
Guidelines for the determination of the long-term strength of geosynthetics for soil reinforcement

*Lignes directrices pour la détermination de la résistance à long terme
des géosynthétiques pour le renforcement du sol*

STANDARDSISO.COM : Click to view the full PDF of ISO/TS 20432:2022



STANDARDSISO.COM : Click to view the full PDF of ISO/TS 20432:2022



COPYRIGHT PROTECTED DOCUMENT

© ISO 2022

All rights reserved. Unless otherwise specified, or required in the context of its implementation, no part of this publication may be reproduced or utilized otherwise in any form or by any means, electronic or mechanical, including photocopying, or posting on the internet or an intranet, without prior written permission. Permission can be requested from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office
CP 401 • Ch. de Blandonnet 8
CH-1214 Vernier, Geneva
Phone: +41 22 749 01 11
Email: copyright@iso.org
Website: www.iso.org

Published in Switzerland

Contents

Page

Foreword	v
1 Scope	1
2 Normative references	1
3 Terms, definitions, abbreviated terms and symbols	1
3.1 Terms and definitions	1
3.2 Abbreviated terms	2
3.3 Symbols	3
4 Design procedure	4
4.1 General	4
4.2 Design lifetime	4
4.3 Causes of degradation	5
4.4 Design temperature	5
5 Determination of long-term creep strain	5
5.1 General	5
5.2 Extrapolation	6
5.3 Time-temperature superposition methods	6
5.4 Isochronous curves	7
5.5 Weathering, chemical and biological effects	7
6 Determination of long-term strength	8
6.1 Tensile strength	8
6.2 Reduction factors	8
6.3 Modes of degradation	8
7 Creep rupture	9
7.1 General	9
7.2 Measurement of creep rupture: conventional method	10
7.3 Curve fitting (conventional method)	11
7.4 Curve fitting for time-temperature block shifting of rupture curves	13
7.5 Strain shifting and the stepped isothermal method	14
7.6 Extrapolation and definition of reduction factor or lifetime	15
7.7 Residual strength	16
7.8 Reporting of results	16
7.9 Procedure in the absence of sufficient data	16
8 Installation damage	17
8.1 General	17
8.2 Data recommended	17
8.3 Calculation of reduction factor	18
8.4 Procedure in the absence of direct data	18
8.4.1 General	18
8.4.2 Interpolation from measurements with different soils	18
8.4.3 Interpolation between products of the same product line	19
8.4.4 Laboratory damage tests	19
9 Weathering, chemical and biological degradation	19
9.1 General	19
9.2 Data recommended for assessment	20
9.3 Weathering	20
9.4 Chemical degradation	21
9.4.1 Causes of chemical degradation	21
9.4.2 Evidence from service experience	21
9.4.3 Accelerated chemical degradation tests	22
9.4.4 Oxidation of polyolefins	26
9.4.5 Hydrolysis of polyesters	27

9.4.6	Procedure in the absence of sufficient data	29
9.5	Biological degradation.....	29
10	Determination of long-term strength	29
10.1	Factor of safety f_s	29
10.2	Design for residual strength	30
11	Reporting	30
	Bibliography	31

STANDARDSISO.COM : Click to view the full PDF of ISO/TS 20432:2022

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.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular, the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

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. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO's adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT), see www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 221, *Geosynthetics*.

This first edition of ISO/TS 20432 cancels and replaces ISO/TR 20432:2007, which has been technically revised. It also incorporates the Technical Corrigendum ISO/TR 20432:2007/Cor 1:2008.

The main changes are as follows:

- Subclause 7.4 has been modified to further detail and clarify the fitting of linear regression curves to time-temperature block shifted creep-rupture test results.

Any feedback or questions on this document should be directed to the user's national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.

STANDARDSISO.COM : Click to view the full PDF of ISO/TS 20432:2022

Guidelines for the determination of the long-term strength of geosynthetics for soil reinforcement

1 Scope

This document provides guidelines for the determination of the long-term strength of geosynthetics for soil reinforcement.

This document describes a method of deriving reduction factors for geosynthetic soil-reinforcement materials to account for creep and creep rupture, installation damage and weathering, and chemical and biological degradation. It is intended to provide a link between the test data and the codes for construction with reinforced soil.

The geosynthetics covered in this document include those whose primary purpose is reinforcement, such as geogrids, woven geotextiles and strips, where the reinforcing component is made from polyester (polyethylene terephthalate), polypropylene, high density polyethylene, polyvinyl alcohol, aramids and polyamides 6 and 6,6. This document does not cover the strength of joints or welds between geosynthetics, nor whether these might be more or less durable than the basic material. Nor does it apply to geomembranes, for example, in landfills. It does not cover the effects of dynamic loading. It does not consider any change in mechanical properties due to soil temperatures below 0 °C, nor the effect of frozen soil. The document does not cover uncertainty in the design of the reinforced soil structure, nor the human or economic consequences of failure.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements 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 10318-1, *Geosynthetics — Part 1: Terms and definitions*

3 Terms, definitions, abbreviated terms and symbols

3.1 Terms and definitions

For the purposes of this document, the terms and definitions given in ISO 10318-1 and the following apply.

ISO and IEC maintain terminology databases for use in standardization at the following addresses:

- ISO Online browsing platform: available at <https://www.iso.org/obp>
- IEC Electropedia: available at <https://www.electropedia.org/>

3.1.1

long-term strength

load which, if applied continuously to the geosynthetic during the service lifetime, is predicted to lead to rupture at the end of that lifetime

3.1.2

reduction factor

factor (≥ 1) by which the tensile strength is divided to take into account particular service conditions in order to derive the long-term strength

Note 1 to entry: In Europe, the term 'partial factor' is used.

3.1.3

characteristic strength

95 % (two-sided) lower confidence limit for the tensile strength of the geosynthetic, approximately equal to the mean strength less two standard deviations

Note 1 to entry: This should be assured by the manufacturer's own quality assurance scheme or by independent assessment.

3.1.4

block shifting

procedure by which a set of data relating applied load to the logarithm of time to rupture, all measured at a single temperature, are shifted along the log time axis by a single factor to coincide with a second set measured at a second temperature

3.1.5

product line

series of products manufactured using the same polymer, in which the polymer for all products in the line comes from the same source, the manufacturing process is the same for all products in the line, and the only difference is in the product mass per area or number of fibres contained in each reinforcement element

3.2 Abbreviated terms

CEG	carboxyl end group
DSC	differential scanning calorimetry
HALS	hindered amine light stabilizers
HDPE	high density polyethylene
HPOIT	high pressure oxidation induction time
LCL	lower confidence limit
MARV	minimum average roll value
OIT	oxidation induction time
PA	polyamide
PET	polyethylene terephthalate
PP	polypropylene
PTFE	polytetrafluoroethylene
PVA	polyvinyl alcohol
SIM	stepped isothermal method
TTS	time-temperature shifting

3.3 Symbols

A_i	time-temperature shift factor
b_a	gradient of Arrhenius graph
d_{50}	mean granular size of fill
d_{90}	granular size of fill for 90 % pass (10 % retention)
f_s	factor of safety
G, H	parameters used in the validation of temperature shift linearity (see 7.4)
m	gradient of line fitted to creep rupture points (log time against load); inverse of gradient of conventional plot of load against log time.
M_n	number averaged molecular weight
n	number of creep rupture or Arrhenius points
P	applied load
R_1	ratio representing the uncertainty due to extrapolation
R_2	ratio representing the uncertainty in strength derived from Arrhenius testing
$f_{R,CH}$	reduction factor to allow for chemical and biological effects
$f_{R,CR}$	reduction factor to allow for the effect of sustained static load
$f_{R,ID}$	reduction factor to allow for the effect of mechanical damage
$f_{R,W}$	reduction factor to allow for weathering
S_{sq}	sum of squares of difference of log (time to rupture) and straight line fit
S_{xx}, S_{xy}, S_{yy}	sums of squares as defined in derivation of regression lines in 9.4.3
σ_0	standard deviation used in calculation of LCL
t	time, expressed in hours
t_{90}	time to 90 % retained strength
t_D	design life
t_{deg}	degradation time during oxidation
t_{ind}	induction time during oxidation
t_{LCL}	LCL of time to a defined retained strength at the service temperature
t_{max}	longest observed time to creep rupture, expressed in hours
t_{n-2}	Student's t for $n - 2$ degrees of freedom and a stated probability
t_R	time to rupture, expressed in hours
t_s	time to a defined retained strength at the service temperature

T	load per width
T_B	batch tensile strength (per width)
T_{char}	characteristic strength (per width) (see 6.1)
T_x	unfactored long-term strength (see 9.4.3)
T_D	long-term strength per width (including factor of safety)
T_{DR}	residual strength
θ_j	temperature of accelerated creep test
θ_k	absolute temperature
T_{LCL}	LCL of T_{char} due to chemical degradation
θ_s	service temperature
x	abscissa: on a creep rupture graph the logarithm of time, in hours
$\bar{x}$	mean value of x
x_i	abscissa of an individual creep rupture point
x_p	predicted time to rupture
y	ordinate: on a creep rupture graph, applied load expressed as a percentage of tensile strength, or a function of applied load
y_0	value of y at 1 h ($\lg t = 0$)
$\bar{y}$	mean value of y
y_i	ordinate of an individual creep rupture point
y_0	value of y at time 0, derived from the line fitted to creep rupture points

4 Design procedure

4.1 General

The design of reinforced soil structures generally requires consideration of the following two issues:

- the maximum strain in the reinforcement during the design lifetime;
- the minimum strength of the reinforcement that could lead to rupture during the design lifetime.

In civil engineering design, these two issues are referred to as the serviceability and ultimate limit state respectively. Both factors depend on time and can be degraded by the environment to which the reinforcement is exposed.

