BLUE ANGEL
The German Ecolabel

Laundry detergent

DE-UZ 202

Basic Award Criteria
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Version 2
The Environmental Label is supported by the following four institutions:

The Federal Ministry for the Environment, Nature Conservation and Nuclear Safety is the owner of the label. It regularly provides information on the decisions taken by the Environmental Label Jury.

The German Environmental Agency with its specialist department for "Ecodesign, Eco-Labelling and Environmentally friendly Procurement" acts as office of the Environmental Label Jury and develops the technical criteria of the Basic Criteria for Award of the Blue Angel.

The Environmental Label Jury is the independent, decision-making body for the Blue Angel and includes representatives from environmental and consumer associations, trade unions, industry, the trade, crafts, local authorities, academia, the media, churches, young people and the German federal states.

The RAL gGmbH is the awarding body for the Environmental Label. It organises the process for developing the relevant award criteria in independent expert hearings – which involve all relevant interest groups.

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1 Introduction

1.1 Preface

In cooperation with the Federal Ministry for the Environment, Nature Conservation and Nuclear Safety, the Federal Environmental Agency and considering the results of the expert hearings conducted by RAL gGmbH, the Environmental Label Jury has set up these Basic Criteria for the Award of the Environmental Label. RAL gGmbH has been tasked with awarding the Environmental Label.

Upon application to RAL gGmbH and on the basis of a Contract on the Use of the Environmental Label to be concluded with RAL gGmbH, the permission to use the Environmental Label may be granted to all products, provided that they comply with the requirements as specified hereinafter.

The product must comply with all the legal requirements in the country in which it is to be marketed. The applicant shall declare that the product meets these conditions.

1.2 Background

Washing and cleaning agents are used on a daily basis for maintaining cleanliness and hygiene. Approx. 480,000 tonnes of these cleaning and care agents are sold each year in Germany, whereby dishwashing detergents account for around 260,000 tonnes. All of these products contain ingredients that find their way into the wastewater system and can have a negative effect on the environment and human health. If these components cannot be completely retained or degraded in sewage treatment plants, they will enter into bodies of water and can be ingested by water organisms and, in some circumstances, enriched. Furthermore, washing and cleaning agents can negatively affect human health when substances hazardous to health such as fragrance allergens and preservatives are used.

An important component of cleaning agents are surfactants. These can be manufactured based on petrochemicals and/or renewable raw materials. The use of sustainably produced raw materials makes a significant contribution to sustainable development. In order to ensure this is the case, the cultivation of the plants is subject to ecological, social and economic requirements. Criteria for sustainable cultivation are currently being discussed in different initiatives and reliable certification systems for recording and labelling this type of cultivation are being developed or are establishing themselves on the market.

The separation and traceability of the raw materials (segregation) is currently only possible to a very limited extent in the case of palm (kernel) oil for the manufacturing of surfactants. An interim solution that currently exists is the possibility of verifying sustainable cultivation using the mass balance of raw materials. In this process, the end product does not necessarily contain the certified palm (kernel) oil.

However, it is important to ensure in the longer term that certified palm (kernel) oil is exclusively used in the product and other natural resources e.g. other oil plants or raw materials for the manufacture of citric acid or bioalcohol are integrated into the certification system. In future updates to the environmental label criteria, the further development of these certification systems for sustainable cultivation will be taken into account.

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1 www.umweltbundesamt.de/themen/chemikalien/wasch-reinigungsmittel/umweltbewusst-waschen-reinigen
1.2.1 Overview of possible future requirements

The following points will be taken into account, where possible, in future revisions of these Basic Award Criteria:

- Inclusion of other renewable raw materials in the requirements for sustainable cultivation;
- The general biodegradability of all substances;
- Other requirements for the use of recycled materials in the packaging and associated design requirements;

1.3 Objective of the environmental label

The following criteria are designed to promote the use of those laundry detergents that in the interests of the environment, climate and nature conservation have been produced in such a way that they have the least possible impact on the environment and human health during their production, use and disposal. In addition, they should also help to reduce and prevent the risks posed to the environment and human health through the use of hazardous substances and to minimise packaging waste. Furthermore, information should be provided that enables consumers to use the product both efficiently and with the least possible impact on the environment.

Preserving natural resources is also an important focus of this environmental label. Cleaning agents with the Blue Angel ecocert should thus make a contribution to the protection of the environment through the use of renewable raw materials in their production that have been cultivated under sustainable conditions or which support sustainable cultivation.

1.4 Definitions

For the purpose of their use in these Basic Award Criteria, the following definitions are valid:

- **Substance**: means a chemical element and its compounds in the natural state or obtained by any manufacturing process, including any additive necessary to preserve its stability and any impurity deriving from the process used, but excluding any solvent which may be separated without affecting the stability of the substance or changing its composition.
- **Impurity**: An unintended constituent present in a substance as manufactured. It may originate from the starting materials or be the result of secondary or incomplete reactions during the manufacturing process. While it is present in the final substance it was not intentionally added.
- **Mixture**: Mix, mixture or solution composed of two or more substances.
- **End product**: Within the scope of these Basic Award Criteria, the end product describes the laundry detergent offered for sale on the market that should be labelled with the Blue Angel ecocert.
- **Microplastic**: Plastic particles with a size of \( \leq 5 \) mm.
- **Plastic**: A macromolecular substance with a water solubility \(< 1 \) mg/L, obtained through:
  a) a polymerisation process such as e.g. polyaddition or polycondensation or a similar process using monomers or other starting substances; or
  b) chemical modification of natural or synthetic micromolecules; or
  c) microbial fermentation.

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2. REACH, Article 3, and CLP Regulation, Article 2
• **Nanomaterial**: means a natural, incidental or manufactured material containing particles, in an unbound state or as an aggregate or as an agglomerate and where, for 50 % or more of the particles in the number size distribution, one or more external dimensions is in the size range 1 nm - 100 nm.⁴

• **Heavy-duty laundry detergent/universal laundry detergent**: Detergent for washing textiles at a temperature range of between 20 and 95 °C and with a reference dosage of 4.5 kg.

• **Colour-safe laundry detergent**: Laundry detergent for washing coloured textiles at a temperature range of between 20 and 60 °C and with a reference dosage of 4.5 kg.

• **Low-duty laundry detergent**: Special detergent for laundry with delicate fabrics with a reference dosage of 2.5 kg.

• **Pre-treatment stain remover**: Stain remover for direct spot treatment of textiles (before washing in the machine) but not including stain removers dosed in the washing machine or stain removers dedicated to other uses besides pre-treatment.

• **Household washing machine**: An automatic washing machine for cleaning and rinsing textiles with water that has a spin function and is primarily designed for non-professional purposes.⁵

• **Sales packaging (in the context of these Basic Award Criteria)**: All of the packaging contained in one sales unit (primary packaging = direct contact with the contents and, if relevant, a secondary packaging) as it is offered to the end user or consumer at a retail outlet in its smallest sales unit.

