

SLOVENSKI STANDARD SIST ISO 8466-1:1996

01-avgust-1996

Kakovost vode - Umerjanje in vrednotenje analitskih metod in določitev delovnih karakteristik - 1. del: Statistično vrednotenje linearne umerjalne funkcije

Water quality -- Calibration and evaluation of analytical methods and estimation of performance characteristics -- Part 1: Statistical evaluation of the linear calibration function

iTeh STANDARD PREVIEW

Qualité de l'eau -- Étalonnage et évaluation des méthodes d'analyse et estimation des caractères de performance -- Partie 1: Évaluation statistique de la fonction linéaire d'étalonnage https://standards.iteh.ai/catalog/standards/sist/caca8d32-fdb2-426b-a3d9-3570ad2c1602/sist-iso-8466-1-1996

Ta slovenski standard je istoveten z: ISO 8466-1:1990

ICS:

13.060.50 Preiskava vode na kemične Examination of water for

chemical substances snovi

SIST ISO 8466-1:1996 en SIST ISO 8466-1:1996

iTeh STANDARD PREVIEW (standards.iteh.ai)

<u>SIST ISO 8466-1:1996</u> https://standards.iteh.ai/catalog/standards/sist/caca8d32-fdb2-426b-a3d9-3570ad2c1602/sist-iso-8466-1-1996 SIST ISO 8466-1:1996

INTERNATIONAL STANDARD

ISO 8466-1

> First edition 1990-03-01

Water quality — Calibration and evaluation of analytical methods and estimation of performance characteristics

iTeh SPart 1: DARD PREVIEW
Statistical evaluation of the linear calibration function (standards.iteh.ai)

Qualité de l'eau 84 Étalonnage et évaluation des méthodes d'analyse et estimation https://standards.itdes.caractères.de.performance.d32-fdb2-426b-a3d9-

Partie 1: Évaluation statistique de la fonction linéaire d'étalonnage

 $IMI \vdash \bowtie MMM \vdash IMMM$

150



Reference number ISO 8466-1: 1990 (E)

C	Contents	Page
	Foreword	iii
1	Scope	1
2	Definitions	1
3	Symbols	2
4	Performance	3
	4.1 Choice of working range	
	4.2 Calibration and characteristics of the method ANDARD	PREVIEW
	4.3 Assessment (standards.ite	eh.aš)
5	Example	₉₆ . 6
	5.1 Choice of working range https://standards.iteh.ai/catalog/standards/sist/ca	aca8d32 <mark>6</mark> fdb2-426b-a3d9
	5.2 Calibration and characteristics of the method	
	5.3 Evaluation	7
Α	nnex A Bibliography	8

© ISO 1990

All rights reserved. No part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from the publisher.

International Organization for Standardization
Case postale 56 • CH-1211 Genève 20 • Switzerland

Printed in Switzerland

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.

Draft International Standards adopted by the technical committees are circulated to the member bodies for approval before their acceptance as International Standards by the ISO Council. They are approved in accordance with ISO procedures requiring at least 75 % approval by the member bodies voting.

(standards.iteh.ai)

International Standard ISO 8466-1 was prepared by Technical Committee ISO/TC 147, Water quality. 8466-1:1996

https://standards.it

teh ai/catalog/standards/sist/caca8d32-fdb2-426b-a3d9-ISO 8466 consists of the following parts, under the general title Water quality — Calibration and evaluation of analytical methods and estimation of performance characteristics:

- Part 1: Statistical evaluation of the linear calibration function
- Part 2: Calibration strategy for non-linear calibration functions
- Part 3: Method of standard addition
- Part 4: Estimation of limit of detection and limit of determination of an analytical basis method.

Annex A of this part ISO 8466 is for information only.

SIST ISO 8466-1:1996

iTeh STANDARD PREVIEW

This page intentionally left blank

SIST ISO 8466-1:1996 https://standards.iteh.ai/catalog/standards/sist/caca8d32-fdb2-426b-a3d9-3570ad2c1602/sist-iso-8466-1-1996

Water quality — Calibration and evaluation of analytical methods and estimation of performance characteristics

Part 1: Statistical evaluation of the linear calibration function

1 Scope

This part of ISO 8466 describes the steps to be taken in evaluating the statistical characteristics of the linear calibration function. It is applicable to methods requiring a calibration. Further parts of this International Standard will cover the determination of limit of detection and limit of determination, the effect of interferences and other performance characteristics.

It is intended especially for the evaluation of the pure analytical method and for the calculation of performance characteristics siteh ai of the calibration function.

In order to derive comparable analytical results and as a basis \$466-1:1996 for analytical quality control the calibration and evaluation order devaluation and evaluation order devaluation and evaluation order devaluation and evaluation order devaluation of the calibration and evaluation of the calibration of the calibra

2 Definitions

For the purposes of this part of ISO 8466, the following definitions apply.

