287 The concept of ‘acceptable risk’ (as applied in risk assessment) is usually  
 288 avoided in CAA in order to support selecting *relatively* less hazardous chemicals  
 289 and materials in products (Whittaker 2015). Despite the current focus on assessing  
 290 chemical hazard, including exposure, life cycle, and social considerations are lately  
 291 also gaining more attention (Jacobs et al. 2016), focusing the CAA discussion  
 292 around using more quantitative and chemical function-based methods and tools  
 293 (Tickner et al. 2015). However, the need for rapid screening of numerous viable  
 294 alternative solutions prevents CAA from simply adopting the use of LCA tools due  
 295 to high complexity and data demand.

296 CAA is mainly used to identify and evaluate solutions to hazardous chemicals in  
 297 products that have been targeted for phase-out, and to inform early product  
 298 development to minimise reliance on hazardous chemicals. With that, CAA takes  
 299 the ‘producer’ perspective similarly to LCA (Fig. 31.2, left-side box), focusing on  
 300 the impact of chemicals and their alternatives on various receptors. The main  
 301 difference between CAA and LCA is that while CAA focuses on seeking for viable  
 302 alternatives to harmful chemicals, LCA considers the life cycle of whole products  
 303 or processes not focusing specifically on the content of one or more chemicals that  
 304 might be considered ‘hazardous’, but instead evaluating the overall product or  
 305 process environmental performance.

## 306 31.2 LCA Applied to Chemicals

307 Chemicals play a central role in the LCA framework for different reasons. Hundreds  
 308 of chemical emission (inventory) flows typically occur along the life cycle of  
 309 products or systems (Fig. 31.3) and are quantified as part of the Life Cycle  
 310 Inventory (LCI; see Chap. 9) phase. Chemicals are also often precursors of product  
 311 materials, and input for manufacturing and disposal processes. Chemical emissions  
 312 associated with energy conversion during manufacturing, transport of goods and  
 313 end-of-life treatment processes often dominate overall emission profiles for many  
 314 product categories resulting in potential environmental impacts that can be char-  
 315 acterised in the Life Cycle Impact Assessment (LCIA; see Chap. 10) phase.

316 Chemicals contribute to nearly all LCIA impact categories affecting human  
 317 health and ecosystem quality as two main areas of protection in LCA, with re-  
 318 sources (e.g. water) being an area that is usually not relevant for chemicals  
 319 (Hauschild et al. 2013). In LCIA, chemicals contribute to global warming, strato-  
 320 spheric ozone depletion, formation of photochemical ozone in the troposphere, air



**Fig. 31.3** Generic life cycle stages and system boundaries for chemical products or materials and LCA-related questions. In some cases, chemical processing may be followed by material production (e.g. polymers) before manufacturing a product (e.g. plastic bottles), while in other cases, chemicals (e.g. solvents) may be directly added to products or product manufacturing processes. *Underlined topics* are mostly lacking methods or not included in environmental LCA studies

321 pollution (via respiratory particles and precursors), aquatic and terrestrial acidifi-  
 322 cation and eutrophication, and last but not least human toxicity and aquatic and  
 323 terrestrial ecotoxicity. Only a handful of chemicals are associated with the majority  
 324 of abovementioned impact categories, such as carbon dioxide, methane and other  
 325 greenhouse gases contributing to global warming impacts or ammonia, nitrogen  
 326 oxides, phosphate and some other nitrogen and phosphorus containing chemicals  
 327 contributing to aquatic eutrophication. In contrast, thousands of chemicals can be  
 328 characterised as potentially toxic to humans and/or ecosystems (Rosenbaum et al.  
 329 2008). This is, however, only a small fraction of the tens of thousands of com-  
 330 mercially relevant chemicals.

331 The generic life cycle stages shown in Fig. 31.3 are applicable to a chemical  
 332 product (e.g. pharmaceutical or dye) or material (e.g. polymer), from raw materials  
 333 extraction to product disposal, often referred to as ‘cradle to grave’ (Fig. 31.3,  
 334 stages A–F). A ‘cradle to grave’ LCA study can provide valuable insight regarding  
 335 which stages dominate the impacts throughout a product life cycle. Some of these



336 life cycle stages, however, may not be relevant depending on the goal (Chap. 7) and  
337 scope (Chap. 8), and the product system under study. For example, the ‘material  
338 processing’ stage may not be relevant in cases where a chemical is directly added  
339 into a product as an ingredient, such as fragrances in cosmetics or detergents in  
340 cleaning products. As another example, the ‘product application/use’ or ‘product  
341 disposal’ stages may not be relevant for comparing the environmental performance  
342 of chemical synthesis or production processes as long as the compared processes do  
343 not influence the chemical amount used in a product or for product disposal.

344 An LCA study from raw material extraction to chemical product manufacturing,  
345 i.e. without considering product use and disposal stages, is referred to as ‘cradle to  
346 gate’ (Fig. 31.3, stages A–D), which refers to the ‘gate’ of the manufacturing or  
347 production facility (which could be the ‘gate’ of a chemical or product ‘factory’,  
348 depending on the focus of the study). In Table 31.1, different assessment scopes for  
349 LCA studies focusing on chemicals in materials, products and processes are con-  
350 trasted and associated with relevant chemicals management questions.

351 LCA can help identify a variety of impacts associated with chemical production,  
352 use, and disposal, that are either intrinsic to a chemical (e.g. toxicity potential) or  
353 related to supporting industrial chemical processes (e.g. water consumption,  
354 greenhouse gas emission). The main uses of LCA for managing chemicals and  
355 chemical processes are to compare impacts between products or services, or to  
356 identify ‘hot spots’ within a life cycle that contributes greatly to the impacts of a  
357 product or service. With respect to chemicals, LCA can be applied to various  
358 combinations of the generic life cycle stages in Fig. 31.3 depending on the LCA  
359 study goal and chosen system boundaries. In some cases, individual life cycle  
360 stages and associated inputs or outputs may be skipped or not considered important  
361 for the defined system. The chemical industry developed a guidance document to  
362 support the assessment of the environmental performance of chemical products  
363 based on attributional LCA, i.e. referring to process-based modelling and excluding  
364 market-mediated effects (WBCSD 2014).

365 In the following sections, an overview is given of how LCA has been applied to  
366 consider these various life cycle stages and the general lessons learnt from these  
367 studies. Thereby, LCA can be used to compare impacts at the level of chemicals in  
368 materials, products and formulations or at the level of chemical synthesis and  
369 production processes.

### 370 **31.2.1 Chemicals in Materials, Products, and Formulations**

371 A subset of materials, products, formulations (combination or mixture of chemicals)  
372 and processes are intrinsically reliant on the functionality of key chemical ingre-  
373 dients. In this section, main trends are summarised in using LCA- or LCA-based  
374 methodologies. This may include also partial LCA studies, e.g. methods only  
375 considering a subset of life cycle stages (i.e. cradle to gate or gate to gate), with  
376 focus on chemicals in materials, products and formulations.