• **Transport packaging**: Transport packaging or delivery packaging is packaging that facilitates the transport of goods, protects the goods against damage during transport or which is used for reasons of safety of the transport.

• **Identity preserved**: Palm (kernel) oil from a specific production location that is sourced from sustainable plantations is kept separate from other palm oils along the whole supply chain.

• **Segregation**: Palm (kernel) oil from different production locations that is sourced from sustainable plantations is kept separate from other non-certified palm oils along the whole supply chain.

• **Mass balance**: In the mass balance model, sustainable palm oil from certified plantations is mixed with conventional, non-certified palm oil in the value added chain. In this process, the proportion of the certified goods is checked and verified so that no more of the end product is labelled as being certified than the amount of certified palm oil before the mixing process. The certified palm oil is recorded and monitored administratively as it is transferred. The mass balance option thus enables sustainable goods to be verified at every stage of the product chain, without having to establish an additional infrastructure for a parallel supply chain. Due to the fact that the certified and conventional goods are not physically separated, it enables the mass balance goods to be really easily traded within the supply chain. This option is especially relevant for the use of palm kernel oil and its derivatives.

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• **Book & claim**: Sustainable plantations are promoted though the sale of certificates. Companies purchase these certificates via a trading platform (e.g. RSPO Credits) based on the quantity of oil required for the production of surfactants.

2 **Scope**

The product group “laundry detergents” comprises all detergents and pre-treatment stain removers in powder, liquid or other form that are marketed and used for the washing of textiles principally in standard household washing machines but not excluding their use in launderettes and their additional use as a hand washing laundry detergent.

Pre-treatment stain removers are stain removers for the direct spot treatment of textiles (before washing in the machine) but not including stain removers dosed in the washing machine or stain removers dedicated to other uses besides pre-treatment. Excluded from the scope of these Basic Award Criteria are:

- Portioned laundry detergent in water-soluble films
- Stain removers combined with carriers such as sheets, cloths or other materials
- Stain removers for use without subsequent washing e.g. for carpets and upholstered furniture
- Products containing microorganisms that have been intentionally added by the manufacturer.

3 **Requirements**

The end products named under Paragraph 2 can be labelled with the environmental label illustrated on the first page of these Basic Award Criteria if they fulfil the following requirements.

If the applicant is required to submit declarations, documentation, analysis reports or other documentation in order to verify compliance with the criteria, these can come from the applicant and/or his/her suppliers and/or their suppliers, etc.

The substance requirements defined by Regulation (EC) No. 648/2004 (Detergents Regulation, DetVO)\(^6\) on detergents and the CLP Regulation (Regulation (EC) No. 1272/2008)\(^7\) are observed.

3.1 **Assessment and testing requirements**

Paragraph 3.5 refers to the "Detergent Ingredient Database" (DID list 2016), which contains the most widely used substances in detergent formulations. The data found in this list shall be used for deriving the calculations for the Critical Dilution Volume (CDV) and for assessing the biodegradability of the substances. In the case of those substances not found on the DID list,

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guidance is given on how to calculate or extrapolate the relevant data. The DID lists are
published as annexes.
In certain cases, RAL gGmbH can request additional verification and carry out independent tests.

3.1.1 Measurement thresholds
Every substance that exceeds a concentration of 0.010% by mass in the final formulation must comply with these Basic Award Criteria. This also applies to the raw materials used in the product, any listed additives and impurities.
In the case of substances dealt with by the following criteria, a deviating measurement threshold of 0.0010% by mass in the final formulation applies:
• 3.4 Biodegradability
• 3.5 Toxicity to aquatic organisms
• 3.7 Exclusion of substances
• 3.8 Requirements for specific substances
There is no lower measurement threshold for colouring agents.
### Table 1: Overview of the measurement thresholds for the requirement criteria

<table>
<thead>
<tr>
<th>Chapter</th>
<th>Criterion</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.2</td>
<td>Renewable raw materials in <strong>surfactants</strong></td>
</tr>
<tr>
<td>3.3</td>
<td>Requirements for renewable raw materials in the production of <strong>surfactants</strong></td>
</tr>
<tr>
<td>3.4.1</td>
<td>Biodegradability of <strong>surfactants</strong></td>
</tr>
<tr>
<td>3.4.2</td>
<td>Biodegradability of organic substances</td>
</tr>
<tr>
<td>3.5</td>
<td>Toxicity to aquatic organisms</td>
</tr>
<tr>
<td>3.6 a)</td>
<td>General exclusion of substances with certain properties – a) Substances of very high concern (SVHC)</td>
</tr>
<tr>
<td>3.6 b)</td>
<td>General exclusion of substances with certain properties – b) Substances classified with the H-phrases listed in accordance with Regulation (EC) No 1272/2008</td>
</tr>
<tr>
<td>3.7</td>
<td>Exclusion of substances</td>
</tr>
<tr>
<td>3.8.1</td>
<td>Requirements for specific substances – <strong>biocides</strong></td>
</tr>
<tr>
<td>3.8.2</td>
<td>Requirements for specific substances – <strong>fragrances</strong></td>
</tr>
<tr>
<td>3.8.3</td>
<td>Requirements for specific substances – <strong>colouring agents</strong></td>
</tr>
<tr>
<td>3.8.4</td>
<td>Requirements for specific substances – <strong>phosphorous</strong></td>
</tr>
<tr>
<td>3.8.5</td>
<td>Requirements for specific substances – <strong>enzymes</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chapter</th>
<th>Criterion</th>
<th>Measurement threshold in percent by mass [% (w/w)]</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.2</td>
<td>Renewable raw materials in <strong>surfactants</strong></td>
<td>≥ 0.010</td>
</tr>
<tr>
<td>3.3</td>
<td>Requirements for renewable raw materials in the production of <strong>surfactants</strong></td>
<td>≥ 0.010, ≥ 0.010</td>
</tr>
<tr>
<td>3.4.1</td>
<td>Biodegradability of <strong>surfactants</strong></td>
<td>≥ 0.010</td>
</tr>
<tr>
<td>3.4.2</td>
<td>Biodegradability of organic substances</td>
<td>≥ 0.0010 (Colouring agents: no lower limit)</td>
</tr>
<tr>
<td>3.5</td>
<td>Toxicity to aquatic organisms</td>
<td>≥ 0.0010 (Colouring agents: no lower limit)</td>
</tr>
<tr>
<td>3.6 a)</td>
<td>General exclusion of substances with certain properties – a) Substances of very high concern (SVHC)</td>
<td>≥ 0.010 (Colouring agents: no lower limit)</td>
</tr>
<tr>
<td>3.6 b)</td>
<td>General exclusion of substances with certain properties – b) Substances classified with the H-phrases listed in accordance with Regulation (EC) No 1272/2008</td>
<td>≥ 0.010 (Colouring agents: no lower limit)</td>
</tr>
<tr>
<td>3.7</td>
<td>Exclusion of substances</td>
<td>≥ 0.0010</td>
</tr>
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<td>≥ 0.0010</td>
</tr>
<tr>
<td>3.8.2</td>
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<td>≥ 0.0010</td>
</tr>
<tr>
<td>3.8.3</td>
<td>Requirements for specific substances – <strong>colouring agents</strong></td>
<td>≥ 0.0010 (no lower limit)</td>
</tr>
<tr>
<td>3.8.4</td>
<td>Requirements for specific substances – <strong>phosphorous</strong></td>
<td>≥ 0.0010</td>
</tr>
<tr>
<td>3.8.5</td>
<td>Requirements for specific substances – <strong>enzymes</strong></td>
<td>≥ 0.0010</td>
</tr>
</tbody>
</table>

