DRAFT EAST AFRICAN STANDARD

Synthetic detergent powder — Specification — Part 2: Machine wash

EAST AFRICAN COMMUNITY
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Foreword

Development of the East African Standards has been necessitated by the need for harmonizing requirements governing quality of products and services in the East African Community. It is envisaged that through harmonized standardization, trade barriers that are encountered when goods and services are exchanged within the Community will be removed.

The Community has established an East African Standards Committee (EASC) mandated to develop and issue East African Standards (EAS) and other deliverables. The Committee is composed of representatives of the National Standards Bodies in Partner States, together with the representatives from the public and private sector organizations in the community.

East African Standards are developed through Technical Committees that are representative of key stakeholders including government, academia, consumer groups, private sector and other interested parties. Draft East African Standards are circulated to stakeholders through the National Standards Bodies in the Partner States. The comments received are discussed and incorporated before finalization of standards, in accordance with the Principles and procedures for development of East African Standards.

East African Standards and other deliverables are subject to review, to keep pace with technological advances. Users of the East African Standards are therefore expected to ensure that they always have the latest versions of the standards they are implementing.

The committee responsible for this document is Technical Committee EASC/TC 074, Surface active agents

Attention is drawn to the possibility that some of the elements of this document may be subject of patent rights. EAC shall not be held responsible for identifying any or all such patent rights.

This second edition cancels and replaces the first edition (EAS 127-2:2013), which has been technically revised. EAS 127 consists of the following parts, under the general title Synthetic detergent powders — Specification

— Part 1: Household hand use
— Part 2: Machine wash
Introduction

There are two main groups of detergents, namely, the soaps and synthetic detergents. In order to guide the production of synthetic detergents of well-defined quality and also to safeguard consumer interests, it has been felt desirable to formulate this standard.

Synthetic detergents are of three types: anionic, cationic and non-ionic. Anionic synthetic detergents of the alkyl aryl type, such as dodecylbenzene sulfonic acid, are now produced in fairly large quantities and hence priority has been given for the standardization of this material.

It has been discovered that equally good cleaning can be achieved at relatively low detergent levels. With the advance of technology, other equally effective, environmentally safe and cheaper builders can be used instead of Sodium Tripolyphosphate (STPP) which is currently being used. Phosphates have been known to bring eutrophication in water bodies and companies world over are doing away with them and encouraging use of alternative builders such as zeolites and carbonates, and therefore for this reason the requirement for levels of STPP are being progressively phased out in the standard with a view of eliminating them in the near future.

The standard is more composition based than performance based and for that reason performance measures for cleaning efficiency and detergency tests have been included. This edition emphasizes on the biodegradability of the surfactants used to manufacture detergent powders and a method for determining the same has been developed.
Synthetic detergent powder — Specification — Part 2: Machine wash

1 Scope
This Draft East African Standard specifies the requirements, sampling and test methods for synthetic detergents powder for machine wash. It does not cover hand wash powders and industrial detergent powders.

2 Normative references
The following referenced documents are indispensable for the application of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

EAS 814, Determination of biodegradability of surfactants — Test method
ISO 672, Soaps — Determination of moisture and volatile matter content — Oven method
ISO 673, Soaps — Determination of content of ethanol-insoluble matter
ISO 862, Surface active agents — Vocabulary
ISO 2870, Surface active agents — Detergents — Determination of anionic-active matter hydrolysable and non-hydrolysable under acid conditions
ISO 2871-2, Surface active agents — Detergents — Determination of cationic-active matter content — Part 2: Cationic-active matter of low molecular mass (between 200 and 500)
ISO 4313, Washing powders — Determination of total phosphorous (v) oxide content — Quinoline phosphomolybdate gravimetric method
ISO 4316, Surface active agents — Determination of pH of aqueous solution — Potentiometric method
ISO 4317, Surface active agents and detergents — Determination of water content — Karl Fischer methods

3 Terms and definitions
For the purposes of this standard, terms and definitions given in ISO 862 and the following shall apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:
— ISO Online browsing platform: available at http://www.iso.org/obp
3.1 detergent
product specially formulated for cleaning through the process of detergency