4.2 Design lifetime

A design lifetime, t_D , is defined for the reinforced soil structure. For civil engineering structures this is typically 50 years to 100 years. These durations are too long for direct measurements to be made in advance of construction. Reduction factors have therefore to be determined by extrapolation of short-term data aided, where necessary, by tests at elevated temperatures to accelerate the processes of creep or degradation.

4.3 Causes of degradation

Strain and strength may be changed due to the effects of the following:

- mechanical damage caused during installation;
- sustained static (or dynamic) load;
- elevated temperature;
- weathering while the material is exposed to light;
- chemical effects of natural or contaminated soil.

4.4 Design temperature

The design temperature should have been defined for the application in hand. In the absence of a defined temperature or of site specific in-soil temperature data, the design temperature should be taken as the temperature which is halfway between the average yearly air temperature and the normal daily air temperature for the hottest month at the site. If this information is not available, 20 °C should be used as the default value.

Many geosynthetic tests are performed at a standard temperature of (20 ± 2) °C. If the design temperature differs, appropriate adjustments should be made to the measured properties.

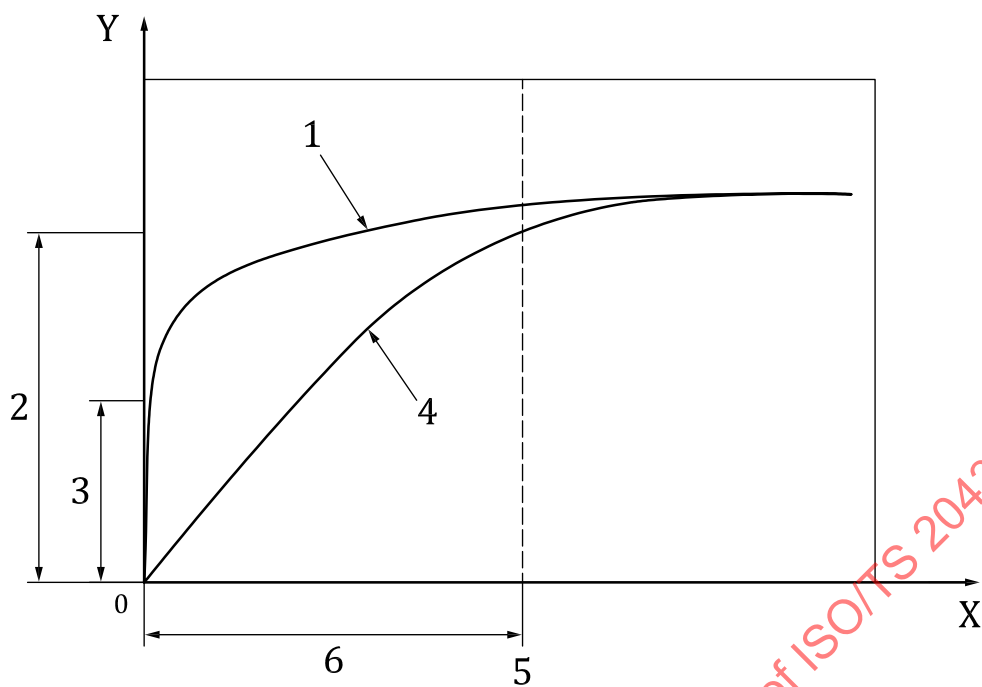
This document does not cover the effects of temperatures below 0 °C (see [Clause 1](#)).

5 Determination of long-term creep strain

5.1 General

The design specification may set a limit on the total strain over the service lifetime of the geosynthetic, or on the strain generated between the end of construction and the service lifetime. In the second case, the time at “end of construction” should be defined, as shown in [Figure 1](#). When plotted against $\lg t$, even a one-year construction period should have negligible influence on the creep strain curve beyond 10 years.

Levels of creep strain encountered in the primary creep regime (creep rate decreasing with time) are thought not to adversely affect strength properties of geosynthetic reinforcement materials.



Key	
X	time
Y	strain
1	laboratory creep test
2	load ramp period on wall
3	load ramp period in creep test
4	loading and creep of reinforcement in wall
5	new time = 0 for post construction creep
6	wall construction time

Figure 1 — Conceptual illustration for comparing the creep measured in walls to laboratory creep data

5.2 Extrapolation

Creep strain should be measured according to ISO 13431 and plotted as strain against the $\lg t$. It may then be extrapolated to the design lifetime. Extrapolation may be by graphical or curve-fitting procedures, in which the formulae applied should be as simple as is necessary to provide a reasonable fit to the data, for example, power laws. The use of polynomial functions is discouraged since they can lead to unrealistic values when extrapolated.

5.3 Time-temperature superposition methods

Time-temperature superposition methods may be used to assist with extending the creep curves. Creep curves are measured under the same load at different temperatures, with intervals generally not exceeding 10 °C, and plotted on the same diagram as strain against $\lg t$. The lowest temperature is taken as the reference temperature. The creep curves at the higher temperatures are then shifted along the time axis until they form one continuous “master” curve, i.e. the predicted long-term creep curve for the reference temperature. The shift factors, i.e. the amounts (in units equivalent to $\lg t$) by which each curve is shifted, should be plotted against temperature where they should form a straight line or smooth curve. The cautions given in 7.6 should be noted.

Experience has shown the strains on loading are variable. Since the increase in strain with time is small, this variability can lead to wide variability in time-temperature shifting (TTS). The stepped isothermal method (SIM) described in 7.5 avoids this problem by using a single specimen, increasing the temperature in steps, and then shifting the sections of creep curve measured at the various temperatures to form one continuous master curve.

If a more accurate measure of initial strain is required, five replicates are recommended at each load. Some of these can be of short duration (e.g. 1 000 s). At a series of loads, fewer replicates at each load will suffice if the data are pooled using regression techniques. One approach is to use regression analysis to develop an isochronous load versus strain curve at 0,1 h. The creep curve should then be shifted vertically to pass through the mean strain measured after 0,1 h.

If the lowest test temperature is below the design temperature, the shift factor corresponding to the design temperature should be read off the plot of shift factor against temperature. The time-scale of the master curve should then be adjusted by this factor.

5.4 Isochronous curves

From the creep curve corresponding to each load, read off the strains for specified durations, typically 1 h, 10 h, 100 h, etc., and including the design lifetime. Set up a diagram of load against strain. For each duration, plot the points of load against strain for the corresponding durations (see Figure 2). These are called isochronous curves. Where a maximum strain is permitted over the design lifetime, or between the end of construction (e.g. 100 h) and the design lifetime, it is possible to read off the corresponding loads from these curves. Where the strain is measured from zero, note that in geosynthetics strains are measured from a set preload (defined in ISO 10319 and ISO 13431 as 1 % of the tensile strength) and that some woven and particularly non-woven materials may exhibit considerable irreversible strains below this initial loading. See Reference [2] for additional details on creep strain characterization.

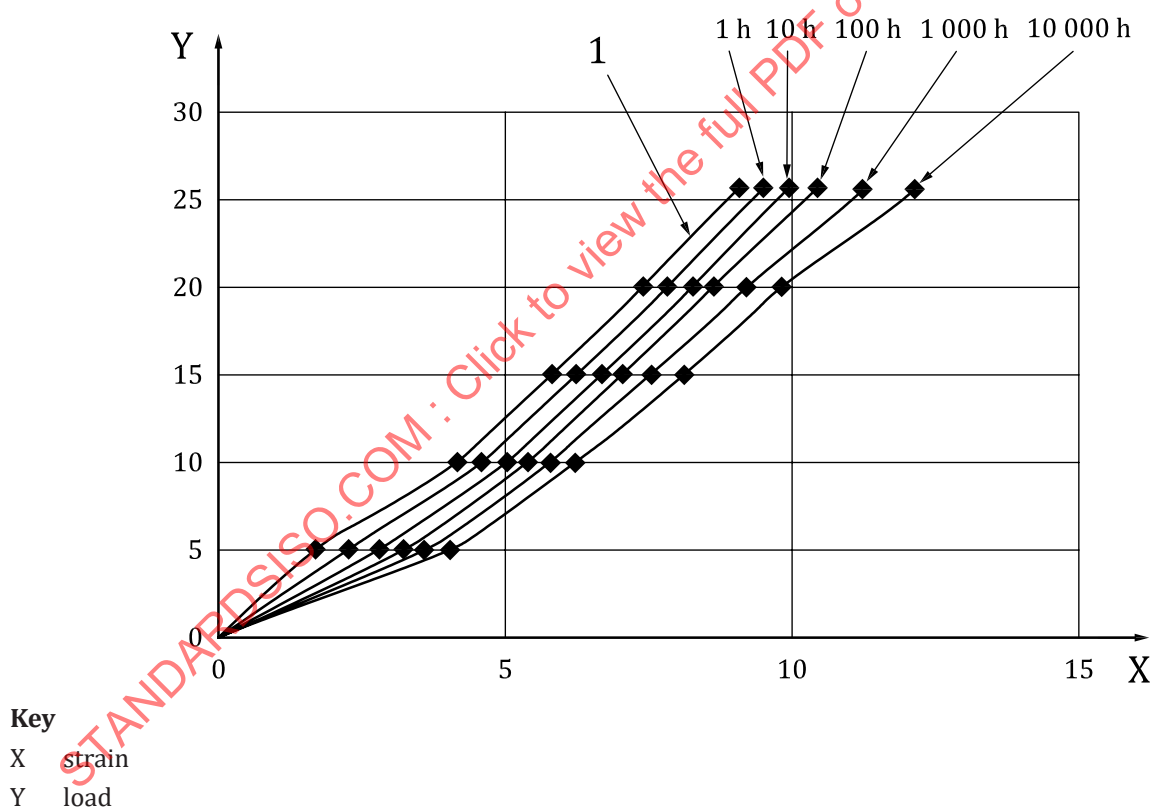


Figure 2 — Isochronous diagram

5.5 Weathering, chemical and biological effects

Creep strain is generally insensitive to limited weathering, chemical and biological effects. In addition, creep strains are in general not affected by installation damage, unless the damage is severe, or unless the load level applied is very near the creep limit of the undamaged material. In most cases, the load level applied is well below the creep limit of the material. See Reference [3] for additional details on this issue. Thus, no further adjustment is generally required beyond the effect of temperature.

