

# codex alimentarius commission



FOOD AND AGRICULTURE  
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Agenda Item 6

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## JOINT FAO/WHO FOOD STANDARDS PROGRAMME

### CODEX COMMITTEE ON METHODS OF ANALYSIS AND SAMPLING

Twenty-seventh Session

Budapest, Hungary, 15-19 May 2006

### CONSIDERATION OF THE METHODS FOR THE DETECTION AND IDENTIFICATION OF FOODS DERIVED FROM BIOTECHNOLOGY

#### GENERAL APPROACH AND CRITERIA FOR THE METHODS

#### BACKGROUND

At the Twenty-fourth Session of the Codex Committee on Methods of Analysis and Sampling, papers giving the methods that had been collated by the *ad hoc* Intergovernmental Task Force on Food Derived from Biotechnology (see CX/MAS 02/8) and outlining general considerations of methods of analysis for the detection and identification of foods derived from biotechnology (see CX/MAS 02/9) were discussed. It was noted that the presence of genetically modified organisms or their derivatives could be assessed by the detection of either DNA sequences present as a result of recombination or the protein coded by the inserted gene. It was pointed out that protein-based methods were cheap, offered high selectivity and sensitivity but that since proteins were denatured during processing these techniques were most suitable for the analysis of raw materials and were not generally, unless antibodies are addressing resistant epitopes as observed in other domains of antigens detection, applicable to highly processed foods. It was also noted that these methods cannot be used when no new protein is expressed in the food, and these methods cannot differentiate between genetic events that produce the same protein.

Methods of detection of DNA markers based on the polymerase chain reaction (PCR) have been used in a variety of food analyses and widely used for detection of GM derivatives in food for many years, and modifications of the PCR method were also widely used. A typical method involved several steps such as sampling, extraction and purification, amplification by PCR and detection/quantification. Specific questions arising in the area of proficiency testing, use of performance criteria and the necessity of quantification due to threshold settings since the results of investigations showed the difficulties in measuring low levels of GM material in processed foods were also discussed. Methods described in the collated documents could only be used successfully if all information about the sequence and certified reference materials were available.

#### GENERAL CRITERIA

In view of the absence of precise provisions for GMOs and of difficulties with the practical application of methodology in this area, the Committee proposed to develop recommendations with respect to criteria for methods of analysis and for quality control measures that should be introduced in laboratories offering GM analyses. It was agreed that a Working Group led by Germany and the United Kingdom would update and further develop the paper for this session and prepare recommendations for quality control measures in laboratories and criteria for methods of analysis for the Twenty-fifth Session of CCMAS.

The paper CX/MAS 05/26/9 was discussed at the Twenty-sixth Session of CCMAS, where the following comments were made or were noted:

- The Committee recalled that at its last session it had agreed that the Delegations of the United Kingdom and Germany with the assistance of a Drafting Group would revise document with a view to the elaboration of Guidelines for consideration at the next session.
- The Delegation of Germany introduced the document and indicated that based on the comments received the following major changes had been made. In the Section on Modular Approach to method validation it was explained how this method could be applied and in Annex V on validation of a Protein-Based Method a new narrative was added.
- The Delegation of the EC supported the development of the paper and expressed the view that it had been elaborated for the endorsement of methods for detection and identification of foods derived from biotechnology in the CCMAS and proposed to send this paper to the Task Force on Biotechnology for their information.
- The Delegation Republic of Korea indicated that still there were some uncertainties in Table 1 on the Criteria for scoring Qualitative PCR analyses especially in expressing of the scoring of test when GM analyte in PCR was positive and endogenous PCR result was negative and proposed that the expression of “±” should be changed to “indeterminate” in the scoring of test expression.
- The Delegation of the United States supported the view expressed by the Delegation of Korea and indicated that it had provided general and detailed written comments presented in CCX/MAS 05/26/9-Add.1. The Delegation proposed that this document should be retained by the Committee until it had been improved and technical issues resolved. This view was supported by several delegations.
- The Delegation of Malaysia proposed to include a wider description of protein based testing as it was less costly and wider applied especially in developing countries.
- The Delegation of Brazil urged the Committee to proceed with this work as the matter of urgency as the trade in GMOs food was growing and there was a need for governments to receive advice on this matter.
- As regards to the status of the document, the Secretariat clarified that the Committee at its 24<sup>th</sup> session, following the request from the Committee on Food Labelling and the Task Force on Foods Derived from Biotechnology, had considered the methods of analysis for foods derived from biotechnology and had concluded that the criteria approach should be applied in the selection of methods of analysis for foods containing genetically modified material and that the selection or endorsement of methods without appropriate provisions was not possible. It was further agreed to prepare recommendations for quality control measures in laboratories and criteria for method of analysis. The Secretariat also indicated that the Intergovernmental Task Force on Biotechnology and the Committee on Food Labelling would be informed about the work of CCMAS in this area.
- The Committee agreed that a Working Group led by Germany and the United Kingdom with participation of all interested parties would revise the paper for consideration by the next Session of the Committee, especially in order to arrive to common understanding on how to proceed on this matter.

The following countries and organisations expressed their willingness to participate in this work: Argentina, Australia, Brazil, Canada, Egypt, France, Iran, Ireland, Italy, Japan, Malaysia, The Netherlands, Norway, Philippines, United States, European Commission, AOAC International, AOCS, Bio, CROPLIFE International, EUROPABIO, and ISO.

These measures are given as Guidelines in the Appendix to this paper.

## **RECOMMENDATIONS**

It is recommended that the draft Guidelines be discussed at the Twenty-seventh Session of CCMAS. If there is sufficient consensus, then the approaches described should be further refined and then sent for approval as a defined Codex “new work item”.

# APPENDIX I: GUIDELINES FOR THE VALIDATION AND QUALITY CONTROL REQUIREMENTS FOR THE ANALYSIS OF FOODS DERIVED FROM BIOTECHNOLOGY

## INTRODUCTION

### Method Criteria

The Codex Alimentarius Commission places an emphasis on the acceptance of methods of analysis which have been “fully validated” through a collaborative trial conforming to an internationally accepted protocol. In a number of sectors, including the foods derived from biotechnology (GMO) sector, there are few methods of analysis which have been fully validated. As a result, Codex is also endorsing by reference single-laboratory validation protocols. In this area there may be pressure to adopt a formal single-laboratory validation as an interim measure in the absence of collaborative trial data. However, methods used for determination of the presence of GMO's are able to be, and intended to be performed at, multiple laboratories and should therefore be validated by multi-laboratory collaborative studies as soon as practicable.

In these Guidelines the term “GMO” has been used for “Foods Derived from Biotechnology”.

Many methods are currently being developed for GMO detection, identification and quantification. Before they are accepted for use by Codex they must be validated to ensure that they are fit-for-purpose.