3.2 active ingredient
organic surface-active material present in the detergent

3.3 builder
material added to the detergent formulation that enhances or maintains the cleaning efficiency of the surfactant, principally by inactivating water hardness either by sequestration, precipitation or ion exchange

3.4 defect
non-conformance with a requirement of the standard

3.5 defective
detergent or package that fails in one or more respects to comply with the relevant requirements of this standard

3.6 package
unit in which the detergent is stored, which is labelled, and distributed for individual sale

3.7 manufacturer
person or organization actually engaged in or being principally responsible for manufacturing or producing the laundry detergent

3.8 sequestering agent
material that suppresses the hardness of water and improves the effectiveness of surface active agents by removing metal ions or molecules from the water

3.9 batch
material from a single mix or, in the case of a continuous production process, the material from a single day’s production

3.10 lot
quantity of a detergent bearing the same batch identification, from one manufacturer, and submitted at any one time for inspection and testing

4 Requirements

4.1 General requirements

4.1.1 Synthetic detergent powder shall:

a) be homogenous, free flowing powder, free from visible impurities, abrasives and organic solvents and shall be readily soluble in water;

b) not be irritating to the normal skin and shall not contain any ingredients in a quantity that is toxic to human beings; and

c) not give an objectionable odour.
4.1.2 The active ingredients used include salt of alkylaryl sulphonie acid, soap and non-ionics. The formulation may contain one or more of the builders or additives given in Annex A.

4.1.3 The active ingredient used shall be biodegradable when tested according to EAS 814.

4.1.4 A solution of the synthetic detergent powder in hot water (at 60 °C ± 2 °C) shall not have objectionable odour.

4.1.5 If perfume is added, it shall not change the product fragrance nor develop an objectionable odour during storage at ambient temperature.

4.1.6 If synthetic detergent powder is coloured, the colouring shall be uniform and shall not change appreciably during storage.

4.2 Specific requirements

4.2.1 Synthetic detergent powder shall also comply with the specific requirements given in Table 1, when tested in accordance with the method prescribed therein.

<table>
<thead>
<tr>
<th>S/No</th>
<th>Characteristic</th>
<th>Requirement</th>
<th>Method of test</th>
</tr>
</thead>
<tbody>
<tr>
<td>i)</td>
<td>Moisture and volatile matter content at 105 °C, % m/m, max.</td>
<td>13</td>
<td>ISO 672, ISO 4317</td>
</tr>
<tr>
<td>ii)</td>
<td>Active ingredient, % m/m, min.</td>
<td>10</td>
<td>ISO 2870, ISO 2871-1, ISO 2871-2</td>
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<td>iii)</td>
<td>Matter insoluble in alcohol, % m/m, max.</td>
<td>90</td>
<td>ISO 673</td>
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<tr>
<td>iv)</td>
<td>Phosphate (expressed as sodium tripoly-phosphate), % by mass of matter insoluble in alcohol, max.</td>
<td>15</td>
<td>ISO 4313</td>
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<td>v)</td>
<td>pH of 1 % solution (m/v), at 25 °C ± 2 °C</td>
<td>9 - 11</td>
<td>ISO 4316</td>
</tr>
<tr>
<td>vi)</td>
<td>Non-detergent organic matter, % m/m, max.</td>
<td>1.0</td>
<td>Annex B</td>
</tr>
<tr>
<td>vii)</td>
<td>Matter insoluble in water, % m/m, max.</td>
<td>5.0</td>
<td>Annex C</td>
</tr>
<tr>
<td></td>
<td>phosphate based synthetic detergent powders</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Non-phosphate based synthetic detergent powders</td>
<td>15.0</td>
<td></td>
</tr>
<tr>
<td>viii)</td>
<td>Available O₂ as sodium perborate, % m/m, max</td>
<td>10</td>
<td>Annex D</td>
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</tbody>
</table>

4.2.2 Storage properties

When stored or transported under normal conditions in its original container, the powder shall not cake into lumps.