### 3.1.2 Functional unit

The functional unit for this product group shall be expressed in g/kg laundry (grams per kilogram of laundry).

### 3.1.3 Reference dosage

For ‘heavy-duty laundry detergents’ and ‘colour-safe laundry detergents’, the dosage recommended by the manufacturer for a water hardness level of 2.5 mmol CaCO₃/l and ‘normally soiled’ textiles is taken as the reference dosage for calculating the environmental criteria and testing the washing performance. For heavy-duty laundry detergents and colour-safe laundry detergents, this refers to the dosage for a 4.5 kg load (dry textiles) in the washing machine.
For ‘low-duty laundry detergents’, the dosage recommended by the manufacturer for a water hardness level of 2.5 mmol CaCO₃/l and ‘lightly soiled’ textiles is taken as the reference dosage for calculating the environmental criteria and testing the washing performance. For low-duty laundry detergents, this refers to the dosage for a 2.5 kg load (dry textiles) in the washing machine.

If the recommended dosage refers to other load sizes than the figures mentioned above, the reference dosage used for calculating the environmental criteria must correspond to the average load size. If the dosage instructions on the packaging state values for a prewash and a main washing cycle (in addition to a single normal washing cycle), the total dosage (prewash and main wash) must also comply with the environmental criteria.

The maximum CDV limit in the case of stain removers is based on an estimated dosage of 2 ml per application and six applications for a 4.5 kg load (2.7 ml/kg laundry; liquid stain remover). Products added in powder or paste form must also comply with these maximum CDV limits. Cloths soaked with liquid laundry detergent are considered to be liquids in the context of these Basic Award Criteria.

3.1.4 Testing institutions

The tests to be submitted to verify compliance with the requirements, with the exception of the tests for the product's fitness for use, shall be carried out by testing institutions that fulfil the following requirements:

- The tests comply with the requirements of Good Laboratory Practice (Annex 1 of German Chemicals Act (ChemG)) or
- The testing institution has been notified or accredited according to DIN EN 17025 and these tests form part of this accreditation in terms of the fields being tested and the processes and specifications used.

Compliance verification

Verification of compliance is to be provided in the form of certification in accordance with Article 19b of the German Chemicals Act (ChemG) and a written declaration from the testing institution that the test was carried out according to the principles of Good Laboratory Practice or through submission of the accreditation certificate from Germany’s National Accreditation Body (DAKKS) or another national accreditation system that has been included in the Multilateral Agreement (MLA).

3.2 Renewable raw materials in surfactants

The proportion of renewable carbon in the total carbon in the surfactant system must be at least 50%.

Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1. The proportion of renewable carbon is calculated based on the organic carbon (Annex 2) and verified with a declaration by the surfactant manufacturer. The calculation shall be based on the annual production volume. (see Appendix A)
3.3 Requirements for renewable raw materials produced from palm oil and palm kernel oil

If raw materials produced from palm oil and palm kernel oil are used, the sustainable cultivation of the oil plants on certified plantations must be verified.

The renewable raw materials for all of the ingredients added to the product that are produced from palm oil and/palm kernel oil must be sourced from plantations cultivated under sustainable conditions. Raw materials produced from palm (kernel) oil must be certified at least in accordance with the “mass balance” model.

Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1.

In the case of RSPO certification, the manufacturing company must verify their membership of the RSPO (in the case of first-time applications, to be applied for after the issuing of the contract).

A supply chain audit via a corresponding certificate shall be submitted to RAL gGmbH at the latest after the Blue Angel ecolabel has been used for the first 15 months and then annually. Verification via the Book & Claim system is not sufficient.

The following certification systems are recognised: RSPO (Roundtable on Sustainable Palm Oil), ISCC+ (International Sustainability & Carbon Certification) or RSB (Roundtable on Sustainable Biomaterial).<sup>8</sup> (Appendix A)

3.4 Biodegradability

3.4.1 Biodegradability of surfactants

All of the surfactants contained in the end product must be readily biodegradable under aerobic conditions and biodegradable under anaerobic conditions.

Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1.

The precise formulation of the end product shall be submitted to RAL gGmbH together with an explanation of the function of every individual substance in Annex 2. Part A of the DID list indicates whether a certain surfactant is aerobically or anaerobically biodegradable (those surfactants with an "R" in the column for aerobic biodegradability are readily biodegradable, while those surfactants with a "Y" in the column for anaerobic biodegradability are biodegradable under anaerobic conditions). The list is not comprehensive, but guidance is given in Part B of the list concerning the determination of the relevant calculation parameters for substances not present on the DID list. For those surfactants which are not included in Part A of the DID list or those surfactants classified with an "O" in the column for anaerobic biodegradability, relevant information from literature or other sources or corresponding test results shall be submitted to verify that these surfactants are biodegradable under anaerobic conditions.

The reference test for anaerobic degradability shall be the OECD test 311, the ISO standard 11734, the ECETOC test No. 28 (June 1988) or an equivalent test method, with the requirement

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<sup>8</sup> It is possible that other certification systems will be accepted after they have been investigated by the Federal Environmental Agency.

<sup>9</sup> According to the regulations in EU Regulation No. 648/2004/EC
of 60 % ultimate degradability under anaerobic conditions. In order to verify at least 60% ultimate degradability under anaerobic conditions, test processes can also be used that simulate the conditions in a corresponding anaerobic environment.

3.4.2 Biodegradability of organic substances

The content of aerobically not readily biodegradable (aNBO) and anaerobically non-biodegradable (anNBO) organic substances in the product must not exceed the following limits for the reference dosage (information in g/kg laundry).