However, the two most common approaches are those based on DNA-based methods and those based on the detection of proteins themselves or their activities. The former is generally performed via PCR, although other methods that achieve measurement without a PCR step may be employed if properly validated. Both DNA and protein-based approaches are considered here, though it is the DNA-based PCR approach which is generally recognised as being the more widely applicable.

The conventional criteria that have been adopted by Codex for the evaluation of methods of analysis are:-

- accuracy
- applicability (matrix, concentration range and preference given to 'general' methods)
- limit of detection
- limit of determination
- precision; repeatability intra-laboratory (within laboratory), reproducibility inter-laboratory (within laboratory and between laboratories)
- recovery
- selectivity
- sensitivity
- linearity

These Guidelines address these requirements in the GMO sector, and anticipates that is likely that these will have to be further expanded (e.g. for PCR) by other items such as:-

for the DNA-based methods.

- amplicon length
- whether the method is instrument specific
- whether there are differences between qualitative and quantitative PCR-based detection methods
- whether single- or multi-plex PCR amplifications are undertaken

and

for the protein based methods

- equivalency of reagents over time

The method validation process accepted by Codex includes the definition of the requirements for the method, testing that the method meets these requirements when carried out, for instance, by different laboratories in different countries, and documentation of the method performance and measurement uncertainty.

### Criteria Approach

Codex Alimentarius Commission has accepted the “criteria approach” for methods of analysis. This approach does not extend to Codex Type I empirical/defining, procedures. It is necessary to ensure that this approach is incorporated into Codex guidelines on the validation of GMO methods of analysis unless it is explicitly stated that all GMO methods of analysis are empirical, both theoretically as well as in practice.

### **Laboratory Quality**

The Codex Alimentarius Commission has adopted guidelines for the “quality” of laboratories involved in the import and export of foods. These quality characteristics are based on accreditation to ISO/IEC Standard 17025, proficiency testing and internal quality control as well as the use of methods of analysis validated according to Codex requirements. These overarching guidelines provide information to and dictate requirements for laboratories working in the GMO sector.

### **Measurement Uncertainty**

Codex has developed guidelines on Measurement Uncertainty. These guidelines, as well as the accreditation requirements cited above, require laboratories to estimate the uncertainty of their quantitative measurements. This is particularly important and has consequences for measurements in the GMO sector where analytical controls may not be as effective as found in other areas of analysis in the food sector. It is frequently not appreciated that the magnitude of the measurement uncertainty is considerably greater in this analytical sector than would normally be expected.

### **INFORMATION TO BE PROVIDED TO CODEX WHEN A METHOD FOR GMOs IS TO BE CONSIDERED FOR ENDORSEMENT BY CCMAS**

The information that should be supplied to CCMAS when a method is to be considered for endorsement is given in Annex I. The annex lists both general considerations and specific requirements.

As GMO methodology becomes more developed the specific requirements will be converted to performance criteria to conform to the “criteria approach” already adopted by Codex.

### **DEFINITIONS**

There are a number of Codex definitions applicable to GMO analysis. Suggested definitions are given in Annex II.

### **METHOD DEVELOPMENT TO FORMAL VALIDATION**

#### **Applicability of the Method**

This is a particularly important criterion in GMO analysis. In principle the method should be applicable to the matrix of concern within the Codex system. If this is a specific product derived from GMO then there is merit in requiring those seeking endorsement to provide information on the method of analysis appropriate to the specific product and, ideally, the matrix in which it is likely to be used. In case of “general purpose” GMO methods, at least one extraction method applicable to a general matrix should be available.

As an example it is required from an extraction method, independent of matrix to which it is to be applied, that it yields DNA of sufficient quantity, structural integrity and purity to allow a proper evaluation of the performance of the subsequent method steps (e.g. adequate amplification of DNA during the PCR step) to be undertaken. This can be tested, for example, by setting up dilution series of the template DNA and determining that the  $\Delta CT$  in a real-time PCR analysis between the dilutions corresponds to the dilution factor, e.g. if DNA is diluted 10X then the  $\Delta CT$  should be approx. 3.32, if the DNA is diluted 4X, the  $\Delta CT$  should be 2, etc. Statistically significant deviations from this relationship may indicate that the extracted DNA contains PCR inhibitors, that the DNA solution is not homogenous or the DNA quantity so low that stochastic variation in copy numbers yield unreliable quantitative estimates.

#### **Validation Process**

Method validation is a process of establishing the performance characteristics and limitations of an analytical method and the identification of the influences, which may change these characteristics - and to what extent. The results of a validation process describe which analytes can be determined in what kind of matrices in the presence of which interference. The validation exercise results in precision and accuracy values of a certain analytical method under the examined conditions.

Formal validation of a method is the conclusion of a long process, which includes the following main steps:

- **Method development and optimisation.** Prior to any pre-validation, the method should be fully optimised so that an inter-laboratory transfer is possible. The protocol should be finalized so that no major changes are needed between the pre-validation and validation.
- **Pre-validation of the method.** Pre-validation should ensure that a method performs in a manner, which allows a successful conclusion of the validation study, i.e. it should provide evidence about the compliance with the requested performance or regulations. Pre-validation should preferably be carried out by involving 2 - 4 laboratories. Statistical analyses (e.g. of “repeatability” and “reproducibility”) should be made according to the validation procedure to be subsequently used.
- **Full validation of the method.** Full validation requires considerable resources and should be conducted only on methods which have received adequate prior testing.

A collaborative trial is expensive to undertake and usually follows only after the method has shown acceptable performance both in a single-laboratory and a pre-validation study.

### **Modular Approach to Method Validation**

The “method” refers to all the experimental procedures needed to estimate the measurand in a particular matrix. For a particular material this may include the methods for DNA extraction and the final quantification in a PCR system. In such a case, the whole chain from extraction up to the PCR-method (or equivalent) constitutes a method, but the different method parts can be considered separately (i.e. modular validation). The theoretical advantage of a modular approach to method validation is that each section of a method or protocol can be validated separately, and once validated, can be combined with other sections in a flexible manner.

A modular approach to method validation assumes that the modules in a method which form the whole are independent of each other.

Where validation for an “official control” method is to be undertaken, a modular approach should not be taken until independence between the modules has been demonstrated and documented. This independence will be more difficult to establish the more variable each module is and the method as a whole will be critically examined when it is presented to Codex for endorsement.

### **METHOD ACCEPTANCE CRITERIA**

In order to evaluate a method prior to full validation, information concerning both the method and the method testing is required. Information on this is given in Annex I.

The method will be evaluated based on the information provided to Codex. The evaluation should verify that the principle preconditions for using the method for Codex purposes are fulfilled. This section describes the method acceptance criteria, which have to be fulfilled by the method in order to conduct further a pre-validation and full collaborative trial.