Note, however, that artificially contaminated soils may contain chemicals, such as organic fuels and solvents, which can affect the creep of geosynthetics. If necessary, perform a short-term creep test according to ISO 13431 on a sample of geosynthetic that is immersed in the chemical or has just been removed from it. If the creep strain is significantly different, do not use this geosynthetic in this soil.

6 Determination of long-term strength

6.1 Tensile strength

The characteristic strength, T_{char} , is taken as the basis for the long-term strength. T_{char} is typically a statistical value generated from the mean strength of production material less two standard deviations sometimes referred to as the minimum average roll value (MARV), unless otherwise defined.

6.2 Reduction factors

T_{char} can then be divided by the following four reduction factors, each of which represents a loss of strength determined in accordance with this Technical Specification, to arrive at the long-term strength T_{D} :

- $f_{\text{R,CR}}$ is a reduction factor to allow for the effect of sustained static load at the service temperature;
NOTE The effect of dynamic loads is not included.
- $f_{\text{R,ID}}$ is a reduction factor to allow for the effect of mechanical damage;
- $f_{\text{R,W}}$ is a reduction factor to allow for weathering during exposure prior to installation or of permanently exposed material;
- $f_{\text{R,CH}}$ is a reduction factor to allow for reductions in strength due to chemical and biological effects at the design temperature (see 4.4).

In addition to the reduction factors, a factor of safety, f_s , takes into account the statistical variation in the reduction factors calculated (see 6.1). It does not consider the uncertainties related to the soil structure and the calculation of loads.

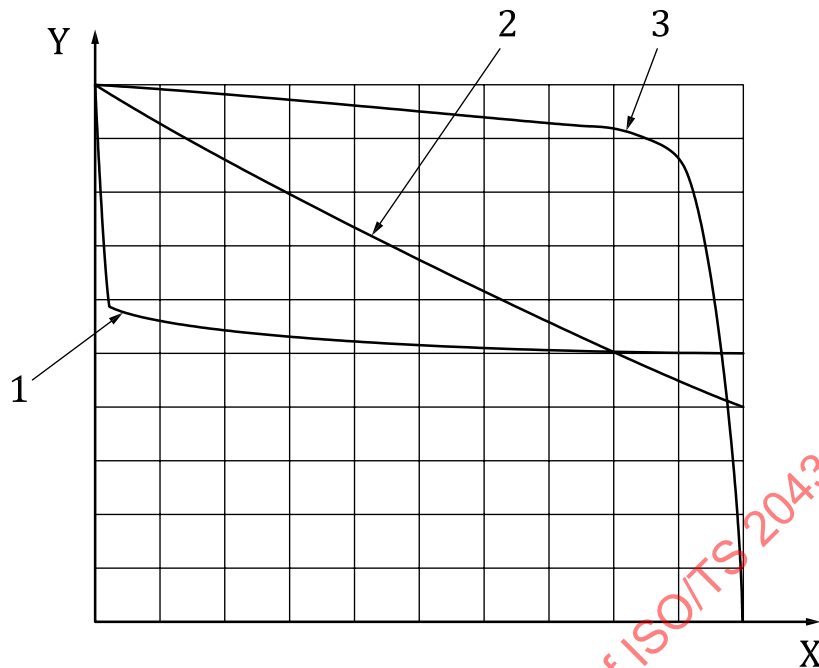
6.3 Modes of degradation

Degradation of strength can be divided into three modes according to the manner in which they take place with time:

- Mode 1: Immediate reduction in strength, insignificant further reduction with time.
- Mode 2: Gradual, though not necessarily constant, reduction in strength.
- Mode 3: No reduction in strength for a long period; after a certain period, onset of rapid degradation.

For Mode 1, of which installation damage is an example, it is appropriate to reduce the tensile strength by an appropriate time-independent reduction factor. For Mode 2, where there is a progressive reduction in strength, the tensile strength will be reduced by a time-dependent reduction factor. For Mode 3, it is not appropriate to apply a reduction factor to the tensile strength but rather to restrict the service lifetime.

These modes are depicted schematically in [Figure 3](#).

**Key**

- X time
- Y retained strength
- 1 mode 1
- 2 mode 2
- 3 mode 3

Figure 3 — Retained strength plotted against time for the three Modes of degradation

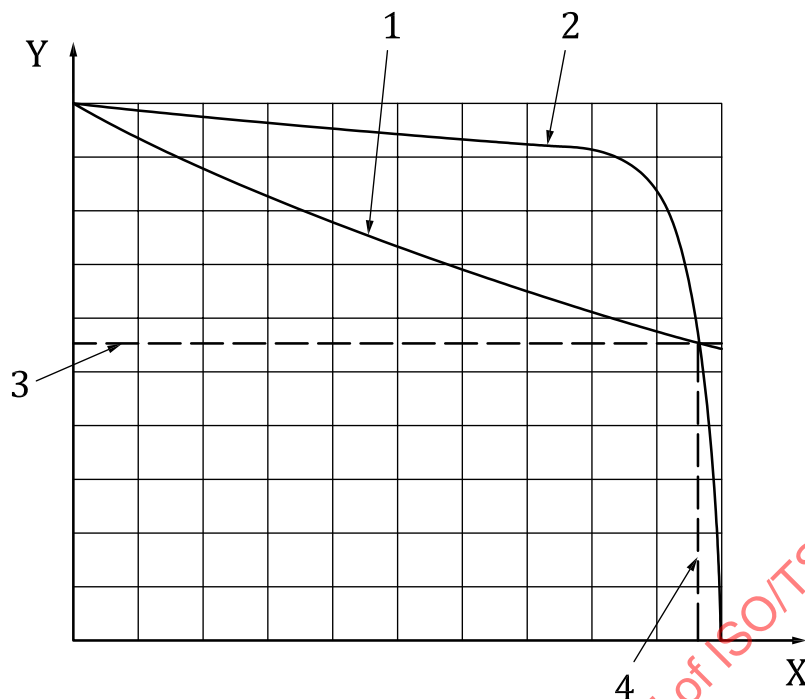
7 Creep rupture

7.1 General

Creep rupture, or lifetime under sustained load, is determined by measuring times to rupture of up to at least 10 000 h. The results are extrapolated to predict longer lifetimes at lower loads and thereby the reduction factor $f_{R,CR}$.

This procedure may be supported by measurements at higher temperatures. Conventional TTS of results obtained on multiple specimens at elevated temperatures provides an improved prediction of the long-term behaviour at ambient temperature. In the SIM, the temperature of a single specimen is increased in steps. The sections of creep strain curve measured at each temperature step are then combined to predict the long-term creep strain and rupture lifetime.

It should be noted that a creep rupture diagram depicts applied load plotted against time to rupture and is not a statement of the loss of strength under continuous load. It has been predicted on the basis of accelerated tests that many geosynthetics exposed to sustained load do not in fact significantly diminish in strength until close to the end of their predicted life. When the strength equals the applied load, the material ruptures (see [Figure 4](#)). Sustained load is therefore a Mode 3 form of degradation.



Key

- X time
- Y applied load, residual strength
- 1 creep rupture
- 2 residual strength
- 3 applied load
- 4 lifetime

Figure 4 — Creep rupture and residual strength as a function of time

The creep rupture curve shows the predicted lifetime corresponding to a particular applied load. During that lifetime, the strength of the geosynthetic follows the residual strength curve, falling to equal the applied load at the moment of rupture.

7.2 Measurement of creep rupture: conventional method

For limit state design, the creep rupture behaviour of the product should be measured according to ISO 13431 with a minimum of 12 measurements. As a guide, at least four of the test results should have rupture times between 100 h and 1 000 h, and at least four of the test results should have rupture times of 1 000 h to 10 000 h, with at least one additional test result having a rupture time of approximately 10 000 h (1,14 years) or more.

Specimens should be tested in the direction in which the load will be applied in use. The tensile strength of the same batch, T_B , of the material in the same direction should be determined according to ISO 10319 using grips similar to those used for creep rupture testing. Loads applied during the creep rupture tests should be expressed as a percentage of T_B . The nature of the failure should be observed and recorded.

It is recommended that creep strain is measured as well as time to rupture, since this can assist with conventional time-temperature strain shifting and in identifying any change in behaviour that could invalidate extrapolation of the results. This practice will also permit laboratory creep data collected at moderate differences (plus or minus 10 °C) in test temperature to be corrected to the desired reference temperature. Similar moderate changes in reference temperature will be facilitated under this practice as well.

The temperature should be as stated in ISO 13431 and ISO 10319; if a different temperature, for example, the design temperature, is used then it should be the same for both tensile and creep rupture measurements. Further tests at elevated temperature may be used for the purposes of TTS.

