RECOMMENDATION

Conformity attestation of product hygiene suitability for drinking water¹

English translation – only the German document version is legally binding

2nd amendment:
important changes to 1st amendment have been shaded

1st amendment:
added annex 7
some adaptions in section “definition of terms”
some adaptions in the main part resulting from addition of annex 7

Notification number: 2018/479/D
1 Foreword

This recommendation serves as a basis for the attestation of conformity of the hygiene suitability of products intended to come into contact with drinking water.

Products that come into contact with drinking water must be both technically and hygienically suitable for this application. The technical requirements on products coming into contact with drinking water and their performance reliability are set out in the relevant product-specific standards, and are therefore not covered in this recommendation.

The hygiene requirements arise from the German Drinking Water Regulation (TrinkwV). This regulation states that the German Environment Agency shall adopt binding evaluation criteria documents further detailing the general requirements of the Drinking Water Regulation for materials coming into contact with drinking water. Due to formal statutory reasons, however, no procedural steps are defined in these evaluation criteria for the attestation of conformity of products with applicable requirements.

This recommendation describes a procedure to certify hygiene requirement conformity for the materials in the product that come into contact with drinking water. This will allow manufacturers to demonstrate that the products they sell meet the requirements of § 17 Section 2 and 3 TrinkwV.

Testing and evaluation of hygiene suitability have not yet been harmonised in Europe, and therefore fall under the scope of national regulations.

Due to the lack of harmonised requirements on hygiene suitability for drinking water, the available EN product standards are not harmonised standards and CE marking based on these standards is not possible. In Commission Decision 2002/359/EC on construction products, the European Commission stipulates that the 1+ system must be used to certify the conformity of hygiene suitability for a future CE labelling.

The attestation of conformity described in this recommendation meets this 1+ system. The procedure, for products to which a conversion factor $F_c \geq 0.5 \text{ d/dm}$ applies, or for metallic materials that are used for of product groups A and B, requires external monitoring by a certification body to obtain a conformity attestation.

2 Scope of application

This recommendation may be used for conformity attestation of hygiene suitability for drinking water for all products coming into contact with drinking water. The certificate of conformity may also be part of a combined certification of hygienic and technical suitability.
3 Normative references


German Environment Agency Evaluation Criteria Document for plastics and other organic materials in contact with drinking water (KTW-BWGL).


German Environment Agency Guideline for hygienic assessment of elastomers in contact with drinking water (Elastomer Guideline) and transitional regulation.

German Environment Agency Recommendation on the provisional hygienic assessment of products made from thermoplastic elastomers in contact with drinking water (TPE Transitional Recommendation)

German Environment Agency Evaluation Criteria Document for metallic materials in contact with drinking water.


DVGW standard W 270: Microbial enhancement on materials to come into contact with drinking water - Testing and assessment.
4 Definition of terms (sequence acc. to alphabetical order in German)

4.1 Starting material
A starting material is a substance used to produce an organic material (monomers, additives, adjuvants; not the same as the definition in Regulation (EU) No 10/2011) or cement-bound materials (e.g. admixtures, additives).

4.2 Component
A component is a part of a traded product or is produced as a piece of equipment for use in one or more products.

4.3 Factory-made product
A factory-made product is a product which is manufactured or applied in a factory as part of a defined production process.

4.4 Family of products or components (product line)
A family of products or components are products or components for which a common attestation of conformity may be issued.

4.5 Functional barrier
A functional barrier is a material layer that delays, but does not prevent, diffusion of the migrating substances.

4.6 Mixture (preparation)
Mixtures or preparations denote a mixture or solution composed of two or more substances (corresponds to definition in REACH regulation). These may contain both intermediate products or pre-products as defined in this recommendation document.

4.7 Trader / Distributor
A trader/distributor is any natural or legal person in the supply chain, other than the manufacturer, that provides a product on the market.

4.8 Manufacturer
A manufacturer is any natural or legal person that manufactures a product or component, and/or arranges their development or manufacturing, and that markets them under its own name or trademark.

4.9 Conformity attestation
A conformity attestation is a certificate from a certification body under the 1+ system (meeting Annex V to Regulation (EU) No 305/2011) attesting to compliance with the requirements on hygiene suitability.
4.10 Material (organic)
Organic material is organic matter from one or more starting materials with a precisely defined formulation and production process.

4.11 Multilayer product
A multilayer product is a product made up of multiple interconnected layers. These may be made from organic or inorganic materials.

4.12 Product
A product is a clearly identifiable manufactured part, with its final shape and surface, that a manufacturer or trader/distributor provides on the market and that is intended to come into contact with drinking water.

4.13 Product group
A product group encompasses various products or components with the same conversion factor or weighing factor (metallic materials) that are comparable in terms of their frequency of use in drinking water distribution and their surface/volume ratios and therefore may be treated the same in the evaluation of hygiene suitability for drinking water.

4.14 Products for on-site-application
A product for on-site application is a product intended for use on the worksite. Intermediate products like cold-curing epoxy resins are a class of on-site application products reacting to the final product only with just the final application step.

4.15 Test specimen
A test specimen is a product or specially produced sample that is tested to represent one or several product(s).

4.16 Testing body (authority)
A body contracted or approved by the certification body that must have a corresponding accreditation and that performs the required tests, calculations or external monitoring.

4.17 Formulation
A formulation is a list and description of the proportionate quantities of the starting materials used to produce an organic material.

4.18 Raw materials
Raw materials are substances or mixtures of substances used by the manufacturer.

4.19 Type testing
A type testing is the basis for every procedure to achieve product hygiene conformity attestation for drinking water. It shall be conducted at the start of conformity testing and repeated once every five years. The type testing shall examine all requirements on the product/component.
4.20 Total barrier

A total barrier is a barrier layer that prevents diffusion of the substance in question to the side in contact with drinking water.

4.21 Pre-product

A pre-product is a polymer which may contain other admixtures or additional components like e.g. glass fibres and which does not undergo further reactions. It serves in manufacturing of a product intended to be used in contact with drinking water (like granulates).

4.22 Material (inorganic, „Werkstoff“)

Inorganic material (also termed basic material) is inorganic matter (metallic, enamel, ceramic materials or cement-bound substances) made from one or more substances of precisely defined composition.