5 Packaging

Synthetic detergent powder shall be packaged in containers that are strong enough to withstand normal handling, storage and transportation and that will prevent leakage and contamination of the product.
6 Labelling

Each container each bulk package shall be in prominent, legibly and indelibly labelled either in English, Kiswahili or French or combination or any other language as agreed between the manufacturer and supplier with the following information:

a) name of the product as "Synthetic detergent powder";
b) indicate that the product is for machine wash;
c) manufacturer’s name and physical address;

NOTE The name, physical address of the distributor/supplier and trade mark may be added as required.
d) batch or lot number;
e) list of ingredients
f) net content;
g) country of origin;
h) instructions for use;
i) date of manufacture and best before date; and
j) indicate mode of disposal of containers/packages

7 Sampling

Sampling shall be done in accordance with Annex F.
Annex A
(informative)

List of suggested builders and additives

The following is a list of suggested builders and additives:

a) trisodium phosphate;
b) sodium carbonate;
c) sodium sulfate;
d) tetraboric acid(pyrophosphoric acid);
e) sodium tripolyphosphate;
f) sodium hexametaphosphate;
g) sodium carboxy methyl cellulose;
h) sodium silicate;
i) optical brighteners;
j) lather boosters;
k) hydrototes;
l) perfum;
m) preservatives;
n) chelating agents (sequestering agents);
o) colours;
p) perborates;
q) enzymes;
r) bactericides;
s) zeolites;
t) common salt;
u) urea;
v) magnesium sulphate;

w) polymers; and

x) any other internationally accepted builder cleared by the national standards body in the respective country.

Annex B
(normative)

Determination of non-detergent organic matter

B.1 Principle

The term non-detergent organic matter includes hydrocarbons, fatty alcohols and perfumes. Using petroleum ether and under the conditions prescribed, non-detergent organic matter only is extracted leaving any alkylolamide present in the material.

B.2 Apparatus

B.2.1 Evaporating basin

B.2.2 Separating funnels, 1 000-ml capacity

B.2.3 Wide mouthed flat-bottomed flask, 200-ml capacity

B.2.4 Buchner flask, 500-ml-capacity fitted with a sintered glass filter funnel (porosity 4)

B.2.5 Vacuum pump

B.3 Reagents

B.3.1 Ethyl alcohol, 50 %, 70 %, 90 % and 96 % (by volume)

B.3.2 Petroleum ether, boiling range 40 °C - 60 °C non-volatile residue at 80 °C maximum 0.001 %

B.3.3 Acetone, non-volatile residue at 80 °C maximum 0.001 %

B.4 Procedure

B.4.1 Weigh accurately about 5 g of the material in a 150-ml squat beaker. Extract with 50 ml of hot 90 % ethanol by heating on the steam bath for about 2 min stirring and breaking up any hard lumps with a glass rod flattened at the end.

B.4.2 Allow the solid matter to settle and decant the hot alcoholic solution through a sintered glass filter funnel (porosity 4) fitted to a 500-ml Buchner flask to which suction is applied. Repeat the extraction in a similar manner with five further consecutive 30-ml quantities of boiling 90 % ethanol. Pass each extract in turn through the filter into the flask.

B.4.3 Transfer quantitatively all the combined filtrate from the Buchner flask to a 1-L separating funnel and rinse the flask four times with 40-ml quantities of distilled water, transferring each wash in turn to the separating funnel. Add 100 ml of petroleum ether, swirl gently to ensure adequate mixing and allow the two phases to separate. Run off the aqueous alcoholic layer into a second separating funnel, and extract with 75 ml of petroleum ether. Repeat the extraction of the aqueous alcoholic phase in the third separating funnel with a further 75 ml of petroleum ether. Combine the three ether extracts in the first separating funnel. Rinse each of the two empty funnels with a few millilitres petroleum ether and add the rinsing to the combined ether extracts.
B.4.4 Wash the combined ether extracts and rinsing (see B.4.3) with four successive 50-ml portions of 70 % ethyl alcohol, shaking and removing the alcoholic phase each time. Transfer the ether layer in stages to a pre-weighed flask and recover the solvent. Add 10-ml of acetone and evaporate off the solvent using a stream of nitrogen. Rotate the flask on a steam bath during the operation. Cool the flask to about 60 °C to 65 °C, gently blow out the last traces of solvent with a current of dry air, cool in a desiccator and weigh.