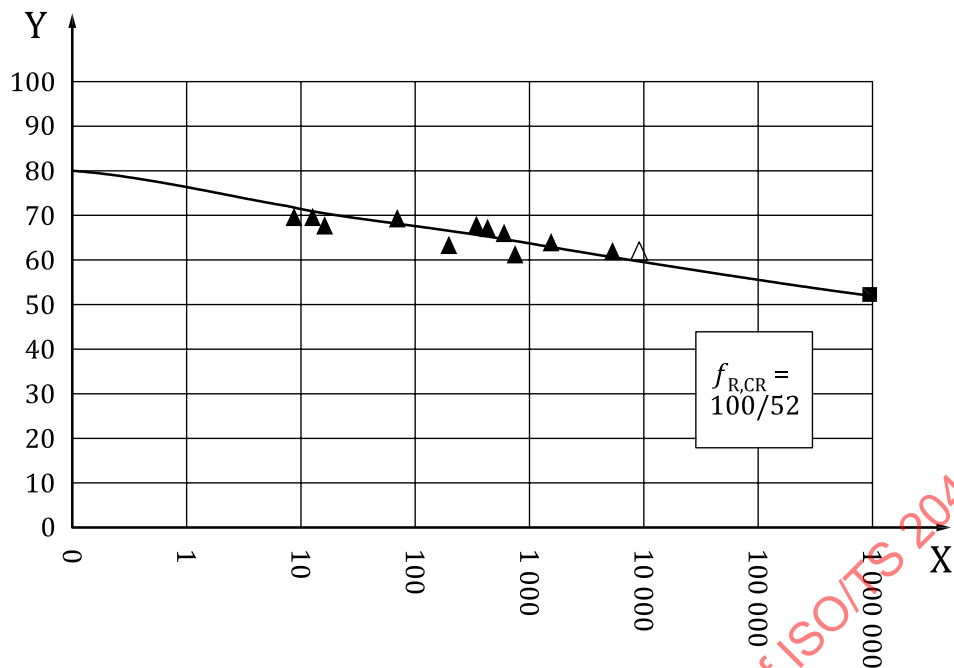
The creep rupture data for the product should be tabulated as:

- load per width T , as percentage of the batch tensile strength, T_B ;
- time to rupture, t_R , in h;
- $\lg t$ to rupture;
- observations on the failure, including the strain at failure or the strain at the point where the rate of creep starts to increase (tertiary creep) and, where visible, the nature of the fracture surface, e.g. ductile, semi-brittle or brittle and smooth;
- creep strain data, if available, particularly if conventional time-temperature strain shifting is applied;
- whether the test was conventional (20 °C), time-temperature accelerated, SIM or was performed on a similar material as supporting data.

Incomplete tests may be included, with the test duration replacing the time to rupture, but should be listed as such. The procedure for handling incomplete tests is described in [7.3](#).

7.3 Curve fitting (conventional method)

The data, including any relevant supporting data, should be plotted as $y = T$ (expressed as a percentage of T_B) against $x = \lg t_R$, which should yield a linear plot (see [Figure 5](#)). This is referred to as a semi-logarithmic plot and has been shown to apply to polyester reinforcements. If the plot is not linear, it may be necessary to plot the ordinate (y) as a function of applied load to achieve a linear plot. The use of the function $y = \lg T$, resulting in a double logarithmic plot, has been shown to apply to polyethylene and polypropylene reinforcements. Where a function of T is used, it should preferably be based on a known physical model.

**Key**X logarithm of time to rupture ($\lg t_R$)Y load per width T , as % tensile strength**Figure 5 — Creep rupture diagram with straight line fit**

Fit a straight line using statistical regression analysis. In the following, x equals $\lg t_R$ and y equals T or a function of P . The creep rupture points, total number n , are denoted as (x_i, y_i) . Note that in contrast to most scientific plots, the independent variable is plotted on the y axis and the dependent variable is plotted on the x axis. Formulae (1) to (3) therefore differ from those conventionally found by having x and y interchanged.

The straight line fit (regression line) is given by the formula:

$$x = \bar{x} + m(y - \bar{y}) \quad (1)$$

where

$$\bar{x} = \frac{\sum x_i}{n} \text{ and } \bar{y} = \frac{\sum y_i}{n}$$

summed over all points (x_i, y_i) .

m is given by the formula:

$$m = \frac{\sum (x_i - \bar{x})(y_i - \bar{y})}{\sum (y_i - \bar{y})^2} \quad (2)$$

Because of the interchange of x and y , the gradient of the graph is equal to $1/m$. For a semi-logarithmic diagram, this should be expressed as percentage tensile strength per decade of time. The gradient should be a negative value.

The intercept y_0 on the line $x = 0$ (i.e. at $\lg t = 0$; $t = 1$ h) is given by:

$$y_0 = \bar{y} - \bar{x} / m \quad (3)$$

The accepted practice for incomplete tests is as follows. The regression should first be performed with the incomplete tests excluded. The time to failure for an incomplete test should then be determined for the corresponding value of T . If the predicted time to failure is less than the duration of the incomplete test, the point may be added and the regression recalculated. If the predicted time to failure is greater than the duration of the incomplete test, the point should continue to be excluded. In [Figure 5](#) the incomplete test shown by an open triangle is included since it lies to the right of the regression line.

Extend the regression line to the design lifetime, for example in [Figure 5](#) where for a design lifetime of 1 000 000 h, $T = 52$ % of tensile strength. $f_{R,CR} = 1/52$ % = $100/52 = 1,92$

Record the duration of the longest test that has ended in rupture, or the duration of the longest incomplete test whose duration has been included in the regression calculation; this duration is denoted as $t_{\max}$.

7.4 Curve fitting for time-temperature block shifting of rupture curves

If data obtained at higher temperatures θ_i are to be included for the purposes of acceleration, tabulate the values of y_i and t_R as in [7.3](#) together with the temperatures θ_i . For each temperature θ_i , assign a nominal shift factor A_i . Assign nominal values to the constants y_0 and m . Include the test points derived at 20 °C for which $A_i = 0$. Then proceed as follows.

For each measured value of t_R , calculate the shifted log time $x_i = \lg t_R + A_i$.

For each value of y_i , calculate the logarithm of the predicted time to rupture $x_p = (y_i - y_0)m$.

For each pair of values, calculate the square of the difference $(x_i - x_p)^2$.

Derive the sum of squares $S_{sq} = \sum i (x_i - x_p)^2$.

Using a spreadsheet optimization programme, minimize S_{sq} as a function of all A_i, y_0 and m .

Using the optimized values of A_i , recalculate the values of x_i .

Plot y_i against the recalculated values of x_i . On the same diagram, using the optimised values of y_0 and m , add the straight line fit as in [7.3](#). This line should be used for prediction of the time to rupture at 20 °C.

Plot the optimized values of A_i against θ_i . Check that the line passes through the point (20 °C, 0) and is then straight or lightly curved, such that if the curve is approximated by Formula (4):

$$A_i = G(\theta_i - 20) + H(\theta_i - 20)^2 \quad (4)$$

then $-0,003 < G/H < 0,003$. If not, the validity of the tests should be reviewed.

This curve may be interpolated or extended to derive the temperature shift factor A_i corresponding to a different service temperature.

For example, in [Figure 6](#), the regression creep rupture lines for 20 °C, 40 °C and 60 °C are assumed to be parallel. The 40 °C and 60 °C lines and associated points have been shifted to the right until they coincide with the 20 °C line to which they form an extension. Temperature steps ≤ 10 °C are recommended for PE and PP.

This procedure assumes that the creep rupture curves at all temperatures are linear and parallel, which has been found empirically to apply to polyester (semi-log plots) and polypropylene (log/log plots). It should be pointed out that the theory of Zhurkov^[4] in the Bibliography, which assumes that the fracture

process is activated thermally with the additional effect of applied stress, predicts that the creep rupture characteristics should be straight when plotted on a semi-logarithmic diagram, and that their gradients should be stress-dependent. This theory has not provided a better fit to experimental creep rupture data than the empirical method used here, but experience has shown that the shift factors can be stress-dependent and block shifting ignores this.

7.5 Strain shifting and the stepped isothermal method

Long-term rupture data can be obtained through the use of the classical TTS of creep strain data. Strain shifting as described in 5.2 can be applied to creep curves terminated in rupture. For example, a creep strain versus $\lg t$ curve obtained under a given load at 60 °C and which terminates in rupture can be shifted to longer times. Needed to accomplish this are creep strain curves at, say, 20 °C and 40 °C under the same load. The lower temperature curves can be terminated before rupture provided that sufficient data are available to effect the TTS procedure properly. Because of the scatter in initial strains mentioned previously, the strain tests should be replicated.

In the SIM, which is a special case of TTS, the temperature of the creep test is raised in a series of steps. The sections of creep curve at the individual temperatures are then combined to form a continuous determination as instructed in ASTM D 6992.

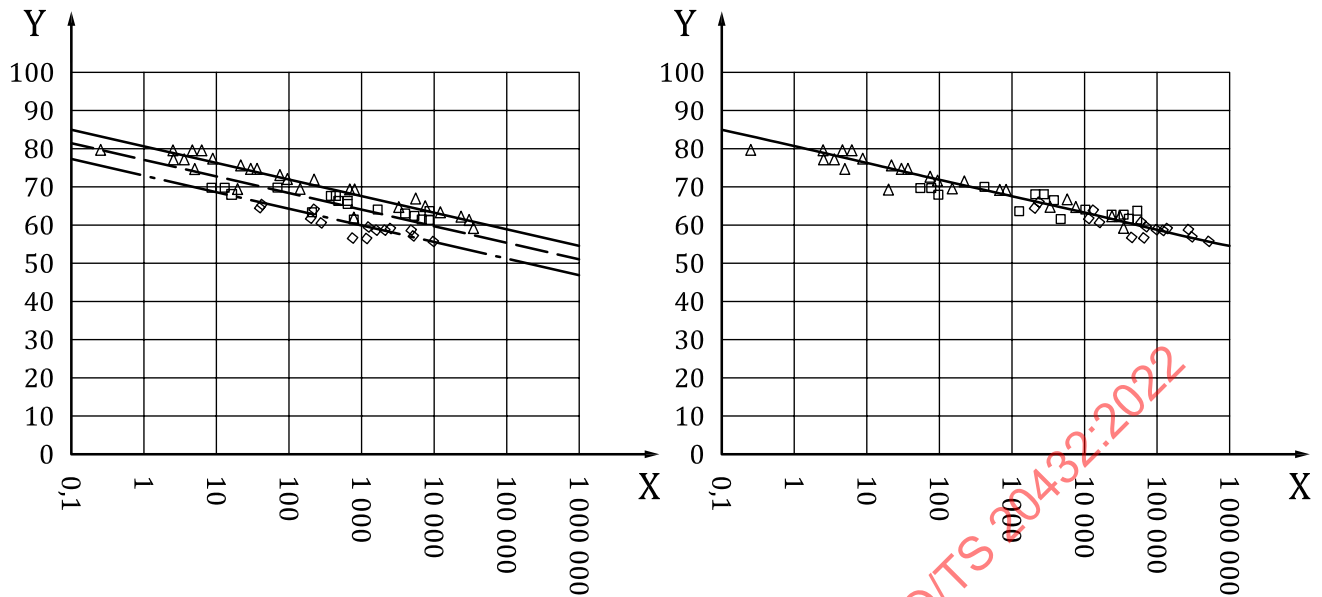
SIM can be considered for use in generating and extrapolating geosynthetic creep rupture data, provided that the predictions are consistent with those based on conventional testing or time-temperature block or strain shifting as described above. To this end, it is recommended that a minimum of 12 data points, time-shifted to the reference temperature, be obtained from accelerated (TTS and SIM) and conventional testing, with a minimum of

- three time-shifted durations between 1 000 and 100 000 h, and
- three time-shifted durations between 100 000 and 10 000 000 h.