4.23 Certification body

A certification body is an independent body with the required competence (demonstrated by a corresponding accreditation as per DIN EN ISO/IEC17065) to assess the characteristics specified in this recommendation. A certification body may authorize notified subcontractors to have them performed defined tasks associated with a conformity assessment procedure, provided this has been settled within the accreditation framework of the certification body.

4.24 Assembled product

Assembled products are products that are made up of different components and that can be disassembled into these components.

4.25 Intermediate product

An intermediate product denotes a substance that is manufactured for and consumed in or used for chemical processing in order to be transformed into another substance (corresponds to definition in REACH regulation).

Other definitions apply from the German Environment Agency Evaluation Criteria Documents for organic materials, for enamels and ceramic materials as well as for metals.
5 Hygiene requirements on materials in contact with drinking water

5.1 General
Pursuant to § 17 Section 3 TrinkwV, the German Environment Agency shall determine the specific hygiene requirements on materials coming into contact with drinking water in the form of legally binding Evaluation Criteria Documents.

Thus far, the German Environment Agency has published Guidelines and Recommendations on materials in contact with drinking water. DVGW regulations also currently provide hygiene requirements (e.g. standard W 347). These Guidelines, Recommendations and Regulations shall be used until legally binding German Environment Agency Evaluation Criteria are determined (as has been accomplished already for metallic materials and enamel/ceramic materials).

The hygiene requirements have been drafted for individual materials and are provided below.

5.2 Metallic materials
Metallic materials must appear on the positive list of metallic materials hygienically suitable for drinking water, which is part of the Evaluation Criteria for metallic materials in contact with drinking water. In addition, the usage restrictions in the positive list shall also apply (product groups, use with specific drinking water types).

Note: For metallic coatings, the general drinking water hygiene suitability of which cannot be determined and for which therefore no entry in the positive list of metallic materials suitable for drinking water exists, no further testing standards are currently in force.

5.3 Organic materials
Organic materials like:
- plastics
- coatings
- lubricants
- elastomers
- thermoplastic elastomers
- silicones

shall meet the requirements of the Evaluation Criteria for organic materials, including the requirements on the results of the test as per DIN EN 16421:2015-05.

2 Until publication of corresponding annexes of the Evaluation Criteria Document on organic materials, the annexes of the relevant transitional provisions shall apply (Elastomer Guideline, TPE Transitional Recommendation, Transitional Recommendation for Silicones).
5.4 Enamels and ceramic materials
Enamels and ceramic materials shall meet the requirements of the Evaluation Criteria Document for enamels and ceramic materials.

5.5 Cementitious materials
Cementitious materials shall meet DVGW standard W 347 \(^3\) (denoted there as cement-bound materials). This standard sets out provisions on the permitted starting materials (positive list) and on a migration test procedure. Cementitious materials with organic content shall also meet DVGW standard W 270.

5.6 Multilayer materials
Multilayer materials shall be subject to the material-specific requirements for the materials in question. In cases of multiple layer material build-up, one layer may constitute a total barrier. If this is the case, the requirements shall only apply to the layers on the side towards the drinking water.

6 Principle of product conformity attestation

6.1 General
The evaluation and testing of hygiene requirement compliance for products in contact with drinking water shall be conducted under the 1+ system, analogously to Regulation (EU) No 305/2011 (see the product-specific testing requirements in Table 2 und Table 4). This entails conformity certification by a certification body. The duties of the certification body in this shall be:

- initial inspection of the manufacturing plant and of factory production control,
- monitoring of test specimen sampling and type testing of the product,
- continuous surveillance, assessment and evaluation of factory production control,
- audit-testing.

Due to difficulties with the realization of initial inspections, external monitoring and test specimen sampling that occurred in 2020 in conjunction with the emerging COVID-19-pandemia, it is possible to obtain conformity attestations based on type testing (simplified procedure) until 21\(^{st}\) March 2023, even in such cases where this recommendation stipulates certification based on the 1+ system.

To accomplish this for organic materials, test reports that had been created for the purpose of obtaining former KTW certificates may be used. As a prerequisite to proceed this way it is required that

- test reports must be issued after 21\(^{st}\) March 2013,
- the evaluation of formulations and of test results have to be performed on the basis of the KTW-BWGL or renewed where necessary.

\(^3\) After entry into force of the Evaluation Criteria Document according to § 17 TrinkwV, the Evaluation Criteria Document for cementitious materials and DIN EN 16421:2015-05 shall be met.
The continuous surveillance, assessment and evaluation of factory production control shall be conducted by the certification body or an inspection body contracted by the certifier as part of external monitoring at the manufacturer site.

The manufacturer shall provide the certification body with the information needed for evaluation (e.g. formulations or descriptions of the production process). Additionally, the manufacturer shall conduct:

- factory production control (FPC).

### 6.2 Conformity attestation of drinking water hygienic suitability

Product conformity attestation of drinking water hygienic suitability has to be performed in a material-specific manner. For products made up of different components (assembled products; see Figure 1), this means that as a general rule from this recommendation, every component produced from a single-layer or multilayer material shall have a separate certificate of hygiene suitability for drinking water. It is thus convenient to readily prepare conformity attestations for all individual components beforehand. For the attestation of conformity for an assembled product, it shall suffice if the corresponding certificates are available for the individual components (see the example in Annex 6).

Figure 1  Tasks for attestation of conformity for a combined product.
See explanatory notes and further details in Chapter 7 and Annexes 3, 5 and 6.
It is possible to issue a common attestation of conformity with hygiene requirements for a family of products or components (e.g. pipes, rubber sealing rings or injection-moulded components of various geometries). The preconditions for this are:

- the production process shall be comparable; and
- the material shall have the same composition and/or the material shall have the same formulation.

This attestation of conformity may also include components used for different products (product groups, product lines; see Figure 2). The conformity attestation must indicate the components or products to which it applies (see Annex 3).

The certification body may also combine the conformity attestations for assembled products if they are made from the same components, but with different geometries.

**Evaluation of formulations of mixtures, intermediate products and pre-products and, under certain circumstances, the conformity attestation for pre-products, may also be conducted independently from the final product, see annex 7.**

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**Figure 2** Attestation of conformity to a supplier for a group of products
6.3 Types of conformity attestation

6.3.1 Conformity attestation following the 1+ -system
Conformity may be certified under the 1+ -System for
- products and
- components
(see Table 2 and Table 4).

