# INTERNATIONAL STANDARD

**ISO** 2322

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# Styrene-butadiene rubber (SBR) — Emulsion- and solution-polymerized types — Evaluation procedures

Caoutchouc butadiène-styrène (SBR) — Types polymérisés en émulsion et en solution — Méthode d'évaluation

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ISO 2322:2009

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### **Foreword**

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

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

ISO 2322 was prepared by Technical Committee ISO/TC 45, Rubber and rubber products, Subcommittee SC 3, Raw materials (including latex) for use in the rubber industry.

This fifth edition cancels and replaces the fourth edition (ISO 2322:1996), which has been technically revised.

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## Styrene-butadiene rubber (SBR) — Emulsion- and solution-polymerized types — Evaluation procedures

WARNING — Persons using this International Standard should be familiar with normal laboratory practice. This standard does not purport to address all of the safety problems, if any, associated with its use. It is the responsibility of the user to establish appropriate safety and health practices and to ensure compliance with any national regulatory conditions.

### 1 Scope

This International Standard specifies:

- physical and chemical tests on raw rubbers;
- standard materials, standard test formulations, equipment and processing methods for evaluating the vulcanization characteristics of emulsion- and solution-polymerized styrene-butadiene rubbers (SBR), including oil-extended rubbers TANDARD PREVIEW

It applies to those rubbers listed in Table 1 which are normally used in vulcanized form.

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Rubber	<del>9db8-c72b36a6c31c/iso-2322-2009</del> <b>Styrene</b>				
(oil-extended or	- · · · · · · · · · · · · · · · · · · ·	Total content	Block content % (by mass)		
non-oil-extended)	Type of copolymer	% (by mass)			
Series A					
Emulsion SBR	Random	≤ 50	0		
Solution SBR	Random	≤ 50	0		
Solution SBR	Partial block	≤ 50	≤ 30		
Series B					
Emulsion SBR	Random	> 50	0		
Solution SBR	Random	> 50	0		
Solution SBR	Partial block	≤ 50	> 30		

#### 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.

ISO 37, Rubber, vulcanized or thermoplastic — Determination of tensile stress-strain properties

ISO 247:2006, Rubber — Determination of ash

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ISO 248, Rubber, raw — Determination of volatile-matter content

ISO 289-1, Rubber, unvulcanized — Determinations using a shearing-disc viscometer — Part 1: Determination of Mooney viscosity

ISO 1795, Rubber, raw natural and raw synthetic — Sampling and further preparative procedures

ISO 2393, Rubber test mixes — Preparation, mixing and vulcanization — Equipment and procedures

ISO 3417, Rubber — Measurement of vulcanization characteristics with the oscillating disc curemeter

ISO 6502, Rubber — Guide to the use of curemeters

ISO 23529, Rubber — General procedures for preparing and conditioning test pieces for physical test methods

### 3 Sampling and further preparative procedures

- 3.1 Take a laboratory sample of approximately 1,5 kg by the method described in ISO 1795.
- **3.2** Prepare test samples in accordance with ISO 1795.

## 4 Physical and chemical tests on raw rubber 1 PREVIEW

### 4.1 Mooney viscosity

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Determine the Mooney viscosity in accordance with ISO 289-1 on a test sample prepared in accordance with the preferred method of ISO 1795 (unmilled test sample). Record the result as ML(1+/4) at 100 °C.

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If ML(1 + 4) at 100  $^{\circ}$ C exceeds 100 Mooney units, the small rotor may be used and the result reported as MS(1 + 4) at 100  $^{\circ}$ C.

Alternatively, the Mooney viscosity may be determined on a test sample prepared by the mill massing procedure of ISO 1795. However, this method gives poorer reproducibility and the results may be different.

#### 4.2 Volatile matter

Determine the volatile-matter content by the hot-mill method or by the oven method as specified in ISO 248.

#### 4.3 Ash

Determine the ash in accordance with method A or method B of ISO 247:2006.

## 5 Preparation of the test mixes

#### 5.1 Standard test formulations

The standard test formulations are given in Table 2.

The materials shall be national or international standard reference materials.

If no standard reference material is available, the materials to be used shall be agreed by the parties concerned.