In addition, a limited programme of conventional creep rupture tests obtained at the reference temperature and therefore un-shifted (except as corrected per 7.2), should be performed in accordance with 7.2. It is recommended that there should be four conventional creep rupture data points between 100 h and 10 000 h and one data point at 10 000 h or more. (The last data point may be an incomplete test). This conventional creep rupture data envelope should then be compared to the envelope determined from the accelerated data.

Linear regression analysis should be performed separately for the conventional and accelerated data in accordance with 7.3 and 7.4. The value of $f_{R,CR}$ determined from the accelerated data at 2 000 h at the reference temperature should differ from the value of $f_{R,CR}$ determined from conventional data at 2 000 h at the reference temperature by no more than 0,15. Also the value of $f_{R,CR}$ determined from the accelerated data at 10 000 h at the reference temperature should differ from the value of $f_{R,CR}$ determined from conventional data at 10 000 h at the reference temperature by no more than 0,15. If both the conditions are fulfilled, the SIM data may be combined with the conventional data and used to determine $f_{R,CR}$. If not, $f_{R,CR}$ should be determined from data from conventional testing alone (additional conventional data will be needed in this case).

The validity of SIM is supported by References [5] and [9].



a) Linear regressions using test results from each 20 °C, 40 °C and 60 °C

b) Block shifted data to a single reference temperature of 20 °C using all data in a)

Key

X	time (h)
Y	percentage tensile strength
◇	60 ruptures
□	40 ruptures
△	20 ruptures
—	regression 20
— —	regression 40
- - -	regression 60

Figure 6 — Block shifting

7.6 Extrapolation and definition of reduction factor or lifetime

Extrapolate the straight line fit to $\lg t_D$. Read off the corresponding percentage y from the formula $y = y_0 - (\lg t_D)/m$ (if y is a different function of load, derive the percentage accordingly).

Calculate $f_{R,CR} = 100/y$. $f_{R,CR}$ should be greater than unity.

A condition of the extrapolation is that there is no evidence or reason to believe that the rupture behaviour will change over this duration. It should be checked that at long durations, and at elevated temperatures, if used:

- there is no abrupt change in the gradient of the creep rupture curve;
- there is no abrupt change in the strain to failure;
- there is no significant change in the appearance of the fracture surface.

Any evidence of such changes, particularly in accelerated tests, should invalidate the extrapolation unless it can be taken into account as described in the following example. Particular attention is drawn to the behaviour of unoriented thermoplastics under sustained load, where a transition in behaviour is observed in long-term creep rupture testing. The effect of this transition is that the gradient of the creep rupture curve steepens at the so-called “knee” such that long-term failures occur at much shorter

lifetimes than would otherwise be predicted. The strain at failure is greatly reduced and the appearance of the fracture surface changes from ductile to semi-brittle. If this is observed, any extrapolation should assume that the “knee” will occur. For the method of extrapolation, reference should be made to ISO 9080:2012.

7.7 Residual strength

Creep rupture is Mode 3 degradation, resulting in little reduction in strength until the duration approaches the design life (see [Figure 4](#)). If the applied load is expected to be lower than $T_{\text{char}}/f_{\text{R,CR}}$, it can be more appropriate to calculate the time to failure corresponding to the applied load and to check that this substantially exceeds t_{D} . On the basis of current measurements, it may then be assumed that the strength remains close to T_{char} over the design life. This is particularly relevant to seismic design and to other cases where a certain reserve strength has to be assured.

7.8 Reporting of results

The results should be reported as a graph of applied load (or a function of applied load) plotted against time to rupture in the manner of [Figure 5](#).

The following should be stated:

- material;
- design lifetime;
- design temperature;
- T_{char} ;
- equation of the regression line $y = y_0 - x/m$;
- $f_{\text{R,CR}}$.

7.9 Procedure in the absence of sufficient data

Long-term creep data obtained from tests performed on older product lines, or other products within the same product line, may be applied to new product lines, or a similar product within the same product line, if one of the following conditions is met.

- The materials and structure of the proposed product are similar to those of the tested product. Data should be provided which shows that the minor differences between the tested and the untested products will result in equal or greater creep resistance for the untested products.
- The results of a limited testing programme on the proposed product are not significantly different from those predicted from the data on the tested product. For creep evaluation, this limited testing programme should include creep tests taken to at least 1 000 h to 2 000 h in length.
- If SIM is accepted for the previously tested product, then SIM can be used RF_{CR} exclusively on the proposed product or products. In this case, the SIM tests should be concentrated in the 100 000 h to 10 000 000 h time window for maximum statistical efficiency. Three SIM tests should be sufficient for each proposed product.

Similarity can be judged on the following.

- Equivalence of polymer structure, molecular weight, carboxyl end group count (CEG) cross-linking, crystallinity and draw ratio. It should be noted that per cent crystallinity is not a controlled property and there is presently no indication of what an acceptable value for percent crystallinity should be. For the method of determining CEG, see [9.4.5.2](#).
- Tensile strength per identifiable unit such as single rib or yarn. Tests performed on single ribs or yarns should, however, be shown to be representative of the material as a whole.

- Polymer additives used (i.e. type and quantity of antioxidants or other additives used).
- Textile (weave, style of non-woven, grid) and yarn structure, and fibre diameter.

NOTE Not all properties apply to all materials.

The data provided should show that the performance of the new or similar product is equal to or better than the performance of the product previously tested. If so, the results from the full testing programme on the older or similar product could be used for the new/similar product. If these conditions are not met, then a full testing and evaluation programme for the new product should be conducted.

Single ribs for geogrids or yarns for woven geotextiles may be used for creep testing for ultimate limit state design provided that it can be shown, for example, by a creep testing programme similar to the conventional creep tests defined in 7.5, that the rupture behaviour and envelope for the single ribs or yarns are the same as that for the full product.

If the procedures described in this section are applied, then this should be noted in the statement of the corresponding reduction factors.

8 Installation damage

8.1 General

Coarse backfills and heavy compaction loads can damage geosynthetics, causing an immediate reduction in strength. The effect is referred to as installation damage and the corresponding reduction factor as $f_{R,ID}$.

Generally, the mechanical damage occurs on installation (Mode 1). If significant further damage is likely to occur in use, there will be an additional time-dependent contribution to this factor.

8.2 Data recommended

Measurement of the effect of installation damage on geosynthetic reinforcement strength and deformation should be determined from the results of installation damage tests. General guidance is given ISO 13437 and BS 8006:1995, Annex D. The installation damage tests should simulate the installation conditions (conditions of service) as closely as practicable to the installation conditions anticipated in the geosynthetic structure. The installation conditions to be simulated should include, as a minimum:

- the nature of the backfill both below and above the sample: particle size distribution, hardness and angularity;
- the depth at which the sample is installed;
- whether the material is driven over by vehicles before compaction;
- method and degree of compaction.

Test results from damaged specimens should be compared to tensile test results obtained from undamaged (i.e., not exposed to installation conditions) specimens taken from the same lot, and preferably the same roll, of material as the damaged specimens.

The specimens should be large enough to be used for wide-width tensile testing (ISO 10319). Consideration should be given to increasing the number of specimens to ensure that they are fully representative of the damaged material. It is desirable that multi-rib tests, with at least four ribs, should be used for installation damage evaluation. With single rib testing it can be difficult to assess the effect of severed ribs on the strength and modulus of damaged materials, and the effect of differences in degree of damage between ribs on the overall tensile strength of the product. Single ribs of geogrids are generally unsuitable for installation damage testing. If this cannot be avoided, for example, for very high

strength materials, then it should be demonstrated that the strength of the single ribs is representative of the full product.

Further information is given in Reference [10].

8.3 Calculation of reduction factor

The reduction factor to allow for the effect of mechanical damage for the site conditions used, $f_{R,ID}$, should be expressed as the ratio of the mean tensile strength of the undamaged material to the mean tensile strength of the damaged material.

8.4 Procedure in the absence of direct data

8.4.1 General

In the absence of site-specific data obtained in accordance with 8.2, one of the approaches in 8.4.2, 8.4.3 or 8.4.4 can be taken.

8.4.2 Interpolation from measurements with different soils

If the $f_{R,ID}$ of the material under consideration is known for other soils with grain size both less than and greater than the soil to be used, then $f_{R,ID}$ should be determined by interpolation using the values of d_{50} or an alternative such as d_{90} for the respective soils to obtain $f_{R,ID}$ for the soil in question. It is recognized that this is only an approximation, particularly for soils with a broad particle distribution, and other soil gradation characteristics may be considered for interpolation purposes if it can be shown that they produce a more accurate correlation than the d_{50} size. An example of this interpolation procedure to obtain $f_{R,ID}$ at a different soil d_{50} is provided in Figure 7, which shows the interpolation of $f_{R,ID}$ for a soil with d_{50} equalling 2 mm from measurements made with soils with d_{50} equalling 0,02 mm, 0,5 mm and 10 mm.