2.1 analytical method: An analytical method is composed of procedural, measuring, calibrating and evaluating instructions (see figure 1).

Whereas the procedural and measuring instructions depend on the method, and are therefore the object of standardization of the respective method, the calibrating and evaluating instructions are valid for any analytical method requiring calibration.

2.2 calibrating instruction: Describes the approach to determine the calibration function from information values, y_i , obtained by measuring given standard concentrations, x_i . The slope of the calibration function, b, as a measure of sensitivity of the analytical method and the standard deviation of the method, s_{xo} , are figures of merit and characteristics which result from the calibration experiment.

The standard deviation, s_{xo} , allows the comparison of independent analytical methods.

For the user of the method, these characteristics present criteria for the internal laboratory quality control.

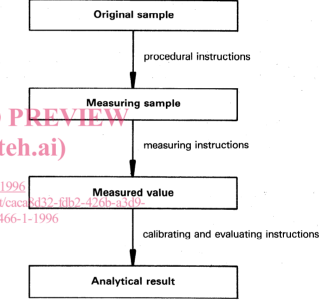


Figure 1 — The analytical method

- **2.3 evaluating instruction:** A calculation guide for the computation of concentrations from the measured values by the use of the calibration function. Additionally, the confidence range permits an objective assessment of the imprecision of the analytical result^[2].
- **2.4** measured values: The concentration-dependent initial values (e.g. extinction) of a measuring system.

NOTE — Information value and measured volume are synonymous.

2.5 residual standard deviation, s_y : The residual standard deviation describes the scatter of the information values about the calculated regression line. It is a figure of merit, describing the precision of the calibration.

For the purpose of this standard, the standard deviation of the method means the standard of deviation of the calibration procedure.