6.3.2 Conformity attestation based on type testing
The process for certifying the conformity of components with only a small portion of surface area coming into contact with drinking water – which therefore have less of an impact on drinking water quality at the tap – has been simplified (see Table 3).

Table 1 gives an overview of the simplified conformity attestation process.

<table>
<thead>
<tr>
<th>Material</th>
<th>Product groups</th>
<th>Conformity attestation process</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organic materials</td>
<td>Product groups with $F_c &lt; 0.5$ $d/dm$ according to the Evaluation Criteria Document</td>
<td>Type testing every five years</td>
</tr>
<tr>
<td>Enamels and ceramics</td>
<td>Product groups with $F_c &lt; 0.5$ $d/dm$ according to the Evaluation Criteria Document</td>
<td>Type testing every five years</td>
</tr>
<tr>
<td>Metals</td>
<td>Product group C according to Evaluation Criteria Document for metallic materials</td>
<td>Company certificate 2.2 for starting material (e. g. bar stock) and type testing of composition every five years</td>
</tr>
<tr>
<td></td>
<td>Product group D according to Evaluation Criteria Document for metallic materials</td>
<td>Type testing every five years</td>
</tr>
</tbody>
</table>

As with the 1+ system, simplified conformity attestation shall be performed by the certification body.

If an attestation of conformity is based solely on a type test, then the respective certificate must clearly indicate this fact.

External monitoring is not conducted in cases of a simplified attestation of conformity. The manufacturer shall bear full responsibility for quality assurance in the manufacture of the component. **With attestation of conformity of pre-products, however, monitoring of manufacture and sampling of test specimens by the certifying body is required (see annex 7).**

**Note:** The provisions for a simplified conformity attestation will be harmonised as part of the 4MS collaboration.
6.4 Test specimens for type testing and external monitoring

Evaluation of drinking water hygiene requirements shall be conducted on the product or component. In exceptional cases however (see Annex 5), representative specimens (e.g. test plates) may also be tested. These shall be produced under the supervision of the certification body or the contracted inspection body, or the certification body shall otherwise verify that the test specimens were produced under conditions comparable to those of the product (e.g. by fingerprint methods for different specimens/products).

For an attestation of conformity for a family of products or components (see 6.2), test specimens shall be taken that may be regarded as representative of the family. The specimens should be expected to exhibit substance release that are high for the product line. During annual monitoring, the certification body or contracted inspection body may collect different test specimens.

Sampling of test specimens by the certification body or contracted inspection body has to be performed at a place in the in-house logistics after in-house clearance for sale has been passed or in a centralised or distribution warehouse of the manufacturer.

7 Material-specific attestation of conformity of hygiene suitability for drinking water

7.1 Overview

Table 2 and Table 3 summarise the material-specific principles of conformity attestation for products or components. The manufacturer shall provide the certification body with the information as per Annex 1. In addition, Table 4 describes the principles for assembled products.

7.2 Metallic materials

The manufacturer shall notify the certification body of all metallic materials used for the product or component.

Hygiene requirement conformity shall be certified for metallic materials coming into direct contact with drinking water.

If metallic materials not appearing on the positive list for metallic materials of suitable hygiene for drinking water are coated, proof shall be submitted that the coating will be resistant for the expected service life of the product and that the coating fully covers the metal surface.

Type testing

The certification body shall verify that the materials coming into contact with drinking water as indicated by the manufacturer appear on the positive list of materials of suitable hygiene for drinking water, for the relevant area of application. In addition, the certification body or contracted inspection body shall take products and components during the initial inspection in
order to verify the composition of the individual basic materials by analysing them or having them analysed by an independent inspection body.

**Factory Production Control (FPC)**
The certification body, together with the manufacturer establish a procedure by which the hygiene suitability of the finished products or components for drinking water is to be verified in the factory production control. An installed, functioning QM system may be used for this purpose.

For factory production control, applicable terms would be:

- acceptance test certificate 3.1 as per DIN EN 10204 by the material supplier; or
- regular inspection of material composition in the finished product or component in the factory.

For foundries, the composition of the cast component shall be checked in the factory.

**External monitoring**
The certification body shall monitor the factory production control regularly (see Table 2). This shall include verifying use of the materials indicated by the manufacturer.

Moreover, the certification body or contracted inspection body shall regularly (see Table 2) take products and components (see 6.4) from the factory and verify the composition of the individual materials by analysing them or having them analysed by an independent inspection body.

### 7.3 Metallic coatings

**Note:** Binding testing requirements are not currently available for metal coatings. The manufacturer shall be responsible for certifying hygiene suitability for drinking water for metal coatings that cannot be included in the positive list of the Evaluation Criteria Document for metallic materials. An attestation of conformity from a certification body must clearly indicate that release of substances from coatings coming into contact with drinking water was not tested. Certification bodies are advised not to issue any certificates of conformity for the hygiene suitability of products with these kinds of non-listed coatings for drinking water.

### 7.4 Organic material

#### 7.4.1 General

The formulation shall be disclosed to the certification body. In general, suppliers are to be involved in this, according to the supply chain. For full disclosure of the finished product formulation, the manufacturer is obliged to provide the certification body with the required information on the supply chain for its product, including details on all suppliers. Any change in the supply chain shall be reported to the certification body immediately, so it can rule out any resulting changes in the finished product formulation.
For substances adding less than 0.02% (w/w, with respect to the finished components), the certification body may waive further disclosure of the formulation if the materials adding less than 0.02% (w/w) do not exceed a total value of 0.1% (w/w). The form according to Annex 2 shall be used to submit the formulation. For multilayer products, the formulation shall be submitted separately for each layer.

For products with a total barrier, the layers on the side not in contact with drinking water shall not be assessed. The term ‘total barrier’ applies e.g. to a continuous aluminium layer with a thickness of at least 9 µm or a continuous glass layer.

In cases of composite materials containing a functional barrier, e.g. a layer of an ethylene vinyl alcohol copolymer, all layers shall be assessed.

The certification body shall ensure that the test specimens used for testing are made from the indicated starting materials (e.g. by using a fingerprint method).