B.5 Calculation

The non-detergent organic matter is expressed as follows:

Non-detergent organic matter (on dry basis), % m/m = \( 100 \frac{m_2 - m_1}{m} \times \frac{100}{(100 - \text{moisture})} \)

where

- \( m_1 \) is the mass, in grams, of empty flask
- \( m_2 \) is the mass, in grams, of the flask with residue
- \( m \) is the mass, in grams, of material taken for the test.
Annex C
(normative)

Determination of matter insoluble in water

C.1 Procedure

C.1.1 Starting with a fresh portion of the material, weigh accurately about 5 g of the material into a beaker, and digest with 50 ml of ethyl alcohol by heating on a steam bath for about 2 min. Stir and break up any hard lump with a glass rod flattened at one end.

C.1.2 Allow the solid matter to settle and decant the hot alcoholic solution through a dried and pre-weighed sintered glass filter funnel fitted to a Buchner flask, to which suction is applied.

C.1.3 Repeat the alcoholic digestion in a similar manner with five further consecutive 30 ml portions of boiling ethyl alcohol. Filter each extract in turn through the same sintered glass funnel and, finally, wash the residue several times with hot ethyl alcohol to remove all the alcohol soluble matter.

C.1.4 Even after digestion with five 30 ml portions of boiling ethyl alcohol, the alcohol insoluble portion may sometimes be found to be sticky. In that case treat it further with more boiling ethyl alcohol until it is free from active matter and the alcohol insoluble portion is no longer sticky.

C.1.5 Dry the sintered glass funnel with the residue in an air-oven at a temperature of 105 °C ± 2 °C for 1 h. Cool in a desiccator and weigh.

C.1.6 Repeat the process in C.3.6 for 30 min until a constant weight is obtained.

C.1.2 Even after digestion with five 30-ml portions of boiling ethyl alcohol, the alcohol insoluble portion may sometimes be found to be sticky. In that case, treat it further with more boiling ethyl alcohol until it is free from active matter and the alcohol insoluble portion is no longer sticky. Do not dry or weigh the matter insoluble in alcohol. After filtering and washing the residue thoroughly with hot ethyl alcohol, change the receiver, extract the residue with successive portions of distilled water at about 60 °C, and wash the residue several times to remove all the water solubles. Dry the sintered glass funnel with the residue in an air-oven at a temperature of 105 °C ± 2 °C until a constant mass is obtained.

C.2 Calculation

The matter insoluble in water is expressed as follows:

\[
\text{Matter insoluble in water, } \% \text{ by mass (on dry basis), } \% \text{ m/m} = 100 \times \frac{m_2 - m_1}{m} \times \frac{100}{(100 - \text{moisture})}
\]

where

- \( m_1 \) is the mass, in grams, of empty sintered glass filter funnel
- \( m_2 \) is the mass, in grams, of sintered glass filter funnel with residue
- \( m \) is the mass, in grams, of material taken for the test.
Annex D
(normative)

Test method for determination of perborate

D.1 Method

Weigh 1.5 g - 2 g of sample, dissolve in water and transfer into a 250 ml volumetric flask and make up to volume with water. Take an aliquot of 25 ml for titration and add 20 ml of 0.1 M sulphuric acid. Titrte immediately with standard 0.1 N potassium permanganate until a faint pink colour persists.

D.2 Calculation

The available oxygen shall be expressed as follows:

\[
\% \text{ available oxygen} = \frac{a \times 8 \times N}{W}
\]

where

- \( a \) is the titre;
- \( N \) is the normality of potassium permanganate;
- \( W \) is the weight of material taken.

NOTE When fresh contains approximately 95 % corresponding to 9.9 % available oxygen.
Annex E
(normative)

Test method for foam

E.1 Foaming

Foaming capacity of detergents is an important property. In some instance, a high foam is required and in others no foam at all or a very low foam is desirable. It is necessary to standardize the following factors:

a) concentration of active water;
b) type of water to be used;
c) temperature of solution; and
d) period of aging.

E.2 Apparatus

E.2.1 500-ml glass measuring cylinder
E.2.2 250-ml glass separating funnel
E.2.3 50-ml glass pipette
E.2.4 250-ml glass cylinder

E.3 Preparation of standard hard water (50 ppm as calcium carbonate)

E.3.1 Reagents

E.3.1.1 Analar calcium chloride dihydrate, CaCl₂·2H₂O
E.3.1.2 Analar magnesium sulphate heptahydrate, MgSO₄·7H₂O
E.3.1.3 Deionized water

E.3.2 Method

Dissolve 0.0440 g Analar calcium chloride dihydrate and 0.0492 g Analar magnesium sulphate heptahydrate in deionized water. Make up the volume to 1 L with deionized water.