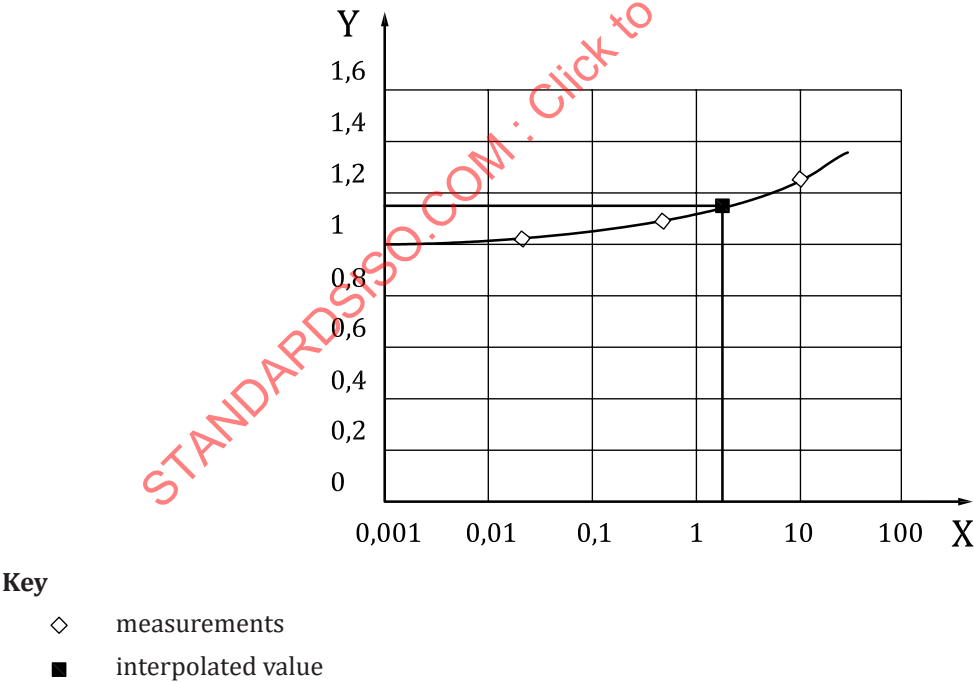
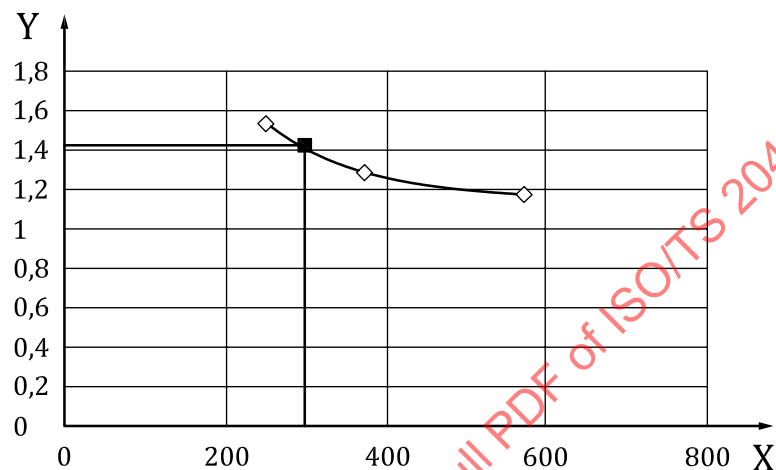


Figure 7 — Interpolation of $f_{R,ID}$

8.4.3 Interpolation between products of the same product line

This interpolation may also be made for other products within the product line of the subject product, provided that a relationship can be established between the weight, tensile strength, etc. of the product and the $f_{R,ID}$ of the product as in Figure 8, and provided that data are available for products which are both lighter (weaker) and heavier (stronger) than the product in question. For products that are heavier (stronger) than the heaviest product tested, the $f_{R,ID}$ for the heaviest product tested may be used. For coated polyester geogrids, the coating thickness or coating mass per area relative to the mass per area of the product should be considered for the purpose of correlating $f_{R,ID}$ between products rather than product weight or tensile strength alone. In Figure 8 for a product of weight 300 g/m^2 , $f_{R,ID} = 1,42$.



Key

X product weight, g/m^2

Y reduction factor $f_{R,ID}$

Figure 8 — Interpolation of $f_{R,ID}$ from damage measurements on products from the same line but with different weights

8.4.4 Laboratory damage tests

It should be noted that ISO 10722 is intended as an index test for comparative purposes and should not be used for the derivation of reduction factors for geosynthetic soil reinforcements.

9 Weathering, chemical and biological degradation

9.1 General

Polymers are susceptible to environmental degradation due to weathering, including exposure to ultraviolet light, to chemical attack and to biological attack. All three effects are further influenced by temperature and, for some polymers, by moisture uptake. The durability of geosynthetic reinforcements is improved by their high degree of orientation and high molecular weights, while for polyolefins in particular the principal reason is the inclusion of special additives.

Environmental degradation can lead to degradation by Modes 1, 2 and 3. Weathering on site before a geotextile is covered can be regarded as Mode 1, while the weathering of geotextiles permanently exposed should be regarded as Mode 2. For chemical degradation, the preferred approach is to restrict the service lifetime to the period over which no significant reduction in strength is predicted. This is, however, not always possible, and for the hydrolysis of polyesters, which takes place continuously (Mode 2) a time-dependent reduction factor should be determined.

Two reduction factors are defined: $f_{R,W}$ for weathering and $f_{R,CH}$ for chemical and biological degradation. Allowances for statistical scatter and uncertainty are made by means of a separate factor of safety, f_s .

9.2 Data recommended for assessment

It is recommended that the following data be provided.

- Statement of principal polymers used.
- Evidence of the resistance of these polymers to weathering (for example EN 12224) and to chemical degradation, in particular to hydrolysis and oxidation in aqueous solutions with or without the presence of oxygen. For polyesters, a statement may be made of the number averaged molecular weight (M_n) and of the carboxyl end group count (CEG).
- A statement that post-consumer recycled material is not used.
- Predicted exposure to daylight: duration, location and season.
- Effective design soil temperature (see 4.4).
- Soil pH.
- A statement of any non-natural contaminants in the soil, e.g. industrial waste.
- Any unusual biological hazards such as termites.

9.3 Weathering

All polymers can degrade when exposed to ultraviolet light, although stabilizing additives will normally have been added to materials intended for outdoor use. In this Technical Specification, “weathering” will be taken as applying solely to the effects of ultraviolet light, either alone or together with temperature and water spray.

The recommendations for weathering are related to the duration of exposure during storage and on site. If the geosynthetic is exposed to ultraviolet light for a maximum of 12 h, no reduction factor need be applied. If the exposure time is longer, then the geosynthetic should undergo an accelerated weathering index test such as EN 12224. If the loss of strength is no greater than 5 % or is not statistically significant, no reduction factor is applicable. This is on condition that the installer covers the geosynthetic within one month.

Any geosynthetic reinforcement showing a greater loss of strength should not be exposed on site for longer than the duration shown in Table 1, and a reduction factor $f_{R,W}$ should be applied.

Table 1 — Installation exposure period

Retained strength after testing according to EN 12224	Maximum exposure time (uncovered) during installation	Reduction factor $f_{R,W}$
> 80 %	1 month ^a	Ratio of tensile strength of unexposed material to that of exposed material
60 % to 80 %	2 weeks	1,25
< 60 %	1 day	1,00
Untested material	1 day	1,00
^a Exposure of up to four months may be acceptable depending on the season and location.		

For a range of products identical except for mass per area, it is sufficient to subject only the product with the lowest mass per area to the test. The results of the test may be applied for the other products in the range, unless they have been tested separately.

If the geosynthetic is to be exposed to light for longer than one month, then it should be tested according to EN 12224 or a similar method for a duration such that extrapolation of the radiant exposure to that expected in service can be justified. The radiant exposure (ultraviolet radiation) in EN 12224 is 50 MJ/

m^2 , corresponding to approximately one summer month's exposure in southern European or central North American latitudes. The strength retained after the full radiant exposure should be predicted. $f_{R,W}$ should be set equal to the ratio of the strength of the unexposed material to that predicted for the exposed material.

9.4 Chemical degradation

9.4.1 Causes of chemical degradation

The principal causes of chemical degradation of polymeric geosynthetics in the soil are described in ISO/TS 13434. The following is a summary.

The principal cause of degradation of polyester geosynthetics (which consist of polyethylene terephthalate (PET)) is by hydrolysis. The rate of hydrolysis is slow at typical soil temperatures but increases rapidly as the temperature is raised. The rate can be less if the polyester is fully coated, but this is discounted since the coating may become damaged by the installation process in the ground. Since PET wicks moisture quite well, any exposure of the fibres due to coating damage could result in hydrolysis at the rate which would occur if the coating was not present. The rate of hydrolysis will be less if the soil is partially instead of fully saturated, but is not zero. Alkaline liquids with $\text{pH} \geq 9$ can, in addition, erode the surface. Polyester reinforced geosynthetics should not be used in natural or industrially polluted soils where $\text{pH} > 9$ is maintained unless proof of their durability can be provided.

The principal cause of degradation of polypropylene and polyethylene is oxidation, also resulting in chain scission, reduced molecular weight and strength loss. Other effects are embrittlement, surface cracking and a change in colour. Oxidation of these materials is a chain reaction whose chemistry is complex but quite well understood. The reaction may be started by ultraviolet light or by heat, and may be accelerated by catalysts such as ions of heavy metals, including iron. The resistance of these materials to oxidation is improved dramatically by the addition of a selection of antioxidant stabilizers which can extend the lifetime by hundreds or thousands of times. Ultimately, the antioxidant is consumed by oxidation, if it has not been lost prematurely by migration, evaporation or leaching. Assessment of the rate of oxidation is complex and is further described in [9.4.4](#).