**Table 31.1** Relevant life cycle assessment scopes and life cycle stages for selected chemicals management questions and example studies

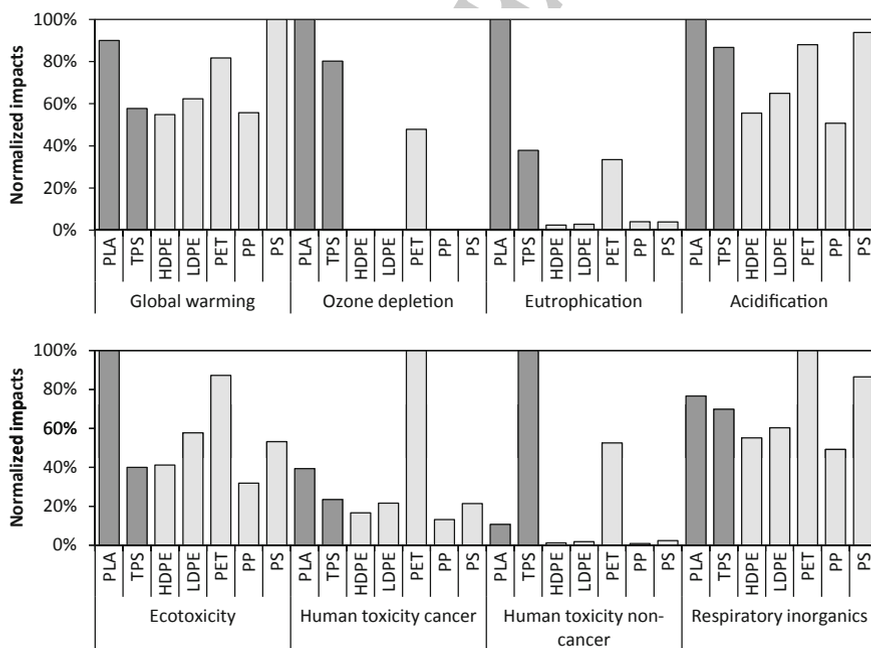
Chemicals management questions	Assessment scopes and considered life cycle stages	Example studies
What is the environmental performance of different products with respect to chemical emissions?	<ul style="list-style-type: none"> <li>• Cradle to grave</li> <li>• Stages A–F (Fig. 31.3)</li> <li>• Focus on chemicals consumption and emissions</li> </ul>	<ul style="list-style-type: none"> <li>• Cleaning products (Van Lieshout et al. 2015)</li> <li>• Textiles (Roos et al. 2015)</li> </ul>
What are the environmental profiles of the production of different chemicals?	<ul style="list-style-type: none"> <li>• Cradle to (factory or consumer) gate</li> <li>• Stages A–D or a subset of these stages (Fig. 31.3)</li> <li>• Focus on chemical manufacturing</li> </ul>	<ul style="list-style-type: none"> <li>• Pharmaceuticals (Wernet et al. 2010)</li> </ul>
Which life cycle stage of a chemical product life cycle contributes most to environmental impacts?	<ul style="list-style-type: none"> <li>• Cradle to grave</li> <li>• Hotspot analysis including stages A–F (Fig. 31.3)</li> <li>• Focus on chemicals as products</li> </ul>	<ul style="list-style-type: none"> <li>• Plant protection products (Geisler et al. 2005)</li> </ul>
Which chemical synthesis and/or manufacturing processes contribute most to environmental impacts?	<ul style="list-style-type: none"> <li>• Cradle to (factory) gate</li> <li>• Hot-spot analysis including stages A–B or A–C (Fig. 31.3)</li> <li>• Focus on chemical manufacturing</li> </ul>	<ul style="list-style-type: none"> <li>• Pharmaceuticals (De Soete et al. 2014)</li> <li>• Nano-materials (Pati et al. 2014)</li> </ul>
Which life cycle stage of a chemical in a product contributes most to human exposure?	<ul style="list-style-type: none"> <li>• Cradle to grave</li> <li>• Partial LCA (only human e.g. exposure estimates) including stages A–F (Fig. 31.3)</li> <li>• Focus on chemicals in products</li> </ul>	<ul style="list-style-type: none"> <li>• Cosmetics (Ernstoff et al. 2016a)</li> </ul>
Which feedstock provides the most environmentally friendly substrate for biochemical synthesis?	<ul style="list-style-type: none"> <li>• Cradle to (factory) gate</li> <li>• Stages A or A–B (Fig. 31.3)</li> <li>• Focus on chemicals and raw materials consumption</li> </ul>	<ul style="list-style-type: none"> <li>• Acrolein (Cespi et al. 2015)</li> <li>• PET (Akanuma et al. 2014)</li> </ul>

377 LCA studies have focused on pharmaceuticals (e.g. De Soete et al. 2014),  
 378 cleaning products (e.g. Van Lieshout et al. 2015) and pesticide formulation products  
 379 (e.g. Geisler et al. 2005) as examples of products where chemicals provide the  
 380 main product functions. Other LCA studies on chemicals with in-product functions  
 381 include studies focusing on flame retardants in electronics (Jonkers et al. 2016),  
 382 nano-materials used in bandages and cosmetics (Botta et al. 2011), and polymers  
 383 used in food packaging (Hottle et al. 2013). Chemicals required for industrial  
 384 processes have also been assessed in LCA studies, including industrial solvents  
 385 (Zhang et al. 2008) and chemicals used for the production of treated water, oil and

386 gas, printing paper and dyed textiles (e.g. Alvarez-Gaitan et al. 2013; Parisi et al.  
 387 2015).

388 When analysing LCA studies on chemical-based functions, a few generalisations  
 389 emerge. For example, it is important to consider life cycle thinking early on in the  
 390 design phase of products and processes whenever possible and it has been shown  
 391 that simplified tools may help in this process (e.g. De Soete et al. 2014).  
 392 Furthermore, it has been demonstrated that hybridised LCA tools or metrics can be  
 393 useful to improve communication and management for specific stakeholders (e.g.  
 394 Alvarez-Gaitan et al. 2013).

395 Several LCA studies indicate that being sceptical of services deemed ‘green’ or  
 396 ‘sustainable’ is crucial, especially when an LCA has not yet been performed. Case  
 397 studies on, e.g. ‘green’ solvents (Zhang et al. 2008) or ‘sustainable’ bio-based  
 398 chemicals and materials (e.g. Hottle et al. 2013) demonstrate that materials and  
 399 products guided by principles of ‘sustainability’, ‘eco-friendliness’ or ‘green  
 400 chemistry’ can have significant, but often disregarded or unassessed, environmental  
 401 impacts. An example is given in Fig. 31.4, where environmental life cycle impacts of  
 402 petro- and bio-based polymers are contrasted based on data from Hottle et al. (2013).



**Fig. 31.4** Impact scores for LCAs of two bio-based polymers (*dark bars*; PLA Poly-lactic acid, TPS Thermoplastic starch) compared to petroleum-based polymers (*light bars*; HDPE High-density polyethylene, LDPE Low-density polyethylene, PET Polyethylene terephthalate, PP Polypropylene, PS Polystyrene) per kg of produced granule, normalised for each category to the polymer with highest impacts (based on data from Hottle et al. 2013)

403 According to this study, bio-based polymers lead to higher impacts than petro-based  
404 polymers for several impact categories, which contradicts assumptions that  
405 bio-based automatically implies ‘green’ or ‘sustainable’ (see also Chap. 30). Higher  
406 impacts for bio-based polymers are mainly associated with feedstock-related agri-  
407 cultural emissions of fertilisers (eutrophication) and pesticides (human toxicity and  
408 ecotoxicity), as well as deforestation (impacts related to changes in land use).  
409 However, the relative importance (i.e. contribution to overall environmental impacts)  
410 of the different impact categories also needs to be considered when evaluating the  
411 overall environmental performance of different polymers or other chemical products  
412 and processes.