### 7.4.2 Factory-made products manufactured from organic materials

#### Type testing

The type testing shall preferably be conducted on test specimens (see 6.4) collected during the initial inspection of the production site. The certification body verifies the conformity of the indicated formulation with the positive list from the Evaluation Criteria Document for organic materials. This shall also include verifying the technological functions of the formulation components and restrictions on the use of starting materials (e.g. purity of the starting materials used, maximum application quantity, residual content). Based on the formulation submitted by the manufacturer and/or the suppliers, the certification body shall specify the test parameters according to the Evaluation Criteria Document for organic materials. The certification body and/or the contracted inspection body shall conduct complete testing of the test specimens collected during the initial inspection based on the Evaluation Criteria Document for organic materials.

#### Factory production control

The certification body, together with the manufacturer establish a procedure by which the hygiene suitability of the finished products or components for drinking water is to be verified in the factory production control. An installed, functioning QM system may be used for this purpose.

For pipes (F: ≥ 5 d/dm), in addition to verification of the starting materials and production procedure at defined intervals closely coordinated with the certification body, the manufacturer shall also conduct migration tests. Migration water testing for the odour threshold value or analysis of a representative substance may be suitable methods for factory production control. As an alternative to migration testing, the manufacturer may apply other methods, such as fingerprint methods, if this is coordinated with the certification body.

#### External monitoring

The certification body shall regularly monitor the factory production control (see Table 2). This includes examination of documents to verify use of the formulation components indicated by
the manufacturer. This shall also encompass verification that upstream products exhibit the required purity and that the specified dosing is actually applied.

The certification body or the contracted inspection body shall regularly take representative samples from all production sites and test them for the basic requirements of the Evaluation Criteria Document for organic materials. In addition, parameters whose levels \( c_{\text{tap}} \) are close to the reference concentration in the type test shall be retested.

The certification body shall take representative samples from the factory at regular intervals (see Table 2) and conduct complete type testing or have this conducted by a contracted inspection body.

7.4.3 Products for on-site application made from organic materials (e.g. coatings)

The attestation of conformity corresponding to this Recommendation is limited to the general suitability of the products: it does not cover certification of individual users.

On-site application has a significant influence over the quality of finished products. Therefore, additional measures shall be required taking into account the formulation details and the defined processing instructions and/or technical bulletins of the manufacturer, to ensure proper on-site application.

A simplified conformity attestations excludes products for on-site application because the specific resulting finished products may also yield applications of extensive areal coverage or high surface/volume ratios. For this reason there is also no distinction between products of risk groups P1 and P2 in this case. For products of risk groups P3 and P4, however, the simplified procedure may be applied since here the wetted surface portion is very small. The respective limitations of use must be observed and shall be notified in the certificate.

It is advisable for the manufacturer to deliver the product for on-site application along with a notice of mandatory adherence to the permitted applications and processing instructions and an indication that failure to comply shall invalidate the existing conformity attestation for the product, i.e. the hygiene conformity of the finished products manufactured by the end user would no longer be guaranteed for drinking water. It is not the manufacturer, but rather the end user that bears responsibility for this. An additional guarantee may be provided if the user has a certification for proper application of the product for on-site application. Because this is difficult to audit, it requires additional certification provisions.

Type testing

The type test shall preferably be conducted with representative test specimens produced during the initial inspection of the production site (e.g. manufacturer of coatings for on-site application) under the supervision of the certification body. The certification body verifies the conformity of the indicated formulation with the positive list from the Evaluation Criteria Document for organic materials. This shall also include verifying the technological functions of the formulation components and restrictions on the use of starting materials (e.g. purity of the starting materials used, maximum application quantity, residual content). In addition, the
conformity shall also be verified for the application instructions/processing conditions and the production of the test specimens. Based on the formulation submitted by the manufacturer and/or the suppliers, the certification body shall specify the test parameters according to the Evaluation Criteria Document for organic materials. The certification body and/or the contracted inspection body shall conduct complete testing of the test specimens collected during the initial inspection based on the Evaluation Criteria Document for organic materials.

**Factory production control**
The certification body, together with the manufacturer and located at the production site establish a procedure by which the hygiene suitability of the manufactured products for drinking water is to be verified by means of representative test specimens in the factory production control. An installed, functioning QM system may be used for this purpose.

In addition to verification of the starting materials and production process, the manufacturer shall conduct regular migration tests. Migration water testing for the odour threshold value or analysis of a representative substance may be suitable methods for factory production control. As an alternative to migration testing, the manufacturer may apply other methods, if this is coordinated with the certification body.

**External monitoring**
The certification body shall regularly monitor factory production control in the production site (see Table 2). This includes examination of documents to verify use of the formulation components indicated by the manufacturer, and it also encompasses verification that upstream products exhibit the required purity and that the specified dosing is actually applied.

The certification body shall have representative samples regularly produced by all production sites and shall test these for the basic requirements of the Evaluation Criteria Document for organic materials. In addition, parameters whose levels ($c_{tap}$) are close to the reference concentration in the type test shall be retested.

The certification body shall have representative samples produced by the factory at regular intervals (see Table 2) and shall conduct complete type testing or have this conducted by a contracted inspection body.

**7.5 Enamels**
The certification body may only issue an attestation of conformity jointly for enamellers and enamel frit manufacturers.

The manufacturer of the enamel frit and enameller shall provide the certification body with the information corresponding to Annex 1.
**Type testing**
The certification body shall conduct a complete test and assessment based on the Evaluation Criteria Document for enamels and ceramic materials. To achieve this, the certification body shall have test plates enamelled at the enamelling factory during the initial inspection, under supervision, and shall subsequently test these.

**Factory production control**
The certification body, together with the enamel frit manufacturer establish a procedure by which the hygiene suitability of the finished products or components for drinking water is to be verified in the factory production control. An installed, functioning QM system may be used for this purpose.

The enamel frit manufacturer shall regularly check the composition of the enamel frit or shall have this checked by a third party.

The enameller shall conduct an incoming goods inspection on the supplied enamel frits. Moreover, it shall check the component enamelling process to ensure consistent quality of the enamelled components. An installed, functioning QM system may be used for this purpose.

**External monitoring**
The certification body shall regularly monitor factory production control by the frit manufacturer and enameller (see Table 2).