Table 2 — Test formulations

Material	Parts by mass			
Material	Series A	Series B		
Styrene-butadiene rubber (SBR) (including oil in oil-extended SBR)	100,00	_		
Standard SBR 1500 <sup>a</sup>	_	65,00		
Series B SBR	_	35,00		
Sulfur	1,75	1,75		
Stearic acid	1,00	1,00		
Industry reference black <sup>b</sup>	50,00	35,00		
Zinc oxide	3,00	3,00		
TBBS <sup>c</sup>	1,00	1,00		
Total	156,75	141,75		

<sup>&</sup>lt;sup>a</sup> The previously used SBR 1500 EST is no longer available. It is therefore necessary to use another commercially available SBR 1500. The type shall be agreed between the interested parties.

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## 5.2 Alternative formulations for oil-extended types 21)

ASTM D 3185 specifies the test formulations given in Table 3 for evaluation of general-purpose, oil-extended SBR, depending on the oil content of the rubber. These test formulations may be used as alternatives to the test formulations given in Table 2. 9db8-c72b36a6c31e/iso-2322-2009

Table 3 — Alternative test formulations for oil-extended types

	Quantity					
	parts by mass					
Formulation number	1B	2B	3B	4B	5B	6B
Parts of oil	25	37,5	50	62,5	75	γa
Oil-extended rubber	125,00	137,50	150,00	162,50	175,00	100 + Y
Zinc oxide	3,00	3,00	3,00	3,00	3,00	3,00
Sulfur	1,75	1,75	1,75	1,75	1,75	1,75
Stearic acid	1,00	1,00	1,00	1,00	1,00	1,00
Industry reference black b	62,50	68,75	75,00	81,25	87,50	(100 + Y)/2
TBBS <sup>c</sup>	1,25	1,38	1,50	1,63	1,75	(100 + Y)/100
Total	194,50	213,38	232,25	251,13	270,00	

Y = parts of oil, by mass, per 100 parts of base polymer in the oil-extended rubber.

b Use the current industrial reference black. Dry the material for 1 h at 125 °C ± 3 °C and store in a tightly closed container.

<sup>&</sup>lt;sup>c</sup> N-tert-butylbenzothiazole-2-sulfenamide. This is supplied in powder form having an initial insoluble-matter content, determined in accordance with ISO 11235, of less than 0,3 %. The material shall be stored at room temperature in a closed container and the insoluble matter checked every 6 months. If this is found to exceed 0,75 %, the material shall be discarded or recrystallized.

b Use the current industrial reference black. Dry the material for 1 h at 125  $^{\circ}$ C  $\pm$  3  $^{\circ}$ C and store in a tightly closed container.

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#### 5.3 Procedure

### 5.3.1 Equipment and procedure

The equipment and procedure for the preparation, mixing and vulcanization shall be in accordance with ISO 2393.

Three alternative mixing procedures are specified:

- method A: mill mixing;
- method B: single-stage mixing using a laboratory internal mixer;
- method C: two-stage mixing using a laboratory internal mixer for initial mixing and a mill for final mixing.

#### 5.3.2 Method A — Mill-mixing procedure

The standard laboratory mill batch mass, in grams, shall be based on four times the formulation mass (i.e.  $4 \times 156,75$  g = 627 g or  $4 \times 141,75$  g = 567 g). Maintain the surface temperature of the rolls at 50 °C  $\pm$  5 °C. Maintain a good rolling bank at the nip of the rolls during mixing. If this is not obtained with the nip settings specified hereunder, small adjustments to the mill openings may be necessary.

	iTeh STANDA	Series A		Se	ries B
	(standar	Duration (1 (min)e h	Cumulative <b>al</b> time (min)	Duration (min)	Cumulative time (min)
a)	Homogenize series B rubbers with the mill opening set at 1,1 mm at a temperature $^{1SO~2}$ 100 °C $\pm$ 5 °C.	322:2009 andards/sist/774 :31e/iso-2322-		<sub>d27-</sub> 1,0	1,0
b)	Band the rubber with the mill opening set at 1,1 mm and make 3/4 cuts every 30 s from alternate sides.	7,0	7,0	_	_
	After banding the SBR 1500, add the rubber [homogenized as in step a) above] and make 3/4 cuts from both sides every 30 s.	_	_	8,0	9,0
c)	Add the sulfur slowly and evenly across the rubber.	2,0	9,0	2,0	11,0
d)	Add the stearic acid. Make one 3/4 cut from each side.	2,0	11,0	2,0	13,0
e)	Add the carbon black evenly across the mill at a uniform rate. When about half the black has been incorporated, open the mill to 1,4 mm and make one 3/4 cut from each side. Then add the remainder of the carbon black. Be certain to add any black that has dropped into the mill pan. When all the black has been incorporated, open the mill to 1,8 mm and make one 3/4 cut from each side.	12,0	23,0	12,0	25,0

f)	Add the zinc oxide and the TBBS with the mill opening still at 1,8 mm.	3,0	26,0	3,0	28,0
g)	Make three 3/4 cuts from each side.	2,0	28,0	2,0	30,0
h)	Cut the batch from the mill. Set the mill opening to 0,8 mm and pass the rolled batch endwise through the rolls six times.	2,0	30,0	2,0	32,0