413 Often products are referred to as ‘green’ or ‘sustainable’ based on a single  
414 environmental issue (e.g. reducing greenhouse gas emissions), or based on fol-  
415 lowing the principles of green chemistry in chemical design only. However,  
416 chemical products that are claimed to be ‘green’ or ‘sustainable’ may in fact lead to  
417 greater impacts on the environment or humans than the conventional alternatives.  
418 For example, ‘eco-friendly’ food packaging made of plant fibres may increase  
419 exposure and environmental emissions of highly hazardous fluorinated chemicals  
420 (Yuan et al. 2016), and ‘green’ solvents can have higher impacts across many  
421 impact categories when compared to conventional solvents (Zhang et al. 2008).  
422 Furthermore, the production of bio-based raw materials (such as corn, sugar cane,  
423 or soy for feedstock) may or may not be associated with lesser greenhouse gas  
424 emissions and consumptions of fossil resources, but may have *equal or greater*  
425 *impacts* in other impact categories (e.g. land use, toxicity related to using pesticides,  
426 eutrophication related to using fertilisers) than fossil-based materials (see Chap. 30  
427 for further details). These phenomena are commonly referred to as burden shifting  
428 (e.g. between environmental issues or compartments). Identifying these is a fun-  
429 damental application principle unique to LCA.

430 LCA is a tool that can be useful for comparing products and processes for  
431 identifying such burden shifting and how to minimise impacts across a variety of  
432 impact categories. However, it is important always to ensure as a practitioner that  
433 all relevant chemical emissions are inventoried and all impact pathways are char-  
434 acterised. These general cautions are also relevant for LCA studies focusing on  
435 chemical synthesis and production processes as discussed in the following section.

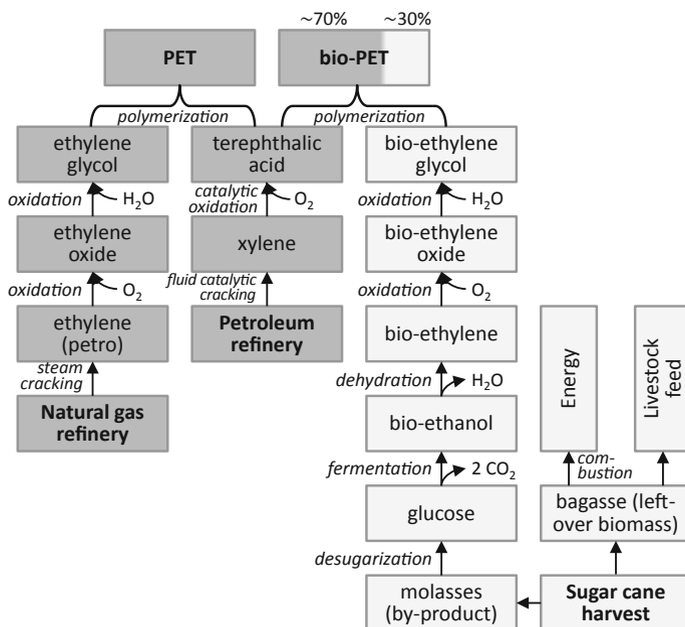
### 436 31.2.2 Chemical Synthesis and Production Processes

437 LCA is a useful tool for improving existing processes and designing new processes  
438 for the synthesis and production (Fig. 31.3, stages A–D) as well as for the  
439 end-of-life treatment (Fig. 31.3, stage F) of chemicals and chemical products, to  
440 inform process systems engineering decisions (Jacquemin et al. 2012). In this  
441 section, LCA case studies focusing on chemical synthesis and production processes  
442 across various economic sectors are discussed.

443 A major issue illustrated by several LCA studies is that management decisions  
444 based on single indicators or criteria can lead to increasing other impacts (that were  
445 not considered in the decisions), thereby indicating the strength of LCA as an  
446 approach to assess multiple indicators and related trade-offs. An example is the  
447 development and application of new plant protection products (pesticides) designed  
448 with the intention to reduce human toxicity and ecotoxicity potentials associated  
449 with emissions after application in agricultural crop protection or elsewhere (e.g.  
450 household pesticides). A related LCA study revealed that the production of a new  
451 and more effective plant growth regulating pesticide with less intrinsic toxicity  
452 (preferable from a risk perspective) than a functionally equivalent earlier marketed  
453 pesticide comes at the expense of increased impacts associated with pesticide  
454 synthesis and production processes (Geisler et al. 2005). The higher impacts for the  
455 new pesticide are mostly explained by the high complexity of its molecular  
456 structure requiring more synthesis and processing steps. In general, impacts related  
457 to the production of chemicals have been attributed to energy consumption which  
458 tends to increase with increasing complexity of a chemical molecule. Highly specialised  
459 chemicals, such as pharmaceuticals, can thereby be associated with higher  
460 energy consumption and related impacts from synthesis and production processes  
461 than other chemicals (Wernet et al. 2010).

462 Not only complexity of chemical synthesis and production processes, but also  
463 the difference in raw materials used drives environmental performance profiles of  
464 chemicals and chemical products. This is shown in another set of LCA studies  
465 contrasting chemical production from fossil fuel-based versus renewable  
466 (bio-based) resources. Synthesising and producing chemicals from biomass (e.g.  
467 sugar cane) instead of from fossil fuels (e.g. petroleum or natural gas) has been  
468 proposed as a 'sustainable technology' with respect to reducing reliance on fossil  
469 resources and greenhouse gas emissions. However, a full sustainability analysis has  
470 typically not been conducted, which is why several LCA studies have focused on  
471 this claim.

472 As an example, a simplified overview of the different chemical synthesis and  
473 processing steps involved in polyethylene terephthalate (PET) polymer production  
474 is given in Fig. 31.5. While terephthalic acid used in the production of the chemically  
475 identical PET and bio-PET is in both cases derived from petroleum, ethylene  
476 glycol can be derived from natural gas as a fossil resource (for PET) or from sugar  
477 cane as a biomass feedstock (for bio-PET). The process of natural gas refinement to  
478 create ethylene glycol alone consists of several steps including cracking (breaking  
479 down) into ethylene and other chemicals, ethylene separation and purification  
480 involving several distillation processes (not shown in Fig. 31.5). Accordingly, LCA  
481 studies have found that bio-based chemical production usually can lead to less  
482 greenhouse gas emissions than fossil-based chemical production, mainly because  
483 less refinement of fossil fuels is required. However, growing, harvesting and processing  
484 bio-based feedstocks may lead to other impacts related to agriculture production  
485 systems, e.g. land use (see Chaps. 29 and 30), which are highly variable  
486 with respect to the type of biomass used (Tabone et al. 2010; Akanuma et al. 2014).  
487 Furthermore, the type of biomass used can influence the energy required, and



**Fig. 31.5** Production process steps for chemical synthesis of polyethylene terephthalate (PET) derived from fossil fuels and bio-PET (partly) derived from bio-resources (modified from Tabone et al. 2010)

488 post-processing of bio-based products and residues greatly influence the overall  
 489 related environmental performance.

490 Other LCA studies have focused on specific aspects of chemical synthesis and  
 491 processing, such as comparing continuous and batch reactor types (e.g. Wang et al.  
 492 2013) or different catalysis and fermentation processes (e.g. Pati et al. 2014). It is  
 493 further important to consider which catalysts are used in other processing steps that  
 494 petro- and bio-based materials like PET have in common, such as antimony trioxide  
 495 found at concentrations of 200–300 ppm in PET or other, metal-free catalysts used  
 496 in the polycondensation process as part of polymerisation. Several studies have  
 497 concluded that processes with higher yields have a lower impact per chemical  
 498 production unit. The use of solvents has additionally been identified as an important  
 499 component influencing environmental performance of chemical products (De Soete  
 500 et al. 2014). Generally and specifically for chemical synthesis and processing it is  
 501 important to be sceptical of processes and products labelled or deemed ‘green’ or  
 502 ‘sustainable’ without performing a full LCA as shown, e.g. for ‘green’  
 503 nano-materials synthesis (Pati et al. 2014). An overview of aspects that are relevant  
 504 for assessing ‘green’ chemical synthesis and production processes is given by  
 505 Kralisch et al. (2015).