<table>
<thead>
<tr>
<th>Product type/product form</th>
<th>Solid (powder, tablet)</th>
<th>Liquid (incl. gel)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>aNBO</td>
<td>anNBO</td>
</tr>
<tr>
<td>Heavy-duty laundry detergent, colour-safe laundry detergent</td>
<td>0.75</td>
<td>1.00</td>
</tr>
<tr>
<td>Low-duty laundry detergent</td>
<td>0.40</td>
<td>0.40</td>
</tr>
<tr>
<td>Stain remover</td>
<td>0.10</td>
<td>0.10</td>
</tr>
</tbody>
</table>

Compliance verification:
The applicant shall confirm compliance with the requirement in Annex 1.
In order to calculate the aNBO and anNBO values, there is an Excel file available for this purpose on the Blue Angel website (Annex 2).
The DID list is authoritative here. In the case of ingredients not included in Part A of the DID list, relevant information from literature or other sources or corresponding test results shall be submitted to verify that these ingredients are biodegradable under aerobic and anaerobic conditions (as described in Part B of this list).

3.5 Toxicity to aquatic organisms

The critical dilution volume toxicity (CDV\textsubscript{chronic}) is calculated for each substance (i) using the following equation:

\[
CDV_{\text{chronic}} = \sum_{i=1}^{n} CDV_{(i)} = CDV_{(1)} + CDV_{(2)} + \ldots + CDV_{(n)} \\
\]

with

\[
CDV_{(i)} = \frac{\text{Weight}_{(i)} \cdot DF_{(i)} \cdot 1000}{TF_{\text{chronisch}}(i)}
\]

where the weight\((i)\) is the weight of the substance (in grams) in the recommended dosage.
DF(i) is the degradation factor and TF\textsubscript{chronic}(i) is the value for the chronic toxicity of the substance (in milligrams/litre).

The parameters DF and TF\textsubscript{chronic} shall be taken from Part A of the Detergent Ingredient Database (DID list) (Appendix). If the substance in question is not included in Part A of the DID list, the applicant shall estimate the values in accordance with the approach stated in Part B of the DID list (Appendix). The sum of CDV\textsubscript{chronic} for the individual substances gives the CDV\textsubscript{chronic} for the end product.

The CDV\textsubscript{chronic} must not exceed the following limits for the reference dosage:

<table>
<thead>
<tr>
<th>Type of product</th>
<th>Maximum permissible CDV\textsubscript{chronic}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heavy-duty laundry detergent, colour-safe laundry detergent</td>
<td>25 000 l/kg laundry</td>
</tr>
<tr>
<td>Low-duty laundry detergent</td>
<td>18 000 l/kg laundry</td>
</tr>
<tr>
<td>Stain remover</td>
<td>3 500 l/kg laundry</td>
</tr>
</tbody>
</table>

**Compliance verification:**

The applicant shall confirm compliance with the requirement in Annex 1. The precise formulation of the end product, together with the individual details of the calculation of the CDV\textsubscript{chronic} demonstrating compliance with this criterion, shall be submitted to RAL gGmbH in Annex 2.

3.6 General exclusion of substances with certain properties

The use of the following substances is not permitted in order to protect the environment and human health. In the case of mixtures e.g. fragrances where it is not possible to obtain information about the individual substances, the classification rules for mixtures shall be applied.

a) Substances of very high concern (SVHC)

It is prohibited to use substances in end products labelled with the Blue Angel ecolabel that have been identified in accordance with Article 57 of Regulation (EC) No. 1907/2006 and listed in accordance with Article 59 of the same regulation on the list of candidates\textsuperscript{10} for inclusion on the Annex of substances subject to authorisation. Impurities in substances added to the end product that correspond to the above named criteria are not permitted. The label holder is obligated to take into account current developments on the list of candidates.

b) Substances which according to the criteria of Regulation (EC) No 1272/2008\textsuperscript{11} are assigned the following H Phrases named in Table 2 or which meet the criteria for such classification.

\textsuperscript{10} \url{http://echa.europa.eu/web/guest/candidate-list-table}

Table 2: Restrictive hazard classifications and their assignment to the categories

<table>
<thead>
<tr>
<th>EC Regulation 1272/2008 (CLP Regulation)</th>
<th>Wording</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Toxic substances</strong></td>
<td></td>
</tr>
<tr>
<td>H300</td>
<td>Fatal if swallowed</td>
</tr>
<tr>
<td>H301</td>
<td>Toxic if swallowed</td>
</tr>
<tr>
<td>H304</td>
<td>May be fatal if swallowed and enters airways</td>
</tr>
<tr>
<td>H310</td>
<td>Fatal in contact with skin</td>
</tr>
<tr>
<td>H311</td>
<td>Toxic in contact with skin</td>
</tr>
<tr>
<td>H330</td>
<td>Fatal if inhaled</td>
</tr>
<tr>
<td>H331</td>
<td>Toxic if inhaled</td>
</tr>
<tr>
<td>EUH070</td>
<td>Toxic by eye contact</td>
</tr>
<tr>
<td>H370</td>
<td>Causes damage to organs</td>
</tr>
<tr>
<td>H371</td>
<td>May cause damage to organs</td>
</tr>
<tr>
<td>H372</td>
<td>Causes damage to organs through prolonged or repeated exposure</td>
</tr>
<tr>
<td>H373</td>
<td>May cause damage to organs through prolonged or repeated exposure</td>
</tr>
<tr>
<td><strong>Carcinogenic, mutagenic and reprotoxic substances</strong></td>
<td></td>
</tr>
<tr>
<td>H340</td>
<td>May cause genetic defects</td>
</tr>
<tr>
<td>H341</td>
<td>Suspected of causing genetic defects</td>
</tr>
<tr>
<td>H350</td>
<td>May cause cancer</td>
</tr>
<tr>
<td>H350i</td>
<td>May cause cancer if inhaled</td>
</tr>
<tr>
<td>H351</td>
<td>Suspected of causing cancer</td>
</tr>
<tr>
<td>H360F</td>
<td>May damage fertility</td>
</tr>
<tr>
<td>H360D</td>
<td>May damage the unborn child</td>
</tr>
<tr>
<td>H360FD</td>
<td>May damage fertility</td>
</tr>
<tr>
<td></td>
<td>May damage the unborn child</td>
</tr>
<tr>
<td>H360Fd</td>
<td>May damage fertility</td>
</tr>
<tr>
<td></td>
<td>Suspected of damaging the unborn child</td>
</tr>
<tr>
<td>H360Df</td>
<td>May damage the unborn child</td>
</tr>
<tr>
<td></td>
<td>Suspected of damaging fertility</td>
</tr>
<tr>
<td>H361f</td>
<td>Suspected of damaging fertility</td>
</tr>
<tr>
<td>H361d</td>
<td>Suspected of damaging the unborn child</td>
</tr>
<tr>
<td>H361fd</td>
<td>Suspected of damaging fertility</td>
</tr>
<tr>
<td></td>
<td>Suspected of damaging the unborn child</td>
</tr>
</tbody>
</table>