At regular intervals (see Table 2), the certification body shall have test plates enamelled, under supervision, and shall conduct complete testing as per the Evaluation Criteria Document for enamels and ceramic materials.

**7.6 Ceramic materials**

**Type testing**
The certification body shall conduct a complete test and assessment based on the Evaluation Criteria Document for enamels and ceramic materials. To achieve this, the certification body shall use test plates that it collected during the initial factory inspection.

**Factory production control**
The certification body, together with the manufacturer shall establish a procedure by which the hygiene suitability for drinking water of the finished products or components is to be verified in the factory production control. An installed, functioning QM system may be used for this purpose.

The manufacturer shall regularly check the composition of the products or shall have this checked by a third party.

**External monitoring**
The certification body shall monitor the factory production control regularly (see Table 2).
At regular intervals (see Table 2), the certification body shall collect components from the factory and conduct complete testing as per the Evaluation Criteria Document for enamels and ceramic materials.

7.7 Cementitious materials

The attestation of conformity procedure for cementitious materials will be supplemented as soon as the Evaluation Criteria Document for cementitious materials or a corresponding European harmonized regulatory document has been defined.
<table>
<thead>
<tr>
<th>Material</th>
<th>Type testing</th>
<th>Self-monitoring (by manufacturer)</th>
<th>External monitoring (by certification body) every year</th>
<th>every 5 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Factory-made products</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Metallic materials (Product groups A and B according to the Evaluation Criteria Document for metallic materials)(^4)</td>
<td>The certification body shall collect test specimens during the initial factory inspection, to test the basic material composition</td>
<td>Analysis of material composition during: Incoming goods inspection or production; or acceptance test certificate 3.1 for the starting material</td>
<td>Verification of self-monitoring, collection of test specimens and analysis of material composition</td>
<td>-</td>
</tr>
<tr>
<td>Organic materials (plastics, coatings, elastomers, thermoplastic elastomers, silicones) ((F_c \geq 0.5 , \text{d/dm}))(^5)</td>
<td>The certification body shall collect test specimens during the initial factory inspection for complete testing according to the Evaluation Criteria Document for organic materials, including under DIN EN 16421</td>
<td>Incoming goods check, testing for conformity of raw materials;</td>
<td>Verification of the raw materials used, collection of test specimens and testing for basic requirements and selected parameters under the additional requirements of the Evaluation Criteria Document for organic materials, verification of self-monitoring</td>
<td>Collection of test specimens and complete testing according to the Evaluation Criteria Document for organic materials and DIN EN 16421</td>
</tr>
</tbody>
</table>

\(^4\) For product groups C and D, an attestation of conformity under the 1+ system is not prescribed (see 6.3)

\(^5\) For products with an \(F_c < 0.5 \, \text{d/dm}\), an attestation of conformity under the 1+ system is not prescribed (see 6.3). This usually also applies to lubricants, which according to the Evaluation Criteria Document for organic materials (KTW-BWGL) are attributed with an \(F_c\) of 0.2 or less
<table>
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<tr>
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<th>every 5 years</th>
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<td><strong>Factory-made products</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>(Contd.: Organic materials)</strong></td>
<td></td>
<td>For pipes ($F_c \geq 5$ d/dm): migration testing on the product and identification of the odour threshold value, or a suitable alternative parameter</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Enamels ($F_c \geq 0.5$ d/dm)</strong></td>
<td>The certification body shall have test specimens (typically enamelled plates) produced in the enamelling system during the initial factory inspection, under its supervision, for complete testing as per the Enamel/Ceramics Evaluation Criteria Document</td>
<td>Analysis of the composition of the enamel frit and enamelling</td>
<td>Verification of self-monitoring</td>
<td>Collection of test specimens and complete testing as per the Enamel/Ceramics Evaluation Criteria Document</td>
</tr>
<tr>
<td><strong>Ceramic materials ($F_c \geq 0.5$ d/dm)</strong></td>
<td>The certification body shall collect test specimens during the initial inspection for complete testing as per the Enamel/Ceramics Evaluation Criteria Document</td>
<td>Regular inspection of the composition of finished products</td>
<td>Verification of self-monitoring</td>
<td>Collection of test specimens and complete testing as per the Enamel/Ceramics Evaluation Criteria Document</td>
</tr>
<tr>
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</tr>
<tr>
<td>----------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------</td>
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<td>--------------------------------------------------------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Products for on-site application</td>
<td>The certification body shall collect test specimens produced during the initial inspection of the starting material manufacturer (e.g. coating plant), under supervision, for complete testing as per the Evaluation Criteria Document for organic materials, including under DIN EN 16421</td>
<td>Incoming goods inspection, testing of conformity of raw materials, odour testing on representative test specimens or suitable alternative processes</td>
<td>Verification of the raw materials used, collection of test specimens produced under supervision and testing for basic requirements and selected parameters under the additional requirements as per the Evaluation Criteria Document, verification of self-monitoring</td>
<td>Collection of test specimens produced under supervision and complete testing as per the Evaluation Criteria Document and DIN EN 16421</td>
</tr>
<tr>
<td>Material</td>
<td>Type testing</td>
<td>Self-monitoring (by manufacturer)</td>
<td>External monitoring (by certification body)</td>
<td></td>
</tr>
<tr>
<td>----------</td>
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<td>---------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Organic materials ($F_c &lt; 0.5 , \text{d/dm}$)</td>
<td>The attestation of conformity is based on the type test, which shall be repeated once every five years using a suitable test specimen</td>
<td>Incoming goods inspection, testing of conformity of the raw materials and finished products</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Organic materials (e.g. repair systems) $F_c &lt; 0.04 , \text{d/dm}$ (risk groups P3 and P4 acc. to KTW-BWGL)</td>
<td>The attestation of conformity is based on the type test, which shall be repeated once every five years using a suitable test specimen</td>
<td>Incoming goods inspection, testing of conformity of the raw materials, odour testing of representative test specimen or other appropriate method</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Enamels and ceramic materials ($F_c &lt; 0.5 , \text{d/dm}$)</td>
<td>The attestation of conformity is based on the type test, which shall be repeated once every five years using a suitable test specimen</td>
<td>Incoming goods inspection, regular inspection of the composition of finished products</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Metals</td>
<td>The attestation of conformity is based on the type test, which shall be repeated once every five years using a suitable test specimen</td>
<td>Company certificate 2.2 for the starting material</td>
<td>N/A</td>
<td></td>
</tr>
<tr>
<td>Assembled product</td>
<td>Type testing</td>
<td>Self-monitoring (by manufacturer)</td>
<td>External monitoring (by certification body) every year</td>
<td></td>
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<td>------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>None</td>
<td>Verification that the attestations of conformity are up-to-date for the components and the corresponding incoming goods</td>
<td>Verification of the attestation of conformity and of self-monitoring</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Annex 1  Information for preparation of an attestation of conformity