### 31.3 Specific and General Methodological Issues for LCA of Chemicals

Applying LCA, specifically in the chemical and pharmaceutical sectors, and in other sectors where chemicals play a central role, comes with several methodological and practical challenges. Generally, gathering chemical inventory data, quantifying impacts, and interpreting results constitute challenges for LCA studies across sectors. In the following sections, some of the most relevant challenges focusing on chemicals in LCA are discussed in relation to the definition of the goal and scope of an LCA study, product system modelling and quantification of life cycle chemical emissions in the inventory analysis, characterisation modelling in the impact assessment, and finally interpretation of LCA results in different contexts.

#### 31.3.1 Goal and Scope Definition

Consistently defining the goal and scope for chemical products or processes (e.g. functional unit of the considered product or service system and related reference flow(s) and system boundaries) is not trivial and needs to be critically considered by a practitioner. Examples of relevant issues when defining functional unit, reference flow(s), and system boundaries are discussed in the following.

##### **Functional Unit (FU)**

LCA (and other types of assessments) can be designed to compare functionally equivalent chemicals and chemical products as classified by *chemical function* (e.g. solvents, catalysts), *material function* (e.g. nanotubes, polymers) or *product function* (e.g. herbicides). It is hence important to define the level of ‘functionality’ based on which a study will be conducted. This functionality must be captured in the definition of the FU of an LCA study as basis for comparing products or systems.

Performing an LCA study is useful for providing valuable insight into which of two alternative, functionally equivalent chemicals or products provides the function with the lowest overall environmental impact profile, thereby focusing on avoiding burden shifting between different types of environmental impacts. To screen multiple alternatives to harmful chemicals in a particular product application, in contrast, the focus often is not mainly on environmental performance, but on a combination of regulatory compliance, economic and technical feasibility, along with considering hazard and human, environmental and social impacts. In such cases, a chemical alternatives assessment (CAA) might be the preferred approach to identify the most viable solution(s).

Chemicals and chemical products can also fulfil more than a single function and, hence, a partial definition of the functional unit could lead to inconsistent

544 comparisons if the appropriate product systems are not considered as demonstrated  
 545 in Example 31.1.

547 **Example 31.1 Functional Unit (FU)** Take a cosmetic product like shampoo,  
 548 where different chemical ingredients provide different functions as part of the  
 549 final shampoo product, e.g. to provide clean, shiny and fragrant hair for one  
 550 person over 24 h. If the **FU is defined** with respect to a single shampoo  
 551 product (one-product system) that **cleans the hair of one person** (by con-  
 552 taining detergents) **and makes it shiny** (by containing siloxanes) **for 24 h**, a  
 553 functionally equivalent service could be also provided by applying two dis-  
 554 tinct products (two-product system), one being a shampoo that only cleans  
 555 hair (and does not make it shiny) and another being a conditioner that makes  
 556 the hair shiny (and does not clean). However, both the one-product and  
 557 two-product systems should not provide fragrance in order to be consistently  
 558 compared via the same FU (underlined text above) that excludes fragrance.

559 Likewise, if the FU is defined to just clean hair for one person over 24 h,  
 560 comparing LCA results of a shampoo that only provides clean hair to a  
 561 shampoo that provides clean, shiny and fragrant hair could yield the mis-  
 562 leading outcome that the former shampoo ‘performs better’, because the  
 563 production and related impacts of additional chemicals of the latter shampoo  
 564 (containing siloxanes for making the hair shiny and terpenes for making the  
 565 hair fragrant) are related to functions not fulfilled by the shampoo that only  
 566 cleans hair. Hence, the comparison would be biased by comparing products  
 567 fulfilling distinct functions.  
 568

569 Defining an appropriate FU for multi-functionality (see Chap. 8) is also  
 570 important. For example, water and propylene glycol are both effective chemical  
 571 solvents and, thus, both would fulfil an FU defined with respect to providing the  
 572 function of a solvent in, e.g. a shampoo product. Propylene glycol, however,  
 573 provides other functions that water does not provide (e.g. stabiliser, humectant,  
 574 emulsifier). Therefore, a comparison of propylene glycol and water in an LCA  
 575 study based on a solvent-based FU would not capture the multi-functionality of  
 576 propylene glycol. Defining the FU with respect to all functionalities and then  
 577 providing system expansion when necessary (e.g. water plus a stabiliser plus a  
 578 humectant plus an emulsifier is functionally equivalent to propylene glycol in  
 579 shampoo) can be an important consideration in any LCA on chemicals or product  
 580 systems. It is, hence, important to ensure the product(s) or chemical(s) investigated  
 581 in an LCA study are functionally equivalent and the FU captures this equivalency  
 582 appropriately.

### 583 **Reference Flow**

584 The reference flows (Chap. 8) in an LCA study reflect the overall amount of goods  
 585 and/or services that are required to fulfil the defined FU. Taking a no-wash  
 586 (dry) shampoo versus a conventional (liquid) shampoo as examples, the reference

587 flows to fulfil an FU of cleaning the hair of one person for one day could be 10 g of  
588 the liquid shampoo product plus the (hot) water used to wash the hair. The  
589 reference flow for the dry, leave-in, no-wash shampoo could be simply 5 g of  
590 powdered product (with no wash-water needed). Furthermore, if functionally  
591 equivalent products or chemicals provide different efficiencies to fulfil a defined  
592 FU, the different efficiencies need to be accounted for in the reference flow. This  
593 issue also points to a problem for cradle to gate LCA studies on chemicals, where it  
594 is possible that a chemical could have greater cradle to gate impacts than another  
595 chemical per unit mass emitted, but far less of the former chemical is required to  
596 fulfil the same FU. Here, pesticides with different efficiencies towards the same pest  
597 offer a typical example.

### 598 *System Boundaries*

599 The system boundaries (Chap. 8) of any defined chemical product or service sys-  
600 tems in an LCA study need to capture all relevant processes for the systems being  
601 compared. For example, if the purpose of an LCA study is to compare bio- with  
602 fossil-based chemical synthesis, the system boundaries must include and differen-  
603 tiate all raw material acquisition processes, namely all refining processes for the  
604 fossil-based chemical and the crop production and processing steps for the  
605 bio-based chemical (see also Fig. 31.5). However, for these systems, it may not be  
606 relevant to include chemical use and disposal stages in the study, whenever these  
607 life cycle stages are equivalent in both cases. Such systems are referred to as ‘cradle  
608 to (factory) gate’ systems and are common in LCA studies on chemical synthesis  
609 and other chemical production processes (Jimenez-Gonzalez and Overcash 2014).  
610 In contrast, if the purpose of the study is to compare two distinct fossil-based  
611 materials fulfilling the same function, the disposal stage could be a relevant driver  
612 of the difference between the compared product systems.

613 For several chemical products and production processes, consistently defining  
614 system boundaries is challenging. An example is the application of plant protection  
615 products containing chemical pesticide active ingredients (e.g. carbamate insecti-  
616 cides) applied in agricultural crop production, where the FU could be defined to  
617 provide a specified amount of crop in a season. Allocating field buffer strips (i.e.  
618 non-agricultural areas that are among other functions introduced to reduce the  
619 impact of applied pesticides on non-treated areas), which may be required by law,  
620 to the technosphere would apparently influence the crop yield per hectare and  
621 amount of pesticide used compared with an equivalent system, where the buffer  
622 strips are defined as part of the environment (Rosenbaum et al. 2015). Including  
623 buffer strips in the considered technosphere system or not will, hence, influence the  
624 related impacts and also defines the scope of the environmental distribution pro-  
625 cesses of pesticides in the LCI and LCIA phases. As a consequence, the definition  
626 of the system boundaries needs to be aligned with the selected pesticide inventory  
627 data and characterisation models to avoid overlaps, double counting of processes  
628 and potential gaps along the pesticide impact pathways.