#### **Principle Conditions**

The provision of the detection method is aimed to serve mainly the requirements for the monitoring and labelling of GMOs, as set out in the specific regulations. To serve these purposes, the method should detect and quantify the specific GM event in the GM product; this may be achieved in most cases using either protein-based or DNA-based methods.

Currently, the DNA-based detection method typically consists of PCR methodology and includes:

- a protocol describing an extraction method which is applicable to a relevant matrix;
- a description of the oligonucleotide primer sequences which uniquely identify the GM event in the GM product<sup>1</sup>;
- a description of the oligonucleotide primer sequences which amplify an endogenous gene sequence applicable to the specific host species;

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<sup>1</sup> Note: the fact that most event-specific sequences are not publicly disclosed should be discussed by CCMAS.

- a protocol describing the conditions, including the apparatus used, under which PCR can be used to detect the GM product;
- appropriate control samples, when available. However, the detection of an unknown GMO does not allow the use control samples since they are generally not available.

The method provider should demonstrate that the method fulfils the principle method requirements:

- (1) **GMO Screening Methods.** In the case of a method used for screening for the presence of multiple events, the method should be specific and allow for unequivocal detection/identification/ quantification of a specific DNA sequence in the case of DNA-based methods. In the case of Protein based methods, the method should be specific and allow for unequivocal detection/identification/ quantification of a specific antigen or epitope.
- (2) DNA-based event-specific methods should allow for unequivocal detection/identification/ quantification of a known target nucleotide sequence.

Currently, the best choice concerning event-specificity of a method, should PCR be the chosen technique, is to target an event-specific genomic region using a set of oligonucleotides (primers) that trigger the amplification of such a region. Among various types of event-specific genomic regions, the one relative to the junction between the transgenic insert and the host genomic DNA will probably be the location of choice. However, when a unique DNA sequence can be found within the transgenic insert, such a sequence (generally called construct specific) can also be targeted by appropriate oligonucleotide primers and amplified through a PCR. Identification of the amplified fragment, by e.g. probe hybridization or any appropriate equivalent method, is recommended.

- (3) All methods should be applicable to the material specified in their scopes, and to appropriate quality control and reference materials when available.

It should be noted that at present only relative quantitation, can be carried out which means that the transgenic material relative to the corresponding ingredient/species is measured.

## **COLLABORATIVE TRIAL REQUIREMENTS**

### **General Information**

The purpose of a collaborative trial is to fully validate the data provided by previous testing in a pre-validation or a single laboratory exercise and to determine methodological precision in terms of repeatability and reproducibility.

The values of any performance parameters reported from validation studies must be interpreted and compared with care. The exact values and their interpretation may depend – besides the performance of the method - on the extent of the method (e.g. a real-time quantitative PCR only versus a method chain ranging from extraction to the real-time PCR quantification), experimental design applied, e.g. calibration *vs.*  $\Delta\Delta Ct$  methods, exact calculation forms used to determine the parameters and the approach used to detect and analyse outliers. In order to have meaningful “minimum performance requirements” the above factors must be treated appropriately and in a standardized manner.

For Codex purposes the ISO/AOAC/IUPAC harmonized protocol<sup>1</sup> has been adopted.

### **Minimum Performance Requirements**

In a collaborative trial, the method performance should comply with the relevant parts of the method acceptance criteria and fulfil the method performance requirements specifically set below for the collaborative trial. Thus, the collaborative trial should confirm the results obtained during the previous method evaluation phases and should provide additional information about the method performance in a multi-laboratory setting. In particular, the compliance with the criteria for sensitivity and repeatability/reproducibility standard deviations should be assessed.

In addition to the method acceptance criteria, at least the method performance requirements listed in Annex I should be evaluated from the experimental data of a collaborative trial. First, the definition and thereafter the requirements are described.

The endorsed methods and their associated validation data will be revised on a regular basis as the scientific knowledge and experience gained in Single-Laboratory validation and collaborative trials evolve. These Guidelines will also be complemented with practical information about the operational steps of the validation process.

### **Collaborative Trial Test Materials**

In principle, the method should be applicable to and tested on the matrix of concern (i.e. on which any specification has been made).

In other fora recommendations have been made that in case of “general purpose” GMO procedures (in contrast to consideration of a specific product derived from GMO) that the validation of the detection module is carried out using genomic, or plasmidic DNA as the analyte (for a PCR-based method) for calibration using control samples). This allows the detection step to be combined with various extraction methods applicable to different matrices. However, real materials/matrix typical of a type/group of matrices are preferred unless the effects of the materials/matrix on DNA quality in the extraction step has been evaluated on a series of matrices prior to applying a modular approach. Otherwise a modular approach is inappropriate when considering Codex specifications.

### **VALIDATION OF PCR METHODS**

Specific information on the validation of quantitative and qualitative PCR methods is given in Annexes III and IV respectively.

Specific information on the validation of quantitative and qualitative protein-based methods is given in Annex V.

### **UNITS OF MEASUREMENT**

Various countries have thresholds established for labelling of food and feed derived from modern biotechnology. These thresholds are explicitly expressed as weight or kernels by relative percentage. However, none of the current detection methods (DNA – or protein-based) are able to measure this directly. Although there are correlations between kernels / weight-% and the amount of DNA or protein, respectively, the very nature of these relationships is influenced by a number of biological factors and thus remains highly variable. This continues to cause considerable misunderstanding and requires significant technical guidance.

Based on the PCR technique used for GMO identification and quantification genome equivalents are measured.

Therefore it is not trivial to consider how the genetically modified material is calculated. For example, if a maize seed lot containing 2% genetically modified seeds with the “new” trait in a hemizygous state (coming from the pollen) is used to prepare a flour sample then, in theory, only 0.29% of the isolated genomic DNA copies will represent the genetically modified status. This is due to the different tissue types, the source from where the genomes in these tissue types are derived (maternal or paternal) and the contribution of the tissue types in the seed kernel. Consequently the amount of genetically material would be underestimated (on a seed basis), or overestimated in case of gene stacking, by a DNA and a protein based approach to express the content of material derived from genetically modified organisms.

Quantitation based on the “newly” expressed protein in the GMO would also lead to a significant contribution to the uncertainty of the analysis. For example the environment in which the material was grown can affect the amount of protein expressed. In addition, it is often the case that the protein is expressed at different levels in different tissue types of the plant as well as in different cultivars of the same transformation event. Consequently foods produced from different parts of a genetically modified plant or from different cultivars would contribute a different amount of the “newly” expressed protein.

This issue needs to be appropriately addressed and conversion factors or any appropriate mean such as agreed consensus GMO units, performance and data reporting criteria established for these methods.

For relative quantification, as far as possible a consensus agreed reference gene should be used for the same plant species.

