# INTERNATIONAL STANDARD

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# Validation methods for fire gas analyses —

# Part 2: Intralaboratory validation of quantification methods

Méthode de validation des analyses de gaz d'incendie — Partie 2: Validation intralaboratoire des méthode de d'analyse





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#### Foreword Foreword

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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).

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For an explanation on 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 the following URL: www.iso.org/iso/foreword.html.

The committee responsible for this document is ISO/TC 92, Fire safety, Subcommittee SC 3, Fire threat to people and the environment.

A list of all parts in the ISO 12828 series can be found on the ISO website.

#### **Introduction** <u>----- - -- -- - -- - --</u>

The reduction of human tenability from fire effluent has long been recognized as a major cause of injury and death in fire. The composition and concentration of the effluent from a large fire are also clearly key factors in determining the potential for harm to the environment. The harmful components of fire effluent can be determined from both large-and small-scale tests of materials and finished products. Equations have been developed for quantifying the effects of the effluent components, for example, to estimate the available safe egress time (ASET). Related documents are also being developed in ISO TC92 SC3 which deal with environmental threats from fire effluent.

These advances in fire science and fire safety engineering have led to an increasing demand for quantitative measurements of the chemical components of the fire effluent. Characterizing these measurements is a key factor in evaluating the quality of the quantitative data produced. Such a characterization is developed over four items.

Item 1: Define the objective of the analysis. Before undertaking a chemical analysis of fire effluent, the final objective of the analysis should be established. For example, the objective might be part of a fire safety engineering design of a building, validation of a numerical fire model, or determination of the toxic potency of the effluent from a particular combus tible item.

Item 2: Determine the degree of accuracy and precision required from the analysis. Accuracy is dependent on a combination of the physical fire model being used, the sampling of the effluent and the analytical chemical technique. Precision means the tolerable uncertainty in the measured result. For example, in an FED (Fractional Effective Dose) calculation, where the individual contribution of a range of different species to the overall toxic potency of a fire effluent is estimated, interest might range from concentrations which might incapacitate people of average sensitivity to the effluent, to concentrations which show negligible toxic effect over a long exposure period.

Item 3: Select the appropriate chemical analytical methods, considering specificity, i.e. the other gases present. Guidance on options for measuring a wide variety of chemical species is provided in ISO 19701 and ISO 19702.

Item 4: Evaluate the suitability of the chosen method considering specificity. For chemical analyses, as with any other measurement, it is important to evaluate a specific methodology for its ability to provide appropriate, sufficient, and adequate data for a particular application. This evaluation normally has to consider a range of factors, including repeatability, reproducibility, and a measurement of uncertainty, especially for laboratories working under ISO 17025 rules. For fire effluent toxicity, these properties are discussed in ISO 19706.

Different methods may be deemed suitable for the particular application and for consistency in the interpretation of results from these different methods, it is also important to be able to compare the validity of the analytical technique used. In the field of fire effluents, many factors can affect the trueness and the fidelity of a measurement technique.

# Validation methods for fire gas analyses —

# Part 2: Intralaboratory validation of quantification methods

# 1 Scope

This document describes tools and techniques for use in validating the analysis of fire gases when an analytical method is developed in a laboratory. It complements ISO 12828-1, which deals with limits of quantification and detection.

The tools and techniques described can be applied to the measurement of quantities, concentrations (molar and mass), volume fractions, and concentration or volume fraction versus time analyses. Fire effluents are often a complex matrix of chemical species, strongly dependent on the materials involved in the fire, but also dependent on fire scenario parameters (see ISO 19706). With such a wide variety of conditions, the analytical techniques available will differ in terms of the influence of the matrix on the methods and on the concentration ranges which can be measured. The analytical techniques available are likely to differ significantly in several respects, such as their sensitivity to the matrix and the range of concentrations/volume fractions which can be reliably measured. For these reasons, a unique reference analytical technique for every fire effluent of interest is, in practical terms, difficult or impossible to achieve. The tools in this document allow verification of the reliable measurement ranges and conditions for the analysis of fire effluents, thereby enabling a comparison among various analytical techniques.

Examples of existing International Standards where the information contained in this document can be used are the analytical chemical methods in ISO 19701, ISO 19702, ISO 5660-1, and the chemical measurements in the methods discussed in ISO/TR 16312-2, ISO 16405, or their application to fire toxicity assessment using ISO 13571 and ISO 13344.

The variable "concentration" is used throughout this document, but it can be replaced in all places NOTE<sub>1</sub> with "volume fraction" without altering the meaning. This does not apply to the Annexes.

NOTE 2 Concentration can be calculated from volume fraction by multiplying by the density of the relevant gas at the relevant temperature and pressure.

#### **Normative references**  $\overline{2}$ ========================

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 12828-1:2011, Validation method for fire gas analysis  $-$  Part 1: Limits of detection and quantification

ISO 5479 , Statistical interpretation of data — Tests for departure from the normal distribution

#### **Terms and definitions** 3 3 Terms and definitions and definitions

For the purposes of this document, the terms and definitions given in ISO 13943, ISO 5725-1, ISO 2854, ISO 2602, ISO 13571 and the following apply.

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

— IEC Electropedia: available at http://www.electropedia.org/

— ISO Online browsing platform: available at https://www.iso.org/obp/

#### 3 .1

#### matrix (of fire effluents)

mixture of fire effluents in which the analyte of interest is present

Note 1 to entry: This includes all other species, solid, liquid and gas phases. It constitutes all components that could affect analysis, such as interfering species.

# 4 Symbols and abbreviated terms



#### **General considerations** 5 5 General considerations

# 5 .1 Actual concentration and measured concentration

The objective of every chemical analysis used in fire science is to approach the actual concentration of an and a ly  $\alpha$  , you is the variance of  $\alpha$  is under a lue of  $\alpha$  is the one concentration is the concentration of  $\alpha$ 

y<sup>m</sup>. The concentration y<sup>m</sup> is affec ted by the measurement trueness and prec is ion (uncerta inty) of the chosen analytical technique

The discussion of the discussion you like the s ing interesting ing is done to the measurement. technique chosen. For fire gas analyses, there could be many alternative analytical techniques available, (see ISO 19701 and ISO 19702 for examples). Stages of the analytical procedure which could affect the measurement are sampling (e.g. probe design and temperature), transportation (e.g. size, length and temperature of sampling lines), conditioning of sample (e.g. filtration, drying), and the analysis efficiency. This last factor could be integrated in the trueness of the analytical technique. The different steps of this analytical process of fire effluents and the associated efficiencies are presented in Figure 1.



Figure 1 — Measurement ratios

#### $5.2$ 5 .2 Selection of analytical methods with respect to the physical fire model used

The selection of a physical fire model has an influence on the composition of the effluent, the concentration of individual components in the effluent and variations of effluent concentration with time. These parameters imply that the choice of an analytical method for fire effluents will depend on the physical fire model that produced the effluent. An analytical method validated by using a given physical fire model may therefore be of limited use with another physical fire model. See ISO 19706 and ISO 16312-1 for further details on the selection of physical fire models.

# 5 .3 Validation of analytical techniques

Fire effluent from accidental fires is typically very specific matrix, characterized by a constantly changing and very wide range of chemical species and their concentrations. Some analytical techniques commonly used for combustion gas analysis are not suitable in the case of accidental fires. The selection of a technique with a wrong selectivity for example could lead to erroneous conclusions in a safety assessment. For example, the measurement of incinerator stack composition using solid-state detection techniques would be too limited in selectivity for use in a fire atmosphere safety assessment.

The conditions under which the analytical method is used in practice shall not differ from the conditions used to validate the method. This document proposes different steps to be followed and different techniques from those used in combustion gas analysis in order to validate that an analytical technique could be applied specifically and meaningfully to fire effluents. The validation is therefore limited to the specific nature of a matrix and range of concentrations within the matrix.

Due to the variety of physical and chemical principles used in the analysis of fire effluents (see ISO 19701 and ISO 19702), the technique and its range of application shall be rigorously defined and selected. Figure 2 illustrates the different steps required to validate an analytical technique. Figure 3 illus trates the different steps required to compare two analytical techniques.



Figure 2 — Steps in validating an analytical technique



Figure  $3$   $-$  Steps in comparing two analytical techniques

# 6 Sampling and measurement effectiveness

# 6 .1 General considerations

Fire gases are a complex mixture of water, reactive/corrosive species, condensable species, aerosols, hygroscopic components and are usually in the presence of solid particles which may adsorb or absorb gases to a varying degree. The gases may be at temperatures between ambient and over 1 000  $\degree$ C at the sampling point. This makes the sampling and analysis of fire effluents generally a difficult process requiring much attention to best practice procedures (e.g. as provided in ISO 19701).