### 31.3.2 Product System Modelling and Inventory Analysis

There are several obstacles that need to be considered in the product system modelling and inventory analysis phase (Chap. 9), after the goal and scope of an LCA study have been defined.

#### *Data Availability and Quality*

All relevant chemical elementary flows from a given product system to the environment need to be quantified in the LCI phase. When using LCA software, emission quantities are often available through an LCI database, for example for processes occurring in Europe or the 'rest of the world.' LCI databases generally rely on typical or average emission inventories or an inventory taken by one industry for a given unit process, which may be outdated or tied to, e.g. a specified electricity mix. Thus, it is always preferred to gather primary data, especially for the foreground system modelling (Chap. 9), of the specific LCA case under study. This poses a particular challenge to LCA practitioners, who may or may not have access to company-specific data to resolve the nuances of a particular supply chain. While in some cases, a particular commissioner of an LCA study might provide such data, while in other cases such data have to be collected from different parties. An example is the application of plant protection products, where pesticide manufacturers will know the concentration of a pesticide active ingredient in a formulation product, but where the different farmers might know the effectively used amount that is applied on agricultural fields and this usually depends on pest-, climate-, soil- and application-specific conditions.

#### *Emission Estimation and Modelling*

Most chemical synthesis and material/product manufacturing processes involve several steps, which can yield usable by-products that have to be considered in an LCA study (see Chap. 9 for further details). As an example, harvesting sugar cane yields refined sugar, but also molasses (sugar refining by-product) and bagasse (dry leftover biomass after extracting the juice from the sugar cane). While molasses can be further used to produce biochemicals, bagasse is usually burned (for energy conversion) or used as livestock feed (Fig. 31.5). In an LCA study, usually only one of these products (sugar, biochemical, energy, livestock feed) is in focus and the other products must be accounted for through subdivision or system expansion or if it cannot be avoided through different types of allocation (see Chap. 9).

When building a product system model, different tools and software packages are available. Specifically for simulating material and energy balances of chemical production and processing, there exist several (open-source and commercial) chemical process simulators, such as Aspen HYSYS for oil and gas process simulation and Aspen Plus for chemical process optimisation ([www.aspentech.com](http://www.aspentech.com)), BatchReactor and BatchColumn for chemical reactor and batch distillation columns simulation, respectively ([www.prosim.net](http://www.prosim.net)), or CHEMCAD software suite for chemical process simulation and optimisation including batch operations ([www.chemstations.com](http://www.chemstations.com)). Such software packages may include proprietary data from the



671 chemical and other industries that are otherwise not accessible and may intrinsically  
672 use different allocation systems. The responsibility of ensuring transparency and  
673 consistency when building a product system including proper consideration of  
674 co-products and by-products lies with the LCA practitioner. However, several  
675 documents exist for LCA practitioners to seek guidance, and working examples of  
676 co-product consideration for the chemical industry can be found elsewhere (e.g.  
677 Weidema 2000; Karka et al. 2015).

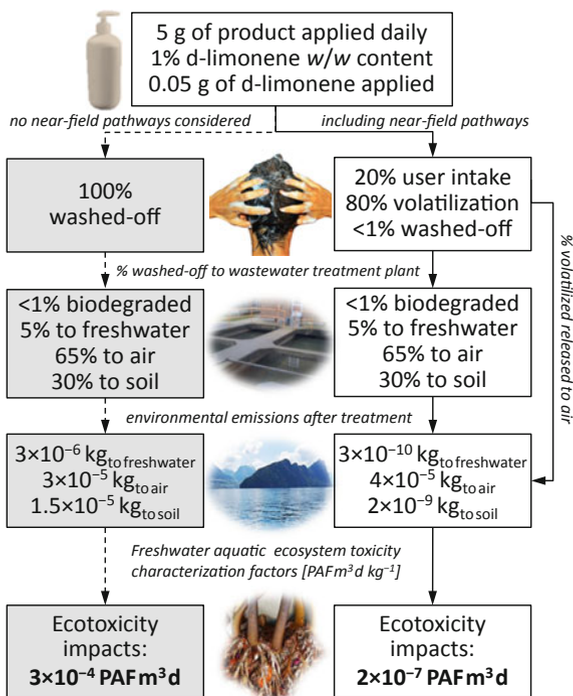
678 In most LCA studies, an inventory covers hundreds of processes and emission  
679 flows but not all chemical emissions are usually able to be covered. Often missing  
680 from LCI databases are, e.g. emissions to the occupational and consumer envi-  
681 ronments, and the ingredients (e.g. chemicals) in a product, which can be emitted  
682 indoors during product use or outdoors post-use as demonstrated in Example 31.2.

684 **Example 31.2 LCI Emission Pathways** When a **consumer product** (e.g.  
685 perfume) or industrial product (e.g. agricultural pesticide) is used, the  
686 chemicals within the product undergo various pathways, thereby exposing for  
687 example the product users and people nearby. Consider that a colleague at  
688 work **applies an air freshener or perfume in the office**. Perhaps you smell  
689 or even taste it in the first minutes after application (indication of exposure),  
690 maybe the scent remains in the office for some days (indicating sorption and  
691 desorption from indoor walls and other surfaces), and maybe you can even  
692 smell it just outside the office building (indicating transport outdoors via  
693 ventilation).

694  
695 In some cases, a large proportion of chemicals within products can be taken in  
696 by humans during and after product use, which is a major concern amongst reg-  
697 ulators and researchers. In LCA, such considerations are currently largely missing,  
698 but first efforts were made to include indoor fate and exposure pathways (referred to  
699 as ‘near-field’) into the toxicity characterisation model USEtox 2.0 (<http://usetox.org>).  
700 Without accounting for near-field fate and exposure pathways, LCA studies  
701 typically may assume a fixed-fraction like 100% of product ingredients being  
702 emitted to the environment. In general, assuming such emission distributions could  
703 lead to an underestimation of resulting human toxicity potentials and an overesti-  
704 mation of environmental or ecosystem impacts. This is illustrated in Fig. 31.6 for  
705 d-limonene as commonly found chemical in a shampoo product, where assuming  
706 100% of the used product being washed-off (left-side pathway in figure) instead of  
707 modelling a more complex yet more realistic distribution (right-side pathway in  
708 figure) yields a difference of more than three orders of magnitude for freshwater  
709 ecotoxicity impacts, which is significantly beyond the uncertainty range for this  
710 impact category.

711 Emissions can also occur from chemical residues in products that are related to  
712 cross-contamination, i.e. such chemicals are not purposefully added to a product  
713 and enter a product from using, e.g. recycled material where not all chemical  
714 ingredients are known. Often, inventory data related to cross-contamination

**Fig. 31.6** Illustrative example of assumptions for emission distributions of chemicals in consumer products in life cycle assessment showing a substantial decrease in the estimated potentially affected fraction (PAF) of freshwater species from the chemical ingredient d-limonene (CAS: 5989-27-5) in shampoo when accounting for indoor fate and exposure of cosmetic products. Adapted from results from Ernstoff et al. (2016a) combined with freshwater ecotoxicity characterisation factors from USEtox 2.0 (<http://usetox.org>). Air emissions were assumed to be to urban air, water emissions to continental freshwater, and soil emissions to continental natural soil



715 pathways are very limited if at all available. Using similar processes or pathways as  
 716 proxy might be a possibility to address this limitation, but also introduces additional  
 717 uncertainty in the emission estimates.

### 718 *Spatiotemporal Variability in Emissions*

719 Time (e.g. year, season or duration) and location can influence variations in  
 720 emissions, referred to as 'spatiotemporal' variability. In many cases, LCI results do  
 721 not capture the time of emissions from systems, e.g. agricultural practices (e.g.  
 722 harvesting, applying fertilisers) can occur according to daily or seasonal cycles  
 723 according to the geographic location of the farm. Likewise, emissions of landfill  
 724 leachate are influenced by changes in environmental conditions (e.g. acidity and  
 725 temperature) which can change through time and according to location (Bakas et al.  
 726 2015).