the residual	lard deviation of the method s_{xo} : The ratio of standard deviation, s_y , to the sensitivity of the function, b . It is a figure of merit for the perfor-	$\hat{y_i}$	Information value of the standard concentration x_i calculated from the calibration function			
	e analytical method, and is valid within the working equation 13).	s_i^2	Variance of the information values for the analyses of standard samples, having the con-			
For the purp	oose of this standard, the standard deviation of the		centration x_i .			
	ans the standard of deviation of the calibration pro-	f_i	Degrees of freedom for the calculation of the variance $(f_i = n_i - 1)$.			
of the stand	icient of variation of the method, V_{xo} : The ratio ard deviation of the method s_{xo} to the appertaining nich is the centre of the working range.	a	Calculated blank (ordinate intercept of the calibration straight line).			
See also no	te to 2.5 and 2.6.	b	Sensitivity of the method (slope of the calibration line; coefficient of regression).			
being experi calibration	ing range (of an analytical method): The interval, imentally established and statistically proved by the of the method, between the lowest and highest mass concentration. The lowest possible limit of a	\overline{x}	Mean of the standard concentrations x_i , resulting from the calibration experiment.			
	ge is the limit of detection of an analytical method.	\overline{y}	Mean of the information values y_i , resulting from the calibration experiment.			
of pooled da	egeneity of variances: Homogeneity of variances ata, such as those resulting from replicate analyses	s_y	Residual standard deviation.			
	levels, is confirmed if these variances are not correlated to their appertaining concentrations.	s_{y1}	Residual standard deviation obtained by linear regression calculation.			
2.10 sens	sitivity of the analytical Teh, STAND	RD PR	EVIEW			
the calibrati	sitivity of the analytical method: The slope of an information of the complete analytical method, in- all procedural steps, within the working range in all	ds.iteh.a	Residual standard deviation obtained by non- linear regression calculation.			
question.	CICT ICO	DS ²	Difference of variances.			
	5151 150	8466-1:1996				
	suring sample (reaction sample): A sample which		22Information value of an analysed sample.			
can be direc	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the dete					
can be direct nand. A me required reat ing and anal	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the dete	da y ds/sist/caca8d3				
can be direct nand. A me required reat ing and anal	suring sample (reaction sample): A sample which city submitted to the measurement of the determination assuring sample is normally obtained by adding the 02/s gents to the analytical sample. Obviously, measur-	da y ds/sist/caca8d3 ist-iso-8466-1-19	Number of replicates on the same analysed			
can be direct nand. A me required reat ing and anal	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the dete	dayds/sist/caca8d2 ist-iso-8466-1-19 n $\hat{\overline{y}}$	Number of replicates on the same analysed sample. Mean of information values, resulting from <i>n</i> replicates.			
can be direct nand. A me required reating and anal added to th	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the dete	dayds/sist/caca8d2 ist-iso-8466-1-19 n	Number of replicates on the same analysed sample. Mean of information values, resulting from n			
can be direct nand. A me required realing and analadded to the	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the dete	dayds/sist/caca8d2 ist-iso-8466-1-19 n $\hat{\overline{y}}$	Number of replicates on the same analysed sample. Mean of information values, resulting from <i>n</i> replicates. Concentration of the analytical sample, calculated from the information value <i>y</i> . Concentration of the analytical sample, calculated from the mean of the information			
can be direct nand. A merequired realing and analaded to the symbol of t	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the dete	dayds/sist/caca8d3 ist-iso-8466-1-19 \hat{y} \hat{x}	Number of replicates on the same analysed sample. Mean of information values, resulting from n replicates. Concentration of the analytical sample, calculated from the information value y . Concentration of the analytical sample, calculated from the mean of the information values \widehat{y} . Tabled value of the t -distribution with f_1			
can be direct nand. A merequired realing and analaded to the symbol symb	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the analytical sample. Obviously, measurelytical sample are identical if no reagents have to be analytical sample. Ols Concentration of the <i>i</i> th standard sample. Subscript of the concentration levels, where <i>i</i> = 1, 2,, <i>N</i> . Number of concentration levels (for this part of	dayds/sist/caca8d3 ist-iso-8466-1-19 \hat{y} \hat{x}	Number of replicates on the same analysed sample. Mean of information values, resulting from n replicates. Concentration of the analytical sample, calculated from the information value y . Concentration of the analytical sample, calculated from the mean of the information values $\widehat{\hat{y}}$.			
can be direct nand. A merequired realing and analaded to the symbol symb	suring sample (reaction sample). A sample which only submitted to the measurement of the determination of the determination of the determination of the analytical sample. Obviously, measurelytical sample are identical if no reagents have to be analytical sample. Concentration of the i^{th} standard sample. Subscript of the concentration levels, where $i = 1, 2,, N$. Number of concentration levels (for this part of ISO 8466, $N = 10$). Concentration of the standard sample at the lower	dayds/sist/caca8d3 ist-iso-8466-1-19 \hat{y} \hat{x}	Number of replicates on the same analysed sample. Mean of information values, resulting from n replicates. Concentration of the analytical sample, calculated from the information value y . Concentration of the analytical sample, calculated from the mean of the information values \widehat{y} . Tabled value of the t -distribution with $f_1 = N - 2$ degrees of freedom and a confidence level of $(1 - \alpha)$ (t -factor of Student's distribution). Tabled value of the F -distribution (Fisher-Snedecor) with f_1 and f_2 degrees of freedom			
can be direct nand. A merequired realing and analadded to the state of	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the analytical sample. Obviously, measuring sample are identical if no reagents have to be analytical sample. Concentration of the i^{th} standard sample. Subscript of the concentration levels, where $i = 1, 2,, N$. Number of concentration levels (for this part of ISO 8466, $N = 10$). Concentration of the standard sample at the lower level of the working range (1st standard sample). Concentration of the standard sample at the upper level of the working range (10th standard	dayds/sist/caca8d3 ist-iso-8466-1-19 \hat{y} \hat{x} \hat{x} $t(f_1, 1-\alpha)$	Number of replicates on the same analysed sample. Mean of information values, resulting from n replicates. Concentration of the analytical sample, calculated from the information value y . Concentration of the analytical sample, calculated from the mean of the information values \widehat{y} . Tabled value of the t -distribution with $f_1 = N - 2$ degrees of freedom and a confidence level of $(1 - \alpha)$ (t -factor of Student's distribution). Tabled value of the F -distribution (Fisher-			
can be direct nand. A merequired realing and analadded to the state of	suring sample (reaction sample): A sample which city submitted to the measurement of the determination of the determination of the determination of the determination of the analytical sample. Obviously, measurelytical sample are identical if no reagents have to be analytical sample. Concentration of the i^{th} standard sample. Subscript of the concentration levels, where $i=1,2,,N$. Number of concentration levels (for this part of ISO 8466, $N=10$). Concentration of the standard sample at the lower level of the working range (1st standard sample). Concentration of the standard sample at the upper level of the working range (10th standard sample).	dayds/sist/caca8d3 ist-iso-8466-1-19 \hat{y} \hat{x} \hat{x} $t(f_1, 1-\alpha)$	Number of replicates on the same analysed sample. Mean of information values, resulting from n replicates. Concentration of the analytical sample, calculated from the information value y . Concentration of the analytical sample, calculated from the mean of the information values \widehat{y} . Tabled value of the t -distribution with $f_1 = N - 2$ degrees of freedom and a confidence level of $(1 - \alpha)$ (t -factor of Student's distribution). Tabled value of the F -distribution (Fisher-Snedecor) with f_1 and f_2 degrees of freedom and a confidence level of $(1 - \alpha)$.			
can be direct nand. A merequired realing and analadded to the state of	suring sample (reaction sample): A sample which city submitted to the measurement of the determination standard sample is normally obtained by adding the gents to the analytical sample. Obviously, measurifytical sample are identical if no reagents have to be analytical sample. Concentration of the i^{th} standard sample. Subscript of the concentration levels, where $i = 1, 2,, N$. Number of concentration levels (for this part of ISO 8466, $N = 10$). Concentration of the standard sample at the lower level of the working range (1st standard sample). Concentration of the standard sample at the upper level of the working range (10th standard sample). j^{th} information value for the concentration x_i . Subscript of the replicate j of level i , where $j = 1$,	dayds/sist/caca8d3 ist-iso-8466-1-19 \hat{y} \hat{x} \hat{x} $t(f_1, 1-\alpha)$ $F(f_1, f_2, 1-\alpha)$	Number of replicates on the same analysed sample. Mean of information values, resulting from n replicates. Concentration of the analytical sample, calculated from the information value y . Concentration of the analytical sample, calculated from the mean of the information values \hat{y} . Tabled value of the t -distribution with $f_1 = N - 2$ degrees of freedom and a confidence level of $(1 - \alpha)$ (t -factor of Student's distribution). Tabled value of the F -distribution (Fisher-Snedecor) with f_1 and f_2 degrees of freedom and a confidence level of $(1 - \alpha)$. Standard deviation of the method.			