The analysis can be performed in situ or with an extractive sampling technique. Quantification can be time-resolved or cumulative, depending on the end-use requirements for the data.

For the purposes of quantification, it should be recognized that in such a mixture, there is much scope for losses from a variety of causes. In any validation process of an analytical method, great care should be exercised to ensure that these losses are properly taken into account especially where there is a chemical or physical modification of the analyte between sampling point and analysis point. This is of particular importance with extractive sampling methods.  $[9]$ 

### 6.2 Sampling probe

Sampling of fire effluents prior to their analysis should be carefully considered to ensure a representative sample is ultimately delivered to the analyser. The first part of a typical sampling system is the sampling probe, positioned in the effluent. The design of a sampling probe for fire gas analysis should allow the required portion of the effluent to be passed on to the sampling line. The probe may be a simple openended tube where the effluent stream is homogenous or may require holes along its length to allow for non-homogeneity. The location and diameter of the holes are supposed to be designed so that the sampling is indicative of the full effluent flow. Since the temperature, density and mixing of the flow can vary during a test, the assumption of representative sampling has limitations.

In general, to limit flow disturbances, the sampling flow rate shall be low in comparison with the effluent flow rate, and shall limit added turbulence. Some bench-scale systems require a complete sampling of the effluent.

Ideally, the sampling point will be in a known position with respect to the fire source in a location where temperature conditions are measured and where the effluent flow is as homogenous and representative as possible. Clearly, these conditions will sometimes not be met. In some cases, the sampling probe may have to be heated to avoid or reduce condensation. It could also be designed to limit soot particulate deposits (e.g. by incorporating a microcyclone device). Where an extractive method is used for aerosol sampling, isokinetic techniques shall be used (i.e. with the sampling velocity made equivalent to the aerosol flow velocity, see ISO 29904).

However, all these systems will have a limited efficiency, and technical choices are made in order to have X<sup>1</sup> close to 1 .

### 6.3 Transportation of effluent from sampling probe to analysis system

Between sampling point and analysis point, effluent may be transported along a sampling line, trapped in a gas bag or passed through trapping solutions or solid adsorbates. The materials in contact with the sample should be carefully chosen to reduce losses, for example, through chemical reactivity, and the temperature conditions in the sampling line should be carefully chosen to avoid losses through condensation and/or further chemical reaction. The flow velocity shall be as high as possible (consistent with an extraction rate which will not disturb the effluent stream) to minimize losses due to adsorption on surfaces. on surfaces .

For example, some species such as HBr have an important tendency to be physically trapped and re leased by surfaces of sampling lines. This affects the kinetics of the analysis, and can result in prolonged delivery of the species to be analysed with a consequent spreading of detector response. Materials such as stainless steel, epoxy-lined stainless steel, glass (not where HF is present) or PTFE are often used.

With sampling line temperatures, the main factors to consider are the temperature of the gas itself and the temperature of the sampling line surfaces. Temperature is often chosen high enough to limit condensation of water, but also of other condensable species such as formaldehyde. Nevertheless, a too high temperature will affect the composition of the transported gas, as it is a reactive mixture, and increases in temperature will accelerate many reactions. A range from 150 °C to 200 °C has been found suitable for the large majority of extractive gas analysis methods used with fire effluents (ISO 19701, ISO 19702 ) , but ana lys is of some non-hygroscop ic gases such as NO , CO or CO<sup>2</sup> cou ld be performed with sampling lines at ambient temperatures.

Because of transportation delay and the thermodynamic conditions, effluent could also react between the sampling point and the analyser and its composition could be modified. This phenomenon is particle in the form it is given bang samples ing the species such as  $\Lambda$  in in international influence in parameters here are temperature and time. It should be appreciated that the validation of various sample transportation methods is only valid where the sampling systems have similar intervals between times between sampling and analysis and are at similar temperatures.

### 6 .4 Conditioning of the effluent

Effluent is often conditioned between the sampling point and the analysis point. The sampling line may be placed either before or after the conditioning procedure, or the conditioning could be performed in various steps, e.g. pre-filtration before the sampling line then final filtration after the sampling line.

Depending on the analytical technique used (See ISO 19701 and ISO 19702), conditioning may consist of filtration to remove soot from the effluent and/or a water trap. The water trap could be based on physical drying (i.e. through cooling) or chemical drying (e.g. calcium chloride, silicone oxide). Other gas traditions traditions as a contract  $\alpha$  remover  $\alpha$  removes (equations in ) . The mass of  $\alpha$ 

Some conditioning systems also include procedures to remove specific species, which could interfere with the analysis technique, but care should be exercised to ensure that other (wanted) species are not affect ted . For example, chem international letters users use converter to convert to convert  $L$  into NO before analysis. This operation has a limited efficiency, depending on the technique and design of the oven. For a successor italy is of the NO2 fraction in a NOX m is  $\Lambda$  m interesting  $\mu$  and the over show in a norm interesting .

All conditioning systems have a limited and variable efficiency. For example, a gas of interest could be partially adsorbed on filters. The filter could be analysed after test (see ISO 19702), but the kinetic information is particle . Hygroscopes procepts (HC l , HE l , HE ) and  $\mu$  , MF,  $\mu$  , MF is and time  $\mu$  is and time  $\mu$ are particularly sensitive to such losses.

Conditioning systems should, therefore, be studied before use to determine the effects on the quantification of each analyte of interest. In addition, it is essential to check how the conditioning system modifies the effluent as a whole. This also includes the effects the conditioning system may have on the sampling flow rate. A quantification of the mass loss in the effluent stream from sampling point to analyser shall be determined.

# 6 .5 Measurement technique

No measurement technique is perfect. Analysers are selective with variable sensitivity depending on the mix of species present. Calibration with "pure" gases may not take into account the effects of the other species in the matrix of compounds in the effluent being measured.

In addition to these effects, the response time from the sampling point to the end of the analysis has to be considered. This response time is an important characteristic of the system. In a dynamic measurement system, the transfer function of the system (i.e. a measure of the time required to achieve a given proportion of the species of interest at the analyser) could be a crucial parameter. A simple way to approach it is the time needed between 10 % and 90 % of the value for a single concentration measurement, as described for FTIR in ISO 19702. However, this parameter is not sufficient to fully characterize the response time in dynamic analysis conditions, as it doesn't cover the transfer function of a particular apparatus.

# 7 Validation steps

# 7.1 General

Details on the different validation techniques outlined in this document are available in References  $[11]$ ,  $[12]$  and  $[13]$ . Figure 4 gives indications on validation sequence and related clauses.



Figure  $4$   $-$  Guidance on validation steps

#### $7.2$ 7 .2 Definition of the range of application and range of calibration

An analytical technique is only applicable over the range of conditions used for calibration – and then only when the tests for calibration validity have been successfully carried out.

The lower limit of the range of application shall be determined according to ISO 12828-1. ISO 12828-1 describes various techniques for use in different applications. Methods described in ISO 12828-1:2011, 6.2 and 6.3 allow the determination of the physical limits of detection and quantification of an analyte. The method described in ISO 12828-1:2011, 6.4 provides a check on whether a given value is within an acceptable range. For example, this method is suitable for validating the lowest value of a set of calibration data, confirming it is a fully quantifiable value.

The highest value of a set of calibration data often provides the upper quantification limit of the analytical device. No measurements higher than this value shall be performed. The upper limit of a range of calibration is fixed by the sensitivity of the analytical method. Some analytical instruments exhibit spurious behaviour with high concentrations of analyte. An example is where the detector becomes progressively saturated leading to a loss of sensitivity and possibly a complete lack of response as the concentration of the analyte increases further. Beyond these points, the calibration model becomes unusable and the tests given in  $7.6$  shall be used to determine if the upper point of the calibration is still suitable.

The calibration points are selected on the basis of the characteristics of the measuring instrument **NOTE** and the practical technique used. In general, a data set consisting of 5 to 10 different analyte concentrations distributed uniformly over the required measurement range is suitable. This distribution can be chosen in the absence of knowledge of the actual calibration model to be used. For analytical instruments whose calibration model is known, e.g. where a perfectly linear relationship between concentration and detector output exists over the calibration range, a data set consisting of low and high values plus checkpoints for intermediate values is statistically more appropriate.