### 727 *Incomplete Emission Inventories*

728 It is important to be aware of the incompleteness of some emission inventories. For  
 729 example, energy conversion processes generally are well detailed in LCI which can  
 730 result in high toxicity related impacts resulting from energy consumption, but other  
 731 processes, e.g. related to chemical processes may have less complete inventory and,  
 732 hence, related toxicity impacts might be underestimated (Laurent et al. 2012).



### 31.3.3 Impact Assessment

Characterising chemical emission flows resulting from the LCI in terms of their impacts on humans and the environment requires a careful consideration of study context (e.g. spatial region), number and relevance of chemicals to be characterised (in many cases, most chemicals contribute marginally to overall impacts, while only few chemicals dominate overall impact profiles). In the following, challenges and pitfalls in the impact assessment of chemical products and processes are discussed with focus on toxicity-related impacts, where special challenges exist mainly due to the countless chemicals to be characterised and the complexity of related impact pathways.

#### *Limited Substance Coverage*

USEtox, a scientific consensus model for characterising human and ecotoxicological impacts of chemicals, presently provides characterisation factors for more than 3000 chemicals, which constitutes the largest list currently available in LCIA (<http://usetox.org>). However, with tens of thousands of chemicals on the market, inventoried chemical emissions either documented in an LCI database or by a practitioner investigating a specific system or process, may in many cases not have existing characterisation factors or the data required to develop new characterisation factors (e.g. toxicity dose-response information). This limitation to substance coverage in LCIA is important when interpreting results, because a lack of data for many chemicals does not preclude their possible impacts.

#### *Chemical Degradation Products*

When a chemical does not degrade, or degrades very slowly, it is considered 'persistent.' Persistent chemicals thereby can be linked to greater impacts because they are not or very slowly removed from the system through degradation. In current LCIA methodologies, abiotic (e.g. where a chemical is transformed via interactions with sunlight) and biotic (e.g. when a chemical is metabolised by soil bacteria) degradation essentially 'removes' organic chemicals from the system and no further impacts are characterised. In reality, degradation processes transform a chemical into various degradation or transformation 'products', including other chemicals or gases, which can also impact the environment. Degradation products can have greater or lesser impacts than their parent compounds, for example aminomethylphosphonic acid (AMPA), which is the main degradation product of the broad-spectrum herbicide glyphosate, is more persistent and more toxic than the glyphosate parent compound. As an example, not considering AMPA in an LCA study that considers agricultural processes where this herbicide is used could underestimate the impacts of using glyphosate. Therefore, an LCA practitioner should include estimates of persistent degradation products and appropriate characterisation factors (in this case for AMPA, not glyphosate) to better capture the impacts of chemicals. While this approach will not be feasible for all chemicals (due to data limitations), it should be performed when the issue is known and data are available.

### ***Impacts from Chemical Mixtures***

Impacts towards humans or different ecosystems, related to chemical emissions, are a function of the simultaneous prevalence of other chemicals, which might have synergistic (enhancing) or antagonistic (counteracting) properties with respect to the effect of a considered chemical. Since information on the site-specific mixture of chemicals in any environmental medium or compartment is not usually available, and the impacts of such a mixture on humans or the environment are not known, synergistic or antagonistic effects are usually not considered, and instead additivity of concentrations and effects is assumed. This means that the effects of all chemicals contributing to the same impact category, e.g. freshwater aquatic ecosystem toxicity or ozone depletion, are summed up to arrive at an overall product system-related impact score. If for any LCA study the emission location and time is known and related background concentration profiles available for all relevant chemicals, this assumption could be evaluated by identifying and quantifying the synergistic or antagonistic effect potentials. However, the potentially added accuracy in an LCA context is most likely not relevant given existing uncertainty attributable to other aspects in the characterisation of chemical emissions. Besides, the large number of chemicals present and emitted into the environment yields an almost limitless amount of possible mixtures, rendering it impossible to quantify the specific effect potentials for each mixture.

### ***Missing Fate and Exposure Processes and Pathways***

In order to reduce the demand put on LCA practitioners, streamline workflows, and allow for science-based and consensus-driven solutions, LCIA often relies on predefined methodologies. However, hundreds or even thousands of chemicals might be inventoried for various processes in an LCA study, but characterisation factors or a LCIA method for a given impact at mid-point or end-point level may be missing, especially for toxicity-related impacts (see Chap. 10). Moreover, certain exposure settings (occupational, consumer) or routes (e.g. dermal exposure) or target organisms (e.g. exposures of bees) may be missing from an LCIA model. Effect factors may also be missing or inconsistent, e.g. in the case of human toxicity, effects of allergy or endocrine disruption (i.e. interaction with the hormone system) are often not included, but may be highly relevant for chemicals in consumer products. Finally, many of the methodological gaps in LCIA are also due to the reliance on simplifying assumptions. The LCA practitioner, who is constrained by resources (time, money, data access), is responsible for compiling the necessary data and for ensuring that the LCIA methodology chosen (or developed) is suitable for the defined goal and scope of an LCA study. Specifically, to characterise a chemical's impact, several assessment factors are required and must be sufficiently scrutinised within the chosen LCIA method, such as the chemical environmental fate, ecosystem and/or human exposure if relevant, and subsequent effects with respect to given impact categories. Each of the related data requirements poses its own challenges. To avoid the misleading conclusion that missing aspects of the chosen LCIA method do not cause impacts because they were not assessed, it is important to be familiar with which processes (e.g. biotransformation),

819 environmental compartments (e.g. indoor air), exposure pathways (e.g. dermal  
820 uptake), and effects (e.g. endocrine disruption), may be missing from the selected  
821 characterisation methods but are relevant for the system under study. In some cases  
822 such missing aspects can be addressed by the practitioner by developing new  
823 methods or by adapting existing methods; if not, it is important to be aware of how  
824 this could influence results.

### 825 *Spatiotemporal Variability of Impacts*

826 LCIA methods are generally based on regional or global averages for various  
827 chemical, environmental and pathway data and processes, e.g. how long it takes  
828 chemicals emitted to freshwater to reach the sea (i.e. residence time), or how many  
829 persons live in an urban area (i.e. population density). Studies have shown, intu-  
830 itively, using a continental average instead of ‘spatially differentiated’ regionalised  
831 models can yield large uncertainty in the estimated impacts (e.g. Kounina et al.  
832 2014). Thus, if the location of the emissions (e.g. from a specific factory) in an LCA  
833 study is known, using a model with characterisation factors specific for that region  
834 can reduce uncertainty of model results. If emission locations are not known (as is  
835 the case for most chemicals in typical LCA studies), characterisation results for  
836 regions can be applied that are parameterised, i.e. averaged for the characteristics of  
837 a particular region. The same rule applies for temporal aspects, where in LCA  
838 mostly steady-state conditions and continuous emissions are assumed, which might  
839 not be true for, e.g. agricultural pesticides that are applied on specific days only (i.e.  
840 pulse emissions). In such cases, accounting for the dynamics of the chemicals in the  
841 modelled environmental system may reduce uncertainty in characterisation results  
842 (e.g. Fantke et al. 2012), but whenever temporal information on emission patterns is  
843 not available, parameterised characterisation results can be applied that account for  
844 the most important temporal aspects of a modelled system.