### **MEASUREMENT UNCERTAINTY**

Analysts using methods which have been validated according to these guidelines will have available to them sufficient information to allow them to estimate the uncertainty of their result.

Guidance on both the estimation and use of any measurement uncertainty estimation has been/are being developed and adopted by Codex<sup>2,3</sup>.

An amount of work has already been carried out and published on the validation of a number of specific methods of analysis. For convenience these are summarised in Annex VI.

## **GUIDANCE ON LABORATORY SET-UP AND OPERATION**

DNA-based methods for the analysis of foods derived from modern biotechnology apply techniques that are not considered as commonly available methods, as they currently require specific apparatus and handling techniques that differ from most chemical-analytical methods. However, the use of DNA based is consistently growing in other detection fields such as microbiology of food pathogens. In the current absence of feedback of other detection domains, it is therefore necessary to provide information and instructions on the essential differences in laboratory set-up and handling techniques. Examples are available<sup>4</sup>.

## **REFERENCE MATERIALS**

There are a number of matrices that can be used to develop reference materials or working standards for methods of detection of GM products. Each has its own advantages and drawbacks for particular purposes. However, in some cases as the detection of an unapproved / unknown GMO, the reference material will be, by nature, not available.

Codex may consider requiring the availability of suitable reference materials as part of the method endorsement procedure. However, it is recognised that there are specific problems with the development of reference materials, e.g. for maize materials should the maize event or the construct specific methods be considered.

A suitable reference material is generally required for validation of a method. Suitable reference materials are becoming available for many commercialized events. Where they are not available, the availability of quality control materials from proficiency testing schemes or from the use of Plasmid or amplicon DNA may be considered. This area of work needs further development.

## **SAMPLING**

In the area of GMO analysis it may be anticipated that sampling error can be expected to contribute significantly – if not dominate - the overall uncertainty of an analytical result, particularly when considering raw commodities. However, this is also found to be the case in other areas of “analysis”, most notably the estimation of the mycotoxin concentration in a bulk, or even the estimation of the nitrate content of a greenhouse. The combination of sampling and analytical uncertainties is now being addressed by a number of International Organisations, most notably EURACHEM which has set up a new Working Group dealing with uncertainty of sampling. Much work has been carried out on sampling generally by CCMAS<sup>5</sup> and of bulk sampling for GMOs by the EU JRC<sup>6</sup>, ISO/CEN<sup>7</sup> and GIPSA.

## **CONCENTRATION DISTRIBUTIONS**

Conventionally concentrations in the GMO area are considered to be normally distributed. However, a study of data from two GMO proficiency schemes has been carried out. This considered data from a total of 29 rounds and 43 test materials over a period of three years. The results from the two schemes are similar and reinforce each other. The amplification process used in quantitative PCR determinations predicts a mixture of normal, binomial, and lognormal distributions dominated by the latter two. As predicted, the study results consistently follow a positively skewed distribution. Log-transformation prior to calculating z-scores is effective in establishing near-symmetric distributions that are sufficiently close to normal to justify interpretation on the basis of the normal distribution<sup>8</sup>. Consideration should therefore be given as to whether all data from quantitative determinations should be log transformed before use. The consequence for proficiency testing schemes is outlined in Annex VII.

## REFERENCES

1. ISO/AOAC/IUPAC harmonized protocol (Protocol for the Design, Conduct and Interpretation of Method-Performance Studies, Ed. Horwitz, Pure & Appl. Chem. 331-343, **67**, 1995)
  2. Guidelines on Measurement Uncertainty (CAC/GL 54-2004)
  3. Codex Draft Guidance Document on “*The Use of Analytical Results: Sampling Plans, Relationship between the Analytical Results, the Measurement Uncertainty, Recovery Factors and Provisions in Codex Standard*”
  4. Draft ISO-standard (ISO/DIS 24276) or the corresponding French standard (AFNOR XP V03-020-2, tabled as room document CRD 5 in its previous version AFNOR XP V03-020-1 by the French Delegation at the 24<sup>th</sup> Session of CCMAS)
  5. Codex General Guidelines on Sampling (CAC/GL 50-2004)
  6. FP5 KeSTE project.
  7. prEN ISO 21568
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8. Scoring in Genetically Modified Organism Proficiency Tests Based on Log-Transformed Results, Michael Thompson, Stephen I. R. Ellison, Linda Owen, Kenneth Mathieson, Joanne Powell, Roger Wood and Andrew P Damant, Journal of AOAC International, 2006, 89(1), 232- 239

## **ANNEX I: INFORMATION TO BE PROVIDED TO CODEX WHEN A METHOD IS TO BE CONSIDERED FOR ENDORSEMENT BY CCMAS**

In order to aid the endorsement of a proposed method of analysis in the GMO sector by Codex, and in particular CCMAS, the following should be provided:

### **DESCRIPTION OF THE METHOD**

A complete and detailed description of all the components of the method should be provided. The use of multiple plates for PCR and protein methods, as an example, should be explicitly addressed. The information should also include information on the following:

#### ***Purpose and relevance of the method***

The objective of the method and the relevance of the method with respect to relevant legislative requirements should be indicated. In particular, the proposer should indicate that the principle conditions for the method are fulfilled.

#### ***Scientific basis***

An overview of the principles of how the method, such as DNA molecular biology based (e.g. for real-time PCR) information should be provided. References to relevant scientific publications are useful.

The prediction model adopted to interpret results and to make inferences must be described in complete detail.

#### ***Specification of the prediction model/mathematical model needed for the method***

If the derivation of the results relies upon a mathematical relationship this must be outlined and recorded (e.g.,  $\Delta\Delta C_t$  method or a regression line or calibration curve obtained by other means). Instructions for the correct application of the model should be provided. These may include, depending on the method, a recommended number and range of levels to be analysed, minimum number of replicates and/or dilutions to be included for routine analyses or the means and confidence intervals to evaluate the goodness-of-fit.

Outline of the experimental design used for validation and for routine analyses, including the details about the number of runs, samples, replicates, dilutions etc. should be stated.

### **INFORMATION ABOUT THE METHOD OPTIMISATION**

#### ***Primer pairs tested***

For PCR methods, sufficient justification should be given of how and why the proposed primer pair has been selected; also for the reference gene (should this be part of the method).

#### ***Specificity testing***

Empirical results from testing the method with non-target transgenic events and non-transgenic plants should be provided. This testing should include closely related events and cases where the limits of the sensitivity are truly tested. In addition it might be appropriate, particularly for reference genes, to test other plants to reduce the potential for obtaining a false positive.

#### ***Stability testing***

Empirical results from testing the methods (to detect reference genes but also GMOs) with different varieties should be provided in order to demonstrate, for instance, the stability of the copy number and sequence conservation of the reference gene.