It is recommended that analysis is confined within the limits of 10 % to 90 % of the calibration range.

### 7 .3 Validation of the independence from the matrix effects

The matrix on which the analyte of interest is present may influence the concentration measured. A simple technique to evaluate the influence of the matrix is to test several different matrixes containing the analyte, and to compare the results obtained.

An example of procedure is as follows.

- $-$  Produce several blank samples i.e. various typical matrices but without the species of interest being present.
- $-$  For each matrix, add a known quantity of the analyte of interest. Repeat so as to produce a range of added quantities.
- The added quantities shall be the same, within acceptable limits, for each matrix.

It shall be emphasized that the blank matrixes shall have no trace of the analyte of interest, but shall retain a similar mix of species apart from the analyte of interest. Such a suitable blank can be difficult to obtain.

An example of determination of the specificity of the chosen method is given in  $A.2$ .

### 7 .4 Validation of the specificity of the chosen method

#### $7.4.1$ General 7 .4.1 General

Fire effluents may contain hundreds of chemical species, some of which are very similar chemically to others, e.g. aldehydes, and ketones. Depending on the analytical method used, some species could have a positive or negative interference on the measured quantities of similar species. The analytical method selected should, therefore, be chosen to be specific to the analyte required. To ensure such specificity, a range of analytical techniques may have to be studied.

An example of determination of the specificity of the chosen method is given in  $A.3$ .

#### 7 .4.2 Simple method

A simple technique to validate the specificity of the method is as follows.

- First, produce a representative sample of fire effluent, e.g. smoke from a physical fire model or in a trapping solution obtained from a fire test.
- Second, analyse it for the species of interest and then separate the sample in two parts.
- $-$  Third, add known quantities of the species of interest in one of these two parts, then analyse these samples for the species of interest. The difference between measurements shall be equal to the quantity added, allowing for experimental error.
- Fourth, add various quantities to the other part of the matrix of chemical species that could also be encountered and that could interfere. The selection of these interfering species shall be done with regard to the application, the expected species present in the smoke, and the limitations of the analytical technique chosen. The influence of these interfering species is evaluated as a sensitivity factor on the variation observed for analysis of the species of interest. Note that interference could be positive or negative.

**NOTE** ISO 19701:2013, Annex A gives a list of analytical techniques that are not suitable with fire effluents, even if they are commonly used for combustion gases.

#### 7 .4.3 Quantitative method

#### 7 .4.3 .1 Quality of separation

For chromatographic methods, such as described in ISO 19701, the quality of separation between two analytes can be expressed as a specific resolution between two adiacent peaks. The resolution for each set of two consecutive peaks shall be calculated according to Formula  $(1)$ . If the resolution is higher than or equal to  $0,6$ , then a qualitative analysis is possible. If the resolution is higher than or equal to 1,5, then a quantitative analysis is possible.  $\boxed{13}$ 

$$
R_s = 1,18 \times \left(\frac{t_2 - t_1}{w_2 - w_1}\right) \tag{1}
$$

where

- $R_{\rm s}$ resolution of the chromatographic method for two consecutive analytes (i.e. adjacent peaks);
- $\cdot$  1 and the  $\mu$  retention times for the two consecutive analytics  $\mu$  analytics ;

wat was well width at hand at height retention times for the peaks given from two consecutive analytes .

#### 7 .4.3 .2 Determination of degree of specificity

The goal of this step is to check that the method has a sufficient specificity for the selected component to be analysed. The study of specificity is carried out with samples representative of the usually analysed samples.

For each analyte, the analytical instrument is first calibrated. Then, a series of samples containing increasing concentrations of the analyte is prepared. This series shall cover the whole range of expected concentrations for the particular suppresses . The best s the best s transport leads  $\{1,1,1\}$  is particular  $\{1,2,3\}$ ana lytica la result for samples in the concentration is equation in a large process in a large result in the  $\cdots$   $\cdots$   $\cdots$   $\cdots$   $\cdots$   $\cdots$  and  $\cdots$   $\cdots$ conditions are simultaneously met, then the selectivity for the analyte is acceptable.

The calculations to be performed are given in Formula (2) for the residual standard deviation  $s(e)$ , in Formu la (3 ) for s(b<sup>1</sup>) , the s tandard deviation on b<sup>1</sup> and in Formu la (4) for s(b0) , the s tandard deviation on b0. p is the the total measurements of measurements performed ...

$$
s(e) = \sqrt{\frac{\sum_{i=1}^{p} (r_i - \hat{r}_i)^2}{p-2}}
$$
\n
$$
s(b_1) = \sqrt{\frac{s^2(e)}{\sum_{i} (v_i - \overline{v})^2}}
$$
\n(2)

$$
s(b_0) = \sqrt{s^2(e) \left( \frac{1}{p} + \frac{\overline{v}^2}{\sum_i (v_i - \overline{v})^2} \right)}
$$
(4)

The test performed to check if  $\mathbf 1$  is sufficiently close to unity is a uniform stationary t-test for 3 ° freedom with a 95 % confidence, according to Formula  $(5)$ .

For an acceptab le resu lt, tob s sha l l be less than the t va lue .

$$
t_{\rm obs} = \frac{|b_1 - 1|}{s(b_1)}\tag{5}
$$

The test performed to check if by a sufficiently close to zero is a universal structure to the form  $\sim$ freedom with a 95 % confidence, according to Formula  $[6]$ .

For an acceptab le resu lt, t′obs sha l l be less than the t va lue .

$$
t'_{\text{obs}} = \frac{|b_0|}{s(b_0)}\tag{6}
$$

### 7 .5 Influence of the measurement technique on results

#### 7.5.1 Generalities 7 .5 .1 Generalities

A species of interest could be suitably measured by different analytical techniques, based on various physical and chemical principles. As an example, ISO 19701 proposes two or three techniques for a large number of species of interest (see ISO 19701:2013, Table 1). In addition, several of these gases could also be analysed with FTIR according to ISO 19702.

For a given analytical species and a given application (e.g. physical fire model), these techniques are not, however, equivalent in terms of scope or response, or may have a different concentration range of application.

Sometimes, a technique is also referred to as "reference technique," usually perceived as the best method for a particular species in terms of specificity and quantitative accuracy. Tools are then needed to validate an alternative technique (which may be easier and/or cheaper to operate), by comparison with the reference technique.

Figure 5 details validation steps for the comparison between analytical techniques. Examples of the influence of the measurement technique on the analytical result according to various techniques presented hereafter are detailed in  $A.4$ .



Figure 5 — Guidance on comparison between analytical techniques

### 7 .5 .2 Simple methods

#### 7.5.2.1 Direct comparison

The following method is suitable for checking the influence of the measurement technique on results.

- $-$  First, select the two analytical techniques to be compared and calibrate the corresponding instruments. It is recommended that the techniques chosen are based on different properties of the analyte. Analytical techniques are normally only capable of measuring different samples in the same phase. A comparison of a gas-phase technique and a solution-based technique using the same equipment is therefore difficult with this method.
- $-$  Second, prepare at least five samples of increasing quantities in a representative matrix. The samples have to cover the range of concentrations capable of being measured by the two analytical techniques
- Third, analyse samples on both instruments. The values shall be the same within experimental error.

Results of both devices could be presented as a graph of measurement from analytical technique 1 vs. measurement from analytical technique 2. The linear regression of these data shall have a slope of  $1$ and an intercept of 0 if the two techniques are to be considered equivalent. Statistical tests proposed in  $7.4.3$  could be used to demonstrate that slope and intercept are statistically 1 and 0, respectively. An example of application is presented in  $A.4.2$ .

### 7 .5 .2 .2 Graphical method of Bland and Altman

Reference  $[14]$  proposes a graphical technique to evaluate the differences between two series of data.

- $-$  Proceed as described in 7.5.2.1 to obtain two series of data for each measurement technique to compare , des ignated as ya, <sup>i</sup> and yb, <sup>i</sup> .
- $\frac{1}{\sqrt{a}}$  is the average of each part of each  $\frac{1}{\sqrt{a}}$  ,  $\frac{1}{\sqrt{a}}$  ,  $\frac{1}{\sqrt{a}}$  ,  $\frac{1}{\sqrt{a}}$  $\sum_{i=1}^{n} a_i$  is one in the interest of  $\sum_{i=1}^{n} a_i$  if  $\sum_{i=1}^{n} b_i$
- Calculate the average value d and its standard deviation  $\sigma_{\rm d}$ .
- $-$  Plot  $d_i$  as function of  $\bar{y}_i$  and add to the graph the lines corresponding to  $d$  and  $d \pm 2\sigma_{d}$ .