### 845 *Impacts Versus Benefits*

846 Life cycle *impact* assessment inherently focuses on quantifying ‘negative’ impacts  
847 on humans and the environment. A stakeholder could in some cases argue that their  
848 product or service offers a benefit to society that is not accounted for, meaning that  
849 LCA yields misleading results. When facing such an argument as LCA practitioner,  
850 it is important to go back to the fundamentals of LCA. The impact assessment phase  
851 of LCA is designed to assess environmental ‘benefits’ in the form of ‘avoiding  
852 environmental impacts.’ For example, a wastewater treatment plant design that also  
853 decreases environmental pollution compared to another design offers a ‘benefit’ that  
854 is quantifiable in an LCA context (see Chap. 34 on LCA of wastewater treatment).  
855 Furthermore, when comparing functionally equivalent products or services, their  
856 benefits (e.g. restoring a wetland to yield a level of biodiversity, or designing a car  
857 with a certain safety rating) is often captured in the functional unit of an LCA study,  
858 which defines a unit of the (beneficial) service being provided. There are special  
859 cases where considering societal benefits that are not captured in the functional unit  
860 or by the assessment methods can be extremely important when guiding  
861 decision-making. In some cases, LCA may not be the appropriate tool to assess  
862 such benefits; however, developing LCA-compatible methods to quantify societal

863 benefits (specifically positive human health outcomes) is a topic of high interest  
864 when assessing human nutrition and dietary patterns, where two functionally  
865 equivalent diets can have very different health impacts (Nemecek et al. 2016).

### 866 **31.3.4 Interpretation**

867 The interpretation of results is fundamental for the findings and reliability of every  
868 LCA study and subsequent guidance provided to stakeholders, and to LCA in  
869 general (see Chap. 12). Robust and transparent interpretation of results from an  
870 LCA study can offer sound council for the stakeholders and when aggregating with  
871 other LCA studies can elucidate generalisable findings important for sustainable  
872 development. As an example of nuances of interpretation, the ‘New Plastics  
873 Economy’ report (WEF 2016) cites interpretation of several LCA studies and  
874 implies that a major shortcoming of LCA is its inability to identify and support  
875 ‘target states’, such as moving towards increased production and use of bio-based  
876 plastics. Indeed, as previously discussed, LCA studies on bio-based versus  
877 fossil-based plastics have demonstrated similar, if not greater impacts (e.g. on land  
878 use and toxicity potentials) for bio-based plastics due to agricultural practices (see  
879 Chap. 30), which is a finding that may be unintuitive or undesirable to some (e.g.  
880 stakeholders in the bioplastics industry). When interpreting such LCA results, it is  
881 important to distinguish what an LCA says about ‘here and now’ versus what it  
882 could mean for future sustainability goals or targets of stakeholders. For example,  
883 LCA results showing bio-based plastics have ‘greater impacts’ than fossil-based  
884 plastics do not discredit bio-based plastics as a sustainability goal, but they do  
885 indicate that bio-based plastics face sustainability challenges given current agri-  
886 cultural practices, which thus must be addressed to avoid impact trade-offs.  
887 Furthermore, LCA results can help indicate which feedstock is the most  
888 eco-efficient (less impacts per kilogram) to work towards a bio-based ‘target’. In  
889 practice, LCA may not be able to easily identify target states often elucidated  
890 according to societal values (which may include socioeconomic or political factors)  
891 or intuitive/consensual sustainability goals, but LCA can be instrumental in  
892 reaching goals and target states in a holistically sustainable manner and shedding  
893 light on challenges faced when working towards such goals. In the following, some  
894 additional challenges in interpreting LCA results are outlined.

### 895 **Contribution to Impact Results**

896 Especially for LCAs on chemical products or processes, it is important to trans-  
897 parently report and document the contribution of different chemicals to impacts  
898 related to product life cycle stages and individual processes. This can help identify  
899 potential problems in the processing of LCI or LCIA results (e.g. if one chemical  
900 dominates results). Interpreting LCA results might be particularly challenging if it is  
901 not clear whether toxicity-related impacts are associated with chemical emissions  
902 occurring along the product life cycle or, in contrast, with chemicals that are

903 product *ingredients* (Roos and Peters 2015). As an example, glass used as food  
904 packaging can show higher potential toxicity impacts compared with plastic  
905 packaging due to transport-related emissions of toxic chemicals from fossil fuel  
906 burning (Humbert et al. 2009), which is linked to the fact that glass is usually  
907 heavier than plastic. However, plastic food packaging can likely lead to greater  
908 exposures to various chemicals through food than glass, but this aspect is not  
909 (yet) considered in current LCIA toxicity models (Ernststoff et al. 2016b). Further, it  
910 might be unclear whether worker and/or consumer exposure pathways are included  
911 as these are currently beyond the scope of LCA studies focusing primarily on  
912 environmental emissions. The covered pathways and exposed populations should  
913 always be clarified in an LCA study to avoid possible misinterpretation of results.  
914 This is of specific relevance for the comparison of chemicals and chemical products  
915 and processes, where such ambiguities can cause confusion regarding the contri-  
916 bution of chemicals and related impact pathways and life cycle stages are lacking.

### 917 ***Identification of Considered Chemicals***

918 In any of the aforementioned contexts, it must be acknowledged that most chemi-  
919 cals have various common names (lindane, CAS RN: 58-89-9, is for example also  
920 commonly known as HCH, hexachlorobenzene, or cyclohexane, etc.). Hence, it is  
921 important to ensure that names for chemicals in the different phases of an LCA  
922 study (e.g. inventory analysis and impact assessment) are consistently chosen based  
923 on using CAS registry numbers or similar as chemical identifier to, e.g. avoid  
924 double counting or neglecting chemicals with ambiguous names. This exercise can  
925 prove to be challenging as LCA software packages often report chemical inven-  
926 tories by chemical name and not by CAS number.

### 927 ***Quality and Uncertainty***

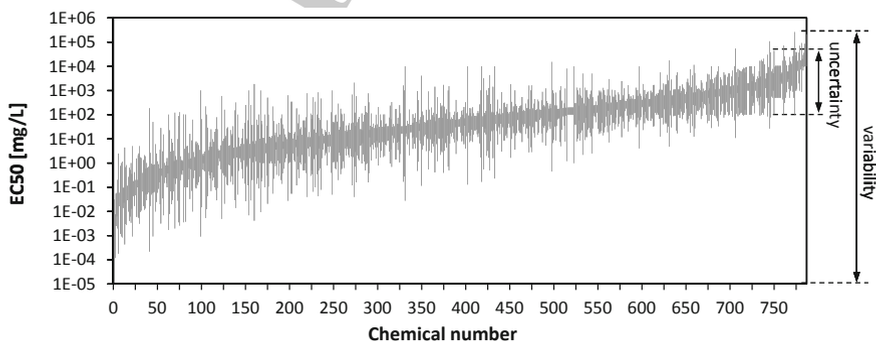
928 Quality checks across the large number of inventoried chemicals is usually difficult,  
929 but inventory results should nevertheless be verified by, e.g. checking the mass  
930 balance of only those chemicals that drive overall impact results, for examples  
931 heavy metals that dominate toxicity impact profiles. Furthermore, it is essential to  
932 report and discuss uncertainties of LCA data and results with respect to each impact  
933 category as integral part of the analysis, and consider such uncertainties in the  
934 interpretation of results and guidance provided to decision makers (see Chap. 11).