### ***Sensitivity testing***

Empirical results from testing the method at different concentrations in order to test the sensitivity of the method. Limits of detection must be defined using samples comprising of single crops only, e.g. “the LOD for Roundup Ready® soy is 0.1 % of total soy *if the product is comprised of 100 % soy*”. For food products made up of multiple ingredients, the actual sensitivity will be reduced, as total extracted DNA will be derived from more than one ingredient so that the starting amount of the actual measurand will be decreased. This dilution effect will depend on how much of the target ingredient (e.g. soy) is in the food product and the total quantity of DNA derived from the other ingredients. Some ingredients will contribute a large amount of DNA, such as wheat or maize flour and eggs, while other ingredients will not contribute any DNA, such as refined sugar, pure water or highly processed oils.

LOD should be determined in terms of haploid genome equivalents for each PCR system separately.

### ***Robustness testing***

Empirical results from testing the method against small but deliberate variations in method parameters.

### ***Cross-reactivity***

The cross-reactivity, interferences and matrix effects should be evaluated, particularly for the protein-based methods of analysis or enzymes based tests such as PCR or LCR.

## **PRACTICAL APPLICATION OF THE METHOD**

### ***Applicability***

Indication of the matrix (e.g., processed food, raw materials, etc.), the type of samples (e.g., seeds, flour, pizza, cookies, etc.) and the range to which the method can be applied. Relevant limitations of the method should also be addressed (e.g. inference by other analytes or inapplicability to certain situations). Limitations may also include, as far as possible, possible restrictions due to the costs, equipment or specific and non-specific risks implied for either the operator and/or the environment.

### ***Operational characteristics and practicability of the method***

The required equipment for the application of the method should be clearly stated, with regards to the analysis *per se* and the sample preparation. An indication of costs would be particularly useful for Codex purposes, as well as other practical difficulties, and of any other factor that could be of importance for the operators should be also indicated.

### ***Operator skills requirements***

A description of the practical skills necessary to properly apply the proposed method should be provided.

## **ANALYTICAL CONTROLS**

The proper use of controls, when available, when applying the method should be indicated. Controls should be clearly specified and their interpretation recorded. These may include positive and negative controls, their detailed contents, the extent into which they should be used and the interpretation of the obtained values. However, it is recognized that control samples may be not available particularly on the case of unknown / unapproved / being approved GMOs.

In particular the following should be stated:

- Positive and negative controls used
- Control samples, plasmids and alike used

- Reference materials used, when applicable.

## METHOD VALIDATION/PERFORMANCE

See the Codex “Check-list” (i.e. accuracy, applicability (matrix, concentration range and preference given to 'general' methods), detection limit, determination limit, precision, recovery, selectivity, sensitivity and linearity) and an assessment that the methods will be fit for purpose.

and in particular the following additional information should be supplied for DNA-based procedures:

- ***amplicon length and sequence***

*The boundaries of the amplified product are formed by the primers at both sides. Therefore the selection of suitable primers is a crucial factor in the PCR analysis. The length of the amplified product does have a direct influence of the PCR performance. By increasing the product length, the PCR efficiency will decrease reciprocal as illustrated below (Fig. 1). In theory in every cycle the target DNA sequence is doubled (amplification factor of 2). In reality the PCR efficiency is less than 100% resulting in a decreased amount of amplified product. Moreover, food processing will generally lead to a degradation of target DNA. Therefore the selection of shorter amplicon sizes (within reason) will increase the possibility to get a positive signal in the analysis of highly processed foodstuffs.*

Amplification factor	2	1.65	1.60	1.55	1.50	1.45
Target copies after:						
10 cycles	10 <sup>3</sup>	150	110	80	58	41
20 cycles	10 <sup>6</sup>	2.2x 10 <sup>4</sup>	1.2x10 <sup>4</sup>	6.4x10 <sup>3</sup>	3.3x10 <sup>3</sup>	1.7x10 <sup>3</sup>
30 cycles	10 <sup>9</sup>	3.3x10 <sup>6</sup>	1.3x10 <sup>6</sup>	5x10 <sup>5</sup>	1.9x10 <sup>5</sup>	7x10 <sup>4</sup>
40 cycles	10 <sup>12</sup>	5x10 <sup>8</sup>	1.5x10 <sup>8</sup>	4.1x10 <sup>7</sup>	1.1x10 <sup>7</sup>	2.8x10 <sup>6</sup>
	100%	82.5%	80%	77.5%	75%	72.5%

Figure 1 PCR efficiency. A decrease of efficiency in PCR leads to lower amounts of amplified products being present after a certain number of cycles.

- ***whether the method is instrument or chemistry specific***

*At the moment a number of different types of real time instruments and chemistries are available. These instruments and chemistries may have different performance such as stability of reagents, heating and cooling characteristics, which affects ramp rates and affects the time necessary for a whole PCR run.*

*Beside the differences in the heating and cooling system there are differences in the technique and software used to induce and subsequently to record the fluorescence. Some real time instruments use laser technique for inducing fluorescence, others are equipped only with a halogen lamp and filters for selecting a specific wave length. The detection and quantification of the fluorescence could also vary according to the recording instruments and software used.*

*Taking all the differences into account it is impossible to change the instrument without adaptation of the PCR method. Thus, because the methods are generally instrument and chemistries dependent they cannot be transferred to other equipment and chemistries without evaluation and/or modification.*

*This is in many ways equivalent to the Codex Type I method and should be considered in the same light.*

- ***whether single- or multi-plex PCR amplifications are undertaken***

*Using more than one primer set in a single reaction is called multi-plex PCR. The aim of using such approach is to reduce costs and time for the analysis of different targets of a single sample (e.g.. a GMO specific system is combined with a target taxon specific for relative quantitation. Unless appropriate optimization of the multiplex has been performed, it must be emphasised that the unbalanced presence of one of the target sequences will lead in a preferred amplification by the polymerase during PCR. Moreover the combination of different primer sets is limited up to 7 to 10 in a single reaction.*

*The information provided should demonstrate the robustness of the method for inter-laboratory transferability. This means that the method should have been tested by at least one other laboratory besides the laboratory which has developed the method. This is an important pre-condition for the success of the validation of the method.*

And for both protein and DNA based methods:

- ***whether there are differences between PCR-based and immunological methods concerning validation criteria***

The DNA and protein-based techniques used to detect and quantify a GMO derived material in foods are based on different principles. In PCR the targeted DNA is amplified in a exponential manner, in which a small difference in the beginning of the PCR process will lead to a big difference in the amplified amount of DNA after 35-45 cycles. In contrast to that, immunological detection assays are based on the direct interaction with the target molecule and do not include an amplification step.

Moreover, the quantitation by real time PCR is often based on two independent PCR systems: one for the genetic modification and one for the taxon specific sequence.