An example of application is presented in  $A.4.3$ .

#### 7.5.3 **Quantitative method**

### 7 .5 .3 .1 Technique and statistical method

To determine whether two sets of analyses performed with two different techniques are equal or different, statistical tests are carried out, which require the mean value, the standard deviation and the variance to be calculated. Two series of data are then obtained, one for each analytical technique to be compared. Two tests shall then be carried out to verify the equivalence of the two techniques by comparing the variances and the mean values for each.

For comparison of variances, several statistic tests can be used.

- $-$  Fisher F-test $\equiv$  is the most Known and simple statistical test for comparison or variances. This test  $\,$ supposes that variables are normal. However, checking normality (see ISO 5479) requires a large number of data, which is difficult to obtain for many fire tests.
- Alternate tests as Levene<sup>[15][16]</sup> or Brown-Forsythe tests<sup>[15][17]</sup> are less sensitive to deviations from normality, and might be used. These tests are recommended when fewer data are available.

For comparison of means, several statistic tests can be used.

- Student's t-tes t[15 ] is the mos t known s tatis tica l tes t for comparison of means . Th is tes t supposes that equality of variances have been previously demonstrated. This test is successive to a variance test and only valid if variances are equal.
- Alternate tests such as Welch's t-test<sup>[15][18]</sup> are independent from any hypothesis on variances. These tests are recommended and may be used in parallel to comparison of variances.
- Wilcoxon's rank sum test<sup>[15]</sup> and Bayesian data analysis might be used as alternatives (not detailed in this document).

Results are sometimes reported as p-values instead of test statistics. The p-value is the probability of observing a difference at least as large as observed due to the vagaries of sampling alone, and that therefore there is no real difference. Small p-values, typically  $0.05$  or less, suggest that there is a statistically significant difference compared to a given limit.

### 7 .5 .3 .2 Comparison of variances

#### Method 1: Fisher F-test  $7.5.3.2.1$

The Fisher F-test is extremely sensitive to deviations from normality. If data are not normal, this test shall not be used. ISO 5479 proposes methods to check the normality of data.

The variances are statistically compared with an  $F$ -test. In this test, the ratio of the two variances (Fobserved) is compared to a theoretica l F-ratio (Ftheory) . The s tatis tics are ca lcu lated accord ing to Formula (7).

In this case, I fheory values are tabulated for uniterent levels or probability.[33] It is then iltersal y to refer to the Fisher-Snedecor tab le at e .g . 1 % or 5 % of s ign i ficance , in order to get the va lue of <sup>F</sup>theory.

I f Fob served < Ftheory, then the var iances are cons idered as equa l . I <sup>f</sup> not, ana lytica l techn iques are significantly different in terms of variances.

$$
F_{\text{observed}} = \frac{s_1^2}{s_2^2} \tag{7}
$$

where

- $S<sub>1</sub>$ standard deviation of series 1;
- $S<sub>2</sub>$ standard deviation of series 2;
- s<sup>1</sup> > s<sup>2</sup> so that Fob served > 1 .

NOTE The Bartlett tes t[15 ] is an a lternative to the Fisher F-tes t, with the same assumption of norma l ity of data .

A strong deviation from a normal distribution may indicate a problem in the data and would suggest that further investigation is needed.

### 7.5.3.2.2 Method 2: Levene and Brown-Forsyth (LBF) tests

Levene's testiffs is a statistical test for the equality or variances calculated for two or more groups. Equal variances across samples are called homogeneity of variance. Analysis of variance assumes that variances are equal across groups or samples. The Levene test can be used to verify that assumption. The Levene test is less sensitive than the Fisher F-test or the Bartlett test<sup>[15]</sup> to departures from normality. If there is strong evidence that data do in fact come from a normal, or nearly normal, distribution, then

bar ticles testi— is more appropriate. The Levene test statistic W is defined according to Formula (8). The variables are transformed to measure the spread in each group.

$$
W = \frac{\left(N - k\right)}{\left(k - 1\right)} \frac{\sum_{i=1}^{k} n_i \left(z_{i.} - z_{..}\right)^2}{\sum_{i=1}^{k} \sum_{j=1}^{n_i} \left(z_{ij} - z_{i.}\right)^2}
$$
\n(8)

where

 $W$ is the result of the test;

 $k$  is the number of different groups to which the sampled case belong;

 $\overline{N}$ is the total number of cases in all groups;

 $n_i$ is the number of cases in the group i;

 $Z_{ij} = |Y_{ij} - Y_i|_{\text{group and } \overline{y}}$ . group and y <sup>i</sup> ;

z. . is the genera l mean of a l l zij;

 $Z_i$ is the mean of the  $z_{ij}$  values for *i*-th group.

The s ign i ficance of Wis tes tes tes ing in 1.5 . 1 . 1 and N − 1 .1 with k − 1 and N − 1 and N − k its degrees of freedom at e.g.  $1\%$  or 5 % of significance.

Brown and Forsythe<sup>[15][17]</sup> extended Levene's test to use either the median or the trimmed mean instead of the mean (LBF test).

<sup>y</sup> <sup>y</sup>  $y'' + y'' = y$  $=$   $\left\{ \begin{array}{ccc} & \circ & \circ \\ & & \circ \end{array} \right.$ ij <sup>i</sup> − − . . a sa n  $||y_{ii} y||$ . . . . , considering  $y_{ij}$  as the value of the measured variable for the  $j$  case from the  $i$  group,  $\overline{y}_i$ 

and  $\bar{v}'$ , respectively as the median and the 10 % trimmed mean of the *i*-th group.

The three cho ices for defin ing zij determ ine the robus tness and power of the d ifferent tes ts . Robus tness means the ability of the test to not falsely detect unequal variances when the underlying data are not normally distributed and the variables are in fact equal. Power means the ability of the test to detect unequal variances when the variances are in fact unequal. Using the mean provides the best power for symmetric, moderate-tailed, distributions. Although the optimal choice depends on the underlying distribution, the definition based on the median is recommended as the choice that provides good robustness against many types of non-normal data while retaining good power. If the underlying distribution of the data is known, this may indicate using one of the other choices.

#### 7 .5 .3 .3 Comparison of mean values

### 7.5.3.3.1 Method 1: Student's t-test

The application of Student's test to compare mean values supposes that variances are equal. If equality of variances has not been demonstrated earlier, or if the data failed comparison of the variances test, then another technique should be used.

The s tatis tica l parameters used to compare meaning the pools the pools the pools the pools the pools in specific to <del>Formula (1)</del> and the value of tobserved accord ing to Formula (10) . The value to see to see to see to see with the Value of the Student's table is tudent at a degree of freedom n = n1 + n2 − 2 . I foodball for the top served to the transfer the mean values are regarded as equal. If not, analytical techniques are significantly different in terms of values.

$$
s_p = \sqrt{\frac{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2}{n_1 + n_2 - 2}}
$$
\n(9)

$$
t_{\text{observed}} = \frac{\left|\overline{y}_1 - \overline{y}_2\right|}{s_p} \sqrt{\frac{n_1 \times n_2}{n_1 + n_2}} \tag{10}
$$

where

 $\overline{\phantom{0}}$ 

 $n_1$  and  $n_2$ number of values in groups 1 and 2;

<sup>y</sup> and y mean values of groups 1 and 2.

### 7.5.3.3.2 Method 2: Welch's t-test

we left is t-testical is an adaptation of Student's t-test, and is intended for use when the two samples have possibly unequal variances. Welch's t-test defines the statistic t by the Formula  $(11)$ . Unlike in Student's  $t$ -test, the denominator is not based on a pooled variance estimate.

$$
t_{\text{observed}} = \frac{\bar{y}_1 - \bar{y}_2}{\sqrt{\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}}} \tag{11}
$$

The degrees of freedom  $v$  associated with this variance estimate is approximated using the Welch– Satterthwaite Formula (12):

$$
v \approx \frac{\left(\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}\right)^2}{\frac{s_1^4}{n_1^2 v_1} + \frac{s_2^4}{n_2^2 v_2}}
$$
(12)

 $v_i = n_i - 1$  is the degrees of freedom associated with the *i*-th variance estimate.

Once  $t_{\rm observed}$  and  $\rm\,v$  have been computed, these statistics can be used with the t-distribution to test the null hypothesis that the two population means are equal.