Polyamides can degrade by either mechanism. Aliphatic polyamides such as PA 6 are susceptible to thermal degradation, oxidation, ultraviolet radiation, acid or alkali attack causing chain scission and by hydrolysis through contact with water at elevated temperatures. They are stabilized by copper salts, aromatic amines and hindered phenolic antioxidants which all act as heat stabilizers. Hindered phenol antioxidants are the most effective as they also resist thermoxidative degradation. Aromatic polyamides such as aramids are more resistant than PA 6 to degradation by oxidation, acids, alkalis and hydrolysis but are susceptible to ultraviolet degradation. Stabilization of these polyamides is effected by adding chlorine and nitro substituents into the recurring structural unit of the polymers.

Service lifetime prediction can be based on evidence from service or from accelerated testing. In some cases, there is sufficient experience to define index tests that will assure a certain minimum level of durability.

9.4.2 Evidence from service experience

The rate of degradation, or evidence for lack of degradation, can be based on results of analysis of specimens of the product exposed to a comparable environment and then exhumed, or of products with a similar physical structure and chemical formulation and including the same additives.

The plan of exhumation and testing should be in accordance with ISO 13437. The following additional points should be noted.

- The observation period should be of sufficient length for extrapolation to the full design life to be justified. This justification is particularly important when rapid degradation follows a long incubation period, or when degradation takes place in a series of separate stages (e.g. [9.4.4.3](#)). Service experience of at least 10 years may be necessary for extrapolation to a service life of 50 to 100 years for PET geosynthetics. Longer periods of time may be needed for polyolefins due to the

presence of antioxidants, as no loss in strength will be observed until the antioxidants are used up. Without knowing how long this will take, it is impossible to predict lifetime.

- Generally, a minimum of three retrievals are to be made (i.e., the first retrieval is taken right after installation, the second retrieval is taken at some time during the middle of this period, and the third retrieval is taken at the end of the study period).
- Enough specimens for each retrieval should be taken into account for statistical variability in the properties measured. For a more detailed description, see Reference [11].
- The polymer and physical characteristics of the exhumed material should meet the recommendations for “similar” products in 7.9.

An assessment and lifetime prediction should then be made on the basis of this evidence. For a more detailed description of the issues that should be considered when evaluating results from service experience, see References [2], [11] and [12] in the Bibliography.

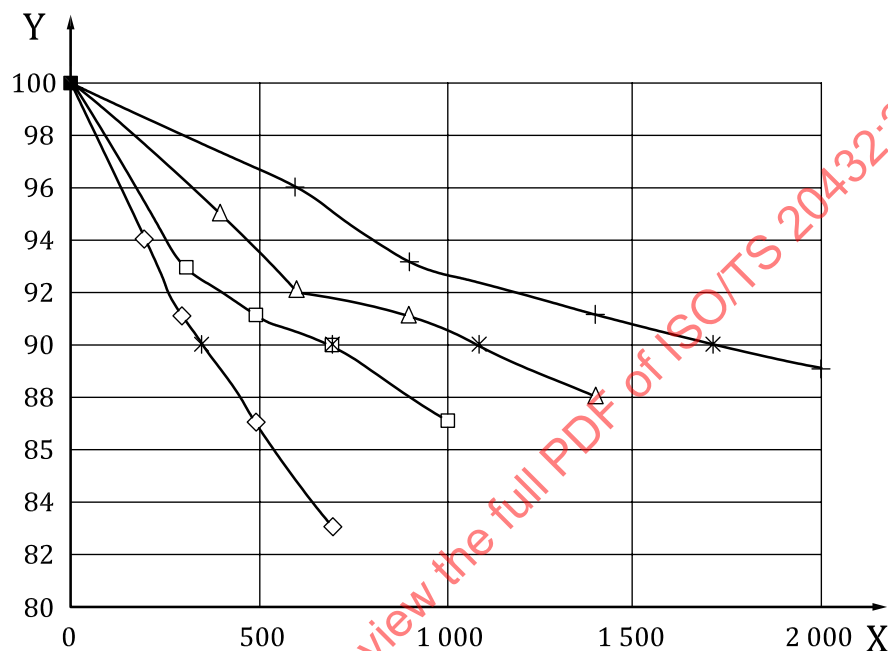
9.4.3 Accelerated chemical degradation tests

The rate of degradation can be estimated using accelerated testing, in which either temperature or chemical concentration, or both, is increased in order to accelerate the rate of reaction. The relation between the rate of degradation under service conditions to that during accelerated testing can be derived from chemical rate kinetics or from Arrhenius’ equation. Take care that the conditions during the accelerated tests are representative of those in service. There should be no change in the mechanism of degradation or in the physical structure of the material; and no barrier layers should form or be present that might retard the degradation process in a manner that does not occur in service.

In such a programme, carry out the following procedures.

- Select the parameter to be measured, for example a level of retained strength such as 90 %, 80 %, 70 %, 60 % or 50 %. If experience has shown that another physical or chemical quantity, such as CEG, gives more precise results, then this may be used instead, provided there is an established relationship between the parameter measured and the strength. It is important that the degradation can be observed and measured; if not, the degree of acceleration cannot be ascertained.
- Decide on the environment: pure water, air or a special chemical environment. If the geosynthetic is to be placed in a natural soil with $\text{pH} < 4$ or > 9 , or in a soil with non-natural contaminants, e.g. industrial waste, immersion tests should be performed in liquids with corresponding chemical composition and extrapolated to the corresponding design soil temperature, chemical composition and service life. ISO 12960 describes a method of immersion. In testing with alkaline solutions, care should be taken to reduce conversion of hydroxide to carbonate ions by reaction with atmospheric carbon dioxide.
- Select a range of at least three to four temperatures, spaced typically at 10 °C intervals. The lowest test temperature should ideally be not more than 25 °C above the service temperature, allowing for the fact that the test duration at this temperature has to lie within the time-scale of the test programme. This can extend for as long as four years. Caution is advised if any transition occurs in the physical state of the polymer or the mechanism of degradation less than 10 °C above the highest test temperature, or between the lowest test temperature and the service temperature. A glass transition occurs in the range of 50 °C to 80 °C in polyester and crystalline melting in high density polyethylene (HDPE) takes place at a range of temperatures, peaking at 128 °C. Furthermore, drawn polyethylene tends to lose its orientation at temperatures of around 70 °C. If a transition is present, then it should be demonstrated that it leads to no significant change in the rate of degradation, for example by confirming that the Arrhenius plot is a straight line.
- Measure the reduction in strength (or other parameter) over time at each of a range of temperatures. To do this, expose groups of samples over a range of times at each temperature. Include spare sets for exposure over longer times in case the rate of reduction in strength is less than predicted. Note that full wide-width specimens are preferred for this testing; however, single rib or yarn specimens can be used if necessary. Plot the retained strength against time and determine either the rate of

change or, by interpolation, the exact times to the desired retained strength (Figure 9). In Figure 9, the durations for 90 % retained strength are interpolated from the lines, noting that these are often irregular in shape. Examine each test sample for any change in the nature of degradation or of failure, for example, the growth of a barrier layer on the surface or circumferential cracking on the fibre surface, or increased ductility as evidenced by the geosynthetic modulus and peak strain at the higher temperatures. Scanning electron microscopy is a useful aid to this purpose. If a change is observed, only those results should be retained which are regarded as being representative of long-term degradation. If process of degradation comprises two or more separate stages, separate extrapolations should be made for each stage.

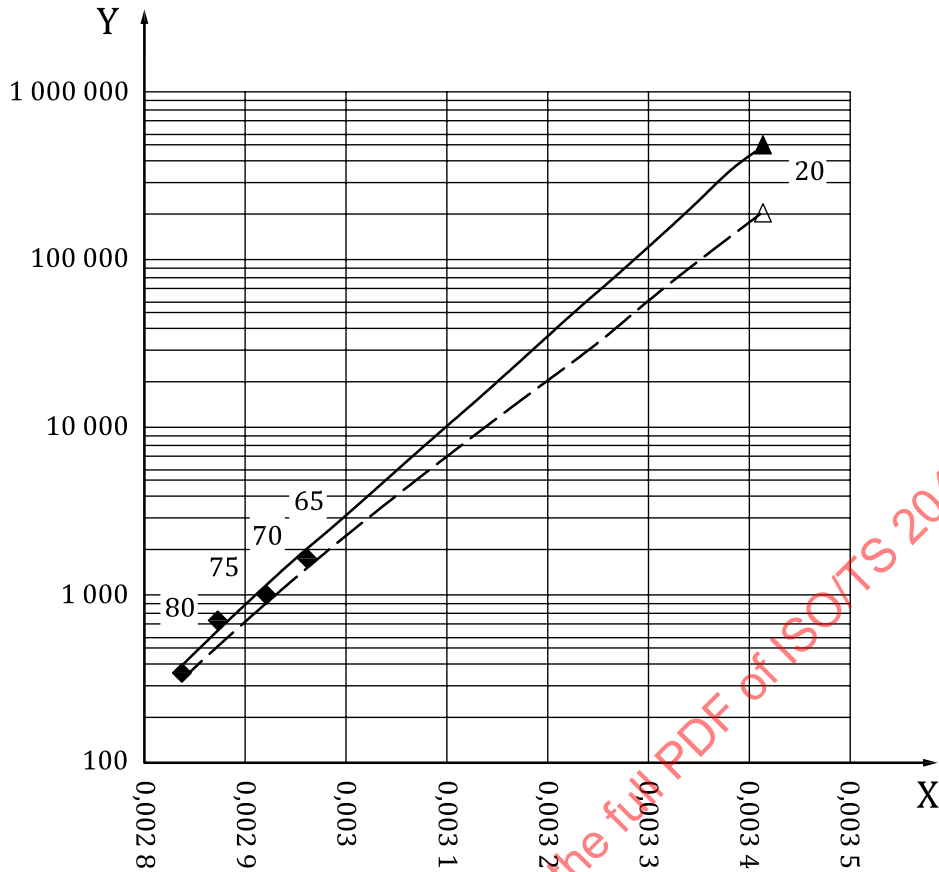


Key

X	time (h)
Y	% retained strength
◇	80 °C
□	75 °C
△	70 °C
+	65 °C
×	durations for 90 °C retained strength

Figure 9 — Reduction in strength at selected temperatures prior to application of Arrhenius' formula

- The number of specimens taken at each retrieval for testing may need to be greater than what is required for testing of the unaged material. This is because the degradation may lead to additional variability in the strength.
- Plot the times to a particular retained strength or other parameter against the inverse of the absolute temperature θ_K in K (see Figure 10). If Arrhenius' formula applies this plot should be a straight line. If it is not a straight line, then the order of the chemical reaction may be different (for the procedure, see Reference [13] in the Bibliography, or a transition may have occurred within the range of test temperatures selected as discussed above. If no straight line is obtained, then Arrhenius' formula does not apply and extrapolation is invalid.