## ANNEX II: CODEX DEFINITIONS APPLICABLE TO GMO ANALYSIS

This Annex is concerned with the definitions needed in GMO analysis. *(Note: a number of definitions have been grouped together in one heading; these may be contradictory and this needs to be resolved. The Codex definition given in the Procedural Manual should be used and amplified as necessary. Codex definitions have not been reproduced here if they need no further qualification for GMO analysis).*

### Accuracy

The closeness of agreement between a reported result and the accepted reference value<sup>2</sup>.

### Applicability

The analytes, matrices and concentrations for which a method of analysis may be used<sup>3</sup>.

The analytes, matrices, and concentrations should be appropriate for the control purposes for which the method has been proposed. The description may also include warnings to known interferences by other analytes, or inapplicability to certain matrices and situations.

It is not feasible to provide reference materials for every one of the many food matrices that are available, so that the use of a representative matrix reference will usually be necessary. The use of the method in a new matrix will need to be validated at a minimum via Single Laboratory validation, usually by spike and recovery experiments, and the reference material, when available, used should be described on the report to the customer.

### Dynamic Range - Range Of Quantification

The interval of concentration within which the analytical procedure has been demonstrated by collaborative trial to have a suitable level of precision and accuracy.

### Limit of Detection (LOD)

Limit of detection is the lowest concentration or content of the analytes that can be detected reliably, but not necessarily quantified, as demonstrated by collaborative trial or single-laboratory validation<sup>4</sup>. Alternatively it may be taken from the last value with reliable data used to determine the LOD. LOD is generally expressed as the amount of analyte at which the analytical method detects the presence of the analyte at least 95% of the time ( $\leq 5\%$  false negative results).

### Limit of Quantification (LOQ)

The limit of quantification of an analytical procedure is the lowest amount or concentration of analyte in a sample, which can be quantitatively determined with an acceptable level of precision and accuracy as demonstrated by satisfactory collaborative trial or single-laboratory<sup>5</sup> validation<sup>6</sup>. Alternatively, it may be taken from the last value with reliable data used for determining the LOQ.

### Practicability

The ease of operations, in terms of sample throughput and costs, to achieve the required performance criteria and thereby meet the specified purpose<sup>7</sup>.

Generally, the method should be practical for its intended purposes.

### Repeatability standard deviation (RSD<sub>r</sub>)

The standard deviation of test results obtained under repeatability conditions. Repeatability conditions are conditions where test results are obtained with the same method on identical test items in the same laboratory by the same operator using the same equipment within short intervals of time.<sup>8</sup>

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<sup>2</sup> Definition adopted from ISO 3534-1.

<sup>3</sup> Slightly modified from the definition provided in Codex CX/MAS 02/4: Proposed draft guidelines for evaluating acceptable methods of analysis. Version November 2002.

<sup>4</sup> Slightly modified from prEN ISO 24276:2002 (E).

<sup>5</sup> E.g. Thompson et al. 2002. IUPAC Technical Report: Harmonised guidelines for single-laboratory validation of methods of analysis. Pure Appl. Chem. 74(5): 835-855.

<sup>6</sup> Slightly modified from prEN ISO 24276:2002 (E).

<sup>7</sup> Adopted from prEN ISO 24276:2002 (E).

### **Reproducibility standard deviation (RSD<sub>R</sub>)**

The standard deviation of test results obtained under reproducibility conditions. Reproducibility conditions are conditions where test results are obtained with the same method on identical test items in different laboratories with different operators using different equipment.<sup>9</sup>

### **Recovery**

Proportion of the amount of analyte, present in or added to the analytical portion of the test material, which is extracted and presented for measurement.

### **Ruggedness (Robustness)**

Robustness refers to variations in the method as performed in different laboratories by different ‘technicians’. The language used here is derived from “Ruggedness” which is the equivalent in the harmonized guidelines. Ruggedness should be demonstrated by the validation of the method in 8-12 laboratories as defined in the harmonized guidelines. It is preferable from a CODEX point of view, that these laboratories be distributed across several continent/trading blocks.

The robustness of an analytical method is a measure of its capacity to remain unaffected by small, but deliberate variations in method parameters and provides an indication of its reliability during normal usage<sup>10</sup>.

### **Sensitivity**

The sensitivity of a method is a measure of the magnitude of the response caused by a certain amount of analyte.

The method should be sensitive enough in order to be able to detect/quantify with respect to the thresholds as provided in the relevant legislation.

Since sensitivity is method- and purpose-dependent it should be specified in the protocol. A reasonable goal for sensitivity is that required to meet levels specified in contracts, with a reasonable certainty that the level does not exceed the required limit.

Sensitivity as a term is used in two different ways - LOD and the slope of a curve. The use of “detection limit”, or “limit of detection” is the preferred term to use as a measure of the ability of a method to detect a small amount of analyte. See also previous comments regarding sensitivity in this document.

### **Specificity**

Property of a method to respond exclusively to the characteristic or analyte of interest.

### **Trueness**

The closeness of agreement between the average value obtained from a large series of test results and an accepted reference value<sup>11</sup>.

The measure of trueness is usually expressed in terms of bias. Trueness has also been referred to as “accuracy of the mean”.

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<sup>8</sup> Definitions adopted from ISO 3534-1.

<sup>9</sup> Definitions adopted from ISO 3534-1

<sup>10</sup> Definition adopted from ICH Topic Q 2 A “Validation of analytical methods: definitions and terminology.” The European Agency for the evaluation of medicinal products. CPMP/ICH/381/95. Version November 1994.  
<http://www.emea.eu.int/pdfs/human/ich/038195en.pdf>

<sup>11</sup> Adopted from ISO 3534.

## **ANNEX III: VALIDATION OF A QUANTITATIVE PCR METHOD**

### **INTRODUCTION**

DNA-based analysis is commonly performed using Polymerase Chain Reaction (PCR). This technique amplifies a specific (short) segment of DNA to the extent that its quantity can be measured instrumentally (e.g., using fluorometric means). As DNA is a molecule that is easily degraded during food processing operations (e.g., due to heat, enzymes and mechanical shearing), we urge that this be considered in the performance criteria assessment of this technique. This is relevant as in most foods raw ingredients are not present, but are in a processed form, which has an effect on proteins and/or DNA present in food. Furthermore, these protein(s) and/or DNA may be degraded, or its total amount may be decreased due to processing. As a result, any current detection method (DNA- or protein-based) is affected.

It is often the case that the results of a determination are expressed in terms of percent of a sample that contains a particular biotechnology-derived sequence. In a quantitative test, this measurement actually involves two PCR-based determinations – that of the primary analyte (e.g. an inserted gene sequence) and that of the endogenous, or comparator sequence (e.g. an endogenous maize gene). Each of these determinations has its own uncertainties, and the two are likely to have different measurement characteristics. In most applications, the primary analyte will be present at low concentrations, and the comparator will be present at concentrations 10 to 1000 times higher. It is thus important that both measurements are properly validated. In cases where the measurement is expressed directly as a percentage (as in the use of  $\Delta CT$ ), these factors must be considered when validating the method.