If the population variances are equal, or if the samples are rather small and the population variances can be assumed to be approximately equal, it is more appropriate to use Student's t-test.

Analytical techniques use a calibration model to compare the concentration of the analyte with the value of a detector. This model is often linear, but not usually for all conditions. The regression coefficient  $r^2$  given for many regression calculations is not sufficient to characterize the capability of the model to represent data: it is often not representative of the error due to the model for small values, as its calculation favours high values.

An example of a calibration study is given in  $A.5$ .

To study the capability of the chosen calibration model to represent data, a calibration curve with  $p$ different levels of concentration is constructed for each analyte. Each solution is prepared several times and consists of a series of n values of  $\nu$ . The study of the calibration model then consists in checking that the model represents the data. The study could start with a linear regression model as the simplest form If this model does not explain the data, then a second degree polynomial model is used. The calculation of global mean and associated standard deviation are detailed in Formulae  $(13)$  to  $(14)$ .

$$
\overline{y}_{i} = \frac{\sum_{i=p}^{i=1} y_{i}}{p}
$$
\n
$$
S_{Y_{i}} = \sqrt{\frac{\sum_{i=p}^{i=1} (y_{i} - \overline{y_{n}})^{2}}{p-1}}
$$
\n(14)

where

 $\overline{\phantom{a}}$ 

 $\overline{y_i}$ is the global mean;

 $\ddot{\phantom{0}}$ 

- $p$  is the total number of measurements performed;
- $S_{Y}$ is the global standard deviation.

The coefficients of the model are calculated according to least-squares models. For a linear model, coefficients are calculated according to Formulae  $(15)$  and  $(16)$ . Formula  $(17)$  calculates the response of the moder  $y_j$ . The non-intearity residuals correspond to  $y_j - y_j$  and the repeatability res idue s to y <sup>y</sup> <sup>−</sup> .

$$
b_0 = \frac{\sum (x_i - \overline{x_i})(y_i - \overline{y_i})}{\sum (x_i - \overline{x_i})^2}
$$
(15)

$$
b_1 = \overline{y_i} - b_1 \times \overline{x_i} \tag{16}
$$

$$
\hat{y}_i = b_1 \times x_i + b_0 \tag{17}
$$

The sum of the squares of the deviations SCE at  $y_1$  is the  $g$  iven by <u>Formu (23)</u> . Then , the medi square (*MS*) is calculated according to **Formula** (19). Various *MS* are calculated as function of various SCEs and degrees of freedom df as defined hereafter.

The model chosen is the one that minimizes a suitable model selection criterion. Several statistical techniques could be used to quantify the quality of the calibration model studied:

- $-$  Fisher statistic;
- $\equiv$  Big (Daycsian information Criterion)  $\equiv$
- $-$  AIGC (AKaIKG 3 Information Gritchion with finite sample size correction) $(21)$

#### 7 .6 .2 Analysis of calibration model using the Fisher statistic

The Fisher s the Fisher s the Fisher street for the ined is  $\lambda_{\rm DM}$  in a individual of  $\lambda_{\rm DM}$  , consequently associated in df Fabiatiica = dfModel – for the calculation of MSE xpiatiica dfresh dipublikation = np = np = np = np = np + p = n . .- . . . . . . . . . . .

The Fisher s tatis tic observed for non-linear ity, Fob s , Non linear ity, is given by Formula (21) , conserved dimon lating and the case the calculation of the MONNING the case in India is the case the calculation of the MS

$$
SCE_{y_i} = \sum (y_i - \overline{y_i})^2
$$
 (18)

$$
MS = \frac{SCE}{df}
$$
 (19)

$$
F_{\rm obs\;Explained} = \frac{MS_{\rm Explained}}{MS_{\rm residual}}
$$
 (20)

$$
F_{\rm obs \; Non\; linearity} = \frac{MS_{\rm Non\; linearity}}{MS_{\rm Total}} \tag{21}
$$

The first tep of the value  $f$  is the value of the value of the regression . For large  $\alpha$  in the start  $\alpha$  is the sequence in the start of th F95 %, the F isher F-s tatis tic at 95 % con fidence .

The second step is the value of the line interesting in OD3 NONNLOALLO shall be shared in the S95 W.

#### 7 .6 .3 The BIC (Bayesian Information Criterion)

The BIC (Bayesian Information Criterion)<sup>[20]</sup> may be used to compare different calibration models. The BIC is defined according to Formulae  $(22)$  and  $(23)$ .

$$
BIC = k \ln(n) - 2 \ln(L) \tag{22}
$$

if the errors are normally distributed:

$$
BIC = k \ln(n) + n \ln(MSE)
$$
\n(23)

where

- $k$  is the number of parameters of the model, e.g. two for a linear one;
- $n$  is the number of data;
- L is the likelihood of the model, see Reference  $[15]$ ;

*MSE* is the Mean Squared Error defined as 
$$
MSE = \frac{1}{n} \sum_{i=1}^{n} (\hat{y}_i - y_i)^2
$$
.

The best regression model is the one that minimizes the BIC value.

#### 7 .6 .4 Analysis of calibration model using the AICc (Corrected Akaike Information Criterion)

The AICc (Aka ike 's Information Cr iter ion with fin ite samp le s ize correc tion) [21] is an a lternative to the Fisher statistic to compare different calibration models. The AICc is defined according to Formulae (24) and (25) with the parameters defined in  $7.6.3$ .

$$
AICc = 2k - 2\ln(L) + \frac{2k(k+1)}{n-k-1}
$$
 (24)

Or, if the errors are normally distributed:

 $\sim 10^{-10}$ 

$$
AICc = 2k + n\ln\left(MSE\right) + \frac{2k\left(k+1\right)}{n-k-1} \tag{25}
$$

The best regression model is the one that minimizes the AICc value. The AICc is recommended if  $n$  is small or  $k$  is large. Degrees of freedom of the model affects the AICc criterion less than it does the BIC.

#### **Determination of uncertainties**  $\mathbf{R}$ 8 Determination of uncertainties

The objective of an analytical method is to try to determine the real concentration of analytes of interest at the sampling point in a fire atmosphere. Appreciation of the factors discussed in Clause 6 can help to reduce the uncertainty of analyte measurement, and the validation methods discussed in Clause 7 can be used to establish effective calibration and to check the influence of the measurement technique on results. resu lts .

Uncertainties are calculated according to ISO/IEC Guide 98-3. The repeatability (R) and the reproducibility  $(r)$  of an analytical method for fire effluents could be determined according to the ISO 5725 series. However, for fire atmosphere analytical methods, this determination shall specify if the repeatability and reproducibility of the physical fire model is included in the final results. In many cases, a lack of repeatability and reproducibility could be due to the physical fire model (i.e. variations in ignition time, effects of sample homogeneity, etc.) instead of the analytical method itself, when results analysed are issued from combustion plus analysis process.

An example of uncertainty calculation according to ISO/IEC Guide 98-3, including repeatability considerations, is presented in  $Annex B$ .

#### **Annex A** Annex A

# (informative)

# Example of application of validation steps: Analysis of hydrogen chloride and hydrogen bromide from trapping solutions

#### A.1 General ---- ---------

The example given here is for the analysis of hydrogen chloride and hydrogen bromide according to the technique presented in ISO 19701:2013, 5.5.2. In the example, hydrogen chloride and hydrogen bromide are trapped in a hydiu phase as hanue ions using impiligers, as described in NF X 70 Too 2(==). Chioriuc and bromide ions are then analysed using Ion-Liquid Chromatography (ILC) with a conductimetric detector. The standard protocol stated in ISO 19701, has been modified for this example by use of a trapping solution of deionised water containing  $3\%$  of hydrogen peroxide. The addition of hydrogen peroxide allows the simultaneous trapping of sulfur dioxide.

This example is detailed in References [23] and [24]. The analytical system used includes the following:

- ion Liquid chromatograph type Dionex DX500;
- ion exchange column AS 14 (silica grafted with quaternary ammonium functional groups) and precolumn AG14;
- suppressor type ASRS Ultra 4 mm;
- detector ED40 (used in conductimetric mode);
- mob is more in the phase of 3 ,5 mmo l  $\mu$  in  $\Delta$  ,  $\mu$  and 1 mmo l  $\mu$  in l name is  $\mu$
- flow rate of the mobile phase:  $1,5$  ml/min;
- autosampler with injection loop volume of 50  $\mu$ l;
- organic filtration (Varian JR-C18 500mg) and mechanical filtration (PTFE 0,2 mm) of samples.