The manufacturer shall provide the certification body with the information below on its product/component.

a) Precise description of the product / component

b) Manufacturing site(s)

c) List of all materials coming in contact with drinking water
   - for metallic materials: standardised and/or precise material designation
     - for organic materials: complete formulation disclosure by all parties throughout the supply chain, with separate submission for each layer in cases of multilayer materials
   - for enamels:
     o Enamel frit manufacturer: enamel frit designation, enamel frit composition with tolerances for the individual enamel frit components and production site
     o Enameller: enamel production site, description of the procedure and indication of the enamelled products
   - for ceramics: composition, description of the production process
   - for cementitious materials: complete formulation disclosure by all parties throughout the supply chain; for mixing plants only for their own production process

d) Total product and/or component surface area coming into contact with drinking water and the relative proportion of surface area of the respective materials

e) Area of application: cold-, warm- or hot water, product group according to Evaluation Criteria Document with conversion factor $F_C$

f) For products for on-site application: application instructions
Annex 2  Declaration of formulation for organic materials

The manufacturer and/or its supplier shall submit the following tables completed in full.

<table>
<thead>
<tr>
<th>No.</th>
<th>Raw material / trade name</th>
<th>Chemical description</th>
<th>CAS number</th>
<th>Function of the raw material</th>
<th>Percentage by weight (in % w/w)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

In the table, the manufacturer shall enter all formulation components, including additional formulation components of the preparation, such as solvents and impurities (see 7.4.1). A current data safety sheet for the starting materials or preparation can generally provide information on starting material purity and on the other substances contained in the formulation. If the manufacturer does not possess complete formulation details, the supplier shall submit the respective information.

For products with several layers, the composition of each layer must be disclosed for assessment of the product formulation.
Annex 3  Contents of an attestation of conformity for factory-made products

Attestation of conformity
by certification body ’...(Title)...’
regarding the hygiene suitability for drinking water of

Product: Name of the product or component / trade name
Material/polymer
Product group
(Certification body registration number, if applicable)
Manufacturer: Company (Name)
(Adress)

☐ Based on a type test (without external monitoring; simplified procedure)
☐ Under system 1+ with external monitoring of the following production sites: (Adresses)

Reference / Attestation:
We hereby certify that the product / component indicated above
• based on the certification programme ’...(Title)...’ of the certification body
  (Name, Adress),
• the submitted product information, including the formulation details
• and evaluation report No (report number)

meets the requirements of the Evaluation Criteria Document / Guideline Document of the
German Environment Agency:
‘...(Title)...’
for the temperature range (degrees Celsius) and is in compliance with the above document.
(Optional: the requirements on product groups with lower conversion factors (F_c) are also met).

or

This attestation of conformity also applies to the following products / components:
Product / component: (Designation / Name)
Product / component: (Designation / Name) etc.

The evaluation reports with numbers (report number) are part of this attestation of conformity.

This attestation of conformity is valid from the date of issue and its validity will endure as long
as the preconditions specified below under "Remarks“ are fulfilled.

Place, date       Name
Certification Body Manager

6 Any applicable transitional provisions must be indicated here.
Remarks:

The attestation of conformity is issued under the precondition that the starting materials used to manufacture the products and/or their composition and/or components, including their manufacturers and supply chain, have been disclosed in full and the product does not contain any further substances. This document shall be invalid in cases of changes to the composition of the product or the processing conditions that have not been agreed upon with the certification body, failure to meet the conformity requirements or termination of the 1+ system monitoring programme by the certification body and manufacturer.

The findings of our tests and the evaluations apply for the test objects examined and the provisions of the law applicable at the time of testing. Without our express written approval, it is only permitted to publish or reproduce this document in full and unedited.
Annex 4 Contents of an attestation of conformity for products for on-site application

Attestation of conformity
by certification body ’...(Title)…’
regarding the hygiene suitability for drinking water of

Product: Name of the product for on-site application / trade name
Material/polymer
Product group
(Certification body registration number, if applicable)
Manufacturer: Company (Name)
(Adress)

☐ for risk groups with \( F_c \geq 0.04 \text{ d/dm} \) (corresponding to P1 and P2 according to KTW evaluation criteria) under system 1+ with external monitoring of the following production sites: ...(Adress)...

☐ for risk groups with \( F_c < 0.04 \text{ d/dm} \) (corresponding to P3 and P4 according to KTW evaluation criteria) based on type testing (without external monitoring; simplified procedure)

Reference / Attestation:
We hereby certify that the product for on-site application indicated above
- based on the certification programme ‘...(Title)…’ of the certification body (Name, Adress),
- the submitted product information, including the formulation details, the permitted applications and processing instructions,
- and evaluation report No (report number)

in cases of proper use, is generally suitable for the production of on-site finished products that meet the requirements of the Evaluation Criteria Document / Guideline Document of the German Environment Agency7:

‘...(Title)…’
for the temperature range (degrees Celsius) and are in compliance with the above document.

The evaluation reports with numbers (report number) are part of this attestation of conformity.

This attestation of conformity is valid from the date of issue and its validity will endure as long as the preconditions specified below under ”Remarks“ are fulfilled.

Place, date Name
Certification Body Manager

---

7 Any applicable transitional provisions must be indicated here.
Remarks:

The attestation of conformity is issued under the precondition that the starting materials and/or their composition and formulation used to manufacture the product for on-site application, including their manufacturers and supply chain, have been disclosed in full and the product does not contain any further substances.

This document shall be invalid in cases of

- changes to the composition of the product or the processing conditions that have not been agreed upon with the certification body or
- failure to meet the conformity requirements or
- termination of the 1+ system monitoring programme by the certification body and manufacturer

This attestation of conformity does not cover use of the product for on-site applications. The user must observe the manufacturer’s application instructions and should hold a separate certification for proper application of this product.