Key

- X inverse absolute temperature (1/K)
- Y time to 90 % retained strength (h)
- ◆ measured time
- regression line
- - - lower confidence limit

Figure 10 — Arrhenius diagram

- Calculate the equation of the straight line, with $y = \log t_{90}$ and $x = 1/\theta_j$, as:

$$y = \bar{y} + b_a (x - \bar{x}) \quad (5)$$

where

$$b_a = S_{xy} / S_{xx};$$

$$S_{xx} = \sum (x - \bar{x})^2;$$

$$S_{yy} = \sum (y - \bar{y})^2;$$

$$S_{xy} = \sum (x - \bar{x})(y - \bar{y}).$$

- Calculate the lower confidence limit (LCL) of the line:

$$y = \bar{y} + b_a (x - \bar{x}) - t_{n-2} \sigma_0 \sqrt{1 + 1/n + (x - \bar{x})^2 / S_{xx}} \quad (6)$$

where

t_{n-2} is the Student's t for $n-2$ degrees of freedom and a stated probability;

n is the number of Arrhenius points;

σ_0 is the $\sqrt{[(S_{yy} - S_{xy}^2 / S_{xx}) / (n-2)]}$.

- Plot these lines as in [Figure 10](#); from the regression line read off the time t_s to the defined retained strength at the service temperature $x = 1/\theta_s$ (noting particularly if this exceeds 25 °C); from the lower confidence limit read off the time t_{LCL} ; in [Figure 10](#), these two values are 516 000 h and 199 000 h respectively. Such large differences are typical of logarithmic scales.
- Using the shape of the observed degradation curves as a guide, plot the shape of the degradation curve such that the defined retained strength is reached after time t_s ([Figure 11](#)). Read off the unfactored long-term strength per width T_x (expressed as a percentage of the batch tensile strength) after the design life t_D . $f_{R,CH} = 100/T_x$. Make a similar plot for t_{LCL} and derive the LCL T_{LCL} . The ratio $R_2 = T_x/T_{LCL}$. In [Figure 11](#), 90 % retained strength is reached at the predicted duration for the service temperature – in this example, 90 % after 516 000 h. The predicted strength after 1 000 000 h is 81,5 % and the $f_{R,CH} = 100/81,59 = 1,23$. A similar derivation is carried out for the LCL for which the predicted strength after 1 000 000 h is 58,8 %. The ratio $R_2 = 81,5/58,8 = 1,39$.

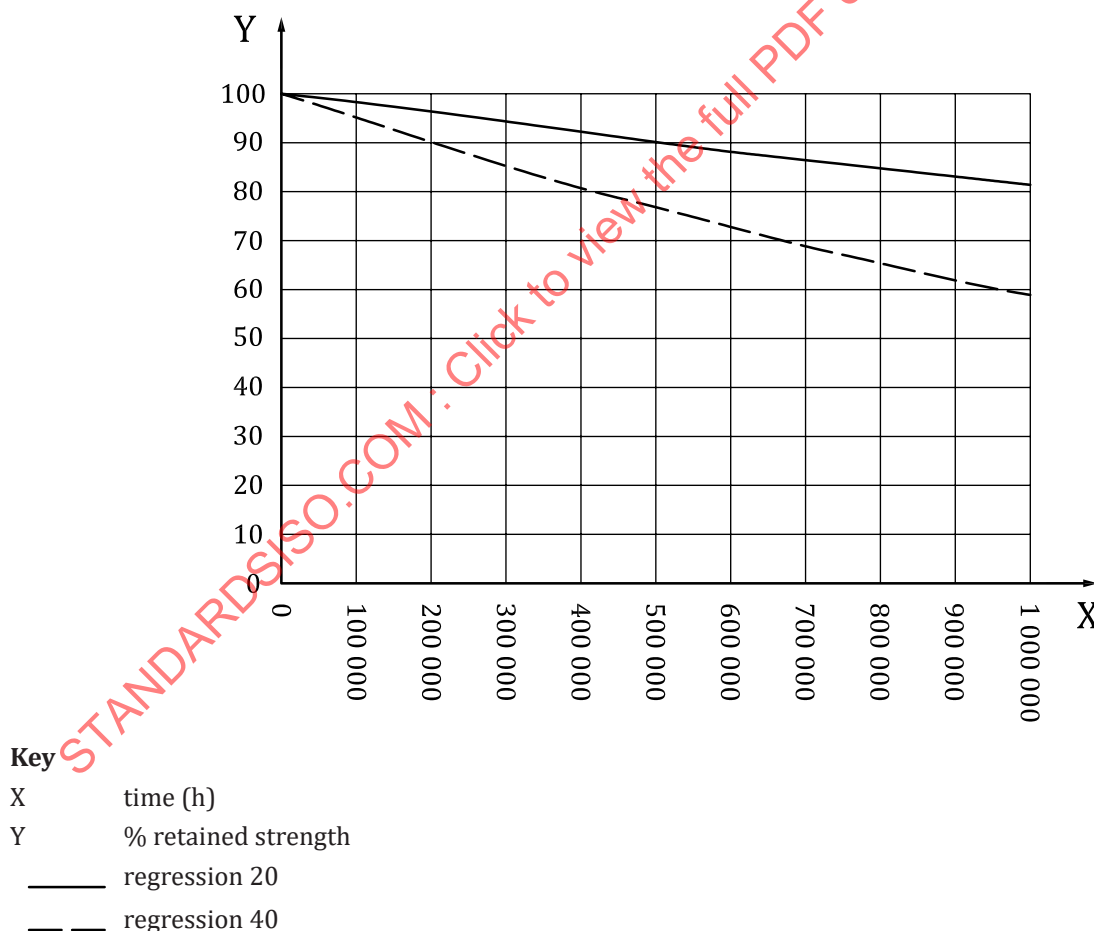


Figure 11 — Degradation curve at the service temperature

Further guidance is given in IEC 60216 and in References [2],[11],[13] and [17] in the Bibliography.

9.4.4 Oxidation of polyolefins

9.4.4.1 General

There are currently three approaches to the assessment of the oxidation resistance of polyolefins: simple Arrhenius testing, multiple Arrhenius testing and pressurized oxygen testing.

9.4.4.2 Simple Arrhenius testing

Historically, overall lifetime has been predicted by treating the entire oxidation process as a single stage. Time to end of life is measured at different temperatures and extrapolated to the service temperature to define the service life, or to the service life to define the operating temperature (e.g. see IEC 60216). The degradation of polypropylene fibre can be extremely sudden and thus give an indication of lifetime prediction. However, the methods have suffered from inaccuracies due to the following.

- The antioxidants present delay the onset of oxidation of the main polymer, causing the reaction rate with oxygen to change over time for the material.
- The oven temperature should stay well below the melting temperature of the polymer, thus restricting the exposure temperatures to a narrow range.
- The mechanism of oxidation can change at higher temperatures, invalidating the extrapolation.
- The different rates of degradation described above can lead to large errors in the measurement of time to end of life and of its extrapolation.
- Surface cracking has been observed in certain grades of polypropylene. This increases the access of oxygen to the polymer and invalidates any prediction based on uncracked material; furthermore, at elevated temperatures, these cracks can heal, which will possibly occur at lower temperatures.
- Diffusion of antioxidants plays a major part at all stages of oxidation. The rate of diffusion of oxygen from the outside, the rate of diffusion of antioxidants and the rate of migration of radicals produced by the chain reaction all increase at higher temperatures and decrease with crystallinity and orientation of the polymer. These effects are accelerated by a high surface-to-volume ratio. Hence, a high surface-to-volume ratio and a low degree of orientation will clearly shorten all stages of oxidation.
- Leaching may also occur in materials having a high surface-to-volume ratio or containing leach-sensitive antioxidants. For these materials, correct selection of stabilizers is essential.

Polyethylene and polypropylene geotextiles cover a wide range of structures from fine, highly oriented fibres to thick and less strongly oriented polymeric geosynthetic barriers. They contain different combinations of antioxidants. Some exhibit surface cracking. In polymeric geosynthetic barriers and the less oriented areas of extruded geogrids, the rate of oxidation should be higher due to the lack of orientation, but simultaneously lower due to the small surface-to-volume ratio.

This explains why it has proved impossible to define a single oven ageing test as a screening test for all geosynthetics. Various attempts to do so have either failed to eliminate poorly stabilized material or conversely have eliminated material which would be expected to be durable. Better results can be obtained by restricting the temperature to 80 °C or below, by dividing the process into stages of oxidation or by raising the oxygen pressure as in ISO 13438.

9.4.4.3 Multiple Arrhenius testing

The degradation of a stabilized polypropylene or polyethylene can be divided into two stages: consumption of the antioxidant, and degradation of the unprotected polymer. To establish which stage has been reached, use is made of oxidation induction time (OIT) measurement or high pressure oxidation induction time (HPOIT) where hindered amine light stabilizers (HALS) are present. OIT cannot be applied universally because it only relates to antioxidants active at the testing temperature, which is in the molten state. In these methods, a sample of material is raised to a high temperature in an