The instrument is calibrated using six standards for both ions. The concentration range for both ions is from 1 mg/L to 20 mg/L.

# A.2 Validation of the non- influence of the matrix

CO2 are measured by ND IR accord in the IR accord ing to ISO 1970. In the theory is the second product of the s other species of interest are adsorbed using two different trapping solutions: deionized water and deionized water containing 3 % of hydrogen peroxide. Deionized water has been previously validated as having no influence on the quantity of CO and CO2 and Journal . A series of tes to performed to valuate that the second trapping solution has no influence either.

Two configurations are successively compared, at several volume fractions:

- impingers filled with deionized water (standard defined solution);
- the same impingers, filled with the alternate solution containing hydrogen peroxide.

As the only difference is the composition of the trapping solution, any difference between the data obtained with the alternate solution, and the data obtained with the standard defined solution, will then be solely due to the difference of trapping solution. The measurements are carried out at three different volume fractions:

- low vo lume fraction in the first  $\sim$  ,  $\sim$  and  $\sim$   $\sim$   $\sim$   $\sim$
- med in the interest and in the fraction of the second second second  $\mu$
- h igh volume fraction is about 5 minutes of 20 and 20 % CO2 . A

All measurements are then compared, in order to determine the influence of the trapping solution. Typ is are used in Fig. are presented in <u>Table Le A .</u> For contents from 0 % to 5 % of Co and 0 % of Co2 , 20 % of C no difference in results is noted. The trapping solution has no influence on the measurement of the amount of CO and CO<sup>2</sup> after the adsorbers .

Table A.1 — data obtained to validate the non-influence of the matrix (in %)

	CO		CO <sub>2</sub>		
	Standard defined Alternative solution solution		Standard defined solution	Alternative solution	
Low volume fraction	$0.91\%$	$0.91\%$	$3.65\%$	3,64 %	
Medium volume fraction	3,00%	$3.00\%$	11,80 %	11,81 %	
High volume fraction	4,54 %	4,54 %	17,38 %	17,38 %	

# A.3 Specificity of the method

### A.3 .1 Quality of separation

In order to have a blank sample representative of the solutions to be analysed, the combustion of a material is carried out. This material has been chosen to produce a similar matrix of compounds in the effluent, but without the presence of chloride and bromide. This is checked using a titrimetric method. One ml of the solution used to trap this effluent, (termed "combustion matrix"), is mixed with 1 ml of a standard poly-annon sondtion containing. F− (10 mg/T), CF− (20 mg/T), NO/− (20 mg/T), Br− (20 mg/T), -NO<sup>−</sup> (20 mg/ l) , PO3− (20 mg/ l) and SO2− (30 mg/ l) . The ana lysed ions are F<sup>−</sup>, C l<sup>−</sup>, NO−, Br<sup>−</sup>, NO−, <sup>3</sup> <sup>4</sup> <sup>4</sup> <sup>2</sup> <sup>3</sup> PO3−, SO2− and CH3COO<sup>−</sup>, wh ich is not quanti fied . A l l these an ions cou ld be present in the trapp ing solution obtained by sampling from a fire test.

Determination of the quality of separation is achieved through the calculation of the resolution as proposed in 7.4.3. The resolution for each set of ions has been calculated as shown in Table A.2. When both acetate and fluoride ions are present, a precise quantitative analysis of these ions is impossible, as peaks overlap. All other ions in solution can be successfully analysed and the resolutions calculated for the bromide and chloride ions are high enough to allow a quantitative analysis of both these ions.

Ion	Retention time (min)	Width at middle height (min)	Resolution with previous ion
Fluoride $(F^-)$	2,33	0,07	
Acetate (CH <sub>3</sub> COO <sup>-</sup> )	2,57	0,16	1,23
Chloride (Cl <sup>-</sup> )	3,27	0,09	3,30
Nitrite ( $NO2$ -)	3,83	0,11	3,30
Bromide (Br <sup>-</sup> )	4,73	0.14	4,25
Nitrate $(NO3^-)$	5,49	0,17	2,89
Phosphate (HPO $4^{2-}$ )	7,12	0,25	<b>ND</b>
$\vert$ Sulfate (SO <sub>4</sub> 2-)	8.53	0.27	3,20

Table  $A.2 -$  Quality of separation in ILC

In fire effluents, a large quantity of hydrogen chloride could be released from materials containing chlorine. In these conditions, it may be difficult to analyse hydrogen bromide as in solution, a large quantity of chloride ions can mask the bromide ions. A solution originating from the combustion of PVC material, containing 1000 mg/L of chloride ions, is used. To this solution,  $2.5$  mg/L of bromide ions are added. Analysis is performed three times and the resulting concentration is  $(2.0 \pm 0.11)$  mg/L. These results show that even the highest concentration of Chloride ions do not prevent weak concentrations of Bromide ions from being correctly measured in these conditions.

## A.3 .2 Determination of specificity

For chloride and bromide ions, five solutions with concentrations from 1 mg/L to 20 mg/L are used. The ions are added over the concentration range to be expected in the matrix from a fire study. For example, for a fire matrix with 5 mg/L of chloride ions, an addition of 5 mg/L of chloride ions is used.

The analytical instrument is first calibrated. Each solution is initially analysed to obtain the analyte concentration. Then, the additions are carried out using the standard solutions. In this way, the added concentration is accurately mixing, which is called the "real" addition  $\left(\theta\right)$  . The resultant source is then ana lysed . S ince the interest in itia later than the interest of the addition is measured (r) . Fig. of data are are are are as in the addition in the addition in the addition is measured (r) . Fig. or are are are are ar ob the ined for the channels in the channels . Results of cancellations described in 1982, and 7.4 . 3 . 3 . 3 <u>Table 1999</u>, is a conserved as  $0$  can be conserved as equal to 0 with a 5 % error. For children  $\mathbb{F}_1$  can be considered as equal to 1 with a 5 % error. This error is 1 % for Bromide. Consequently, the method is specific for the measurement of chloride and bromide ions in such a matrix.

Ion	Chloride	Bromide
S(e)	$4,54 \times 10^{-1}$	$2,03 \times 10^{-1}$
$B_1$	1,039	1,111 2
$S(b_1)$	$3,37 \times 10^{-2}$	$3,32 \times 10^{-2}$
B <sub>0</sub>	0.34	$-0,1516$
$S(b_0)$	0,32	0,16
$t_{\rm obs}$	1,18	3,35
$t'_{\text{obs}}$	1,06	0,94
$t_{student}$ (95 %)	3,18	3,18
$t_{student}$ (99 %)	5,84	5,84

Table  $A.3$   $-$  Results of selectivity test

# A.4 Influence of the measurement technique

The ILC method is then compared to the titrimetric method described in ISO 19701:2013, 5.5.3.

### A.4.1 Simple methods

The two techniques are compared using a wide range of concentrations in solutions obtained from the combustion of polymeric materials. Results are presented in Table A.4 and Table A.5 respectively for hydrogen chloride yield and hydrogen bromide yield, obtained from the combustion of a series of materials. mater ia ls .

Material	HCl yield (mg/g) with titrimetry		$HCl$ yield $(mg/g)$ with ILC		Deviation between mean values		
	Mean value	Std. dev.	Mean value	Std. dev.	(mg/g)	(%)	
A <sub>1</sub>	4,20	1,18	4,18	0,93	0,02	0,5	
<b>B1</b>	8,34	1,01	8.10	1.09	0,24	3,0	
C <sub>1</sub>	14,4	1,69	13,9	1,44	0,47	3,4	
D <sub>1</sub>	134	7,83	133	9.14	0,91	0,7	
F <sub>1</sub>	162	5,67	152	11,72	10,7	7,0	
G <sub>1</sub>	249	1,10	232	3,53	16,6	7,1	
H1	575	8.90	523	9,12	51,6	9.9	

Table A.4 – Comparison of methods (concentration in liquid for hydrogen chloride)

Table A.5 — Comparison of methods (concentration in liquid for hydrogen bromide)

Material	HBr yield (mg/g) with titrimetry		HBr yield (mg/g) with ILC		Deviation between mean values	
	Mean value	Std. dev.	Mean value	Std. dev.	(mg/g)	(%)
A1	33,5	2.93	33,2	2.56	0,35	1,1
B1	59.5	3,21	56,3	3.07	3,19	5,7
C <sub>1</sub>	9,52	1,82	7,84	1,46	1,69	21,6

For all materials releasing HBr and those releasing HCl between  $4.12 \text{ mg/g}$  and  $157 \text{ mg/g}$ , calculations show that there is no influence of the technique on the results. For the two materials releasing a higher amount than 157 mg/g for HCl, amounts measured by ILC are approximately 10 % lower than those measured by titrimetry. For one material, the two sets of results have unequal variances and mean values. For one material  $(H1)$ , the variances are comparable but the mean values are unequal.