The consequence is that the analysis of DNA, especially in processed foods, aims at detecting a very small amount of analyte. Although the result of a PCR analysis is often expressed in % as the relative amount of DNA specific for foods derived from modern biotechnology relative to the total amount of DNA for a specific species, the actual amount of DNA specific for foods derived from modern biotechnology is often in the nanogram/gram range or lower. Analysis of those low amounts of analyte is accompanied by a considerable measurement uncertainty; this needs to be appreciated by the users of analytical results.

### **VALIDATION**

A quantitative PCR assay should be validated for the intended use or application. A harmonized ISO/IUPAC/AOAC protocol was developed for chemical analytical methods. This defines the procedures necessary to validate a method (Horwitz W; Protocol for the design, conduct and interpretation of method-performance studies. Pure and Applied Chemistry, 67, 331 (1995)). It is important to emphasize that all the principles and rules of the harmonized protocol are applicable to quantitative PCR methods.

A number of the parameters involved in validation of the performance of a quantitative PCR assay will be discussed in detail. These are scope, LOD and LOQ, accuracy, precision, sensitivity and ruggedness (robustness). Other important factors are acceptance criteria and interpretation of results, and the issue of the units in which results are expressed.

It is also the case of samples versus reference material which have been are generally different matrices extracted separately. Thus, all parameters listed below, including specificity and sensitivity, have to be assessed individually for each of the assays involved. These are given alphabetically, not necessarily in order of importance.

#### **Accuracy**

As for any method, the accuracy of a method should be compared to known values derived from reference materials, ideally the best characterised. Precision will be determined in the usual way from single laboratory (repeatability) and multi-laboratory (reproducibility) studies. However, the impact of the difference of matrices between sample to be analyzed and reference material has to be considered.

*Recommendation: The accuracy should be within [ $\pm 35\%$ ] of the accepted reference value over the whole dynamic range.*

## **Applicability**

The analytes, matrices and concentrations for which a method of analysis may be used must be stated.

## **Dynamic Range - Range Of Quantification**

The scope of the methods defines the concentration range over which the analyte will be reliably determined. Typically the range for a GM product will range from a tenth of a percent up to a few percent and for the endogenous control the range will be close to 100%, unless the testing of complex mixtures is envisioned. This desired concentration range defines the standard curves and a sufficient number of standards must be used, when applicable e.g. with calibration curves, to adequately define the relationship between concentration and response. The relationship between response and concentration should be demonstrated to be continuous, reproducible and should be linear after suitable transformation.

The range of a quantitative method is typically designed to be in the range 0.1% - 100% (DNA %/ molecules / molecules, haploid genomes / haploid genomes). However, it is common to validate a method for a range of concentrations that is relevant to the scope of the application. If a method is validated for a given range of values, the range may not be extended without further validation. For certain applications (e.g. food or grain analysis) the use of genomic DNA for the preparation of the standard curve (see discussion on the use of plasmid DNA below) may be considered. While it is easy to establish a nominal 100% standard (limited only by the purity of the plant materials used) it is difficult to reliably produce standard solutions below 0.1%. This is due to the uncertainties involved in measuring small volumes and the error propagation if serial dilution steps are applied. Additionally, the number of target sites (DNA sequence to be amplified) becomes so small that stochastic errors will begin to dominate and less reliable analysis is possible<sup>1, 2</sup>. If DNA (genomic or plasmidic) is chosen to be used as calibrator, it is important that this calibrator needs to be traced back (in its metrological meaning) to a reference of highest metrological order, e.g. a certified reference material. The range will be established by confirming that the PCR procedure provides an acceptable degree of linearity and accuracy when applied to samples containing amounts of analyte within or at the extremes of the specified range of the procedure.

*Recommendation: The dynamic range of the method should cover at least [20% and 500%] times the target concentration, where practicable. Target concentration should be understood here as the threshold relevant for a certain regulation.*

*Example: 0.1% and 2.0% for a 1% GMO concentration or 50 and 1000 genome copies if the target is 500 copies.*

There is a general scientific discussion still going on about the interpretation of the percentage values (e.g. dynamic range from 10% to 500% the target value). Although the experts agreed that – at least for PCR – copy number is desired over weight/weight percentage, it was recognised that so far there is no reliable weight/copy number relationship because of inter-variety fluctuation of the 1C value (haploid genome) and because of uncertainty in the correlation of weight of ingredient to number of molecules of DNA. For the time being, both the w/w and copy number/copy number calculations are acceptable.

The unique characteristics of quantitative PCR impose particular restrictions on the low end of the dynamic range of a quantitative PCR. This is due to the difficulty in determining LOD and LOQ values due to the non-normal distribution of variances in the values in this range. Thus it may not be appropriate to require a range extending to 10% of the measured value. The suggestion of a dynamic range that ranges from 10 to 200% can be problematic. For example, capability to analyse a foodstuff composed of more than 50% (w/w) of a biotechnology-derived material (as might be the case for a nutraceutical) would require a dynamic range exceeding 100% (w/w). This is clearly not possible.

## **Limit of Detection (LOD)**

If the validation of the quantitative PCR assay shows that the assay can measure transgenic plant DNA at (for example) 0.1% with acceptable trueness and precision, then it is often not necessary to determine the LOD

and LOQ, as the method is only being applied above the range where these are relevant. However, if the method is being used at concentrations close to the limit of detection and limit of quantification (typically 0.01-0.05%), then the assessment of the LOD and LOQ will become part of the validation procedure.

It is worth noting that a determination of an LOD or LOQ is not necessarily needed to establish the validity of a method for a given application. For example, it does not add much value if an LOD is determined to be 1ng/kg, while the scope of the method validation extends only for concentrations ranging in g/kg. In this and similar cases the reliability of the method will be proven by the other parameters and no efforts are included in the method validation to assess the LOD. However, the LOQ shall always be established and included in the validation study.

If the LOD is required, it is common practice to assume that it is the signal strength of a blank increased by three times the standard deviation of the blank. However, this method gives at best an estimate, relies on normal Gaussian distribution of the blank measurements around zero, and may give a lower value than the actual LOD. Its use is not valid in methods such as Quantitative PCR, in which the distribution of measurement values for blanks is typically truncated at zero and is thus not normally distributed. Thus the LOD need to be experimentally determined unless the targeted concentrations are well above the LOD and the LOD therefore becomes irrelevant. For quantitative methods the LOD is the amount of analyte at which the analytical method detects the presence of the analyte at least 95% of the time ( $\leq 5\%$  false negative results). This, and the false positive rate, are the only parameters required for a qualitative method other than specificity.