E.3.3 Preparation of test solution

Prepare a solution containing 6 g of the detergent per litre in standard hard water.

Warm the solution to the temperature of 60 ºC and allow to ‘age’ at this temperature for 10 min.

E.4 Procedure

Draw 50 ml of the solution into a 50-ml pipette.
Hold the pipette against the upper inside portion of the 500-ml glass measuring cylinder and allow the solution to run into the cylinder, moving the pipette so that it revolves around the complete circumference of the cylinder so that the solution wets the walls of the cylinder. The solution remains in the cylinder.

Into the 250-ml glass cylinder, pour in 200 ml of the solution and pour this into the separating funnel. Clamp into position the separating funnel so that its spout points to the centre of the 500-ml cylinder and is level with the rim of the 500-ml measuring cylinder (see Figure E.1). The positioning is such that the stream of liquid will impinge only on the surface of the liquid.

Open the stopcock fully to allow the solution to fall onto the liquid

The moment the flow ceases, read the volume of the foam ($V_1$).

![Image of measurement of foam]

**Figure E.1 — Measurement of foam**

After 5 min read the foam volume again ($V_2$).

- foaming power = $V_1 - 250$ ml (the amount of liquid in cylinder); and
- foaming stability = $- V_2 - 250$ ml;

replace A and B with $V_1$ and $V_2$

For example, foaming power = $350 - 250$

= 100

For comparison of efficacy in hard and soft water, the whole method can be repeated using deionized water instead of the standard hard water.
Annex F  
(normative)

Sampling procedure for detergent powder

F.1 Requirements

F.1.1 In drawing preparing, storing and handling samples, the precautions in F.1.2 – F.1.7 shall be observed.

F.1.2 Samples shall not be taken from places exposed to damp air, dust or soot.

F.1.3 The sampling instruments shall be clean and dry when used.

F.1.4 The samples, the material being sampled, the sampling instruments and the containers for samples shall be protected from adventitious contamination.

F.1.5 The samples shall be placed in clean and dry glass containers. The sample containers shall be of such a size that they are almost completely filled by the sample.

F.1.6 Each container shall be sealed airtight after filling, and marked with full details of sampling which include, date of sampling, batch or code number, name of manufacturer, and other important particulars of the consignment.

F.1.7 The samples shall be stored in such a manner that the temperature of the material does not vary unduly from the normal temperature, and that they are protected from light.

F.2 Scale of sampling

F.2.1 In a single consignment, all the packages containing the product of the same type and form, and drawn from the same batch of manufacture, shall constitute a lot. If the consignment consists of packages containing products of different types and forms, then the packages containing products of the same type, form and batch of manufacture shall be grouped together, and such group shall constitute a separate lot.

F.2.2 For ascertaining the conformity of the lot to the requirements prescribed in this standard, tests shall be carried out on each lot separately. The number (n) of packages to be selected for drawing the samples shall depend upon the size (N) of the lot and shall be in accordance with Table F.1.

<table>
<thead>
<tr>
<th>No. of packages in the lot (N)</th>
<th>No. of packages to be selected (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4 - 15</td>
<td>3</td>
</tr>
<tr>
<td>16 - 40</td>
<td>4</td>
</tr>
<tr>
<td>41 - 65</td>
<td>5</td>
</tr>
<tr>
<td>66 - 110</td>
<td>7</td>
</tr>
<tr>
<td>111 and above</td>
<td>10</td>
</tr>
</tbody>
</table>

NOTE When the size of the lot is three packages or less, the number of containers to be selected and the criteria for judging the conformity of the lot to the specifications should be as agreed on between the purchaser/inspector and the supplier.