- i) Sheet the batch to an approximate thickness of 6 mm by opening the mill and passing the stock through the mill four times, folding it back on itself each time. Determine the mass of the batch (see ISO 2393). If the mass of the batch differs from the theoretical value by more than + 0,5 % or 1,5 %, discard the batch and remix. Remove sufficient material for curemeter testing.
- j) Sheet the batch to approximately 2,2 mm for preparing test sheets or to the appropriate thickness for preparing ISO ring test pieces in accordance with ISO 37.
- k) Condition the batch for 2 h to 24 h after mixing and prior to vulcanizing, if possible at standard temperature and humidity as defined in ISO 23529.

#### 5.3.3 Method B — Single-stage mixing using a laboratory internal mixer

For a laboratory internal mixer having a nominal capacity of 65 cm<sup>3</sup> to about 2 000 cm<sup>3</sup>, the batch mass shall be equal to the nominal mixer capacity, in cubic centimetres, multiplied by the density of the compound. For each batch mixed, the laboratory internal mixer conditions shall be the same during the preparation of a series of identical mixes. At the beginning of each series of test mixes, a machine-conditioning batch shall be mixed using the same formulation as the mixes under test. The laboratory internal mixer shall be allowed to cool down to 60 °C between the end of one test batch and the start of the next. The temperature control conditions shall not be altered during the mixing of a series of test batches.

The mixing technique shall be such as to obtain a good dispersion of all the ingredients.

The temperature of the batch discharged on completion of mixing shall not exceed 120 °C. If necessary, adjust the batch mass or the mixer head starting temperature so that this condition is met.

NOTE 1 The mixing conditions given in Table A.6 for various sizes of laboratory internal mixer might be helpful.

Compounding materials other than rubber, carbon black and oil can be added to laboratory internal mixer batches more precisely and with greater ease if they are previously blended together in the proportions required by the formulation. Such blends may be made using a mortar and pestle, by mixing for 10 min in a biconical blender with the intensifier bar turning, or by mixing in another type of blender for five 3 s periods, scraping the inside of the blender to dislodge material stuck to the sides after each 3 s mix. A Waring blender has been found suitable for this method. Caution: if mixed longer than 3 s, the stearic acid may melt and prevent good dispersion.

NOTE 2 A general mixing procedure for the laboratory internal mixer is as follows:

		Duration (min)	Cumulative time (min)
a)	Load the rubber, lower the ram and allow the rubber to be masticated.	1,0	1,0
b)	Raise the ram and add the pre-blended zinc oxide, sulfur, stearic acid and TBBS, taking care to avoid any loss. Then add the carbon black, sweep the orifice and lower the ram.	1,0	2,0
c)	Allow the batch to mix.	7,0	9,0

d) Turn off the motor, raise the ram, remove the mixing chamber and discharge the batch. Record the maximum batch temperature.

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After discharging the mixed batch, pass it through a mill set at 50  $^{\circ}$ C  $\pm$  5  $^{\circ}$ C once at a 0,5 mm mill opening and then twice at a 3,0 mm mill opening.

Determine the mass of the batch and record it. If it differs from the theoretical value by more than +0.5% or -1.5%, discard the batch and remix.

Prepare a test piece for determining the vulcanization characteristics in accordance with ISO 3417 or ISO 6502, if required. Condition the test piece for 2 h to 24 h, if possible at a standard temperature and humidity as defined in ISO 23529, before testing.

If required, sheet the batch to approximately 2,2 mm for preparing test sheets or to the appropriate thickness for preparing ring test pieces in accordance with ISO 37. To obtain the effects of mill direction, pass the folded batch four times between mill rolls set at the appropriate opening and a temperature of 50 °C  $\pm$  5 °C. Allow the sheet to cool on a flat, dry surface.

Condition the batch for 2 h to 24 h after mixing and prior to vulcanizing, if possible at standard temperature and humidity as defined in ISO 23529.

## 5.3.4 Method C — Two-stage mixing using a laboratory internal mixer for initial mixing and a mill for final mixing

#### 5.3.4.1 Stage 1 — Initial mixing procedure

The mixing technique shall be such as to obtain a good dispersion of all the ingredients.