# A.4.2 Graphical representation

The results are plotted as Figure A.1 for hydrogen chloride data.



Figure  $A.1$  — Graphical representation of results (case of hydrogen chloride)

# A.4.3 Bland and Altman graphical method

The results of the Bland and Altman method are plotted as Figure A.2 for hydrogen chloride data. It confirms the observation of a problem with data corresponding to H1 material.



### Figure  $A.2$  — Bland and Altman graphical method results (case of hydrogen chloride)

### A.4.4 Quantitative method

### A.4.4 .1 Comparison of variances

### A.4.4.1.1 Using the Fisher F-test

The quantitative technique using the Fisher F-test has been applied to the data shown in Table  $A.4$  and Table A.5. Results of the comparison of variances are given in Table A.6 and Table A.7 respectively for hydrogen chloride and hydrogen bromide. One single case of non-equivalence of variances is observed for hydrogen chloride for the material G1. Equivalence is established for this point at the level of  $1\%$  $\mathbf{v} = \mathbf{u} \cdot \mathbf{u} \cdot \mathbf{v}$  ,  $\mathbf{u} = \mathbf{v} \cdot \mathbf{v} \cdot \mathbf{v}$  . The set of  $\mathbf{v} = \mathbf{v} \cdot \mathbf{v}$ 

The Fisher test is performed using a low number of observations. This yields to a low power of the test, **NOTE** i.e. a low capacity for the test to detect any difference between the variances, unless the difference is very large.



### Table  $A.6 -$  Comparison of variances using the *F*-test  $-$  Results obtained for the yield of hydrogen ch loride



### Table  $A.7 -$  Comparison of variances using the *F*-test  $-$  Results obtained for the yield of hydrogen bromide

#### A.4 .4.1 .2 Using the Levene test as modified by Brown-Forsyth (LBF)

Results of the comparison of variances using the LBF test are given in Table A.8 and Table A.9 respectively for hydrogen chloride and hydrogen bromide. The results indicate that variances are homogeneous for all materials and for both HCl and HBr. Comparison of variances using LBF highlights no significant deviation at the 5 % level (no p-value inferior to 0,05).

#### Table  $A.8 -$  Comparison of variances using the LBF test  $-$  Results obtained for the yield of hydrogen ch loride

	Number of		Comparison of variance		
Material	tests $n_1 = n_2$	W	$F_{\rm theory}$ , 5 %	<i>p</i> -value	
A <sub>1</sub>	5	0,206	6,39	0,66	
B1	3	0,004	19,00	0,95	
C <sub>1</sub>	3	1,014	19,00	0,37	
D <sub>1</sub>	3	0,046	19,00	0,84	
F1	5	0,770	6,39	0,41	
G <sub>1</sub>	5	0,872	6,39	0,38	
H1	5	0,037	6,39	0,85	

Table A.9 — Comparison of variances using the LBF test — Results obtained for the yield of hydrogen bromide



#### A.4 .4.2 Comparison of means

#### A.4 .4.2 .1 Using Student's t-test

Results of the comparison of means are given in Table A.10 and Table A.11 respectively for hydrogen chloride and hydrogen bromide. In the case of hydrogen chloride, the results highlight that the two techniques are not equivalent for higher concentrations in terms of mean values. They are equivalent for the determination of yields up to  $160$  mg/g.



### Table  $A.10$  – Comparison of means – Results obtained for the yield of hydrogen chloride

### Table  $A.11$   $-$  Comparison of means  $-$  Results obtained for the yield of hydrogen bromide



### A.4.4 .2 .2 Using Welch's t-test

Results of the comparison of means using Welch's test are given in Table A.12 and Table A.13 respectively for hydrogen chloride and hydrogen bromide. The results indicate that means are different for materials C1,  $G1$  and H1 in the case of hydrogen chloride (p-values inferior to 0,05).

#### Table A.12 – Comparison of means using the Welch's test – Results obtained for the yield of hydrogen ch loride



### Table A.13 – Comparison of means using the Welch's test – Results obtained for the yield of hydrogen bromide



# A.5 Calibration study

The calibration study of ILC for chloride ion is carried out according to techniques presented in 7.6. The calibration data set is made from 25 solutions: 5 standards are analysed at 5 concentrations, ranging from 1 mg/L to 20 mg/L. The analysis uses the area under the peak  $v$ , vs. concentration x.

As the model uses data from a detector based on a specific physical measurement principle, the choice of the model's mathematical form has to be driven by the physics of the detector principle. For example, an ILC (Ion Liquid Chromatography) measurement with a conductimetric detector, as proposed for chloride or bromide ions in ISO 19701 corresponds to a resistor-capacitor (RC) equivalent alternating current (AC) electric circuit dependent on ion mobility. When concentration increases, ion mobility decreases so the linear model is not applicable to concentrated solutions or to a wide range of concentrations. It is nevertheless possible to use such a model for diluted concentrations and with a narrow range of concentrations . When the range increases , another model such as a second degree polynomial relation is more appropriate to represent the data. See Reference [19] for more details.

Figure A.3 presents graphically the data and both regression models. Figure A.4 presents the residues for both models.

Table A.14 and Table A.15 present results obtained using a linear regression model. As a first step, the regress is value in the mode  $\sim$  Fexplatified  $\sim$  F1 M. The model is the data in the data in the data in the data  $\sim$ s the second step , let it is not ver in the research  $\mu$  in the results  $\mu$  is the resonance of the results which can not be experimented as the experiment be experimented as the experiment of the large deemed excessive. A model of second order is therefore tested.

Model	$y = b_1.x + b_0$
Slope $b_1$	$1,74 \times 10^{5}$
Standard deviation on slope $s(b_1)$	$1,87 \times 10^{4}$
Intercept $b_0$	$-1,03 \times 10^{5}$
Standard deviation on intercept $s(b_0)$	$1,92 \times 10^{5}$
Regression coefficient $r^2$	0.997
Sum of residuals	$6,48 \times 10^{4}$
F-statistic	$8,69 \times 10^{4}$
Degrees of freedom	23
Sum of explained squares SCexp	$3,65 \times 10^{13}$
Sum of residuals SCres	$9,67 \times 10^{10}$

Table A.14 – Results of a regression calculation for the case of a linear model

#### Table A.15 – Results of a Fisher test for calibration for the case of a linear model



Table A.16 and Table A.17 present results obtained for a second-order model. As a first step, regression is valued as Ferry - Feature of the model is the model is the second the second it fits the data second s tep , l inear ity is ver i fied , as Fnon linearity <sup>&</sup>lt; <sup>F</sup>1 %.



### Table A.16 - Results of regression calculation for the case of a 2nd order model

## Table A.17 – Results of a Fisher test for calibration for the case of a 2nd order model



Table A.18 presents results of BIC and AICc analyses for linear and second-order models, assuming  $\overline{\text{normality}}$ . For both methods of analysis, the second order model gives a better fit to the data.







Figure A.3 - Calibration data and regression models tested



Figure A.4 - Residuals from regression models tested

#### **Annex B** Annex B

# (informative)

# Example of an uncertainty calculation: Analysis of hydrogen chloride in trapping solutions

#### **B.1** General --- ------

The example presented here corresponds to the uncertainty calculation applied to measurement of hydrogen chloride yield from three replicates of a single material. Analysis is performed on chloride ions in trapping solution with Ion Liquid Chromatography, as described in Annex A. The physical fire model used to generate the effluent is the NF X 70-100-2 tube furnace and the analytical technique is presented in ISO 19701:2013, 5.5.2.

The material for test is weighed, then introduced in the tube furnace. The effluent passes through impingers, to absorb hydrogen chloride as chloride ions. After the test, the impinger solution is filtered and the volume is adjusted using a volumetric flask. This solution is then analysed by ion liquid chromatography previously calibrated using standard solutions (see  $A.5$  for the details of this calibration). Depending on the concentration of the solution, an additional dilution can be performed. The calibration model used is a second-order polynomial, as validated in  $A_4$ . The result is expressed as a concentration in the volumetric flask, or as a gas yield from the tested material.