935 Particularly uncertainty associated with toxicity characterisation results is high  
936 compared with other impact categories and results can furthermore differ between  
937 toxicity characterisation methods, which can in some cases influence the overall  
938 ranking of compared product systems. Uncertainty (lack of data or understanding)  
939 and variability (data heterogeneity) are distinct concepts, but are sometimes (in-  
940 correctly) aggregated. For example, often ‘high uncertainty’ is perceived negatively  
941 or seen to discredit a particular LCIA method. However, such ‘uncertainty’ can be a  
942 direct reflection of reality and variabilities in temporal and spatial chemical fate and  
943 organism disease responses (see Chap. 11 for further details). Likewise, if an impact  
944 category has low or no associated uncertainty, this is perceived as positive but  
945 should in fact be a warning sign that there may be a lack of understanding of what  
946 uncertainties and/or variabilities exist or that the environmental relevance (or

947 representativeness) of an indicator may be low (which introduces an uncertainty in  
 948 the interpretation phase, but this is usually not quantified). To begin transparently  
 949 addressing this issue, impacts should ideally be cross-compared using different  
 950 LCIA characterisation methods with particular focus on identifying which chemi-  
 951 cals contribute the most to impacts in each LCIA method (which are often not the  
 952 same). Moreover, uncertainty ranges for toxicity-related impacts should be reported  
 953 in logarithmic scale to put average uncertainties of two to three orders of magnitude  
 954 into perspective of more than 15 orders of magnitude in the variability across  
 955 chemicals. This is shown in Fig. 31.7 for 786 chemicals with available measured  
 956 ecotoxicity effective concentrations for 50% of the exposed species (EC50; mg/L)  
 957 for aquatic ecosystems. EC50 values are used to calculate effect factors as part of  
 958 toxicity characterisation in LCIA (see Sect. 10.11). The relation between uncer-  
 959 tainty and cross-chemical variability is not much different for toxicity impacts than  
 960 for other impacts, where uncertainty in characterisation results (of usually only a  
 961 handful of contributing chemicals) and related variability across contributing  
 962 chemicals are both less broad. However, uncertainty ranges vary widely between  
 963 chemicals, but chemical-specific uncertainty around characterisation factors is  
 964 usually not available in LCA, except for specific pathways, e.g. exposure to pes-  
 965 ticide residues in food crops (Fantke and Jolliet 2016), where also the underlying  
 966 method to quantify chemical-specific uncertainty is outlined.

### 967 *Comparison with Results from Other Methods*

968 Comparing results from an LCA study with results from a different method can help  
 969 identify methodological inconsistencies that require further inspection. As an  
 970 example, it might be desired to compare the ranking of chemicals in terms of their  
 971 potential toxicity impacts on humans and/or ecosystems in an LCA study with the  
 972 ranking of chemicals based on persistence, bioaccumulation and toxicity or other  
 973 criteria used, e.g. by risk regulators. In this context, it is important to acknowledge  
 974 that inconsistencies can result from the primary data used in an LCA versus another



**Fig. 31.7** Ranges of measured chemical-specific ecotoxicity effective concentrations (50% of exposed species affected), EC50, for aquatic ecosystems collected and indicated as reliable for 786 chemicals based on REACH ([echa.europa.eu/regulations/reach](http://echa.europa.eu/regulations/reach))

975 method, or assumptions and cut-offs may be based on different criteria, e.g. worst  
976 case versus best estimate or most sensitive species versus average ecosystem sen-  
977 sitivity (Harder et al. 2015). This might lead to problems when comparing chemical  
978 rankings based on different assessment methods and data sources. Chemical toxicity  
979 results may furthermore differ between regions, countries or assessment methods,  
980 and thereby the consideration of chemicals as, e.g. ‘non-carcinogenic’ in LCA  
981 toxicity characterisation models may not be consistent with a specific regulatory  
982 context, such as the Registration, Evaluation, Authorisation, and Restriction of  
983 Chemicals (REACH) legislation framework of the European Union.

## 984 31.4 Conclusions

985 Stakeholders commissioning an LCA study can drive the goal and scope, the  
986 selection of inventory processes, and the selection of impact categories. In many  
987 cases, this can lead to an assessment that is restricted, for example to greenhouse  
988 gas emissions and a focus on climate change. The limited scope of such studies  
989 must be considered in the interpretation and application of their results, whenever  
990 other important impact pathways for chemical production, use, and disposal are not  
991 assessed. It is always important to be critical towards LCA outcomes and under-  
992 stand their limitations and scope, and respect that no tool (including LCA) can  
993 answer all questions related to chemicals and sustainability.

994 Not only can the scope of an LCA study be intentionally restricted according to  
995 its goal and scope, but there are several remaining challenges that also limit LCA,  
996 such as partial coverage of chemical inventory data, fate modelling (e.g. regional  
997 variation), exposure pathways (e.g. dermal exposure of consumers), and charac-  
998 terisation of potential human and ecosystem toxicity impacts. Given that there are  
999 tens of thousands of commercially used chemicals, and often little data on their  
1000 properties or effects, the challenge of addressing chemical risks and impacts is not  
1001 unique to LCA. Generally, the various methods for characterising risk and impacts  
1002 of chemicals face similar challenges of data availability, but they also face  
1003 methodological challenges and intentional differences. For example, results of an  
1004 LCA addressing several impact categories and hundreds of chemicals, where often  
1005 the exact emission location and timing is unknown, are difficult to cross-compare  
1006 with results of a toxicity-focused risk assessment considering specific (e.g.  
1007 worst-case) conditions and only one or several chemicals of concern (Harder et al.  
1008 2015).

1009 Attempts of combining LCA with principles of green and sustainable chemistry,  
1010 integrating LCA- and risk-based approaches, and including life cycle impacts in  
1011 chemical alternatives assessment frameworks demonstrate the growing comple-  
1012 mentarity and relevance of the life cycle approach in other science-policy fields  
1013 (Jimenez-Gonzalez and Overcash 2014; Harder et al. 2015; Jacobs et al. 2016).  
1014 Overall, the number of LCA studies focusing on chemicals or chemical products or

processes is growing; thus, increasing discourse and trust in LCA methods as well as improving existing inventory and impact characterisation approaches.

Over the past years of research, LCA has developed into a powerful tool to identify and assess trade-offs and burden shifting between different environmental issues, identify hotspots and minimise overall environmental impacts of chemicals in the life cycle of products and processes. With rising interest in creating ‘environmentally friendly’ chemicals and products, LCA is particularly important to help avoiding ‘green washing’ and unsupported claims. A common example is the comparison of products that can be developed purely from petrochemicals and also from a combination of petro- and biochemicals. Larger potential greenhouse gas emissions in the petrochemical production are confronted with often larger land use and pesticide-related toxicity impacts from agricultural crop production when serving as feedstock for biochemical production (Tabone et al. 2010; Cespi et al. 2015). Only comparing climate change impacts in this context would lead to false conclusions (i.e. that bio-based chemicals are always ‘greener’) and does not help identify how to optimise production processes and resource use when moving from petrochemicals to biochemicals in, e.g. plastics production. This is especially relevant when assessing emerging technologies, where there is a high level of optimisation potential in the years to come for upscaling lab-level processes to a commercial level.

Future research related to chemicals and LCA should focus on identifying and resolving areas of high uncertainty (such as changes through space and time), filling data gaps (for example with high-throughput exposure and toxicity modelling approaches), and addressing issues of high concern such as consumer and occupational exposure and other toxicity endpoints (e.g. toxicity to bees). Furthermore, applying LCA in case studies and analyses to address issues of existing and emerging technologies can help pinpoint and corroborate solutions towards more sustainable production and consumption of synthetic and naturally occurring chemicals.

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1201 **Peter Fantke** develops methods for LCIA, health impact assessment and chemical alternatives  
1202 assessment since 2006. Has contributed to UNEP/SETAC LCIA working groups and is USEtox  
1203 Manager. Interested in quantifying and characterising chemical emissions, uncertainty analysis,  
1204 consumer exposure, chemical substitution and model parameterisation.

1205 **Alexi Ernstoff** Studied various aspects of chemical fate, transport, and exposure since 2007.  
1206 Recent focus is modelling human exposure to chemicals in products for LCIA. Main interest is  
1207 ensuring human health impacts, mediated by using consumer and food products, is consistently  
1208 considered in quantitative sustainability assessments.