The findings of our tests and the evaluations apply for the product samples examined, the test samples made from these according to the application instructions and the provisions of the law applicable at the time of testing. Without our express written approval, it is only permitted to publish or reproduce this document in full and unedited.
### Annex 5  Test specimens for type testing and external monitoring

<table>
<thead>
<tr>
<th>Material</th>
<th>Test specimens for testing as per DIN EN 16421</th>
<th>Test specimens for other tests</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Factory-made products</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Metallic materials</td>
<td>N/A</td>
<td>Product/component</td>
</tr>
<tr>
<td>Plastics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pipes</td>
<td></td>
<td>Product/component</td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td>Product/component or representative test specimen</td>
</tr>
<tr>
<td>Silicones</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hoses</td>
<td></td>
<td>Product/component</td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td>Product/component or representative test specimen</td>
</tr>
<tr>
<td>Organic coatings</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Product/component or representative test plates</td>
<td></td>
</tr>
<tr>
<td>Silicone hoses</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Elastomers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hoses</td>
<td></td>
<td>Product/component</td>
</tr>
<tr>
<td>Other</td>
<td></td>
<td>Product/component or representative test specimen</td>
</tr>
<tr>
<td>Lubricants</td>
<td>N/A</td>
<td>Applied on plates (where possible)</td>
</tr>
<tr>
<td>Enamels</td>
<td>N/A</td>
<td>Product/component or representative test specimen</td>
</tr>
<tr>
<td>Ceramics</td>
<td>N/A</td>
<td>Product/component</td>
</tr>
<tr>
<td><strong>Products for on-site-application</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Organic materials</td>
<td>Representative test plates</td>
<td>Representative test specimen</td>
</tr>
</tbody>
</table>

The further specifications in Table 2 of the Evaluation Criteria Document for organic materials in contact with drinking water shall apply.
Annex 6  Examples of attestation of conformity under the 1+ -system

Example 1  Finished product manufacturer has injection-moulded components produced from a defined material (attestation of conformity under contract from the finished product manufacturer)

Scenario:
For various injection moulders, a manufacturer of assembled products has components made from a defined material with a prescribed formulation with the same process parameters. The finished product manufacturer incorporates the components produced by the injection moulders into the assembled products.

Conformity attestation for the injection-moulded component:
The certification body may combine the attestation of conformity for components of comparable geometry and the same composition, even if produced by different injection moulders or production sites for a finished product manufacturer. The preconditions on this are that the same raw materials must be used and the production process must be comparable. In addition, at the end, the attestation of conformity must clearly indicate the components to which it applies.

The certification body, along with the manufacturers involved, shall determine the test specimen expected to exhibit the maximum possible substance release for the group of components on the attestation of conformity. Test specimen collection for the type test may be performed at the finished product manufacturer. External monitoring of factory production control shall also be carried out at the finished product manufacturer. The precondition here that the finished product manufacturer must have a clear quality control procedure at incoming goods and must also have the injection moulder document the process parameters.

Conformity attestation of the injected-moulded components shall be certified under contract from the finished product manufacturer.

Conformity attestation for the assembled product:
In the factory production control, the finished product manufacturer must ensure that valid attestations of conformity are available for all components. In addition, it shall also use the documents to ensure that the goods supplied are in accordance with the components for which the attestation of conformity was issued. The certification body shall conduct external monitoring based on documents and verify the validity of the attestations of conformity for the individual components as well as the factory production control of the finished product manufacturer.
Figure 3  Diagram of conformity attestation under system 1+ for a finished product manufacturer (example 1)
Example 2  Manufacturer of elastomer components supplies multiple clients who incorporate the elastomer components into assembled products (conformity attestation under contract from the component manufacturer)

Scenario:
The manufacturer of assembled products produces one component itself and purchases the rest. One of the purchased components is an elastomer component. The elastomer component manufacturer also produces other elastomer components of different dimensions but with the same process, process parameters and raw materials. It supplies these to other clients as well.

Conformity attestation for the elastomer component:
The attestation of conformity may be combined for all elastomer components made with the same process, process parameters and raw materials. The attestation of conformity must clearly indicate the components to which it applies. The certification body, along with the elastomer component manufacturer, shall determine the test specimens expected to exhibit the maximum possible substance release for the group of components on the attestation of conformity.

Sampling for the type test and for the regular product tests as well as external monitoring on the factory production control shall be performed at the elastomer component manufacturer.

Conformity attestation for the component manufactured in-house by the finished product manufacturer:
Test specimen collection for the type test and for the regular product tests as well as external monitoring on the factory production control shall be performed at the manufacturer. The attestation of conformity may be combined for multiple components here as well if they are manufactured with the same raw materials and the same process (see above).

Conformity attestation for the assembled product:
In the factory production control, the finished product manufacturer must ensure that valid attestations of conformity are available for all components. In addition, it shall also use the documents to ensure that the goods supplied are in accordance with the components for which the attestation of conformity was issued.

As part of external monitoring for the component manufactured in-house, the certification body may also verify the attestations of conformity for other components and the factory production control related to the combined product.
Figure 4  Diagram of conformity attestation under system 1+ for a component manufacturer (example 2)
Example 3  Manufacturer of enamelled products purchases enamel frit to make its products (certificate of conformity for frit manufacturer and enameller)

Scenario:
The frit manufacturer produces special frit for an enameller. This enameller uses the frit to make various enamelled products.

Conformity attestation for the enamelled component:
The attestation of conformity may only be issued for the frit manufacturer along with the enameller.

Test specimen collection for the type test and for the regular product tests as well as external monitoring on the factory production control shall be performed at the enameller. For this purpose, the enameller must produce test plates under the supervision of the certification body.

In addition, external monitoring shall be conducted on the factory production control at the frit manufacturer. For the factory production control, the frit manufacturer shall regularly test the frit composition or have it tested by a third party.

The attestations of conformity may be combined for enamelled components produced with the same frit and process. The attestation of conformity shall clearly indicate the components to which it applies.
A7.1 Introduction
Evaluating formulations of organic materials that come into contact with drinking water is of crucial importance to the conformity attestation of end products. This can be very time-consuming as in most cases different sub-formulations for mixtures (e.g. glass fibre sizing agents), intermediate products (e.g. coating hardeners) and pre-products (e.g. plastic granulates) have to be obtained and evaluated.