Preparations Directive). According to the said regulation, substances are classified, labelled and packed until 1 December 2010 according to Directive 67/548/EEC while mixtures (formerly preparations) are classified, labelled and packed until 1 June 2015 according to Directive 1999/45/EC. Thereafter, the GHS Regulation shall be applied. The new indications of danger (H Phrases) as well as the hitherto applicable risk phrases (R Phrases) shall be indicated for substances until 1 June 2015.
<table>
<thead>
<tr>
<th>EC Regulation 1272/2008 (CLP Regulation)</th>
<th>Wording</th>
</tr>
</thead>
<tbody>
<tr>
<td>H362</td>
<td>May cause harm to breast fed children</td>
</tr>
</tbody>
</table>

**Water-hazardous substances**

<table>
<thead>
<tr>
<th>Wording</th>
</tr>
</thead>
<tbody>
<tr>
<td>H400 Very toxic to aquatic life</td>
</tr>
<tr>
<td>H410 Very toxic to aquatic life with long-lasting effects</td>
</tr>
<tr>
<td>H411 Toxic to aquatic life with long-lasting effects</td>
</tr>
<tr>
<td>H412 Harmful to aquatic life with long lasting effects</td>
</tr>
<tr>
<td>H413 May cause long lasting harmful effects to aquatic life</td>
</tr>
</tbody>
</table>

**Other Health and Environmental Effects**

<table>
<thead>
<tr>
<th>Wording</th>
</tr>
</thead>
<tbody>
<tr>
<td>H420 Hazardous to the ozone layer</td>
</tr>
</tbody>
</table>

**Sensitizing substances**

<table>
<thead>
<tr>
<th>Wording</th>
</tr>
</thead>
<tbody>
<tr>
<td>H334 May cause allergy or asthma symptoms or breathing difficulties if inhaled</td>
</tr>
<tr>
<td>H317 May cause an allergic skin reaction</td>
</tr>
</tbody>
</table>

The use of substances or mixtures which upon processing change their properties (e.g. become no longer bioavailable, undergo chemical modification) in a way that the identified hazard no longer applies are exempted from the above requirement.

**Deviations:** The following substances or mixtures are specifically exempted from the above requirement:

<table>
<thead>
<tr>
<th>EC Regulation 1272/2008 (CLP Regulation)</th>
<th>Wording</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surfactant (*)</td>
<td>H400 Very toxic to aquatic life</td>
</tr>
<tr>
<td></td>
<td>H412 Harmful to aquatic organisms with long lasting effects</td>
</tr>
<tr>
<td>Enzymes (**)</td>
<td>H334 May cause allergy or asthma symptoms or breathing difficulties if inhaled</td>
</tr>
<tr>
<td></td>
<td>H317 May cause an allergic skin reaction</td>
</tr>
<tr>
<td>Proteases (e.g. subtilisin)</td>
<td>H400 Very toxic to aquatic life</td>
</tr>
<tr>
<td></td>
<td>H411 Toxic to aquatic organisms with long lasting effects</td>
</tr>
<tr>
<td>NTA as an impurity in MGDA and GLDA(***)</td>
<td>H351 Suspected of causing cancer</td>
</tr>
</tbody>
</table>

(*) This is also valid for impurities from the starting substances.

(**) Including stabilisers and other auxiliary substances in the preparations.

(*** In concentrations lower than 0.2 % in the raw material as long as the total concentration in the end product is lower than 0.10 %.)
Compliance verification

The applicant shall declare compliance with the requirements in Annex 1.
In the event of changes to the list of candidates, the applicant shall inform RAL gGmbH within one month in the event that the end product does not comply with this criterion.
The applicant shall submit the exact formulation of the end product in Annex 2. The applicant shall verify that the substances contained in the end product comply with this criterion by providing information that fulfils at least those requirements according to Annex VII of Regulation (EC) No. 1907/2006. Such information shall be specific to the particular form of the substance, including nanoforms, used in the end product. For that purpose, the applicant shall submit a declaration of compliance with this criterion, together with information on the type (IUPAC nomenclature and CAS number) and content (% by mass) of all substances added to the product and the related safety data sheets in accordance with Annex II to Regulation (EC) No 1907/2006 for the end product, as well as for all substances or mixtures listed in the formulation(s). Concentration limits shall be specified in the safety data sheets in accordance with Article 31 of Regulation (EC) No. 1907/2006. The safety data sheets may not be older than two years. The manufacturer shall verify that he/she has requested that the suppliers of primary/intermediate products submit information on the content of substances up to 0.010 % by mass.

3.7 Exclusion of substances

The following substances are not permitted in the end product, either as part of the formulation or as part of any preparation included in the formulation:
- Alkyl phenol ethoxylates (APEOs) and derivatives thereof
- EDTA (ethylenediaminetetraacetic acid) and its salts
- DTPA (diethylenetriaminepentaacetic acid) and its salts
- Inorganic phosphate(*) (e.g. monophosphoric, diphosphoric, triphosphoric and polyphosphoric acids and their salts)
- Reactive chlorine compounds (e.g. hypochlorite)
- Borate and perborate
- Perfluorinated organic compounds
- Halogenated hydrocarbons
- Aromatic hydrocarbons
- Triclosan
- 3-Jod-2-propinylbutylcarbamate
- Glutaraldehyde
- Quaternary organic ammonium compounds that are not readily biodegradable
- Formaldehyde and formaldehyde releasers(**), e.g. (INCI designations):
  - 2-Bromo-2-Nitropropane-1,3-Diol
  - 5-bromo-5-nitro-1,3-dioxane
  - Diazolidinyl Urea
  - Sodium Hydroxyethylglycinate
  - Dimethylol Glycol
  - Dimethylol Urea
  - DMDM-Hydantoin
  - Quaternium-15
• Tetramethylolglycoluril
• Nitromusks and polycyclic musks including e.g.:
  • Musk xylene: 5-tert-butyl-2,4,6-trinitro-m-xylene,
  • Musk ambrette: 4-tert-butyl-3-methoxy-2,6-dinitrotoluene,
  • Moskene: 1,1,3,3,5-Pentamethyl-4,6-dinitroindan,
  • Musk tibetine: 1-tert-butyl-3,4,5-trimethyl-2,6-dinitrobenzene,
  • Musk ketone: 4’-tert-Butyl-2’,6’-dimethyl-3’,5’-dinitroacetophenol,
  • HHCB (1,3,4,6,7,8-Hexahydro-4,6,6,7,8,8-hexamethylcyclopenta(g)-2-benzopyran),
  • AHTN (6-acetyl-1,1,2,4,4,7-hexamethyltetrali)
• Nanosilver
• Hydroxyisohexyl 3-cyclohexene carboxaldehyde (HICC)
• Atranol
• Chloratranol
• Rhodamine B
• Microplastics

(*) Except for impurities or stabilisers with concentrations lower than 1.0% in the raw material and a total concentration in the end product lower than 0.010%.