Performance

Choice of working range

Each calibration experiment is started with the choice of a preliminary working range^[3].

The working range depends on

a) the practice-related objective of the calibration.

The working range shall cover, as far as possible, the application range for water, waste water, and sludge analysis. The most frequently expected sample concentration should lie in the centre of the working range.

feasibilities of technical realizability.

The measured values obtained must be linearly correlated to the concentrations. This requires that the measured values obtained near the lower limit of the working range can be distinguished from the blanks of the method. The lower limit of the working range should therefore be equal to or greater than the limit of detection of the method. Dilution and concentrating steps should be feasible without the risk of bias.

4.1.2 Test for homogeneity of the variances

Both data sets of the concentrations x_1 and x_{10} are used to calculate the variances s_1^2 and s_2^2 , as given in equation (1):

$$s_i^2 = \frac{\sum_{j=1}^{10} (y_{i,j} - \overline{y_i})^2}{n_i - 1} \qquad \qquad \dots$$
 (1)

with the mean

$$\overline{y}_{i} = \frac{\sum_{j=1}^{10} y_{i,j}}{n_{i}} \quad \text{for } i = 1 \text{ or } i = 10 \qquad ... (2)$$

The variances are tested (F-test) for significant differences at the limits of the working range^[5, 6].

The test value PG is determined for the F-test from equation (3)

$$PG = \frac{s_{10}^2}{s_2^2} \quad \text{for } s_{10}^2 > s_1^2 \qquad \qquad . . . (3)$$

c) the variance of the information values must be independent of the concentration. (standards.iteh. s_{10}^2) for $s_1^2 > s_{10}^2$

The independence is verified by a statistical test on the linearity[6, 8]. **SIST ISO 8466-**

PG is compared with the tabled values of the *F*-distribution^[5].

variances s_1^2 and s_2^2 is not significant.

https://standards.iteh.ai/catalog/standards/sipt/casa8d32-fdb2-426b-a3d9-

4.1.1 Preparation of the calibration

After establishing the preliminary working range, measured values of at least five (recommended N = 10) standard samples are determined. The concentrations, x_{ii} of these standard samples shall be distributed equidistantly over the working range. In order to check for the homogeneity of the variances, ten replicates of each of the lowest and the highest concentrations $(x_1 \text{ and } x_{10})$ of the working range are determined. Ten information values, $y_{i,j}$ result from these series of measurements (see table 1).

3570ad2c1602/sist-iso-8466-1-1996 a) If PG $\stackrel{<}{<} F_{f_1;\ f_2;\ 0,99}$ the difference between the

> b) If PG > $F_{f_1;\ f_2;\ 0,99}$ the difference between the variances s_1^2 and s_2^2 is significant.

If the difference between the variances is significant, the preliminary working range should be made smaller until the difference between the variances is found to be random only.

Table 1 — Data sheet for the calibration

i	x_i	<i>y</i> _{i,1}	<i>y</i> _{i,2}	y _{i,3}	y _{i,4}	<i>y</i> _{i,5}	y _{i,6}	.v _{i,7}	y _{i,8}	<i>y</i> _{i,9}	y _{i,10}
1											
2										_	
3											
4											
5											
6											
7											
8											
- 9											
10											