F.2.3 The packages shall be selected at random and to ensure randomness of selection, a random number table shall be used. In case such tables are not available, the procedure given below may be adopted.
Starting from any package, count all the packages in one order as 1, 2, 3, ..., up to \( r \) and so on, where \( r \) is the integral part of \( N/n \), \( N \) being the lot size and \( n \) the number of packages to be selected. Every \( r \)th package thus counted shall be withdrawn to give a sample for the purposes of test.

**F.3 Preparation of gross samples, test sample and reference sample**

**F.3.1 Gross sample**

From each one of the packages selected as in F.2, draw at random one or more containers. The material in the containers so chosen shall be nearly thrice the quantity required for purpose of test as indicated in F.4.

The powder from the containers selected shall be disintegrated, if aggregated, and mixed thoroughly to give the gross sample for the package.

**F.3.2 Test sample**

**F.3.2.1** Segregate carefully the gross samples of powders. From the gross representing each form of synthetic detergent take a small but equal quantity of material and mix thoroughly into a composite sample which should be of a size sufficient to carry out triplicate testing for all the characteristics specified under F.4. The composite samples representing each form and type of synthetic detergent shall be divided into three equal parts, one for the purchaser/inspector, another for the supplier, and the third for the referee.

**F.3.2.2** The remaining portion of the material in each of the gross samples shall be divided into three equal pans, each forming an individual sample. One set of individual samples, representing the \( n \) selected packages shall be for the purchaser/inspector, another for the supplier, and the third for the referee.

**F.3.2.3** All the composite and individual samples shall be transferred to separate containers. These containers shall then be sealed airtight with stoppers, and labelled with full particulars of identification given in F.1.6.

**F.3.3 Reference samples**

**F.3.3.1** The reference samples shall consist of a composite sample and a set of individual samples. All the containers shall bear the seals of both the purchaser/inspector and the supplier, and shall be kept at a place agreed to between the two parties.

**F.3.3.2** Reference samples shall be used in case of any dispute between the purchaser/inspector and the supplier.

**F.4 Number of tests**

**F.4.1** Tests for the determination of active ingredient shall be performed on each of the individual samples.

**F.4.2** Tests for the determination of other requirements specified in Table 1 shall be conducted on the composite sample.

**F.5 Criteria for conformity**

**F.5.1 For individual samples**

**F.5.1.1** For the characteristic, which has been determined on the individual sample, the mean (\( X \)) and the range (\( R \)) of test results shall be calculated as follows:

\[
\text{Mean}(X) = \frac{\text{Sum of test results}}{\text{Number of test results}}
\]
Range \((R)\) is the difference between the maximum and the minimum value of test results.

**F.5.1.2** If the specification limit for the characteristic is given as a minimum, the value of the expression \((\bar{X} - KR)\) shall be calculated from the relevant test results (see F.5.1.5). If the value so obtained is greater than or equal to the minimum limit, the lot shall be declared as conforming to the requirement for the characteristic.

**F.5.1.3** If the specification limit for the characteristic is given as a maximum, the value of the expression \((\bar{X} + KR)\) shall be calculated from the relevant test results (see F.5.1.5). If the value so obtained is less than or equal to the maximum limit, the lot shall be declared as conforming to the requirement for the characteristic.

**F.5.1.4** If the characteristic has two-sided specification limits, then the values of the expression \((\bar{X} \pm KR)\) shall be calculated from the relevant test results (see F.5.1.5). If the value so obtained lies between the two specification limits, the lot shall be declared as conforming to the requirement for the characteristic.

**F.5.1.5** The value of the factor \(K\) referred to in F.5.1.2 - F.5.1.4 shall be chosen in accordance with Table F.2, depending upon the acceptable quality level, that is, the percentage, of non-conforming packages that may be tolerated reasonably.

<table>
<thead>
<tr>
<th>Acceptable quality level</th>
<th>Value of 'K'</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not more than 3.0 % defectives</td>
<td>0.4</td>
</tr>
<tr>
<td>Not more than 1.5 % defectives</td>
<td>0.5</td>
</tr>
<tr>
<td>Not more than 0.5 % defectives</td>
<td>0.6</td>
</tr>
</tbody>
</table>

**F.5.2** For composite sample

For declaring the conformity of the lot to the requirements of all the remaining characteristics determined on the composite sample, the test results for each one of the characteristics shall satisfy the relevant requirement given in Table 1 of this standard.
Bibliography