(**) Except for impurities of formaldehyde in surfactants based on polyalkoxy compounds up to a concentration of 0.010% by mass in the ingredient.

Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1. The applicant shall submit a declaration, which is also supported by declarations from the manufacturers of the substances, that the listed substances are not contained in the end product. In the event that quaternary ammonium salts have been added to the product, the applicant shall submit documentation demonstrating their biodegradability.

3.8 Requirements for specific substances

These special requirements for specific substances are valid in addition to the general requirements for substances.

3.8.1 Biocides

a) The end product may only include biocides in order to preserve the product and in the appropriate dosage for this purpose. This does not refer to surfactants, which may also have biocidal properties.

Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1. The applicant shall submit the safety data sheets for every preservative added to the product, as well as information about the exact concentrations of these substances in the end product. The manufacturer or supplier of the preservatives shall submit information about the dosage required to preserve the end product.

b) It is prohibited to claim or suggest on the packaging or by any other means that the product has an antimicrobial action.
Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1. The applicant shall submit the texts and layouts used for each individual type of packaging and/or a sample copy of each individual type of packaging to RAL gGmbH.

c) Biocides, either as part of the formulation or as part of any mixture included in the formulation, that are used to preserve the end product are permitted, but only if their log Pow (octanol-water partition coefficient) is < 3.0 or their experimentally determined bioconcentration factor (BCF) is < 100.

Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1. The applicant shall submit the log Pow or BCF value for the biocides (Annex 2).

d) Isothiazolinone

2-methyl-4-isothiazolin-3-one (MIT) and 5-chlor-2-methyl-4-isothiazolin-3-one/2-methyl-4-isothiazolin-3-one (CIT/MIT) may only be added up to the maximum permitted limits stated in Annex V (List of preservatives allowed in cosmetic products) of Regulation (EC) No. 1223/2009 of the European Parliament and of the Council of 30 November 2009 on cosmetic products.

- 5-chlor-2-methyl-4-isothiazolin-3-one/2-methyl-4-isothiazolin-3-one: 0.0015 % by mass.
- 2-methyl-4-isothiazolin-3-one: 0.0015 % by mass.

If the permissible value according to Annex V of Regulation (EC) No. 1223/2009 at the time of application is lower, this lower value applies.

The content of 1,2-benzisothiazol-3(2H)-one (BIT) must not exceed the following content in the product:

- 1,2-benzisothiazol-3(2H)-one: 0.0050 % by mass

Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1.

3.8.2 Fragrances

a) All of the substances added to the end product as fragrances must have been manufactured and/or handled in accordance with the code of practice of the International Fragrance Association (IFRA)\textsuperscript{12}.

b) Fragrances that are subject to labelling as detergents in accordance with Annex VII of Regulation (EC) No. 648/2004 and which are not already excluded by criteria 3.6, as well as (other) fragrances classified as H317 (May cause an allergic skin reaction) and/or H334 (May cause allergy or asthma symptoms or breathing difficulties if inhaled) must not be present in the end product in concentrations $\geq 0.010$ % ($\geq 100$ ppm) per substance.

c) Products specially designed for or advertised as being suitable for allergy sufferers are not permitted to contain any fragrances.

\textsuperscript{12} The code of practice is available on the IFRA website: \url{http://www.ifraorg.org}.
**Compliance verification**

The applicant shall confirm compliance with the requirement in Annex 1.
The applicant shall submit a declaration of compliance with every requirement. For criterion (b), the applicant shall submit a declaration about compliance with this criterion with information about the amount of fragrances contained in the end product. In addition, the applicant shall also submit a declaration from the fragrance manufacturer specifying the content of each of the substances contained in the fragrance which are listed in Annex III of Regulation (EC) No. 1223/2009\(^\text{13}\), as well as the content of (other) substances which have been assigned the risk phrases H317 and/or H334.

### 3.8.3 Colouring agents

The end product must not contain any colouring agents that are bioaccumulating.
A colouring agent is considered to be non-bioaccumulating if it has a bioconcentration factor (BCF) < 100 or a log Pow < 3.0. If the values for both the BCF and the log Pow are available, the highest measurement for the BCF is valid. If using colouring agents that have been approved for use in foodstuffs, no documentation about the bioconcentration factor needs to be submitted.

**Compliance verification**

The applicant shall submit a signed declaration of conformity and, if relevant, supplier declarations or safety data sheets about all colouring agents added to the product and the values for their BCF or log Pow or documentation that verifies that the colouring agent is approved for use in foodstuffs.

#### 3.8.4 Phosphorous

The total phosphorous content (P) of elemental phosphorous must not exceed the following values for the reference dosage.

<table>
<thead>
<tr>
<th>Type of product</th>
<th>Phosphorous content</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heavy-duty laundry detergent, colour-safe laundry detergent</td>
<td>0.040 g/kg laundry</td>
</tr>
<tr>
<td>Low-duty laundry detergent</td>
<td>0.040 g/kg laundry</td>
</tr>
<tr>
<td>Stain remover</td>
<td>0.0050 g/cleaning cycle</td>
</tr>
</tbody>
</table>

**Compliance verification**

The applicant shall confirm compliance with the requirement in Annex 1. In order to calculate the total phosphorous content (P), there is an Excel file available for this purpose on the Blue Angel website (Annex 2).

### 3.8.5 Enzymes

It is only permitted to add encapsulated enzymes (solid) and enzymes in liquid form or as a suspension.

\(^{13}\) OJ L 342 from 22/12/2009
Compliance verification:
The applicant shall submit a signed declaration of conformity and, if relevant, supplier declarations or safety data sheets for the enzymes added to the product.

3.9 Labelling of the end product

a) It is not permitted for the end product to be assigned a H phrase named in Table 2 of Paragraph 3.6 b) in accordance with the CLP Regulation (1272/2008/EC)\(^{14}\).

Compliance verification
The applicant shall confirm compliance with the requirement in Annex 1.
In addition, the applicant shall submit the exact formulation of the end product in Annex 2, as well as the product label and safety data sheet for the end product.

3.10 Dosage requirements
The dosage corresponds to the recommended dosage in g/kg laundry (powder/tablets) or ml/kg laundry (liquids). The recommended dosage for a water hardness level of 2.5 mmol CaCO₃/l and for normally soiled textiles (heavy-duty laundry detergents, colour-safe laundry detergents) and lightly soiled textiles (low-duty detergents), respectively, shall be used.
The dosage must not exceed the following maximum limits:

<table>
<thead>
<tr>
<th>Type of product</th>
<th>Dosage solid/liquid (incl. gel)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heavy-duty laundry detergent, colour-safe laundry detergent</td>
<td>16.0 g/kg laundry</td>
</tr>
<tr>
<td>Low-duty laundry detergent</td>
<td>16.0 g/kg laundry</td>
</tr>
</tbody>
</table>

If using the recommendations for a prewash and a main washing cycle, the total recommended dosage (prewash and main wash) must not exceed the stated maximum limit.