The safety data sheet for the individual products does not usually indicate the various ingredients. In addition, end product manufacturers in most cases have no knowledge of the formulation of purchased mixtures, intermediate products and pre-products. For this reason, the various upstream suppliers must disclose the respective formulations to the end product certification body.

If the certification body can evaluate the formulation of a mixture, an intermediate product or pre-product, then this is one way of simplifying the evaluation of end products. This is especially true if mixtures, intermediate products and pre-products are used for different end products. In this case, the sensitive formulation information has to be fully disclosed to only one certification body.

However, a prerequisite for this is that the manufacturer of the mixture, intermediate product or pre-product has to allow his certification body to disclose the relevant constituents to the certification body of the final product. The relevant constituents are those substances that have to be checked in the final product’s migration test.

Utilization for a conformity attestation of a pre-product
In addition to evaluating a formulation, a conformity attestation conforming to the KTW evaluation criteria (including microbiological requirements) by a certification body is possible for intermediate products (e.g. plastic granulates) if a component manufactured from the pre-product has been tested. This conformity attestation is fully sufficient for components of risk group P2 according to the KTW evaluation criteria.

For components of risk group P1, this conformity attestation is sufficient for formulation evaluation. For P1 products that are not pipes it is also sufficient to confirm conformity with the microbiological requirements.
The procedure described here encompassing a conformity attestation after having done a formulation evaluation is no option for intermediate products of risk group P2, because intermediate products – in contrast to pre-products – chemically react to their final state only at a late stage on their designated use.

A7.2 Evaluation of a formulation

Procedure

Evaluation of formulations shall be carried out by an accredited certification body. This is the only way to ensure that an end product that contains the mixture, intermediate product or pre-product as a source ingredient obtains a conformity attestation as per this recommendation.

Within the scope of the evaluation, the certification body shall check the requirements according to 5.2 of the KTW evaluation criteria and confirm the conformity. The provisions of 7.4.1 of this recommendation apply.

In addition to the formulation, the manufacturer must provide the certification body with the following information:

- Trade name of the product;
- Exact description of the application;
- Maximum input quantity of the mixture, intermediate product or pre-product in the end product;
- Type of end product according to the product groups in Table 7 of the KTW evaluation criteria.

First, the certification body checks the formulation’s information for plausibility. It also checks purity requirements and other specifications if these are required for listed source materials.

The certification body stipulates in the evaluation which substances shall be determined as additional and formulation-specific individual substance requirements in the end product migration test.

The number of substances to be determined in the migration test might be reduced by the fact that the input of substances falls below the formulation cut-off limit when the intended use in the end product is taken into account. The certification body will record the relevant consideration performed.

They can also be reduced by estimating the potential mass transfers by taking into account the intended application of the end product. The certification body will record the relevant estimation performed.
Conformity attestation of formulations

The certification body can attest the conformity of the composition of a mixture, intermediate product or pre-product with the relevant positive lists and their restrictions as a result of the formulation evaluation. This attestation must clearly indicate the following:

- Trade name of the evaluated mixture, intermediate product or pre-product and the manufacturer;
- Description/specification of the mixture, intermediate product or pre-product;
- Maximum input quantity of the mixture, intermediate product or pre-product in the end product;
- Type of end product according to the product groups in Table 7 of the KTW evaluation criteria;
- Specification of the positive list (version/date) used for the evaluation.

If an additional migration evaluation has been carried out assuming complete transfer, the following data is also required:

- absolute quantity of the source material used for a product in contact with one litre of water.

For migration modelling of a product or component that contains the source material further information on its use is required:

- Structure of the product (monolayer or multilayer);
- Thickness of the individual layers;
- Polymer in the individual layers including density;
- Surface/volume ratio considered;
- Conversion factor used.

The following note is needed in all cases:

- **Additional substances must be tested in the end product migration test. The attesting body communicates them to the end product certification body, provided the relevant requirements for confidentiality are met.**

The certification body shall keep a list of those mixtures, intermediate products and pre-products with trade names and the corresponding manufacturers for which it has issued a conformity attestation for the formulation.

The certification body must ensure contractually that it will be immediately informed of any changes in the formulation and will take these into account for the attestation.

In addition to the above listed attestation information, the certification body shall establish a procedure with their client as to how to communicate the substances to be determined in the end product migration test to the certification bodies of end products. If necessary, appropriate confidentiality declarations shall be made by the parties concerned. The passing on of knowledge regarding the formulation components to end product manufacturers is not envisaged and is not necessary.
A7.3 Evaluation of a pre-product

Procedure

The certification body can carry out a complete test of the requirements of the KTW evaluation criteria for pre-products (especially plastic granules that are not involved in chemical reactions). A prerequisite is the formulation evaluation (see above). Additional migration tests shall be carried out on test specimens manufactured according to the manufacturer’s specifications. The test specimens for the migration test shall be as close to actual manufactured components for end products as possible and shall be manufactured and sampled under external monitoring.

The microbiological test as per DIN EN 16421 can be carried out on representative test specimens according to 6.4 of the KTW evaluation criteria for various components (except pipes). The evaluation of the test report according to DIN EN 16421 with the requirement according to 5.6 of the KTW evaluation criteria is part of the evaluation of pre-products.

Conformity attestation of pre-products

The certification body can attest the conformity of components manufactured from pre-products to the requirements of the KTW evaluation criteria including microbiological requirements as a result of the formulation evaluation. This attestation must clearly indicate the following:

- Trade name of the evaluated pre-product and the manufacturer;
- Details of use and reference to the manufacturer’s instructions for use;
- Type of end product according to the product groups of Table 7 of the KTW evaluation criteria used for the evaluation;
- Indication of the positive list (version/date) used for the evaluation;
- Surface-to-volume ratio considered;
- Conversion factor used;
- A note that this attestation only applies to components of product groups P2 to P4 according to Table 2 of the KTW evaluation criteria.

The certification body shall keep a list of the pre-products with trade names and the relevant manufacturers for which it has issued a conformity attestation.

The certification body must ensure contractually that it will be immediately informed of any changes in the formulation and will take these into account in the attestation.