The temperature of the batch discharged on completion of mixing shall be between 150 °C and 170 °C. If necessary, adjust the batch mass or the mixer head starting temperature so that this condition is met.

NOTE 1 The following mixing conditions have been found to be suitable for a laboratory internal mixer with a nominal capacity of 1 170 cm $^3$   $\pm$  40 cm $^3$ :

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- batch mass: 8,5 times the formulation mass (8,5 × 156,75 g = 132,37 g) for series A rubbers,
  - 9,5 times the formulation mass  $(9,5 \times 141,75 \text{ g} = 1 \text{ 346,62 g})$  for series B rubbers;
- rotor speed: 77 rpm  $\pm$  10 rpm.

NOTE 2 A general mixing procedure for the initial mixing using a laboratory internal mixer is as follows:

		Duration (min)	Cumulative time (min)
a)	Adjust the temperature of the laboratory internal mixer to a starting temperature of 50 °C $\pm$ 3 °C. Close the discharge door, set the rotor speed and raise the ram.	_	_
b)	Load the rubber, lower the ram and allow the rubber to be masticated.	0,5	0,5
c)	Raise the ram and load the zinc oxide, stearic acid and carbon black. Lower the ram.	0,5	1,0
d)	Allow the batch to mix.	2,0	3,0
e)	Raise the ram and clean the mixer throat and the top of the ram. Lower the ram.	0,5	3,5
f)	Allow the batch to mix.	1,5	5,0
g)	Discharge the batch.		

After discharging the batch, immediately check the temperature of the batch with a suitable temperature-measuring device. If the temperature as measured falls outside the range 150 °C to 170 °C, discard the batch. Pass the batch three times through a mill with a mill opening of 2,5 mm and a roll temperature of 50 °C  $\pm$  5 °C. Sheet the batch to a thickness of approximately 10 mm and determine the mass of the batch. If the mass differs from the theoretical value by more than  $\pm$  0,5 % or  $\pm$  1,5 %, discard the batch and remix.

Leave the batch for at least 30 min and up to 24 h, if possible at standard temperature and humidity as defined in ISO 23529.

The smaller laboratory internal mixers do not provide enough compound for the final mill mixing, as a batch mass of three times the formula mass is required. In such cases, the laboratory internal mixer may be used for the final mixing. It may be necessary to adjust the head temperature or the batch mass so that the final temperature of the discharged batch does not exceed 120 °C.

#### 5.3.4.2 Final mill-mixing procedure

During final mixing, maintain a good rolling bank at the nip of the rolls. If this is not attained with the nip settings specified, small adjustments to the mill openings may be necessary.

		Duration (min)	Cumulative time (min)
a)	The standard laboratory mill batch mass, in grams, shall be based on three times the formula mass.	_	_
b)	Set the mill temperature at 50°C±5°C and the mill opening to 1,5 mm. (standards.iteh.ai)	_	_
c)	Band the masterbatch on the slow roll.  ISO 2322:2009	1,0	1,0
d)	Add the sulfur and the accelerator Do not cut the band until the sulfur 27 and accelerator are completely dispersed a6c31e/iso-2322-2009	1,5	2,5
e)	Make three 3/4 cuts from each side, allowing 15 s between each cut.	2,5	5,0
f)	Cut the batch from the mill. Set the mill opening at 0,8 mm and pass the rolled batch endwise through the rolls six times, introducing it from each end alternately.	2,0	7,0

- g) Sheet the batch to approximately 2,2 mm for preparing test sheets or to the appropriate thickness for preparing ISO ring test pieces in accordance with ISO 37. Determine the mass of the batch and record it. If it differs from the theoretical value by more than + 0,5 % or 1,5 %, discard the batch and remix.
- h) Condition the batch for 2 h to 24 h after mixing and prior to vulcanizing, if possible at standard temperature and humidity as defined in ISO 23529.

#### 6 Evaluation of vulcanization characteristics by a curemeter test

#### 6.1 Using an oscillating-disc curemeter

Measure the following standard test parameters:

 $M_{\rm L}$ ,  $M_{\rm H}$  at defined time,  $t_{\rm S1}$ ,  $t_{\rm C}'(50)$  and  $t_{\rm C}'(90)$ ,

in accordance with ISO 3417, using the following test conditions:

- oscillation frequency:
   1,7 Hz (100 cycles per minute);
- amplitude of oscillation: 1° of arc;

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