Compliance verification
The applicant shall declare compliance with the requirement in Annex 1 and submit a product label. The density (g/ml) shall be stated for all products (either on the packaging or in a safety data sheet).

3.11 Fitness for use
The end product must be fit for use and meet the requirements of the consumer. To ensure that this is the case, the tests for assessing the product's fitness for use described in Appendix B shall be carried out.

Compliance verification
The applicant shall declare compliance with the requirement in Annex 1 and submit the test results in accordance with the guidelines stated in Appendix B to verify compliance.

3.12 Packaging requirements

a) The weight utility ratio (WUR) of the primary packaging must not exceed the following values:

<table>
<thead>
<tr>
<th>Type of product</th>
<th>WUR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solid laundry detergent (e.g. powder)</td>
<td>1.2 g/kg laundry</td>
</tr>
<tr>
<td>Liquid or gel laundry detergent</td>
<td>1.2 g/kg laundry</td>
</tr>
<tr>
<td>Stain remover</td>
<td>1.2 g/kg laundry</td>
</tr>
</tbody>
</table>

Primary packagings consisting of more than 80% renewable materials are exempt from this requirement.

The WUR is only calculated for the primary packaging (including caps, stoppers and label) based on the following formula:

\[
WUR = \sum \left( \frac{(W_i + U_i)}{(D_i \times R_i)} \right)
\]

Key for the calculation formula:
- \( W_i \): the weight [g] of the primary packaging (i);
- \( U_i \): the weight [g] of non-recycled materials in the primary packaging (i). \( U_i = W_i \), unless the applicant can verify a different number;
- \( D_i \): the number of reference doses in the primary packaging (i). In the case of ready-to-use products, \( D_i = \) product volume (in litres);
- \( R_i \): recycling figure. \( R_i = 1 \) (if the packaging is not reused for the same purpose) or \( R_i = 2 \) (if the applicant can verify that the packaging components are used for the same purpose and he/she sells refill packs).

b) Packagings, sleeves, labels or closures made of halogenated polymers e.g. PVC are not permitted.

c) Paper/cardboard used in the sales packaging must be manufactured using at least 80% recycled materials. In the case of secondary packaging that also serves as transport packaging, the proportion of recycled materials must be at least 70%. Packaging materials are considered recycled if product waste (post-consumer waste) has been subjected to a material recycling process.

d) Recycling-oriented design

Plastic packaging must be designed for the purpose of easy recycling i.e. where possible no potentially hazardous materials and incompatible materials should be used that are known to hinder the separation or recycling of the materials or reduce the quality of the recycled materials. No individual or combination of materials or components listed in Table 3 may be contained in the labels or sleeves, closures and, if relevant, barrier coatings. Pump mechanisms (including in sprays) are exempt from this requirement.
Table 3: Materials and components that are excluded from use as a packaging component:

<table>
<thead>
<tr>
<th>Packaging component</th>
<th>Excluded materials and components (*)</th>
</tr>
</thead>
</table>
| All components      | - Components in the EuPIA list (exclusion list for printing inks and related products)  
                      - PS label or PS sleeve in combination with a PP, HDPE or PET bottle  
                      - A PETG or PETC label or a PETG sleeve or PETC sleeve in combination with a PET bottle  
                      - Other plastic materials for sleeves/labels with a density > 1 g/cm³ in combination with a PET bottle  
                      - Other plastic materials for sleeves/labels with a density < 1 g/cm³ in combination with a PP or HDPE bottle  
                      - Labels or sleeves that are metallised or labels or sleeves that are welded without a seam to a packaging body (in mould labelling) with PET bottles  
                      - Non water-soluble adhesives in combination with moisture-resistant labels with a PE or PP bottle, non-soluble adhesive (in water or alkaline at 80°C) for PET bottles  
| Label or sleeve     | - A PS closure in combination with a PP, HDPE or PET bottle  
                      - PETG closures and/or PETG closure material and other plastic closure components with a density of above 1 g/cm³ in combination with a PET bottle  
                      - Closures made of metal, glass, EVA  
                      - Closures made of silicone. Exempted are silicone closures with a density < 1 g/cm³ in combination with a PET bottle and silicone closures with a density > 1 g/cm³ in combination with a PP or HDPE bottle  
                      - Silicon components with PE and PP bottles  
                      - Components made out of foamed elastomers with a PE and PP bottle  
                      - Elastomer components with a density > 1 g/cm³ with a PET bottle  
                      - Metallic foils or seals which remain fixed to the bottle or its closure after the product has been opened  
| Closures            | - (*) EVA — Ethylene vinyl acetate, EVOH — Ethylene vinyl alcohol, HDPE — High-density polyethylene, PET — Polyethylene terephthalate,  
                      - PETG — Polyethylene terephthalate glycol-modified, PETC — Crystalline polyethylene terephthalate,  
                      - PP — Polypropylene, PS — Polystyrene, PVC — Polyvinyl chloride  
                      Please note: the combinations PP and HDPE and also PE and LLDPE, LDPE or HDPE are permitted.  

(*)
Compliance verification

The applicant shall confirm compliance with the requirement in Annex 1.
The applicant shall submit a calculation of the WUR for the end product in Annex 2.

3.13 Consumer information

3.13.1 Advertising claims

If the product is classified and labelled as hazardous to human health according to Regulation (EC) No. 1272/2008, advertising claims such as “less environmentally damaging”, “less hazardous to water”, “less hazardous substances” or comparable statements which could be considered to play down the risks are prohibited.

Compliance verification

The applicant shall declare compliance with the requirement in Annex 1 and submit a product label to RAL gGmbH.

3.13.2 Dosage instructions for laundry detergent

The recommended dosages shall be stated for “lightly”, “normally” and “heavily” soiled laundry and the different water hardness levels, as well as the weight of the laundry per load.
The reference dosage used for the washing performance test and the evaluation of compliance with the environmental criteria for the ingredients for heavy-duty laundry detergents and colour-safe laundry detergents must be the same as the recommended dosage on the packaging for “normally soiled” laundry at a water hardness level of 2.5 mmol CaCO₃/l.
The reference dosage used for the washing performance test and the evaluation of compliance with the environmental criteria for the ingredients for low-duty laundry detergents must be the same as the recommended dosage on the packaging for “lightly soiled” laundry at a water hardness level of 2.5 mmol CaCO₃/l.
A dosing aid must be provided free of charge on request for laundry detergents that need to be measured out by the end consumer. If the dosing aid is not enclosed with every packaging unit or it cannot be stocked at the retail outlet, it must be possible to request it via a free hotline, via email or via the Internet and the subsequent delivery must be completed free of charge.

Compliance verification

The applicant shall declare compliance with the requirement in Annex 1 and submit a product label to RAL gGmbH.