Tests are performed on three samples. The uncertainty calculation estimate presented hereafter corresponds to uncertainty due to analysis plus the uncertainties inherent to combustion, physical fire model and material homogeneity. This example is detailed in References  $[23]$  and  $[24]$ .

# B.2 Analysis of the uncertainty

The concentration variety , Cflask, are obtained to an 2nd order cannot cannot cannot cannot the model  $\alpha$  . The the yie light in the most energy as an in yields in mg/g are presented in <u>the second</u>

$$
Y_{\text{HCl}} = \frac{C_{\text{flask}} \times V_{\text{flask}} \times M_{\text{HCl}} \times d}{m_{\text{sample}} \times M_{\text{Cl}^-}}
$$
(B.1)

where

 $V_{\text{flask}}$ is volume of the volumetric flask used to collect solution from impingers;

- $m_{\text{sample}}$ is mass of the material sample introduced in the tube furnace (about 1 g according to NF X 70-100);
- $d \sim$  is the dilution required to give concentration within the calibration range;

 $M<sub>HCl</sub>$ is the molar mass of hydrogen chloride;

is the molar mass of chloride ion.

The uncerta inty ca lcu lation cons iders a s tandard uncerta inty u(x) assoc iated with each parameter xi <u>representa prop</u>i reduce of these terms include a repeatably include <u>a repeated bid</u> gives the analytica ly uncerta international contracts with a distribution and with a distribution of 20 . Variation of 20 . Var international and the local late sum of variances for each parameter. Results are presented in Table B.2 and Table B.3 and details available in References  $[23]$  and  $[24]$ .

⊺ ci−

<b>Results</b>	Test 1	Test 2	Test 3	Mean
$m_{\text{sample}}(\text{g})$	0.96	1,02	1,00	0,99
$V_{\text{flask}}$ (mL)	500	500	500	500
$C_{\rm{flask}}$ (mg/l)	259	275	264	266
$Y_{\text{HCl}}$ (mg/g)	138	139	136	138

Table B.1 - Hydrogen chloride yield results of a series of tests



Figure B.1 — Uncertainty from the determination of the analyte concentration in solution, as function of concentration

Parameter $x_i$	Value of $x_i$	Standard uncertainty $u(x_i)$	Relative standard uncertainty $u(x_i)/x_i$	Repeatability uncertainty
$m_{\text{sample}}(\text{g})$	0,9932	$2,31 \times 10^{-4}$	$2,33 \times 10^{-4}$	$1,16 \times 10^{-4}$
$V_{\text{flask}}$ (mL)	500	0.264	$5,28 \times 10^{-4}$	$2,64 \times 10^{-4}$
d	20	0,136	$6,78 \times 10^{-3}$	
$M_{\text{HCl}}\left(\text{g/mol}\right)$	36.46	$5,79 \times 10^{-4}$	$1,59 \times 10^{-5}$	
$M_{Cl^-}$ (g/mol)	35.5	$1,00 \times 10^{-3}$	$2,82 \times 10^{-5}$	
$C_{\rm flask}$ (mg/l)	13,3	$6,13 \times 10^{-2}$	$4,61 \times 10^{-3}$	$2,30 \times 10^{-3}$

Table B.2  $-$  Determination of uncertainty from the analysis





# B.3 Uncertainty from a series of tests

The result reported from a series of tests is the mean of several replicates, three taken as example. The dispersion of these three replicates takes into account the uncertainty of the measurement repeatability and the uncertainties associated with the heterogeneity of the material, thermal decomposition, repeatability, etc. The combination of these uncertainties is calculated using Formula  $(B.2)$ . The final resume is yie for the later and an ing compared factor in the state factor  $\mathcal{A}$  , and it is of any led analysis of  $\mathcal{A}$ contributors to uncertainty indicates that the dispersion due to the material is equivalent to the measurement uncertainty, contributing respectively at 42 % and 58 % of the total uncertainty.

$$
u_h = \sqrt{S_{\text{gas}}^2 + u(Y)^2}
$$
 (B.2)

where  $\dots$ 

> $u<sub>h</sub>$ is the standard uncertainty due to material heterogeneity and fire model;

Sgas is the s tandard deviation between va lues of the three tes ts .

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# **Bibliography**

- [1] ISO 19701:2013, Methods for sampling and analysis of fire effluents
- [2] ISO 19702, Guidance for sampling and analysis of toxic gases and vapours in fire effluents using Fourier Transform Infrared (FTIR) spectroscopy
- [3] ISO 19706, Guidelines for assessing the fire threat to people
- [4] ISO 5660-1, Reaction-to-fire tests Heat release, smoke production and mass loss rate Part 1: Heat release rate (cone calorimeter method) and smoke production rate (dynamic measurement)
- [5] ISO/TR 16312-2, Guidance for assessing the validity of physical fire models for obtaining fire effluent toxicity data for fire hazard and risk assessment — Part 2: Evaluation of individual physical fire models
- [6] ISO 16405, Room corner and open calorimeter Guidance on sampling and measurement of effluent gas production using FTIR technique
- [7] ISO 13344, Estimation of the lethal toxic potency of fire effluents
- [8] ISO 16312-1, Guidance for assessing the validity of physical fire models for obtaining fire effluent toxicity data for fire hazard and risk assessment - Part 1: Criteria
- [9] FARDELL P., & GUILLAUME E. Sampling and analysis of fire effluents, in Fire Toxicity (chapter 11) 1st edition, ISBN 1 84569 502 X, Woodhead publishers / CRC ed., 2010
- [10] ISO 29904, Fire chemistry Generation and measurement of aerosols
- [11] D'AGOSTINO R., & STEPHENS M.A. Goodness-of-fit techniques. Marcel Dekker Inc, 1986
- [12] STEPHENS M.A. EDF statistics for Goodness of Fit and some comparisons. J. Am. Stat. Assoc. 1974, 69 pp. 730-737
- [13] GUIDE EURACHEM The fitness for purpose of Analytical Methods A laboratory Guide to Method validation and related Topics
- [14] BLAND J.M., & ALTMAN D.G. Statistical methods for assessing agreement between two methods of clinical measurement. Lancet. 1986, 327 (8476) pp. 307-310
- [15] NIST/SEMATECH e-Handbook of Statistical Methods. http://www.itl.nist.gov/div898/ handbook/
- [16] Levene H. In: Contributions to Probability and Statistics: Essays in Honor of Harold Hotelling. (OLKIN I. eds.). Stanford University Press, 1960, pp. 278-92.
- [17] BROWN M.B., & FORSYTHE A.B. J. Am. Stat. Assoc. 1974, 69 pp. 364-367
- [18] WELCH B.L. The generalization of "Student's" problem when several different population variances are involved. Biometrika.  $1947, 34$  (1-2) pp. 28-35
- [19] ISO/IEC Guide 98-3:2008, Uncertainty of measurement Part 3: Guide to the expression of uncertainty in measurement (GUM:1995)
- [20] SCHWARZ G. Estimating the Dimension of a Model. Ann. Stat. 1978, 6 (2) pp. 461-464
- [21] HURVICH C.M., & TSAI C.-L. Model selection for extended quasi-likelihood models in small samples. Biometrics. 1995, 51 pp. 1077-1084
- [22] NF X 70-100-2:2006, Fire tests Analysis of gaseous effluents Part 2: tubular furnace thermal degradation method. French standard
- [23] Gu illaume illaume E . , Ath include Line and Society of Hardin Community of Hardin C . Analysis of Hardin during thermal degradation of materials. Adaptation of a standardized method. Proceedings of 13rd International Metrology Congress, Lille, France, June 18-21, 2007
- [24] Yardin C . , & Gu illaume E . Ana lyse de la produc tion de HC l , HBr et SO<sup>2</sup> lors de la dégradation thermique des matériaux - Adaptation d'une méthode standardisée. Traitements & Matériaux. 2010, 403 pp. 43-52
- [25] ISO 2602, Statistical interpretation of test results Estimation of the mean Confidence interval
- [26] ISO 2854, Statistical interpretation of data Techniques of estimation and tests relating to means and variances
- [27] ISO 5725-1, Accuracy (trueness and precision) of measurement methods and results  $-$  Part 1: General principles and definitions
- [28] ISO 13571, Life-threatening components of fire  $-$  Guidelines for the estimation of time to compromised tenability in fires
- [29] ISO/IEC 17025, General requirements for the competence of testing and calibration laboratories

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