3.13.3 Information on the packaging

a) The type of enzyme contained in the product must be stated on the packaging.
b) The packaging must also contain the following or comparable instructions:
   • Wash at the lowest possible temperature.
   • Fill the drum with maximum possible load for the type of textile.
   • Dose the laundry detergent according to the level of soiling and the level of water hardness, follow the dosage instructions to achieve the best possible washing performance with the least amount of detergent.
   • Information on obtaining a dosage aid (if relevant).
• Instructions on where information about water hardness can be found.

c) The primary packaging must contain information on reuse, recycling and the proper disposal of the packaging.

**Compliance verification**

The applicant shall declare compliance with the requirement in Annex 1 and submit a product label to RAL gGmbH.

### 3.13.4 Safety Instructions

End products must carry the following safety instructions (or an equivalent text) in either text form or as a pictogram: "Keep away from children!"

**Compliance verification**

The applicant shall declare compliance with the requirement in Annex 1 and submit a product label to RAL gGmbH.
4 Applicants and parties involved

Manufacturers or distributors of products according to Paragraph 2 shall be eligible for application.

Parties involved in the award process are:

- RAL gGmbH to award the Blue Angel environmental label,
- the federal state being home to the applicant’s production site,
- Umweltbundesamt, (Federal Environmental Agency) which after the signing of the contract receives all data and documents submitted in application for the Blue Angel in order to be able to further develop the Basic Award Criteria.

5 Use of the Environmental Label

The use of the environmental label by the applicant is governed by a contract on the use of the environmental label concluded with RAL gGmbH.

Within the scope of such contract, the applicant undertakes to comply with the requirements under Paragraph 3 while using the environmental label.

Contracts on the Use of the Environmental Label are concluded to fix the terms for the certification of products under Paragraph 2. Such contracts shall run until December 31, 2022. They shall be extended by periods of one year each, unless terminated in writing by December 31, 2022 or March 31 of the respective year of extension.

After the expiry of the contract, the Environmental Label may neither be used for labelling nor for advertising purposes. This regulation shall not affect products being still in the market.

The applicant (manufacturer) shall be entitled to apply to RAL gGmbH for an extension of the right to use the ecolabel on the product entitled to the label if it is to be marketed under another brand/trade name and/or other marketing organizations.

The Contract on the Use of the Environmental Label shall specify:

- Applicant (manufacturer/distributor)
- Brand/trade name, product description
- Distributor (Label User), i.e. the marketing organization.

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Appendix A  Renewable raw materials in surfactants

The proportion of renewable carbon in the total carbon in the surfactant system must be at least 50%.

A declaration from the manufacturer or supplier of the surfactant shall be enclosed with the application as verification for every surfactant (or surfactant raw material) added to the product. (Declaration from the raw material manufacturer).

In Section 16 of the declaration, the manufacturer or supplier of the surfactant shall certify the proportion of renewable carbon in the total carbon for the surfactant or surfactant raw material. In Annex 2 (Excel file), this value shall be entered in the sheet "Result-3" in column E for every surfactant or surfactant raw material added to the product (value between 0 and 100). The subsequent calculation is carried out in Annex 2 as follows:

For every surfactant/surfactant raw material added to the product:

\[ G(i) = \text{Proportional weight of the surfactant/surfactant raw material } i \text{ (column C)} \]

\[ R(i) = \text{Proportion of renewable carbon in the total carbon for the surfactant or surfactant raw material } i \text{ (column E)} \]

The proportion of renewable carbon in the total carbon in the surfactant system (value in cell F62) is calculated using the following formula:

\[ \frac{\sum G(i) \times R(i)}{G(i)} \]

If the surfactant system in the formulation changes during the term of the contract, an amended Annex 2 and a declaration from the new manufacturer of the surfactant shall be submitted. In the annual production volume, the proportion of renewable carbon in the total carbon in the surfactant system must be at least 50%.

\[ P_1 = \text{Production volume in the calendar year with the "old" surfactant system} \]

\[ P_2 = \text{Production volume in the calendar year with the "new" surfactant system} \]

\[ R_1 = \text{Proportion of renewable carbon in the total carbon in the "old" surfactant system} \]

\[ R_2 = \text{Proportion of renewable carbon in the total carbon in the "new" surfactant system} \]

\[ R_J = \text{Average yearly proportion of renewable carbon in the total carbon (must be } \leq 50\% \text{)} \]

\[ R_J = \frac{(P_1 \times R_1) + (P_2 \times R_2)}{(P_1 + P_2)} \]
Requirements for renewable raw materials produced from palm oil and palm kernel oil

If surfactants produced from palm oil and palm kernel oil are used, the sustainable cultivation of the oil plants on certified plantations must be verified.

In section 16 of the “Declaration from the raw material manufacturer”, the manufacturer or supplier shall confirm whether the raw material contains palm/palm kernel oil. If raw materials produced from palm oil and palm kernel oil have been added to the product, this shall be stated in Annex 2 (Excel file) in the sheet "Ingoing substances" in column P.

In the case of RSPO “Mass Balance”, “Segregation” or “Identity Preserved” certification, the manufacturing company must verify their membership of the RSPO (in the case of first-time applications, to be applied for after the issuing of the contract). Normally, the manufacturing company will either be a “Supply Chain Associate” (purchasing less than 500 tonnes of palm oil products) or an “Ordinary Member” (purchasing more than 500 tonnes of palm oil products). A list of RSPO members is published here: https://www.rspo.org/members/all

The following shall be submitted to RAL gGmbH at the latest after the Blue Angel ecolabel has been used for the first 15 months and then annually: RSPO membership number and proofs of purchase (delivery notes/invoices) for the corresponding raw materials. The documents must state the RSPO certificate number for the raw material manufacturer. To verify that sufficient raw materials have been purchased, the annual production volume (for the formulation included in the application) must be stated in the sheet “Results-2” of the Excel table and the form of verification “Delivery notes/invoices (segregated or MB)” must be selected in column E. For raw materials produced from palm oil and palm kernel oil, the amount of the raw material required can then be calculated for a defined period. If multiple products are certified with the Blue Angel ecolabel, the calculations must be carried out for all products and the results (for identical raw materials) added together.

Alternatively, a supply chain audit can be carried out. This must be verified by submitting a corresponding certificate and the audit report. The audit must be carried out by a certification body accredited by the RSPO: https://www.rspo.org/certification/bodies/page/

A supply chain audit is obligatory:
- for ordinary RSPO members
- if using “MB claim transfer cross referencing”
- if products are produced by contract manufacturers who are not RSPO members themselves and are bound to the applicant via “outsourcing agreements”.

Verification via the Book & Claim system (RSPO credits) is not sufficient.
Appendix B  Fitness for use of laundry detergents

The performance test is equal to the „Revised EU Ecolabel protocol for testing laundry detergents“.