For a quantitative method, it is important to know whether the LOQ for a particular matrix is close to the values to be measured. Using the traditional approach, the LOQ can be expressed as the signal strength of a blank equal to the LOD increased by 6-10 times the standard deviation of the blank, unless it is known from other sources that the measured values range so high above the LOQ that its knowledge becomes irrelevant. However, this method to determine the LOQ leads only to an estimate of the true LOQ that may be an artificially high or low approximation.

In practice, two procedures have been employed to determine the LOQ. The first approach is to assay a number of negative samples that have been supplemented (spiked) with known amounts of analyte. The LOQ is then the level at which the variability of the result and percent recovery of the analyte meet certain preset criteria. For small molecules, these criteria have typically been a CV of  $<20\%$  and 70-120% recovery<sup>12</sup>. DNA recovery, however, may be difficult from some matrices, e.g. starches or ketchup, and lower recovery efficiencies may have to be accepted. When recovery efficiencies are low, this must be stated in the validation data and in the analytical report. A more complete approach is to test the method using a number of samples that contain known amounts of the GM material. This is more complicated as it requires access to significant quantities of reference materials that contain a known range of concentrations of the GM event of interest. Procedures for assessing LOD and LOQ during the validation of quantitative PCR methods are discussed in Annexes III and IV..

Validation of methods consists of two phases. The first is an in-house validation of all of the parameters above except reproducibility. The second is a collaborative trial, the main outcome of which is a measure of the repeatability and reproducibility together with detailed information on the transferability of methods between laboratories. It is strongly recommended that a small-scale collaborative trial be performed to test the general ruggedness of a particular method before the expense of organizing a large-scale trial is incurred. In case any improvement of the method or the method description are needed, only limited expenses are incurred through the pre-trial, while a failure of a full interlaboratory method validation due to a ambiguous method description is a very costly failure. Additionally, it may be pointed out that the implementation of an already validated method in a laboratory needs to include necessary experiments to confirm that the implemented method performs as well under local conditions as it did in the interlaboratory method validation. It is important to note that a method should be validated using the conditions under which it will be performed.

*Recommendation: Limit of detection is to be  $< 10\%$  of the value of specification The value of specification should be understood here as the threshold relevant for a certain application.*

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<sup>12</sup> Residue Chemistry Test Guidelines OPPTS 860.1340 "Residue Analytical Method" United States Environmental Protection Agency, August 1996, (Mihaliak & Berberich, 1995).

*Note: limits of detection must be defined using samples comprising of single crops only, e.g. “the LOD for Roundup Ready® soy is 0.1 % of total soy if the product is comprised of 100 % soy”. For food products made up of multiple ingredients, the actual sensitivity will be reduced, as total extracted DNA will be derived from more than one ingredient so that the starting amount of the actual measurand will be decreased. This dilution effect will depend on how much of the target ingredient (e.g. soy) is in the food product and the total quantity of DNA derived from the other ingredients. Some ingredients will contribute much DNA, such as wheat or maize flour and eggs, while other ingredients will not contribute any DNA, such as sugar, water or highly processed oils.*

### **Limit of Quantification (LOQ)**

See introduction above for limit of detection.

*Recommendation: The limit of quantification is to be < 20% of the value of specification with an  $RSD_r \leq [25\%]$  or as close as is practicable. The value of specification should be understood here as the threshold relevant for a certain regulation*

*Example: For a 1 % nominal value  $LOQ_{min} = 0.1 \%$  or for 500 copies  $LOQ_{min} = 50$  copies.*

*For a quantitative method, it is important to know whether the LOQ for a particular matrix is close to the values to be measured. Traditional methods of approximating the LOQ (zero value plus 6-10 standard deviations) rely on normal Gaussian distribution of the blank measurements around zero. This approach is not valid in methods such as Quantitative PCR, in which the distribution of measurement values for blanks is typically truncated at zero and is thus not normally distributed. Thus the LOQ needs to be experimentally determined.*

### **Practicability**

The practicability of the method must be demonstrated.

### **Repeatability standard deviation ( $RSD_r$ )**

*Recommendation: The relative repeatability standard deviation should be  $\leq 25\%$  over the whole dynamic range of the method.*

### **Reproducibility standard deviation ( $RSD_R$ )**

*Recommendation: The relative reproducibility standard deviation should be below 35% at the target concentration and over the majority of the dynamic range.  $RSD_R \leq 50\%$  at the limit of quantification lower end.*

### **Ruggedness (Robustness)**

The evaluation of ruggedness (robustness) demonstrates the reliability of a method with respect to inadvertent variation in assay parameters. Variations that may be included are reaction volumes (e.g., 29 vs. 30 $\mu$ l), annealing temperature (e.g., plus and minus 1°C) and/or other relevant variations. The experiments need to be performed at least in triplicates. The response of an assay with respect to these small changes should not deviate more than  $\pm 35\%$  in reproducibility experiments from the response obtained under the original conditions.

The adequacy of the robustness testing needs to be demonstrated on a method-by-method basis. For instance, for a real-time PCR method, the following factors and their origin / source should ideally be taken into account: different thermal cycler models, DNA polymerase, uracyl-n-glycosylase, magnesium chloride concentration, primer forward and reverse concentration, probe concentration, temperature profile, time profile, dNTP including dUTP concentrations.

### **Sensitivity**

For a quantitative PCR method, a linear relationship of the Ct as a function of the logarithm of the concentration of the target of the individual target should be obtained across the range of the method. The correlation coefficient, y-intercept, slope of the regression line and % of residual should be reported. The % of residual for each of the calibrators should preferably be  $\leq 30\%$ .

In order to obtain a standard curve for event specific quantitative assays, standard DNA mixtures can be prepared by combining purified genomic DNA from transgenic and non-transgenic plants material such as

seed or leaves. The content of transgenic plant DNA in the mixtures might be 100, 50, 10, 5, 1, 0.5, 0.1, and 0% or as appropriate for a smaller concentration range. At least three replicates must be analysed for each point on the standard curve. It has to be pointed out that matrix effects can impact the results as sample to be analyzed and reference material used for calibration might be different matrices and are extracted separately.

For quantitative assays on plant endogenous genes, standard DNA mixtures can be prepared by combining purified genomic DNA from the target plant species and that of a non-target plant species, provided the DNA purity has been demonstrated not to interfere due to the extraction method(s) used. For example, for validation of a maize Adh1 quantitative assay, the target plant species is maize and the non-target plant species could be soybean or another species. The content of DNA molecules of the target plant species in the mixtures is typically 100, 50, 25, 10, 5, 1 and 0% or as appropriate. At least three replicates must be analysed for each point on the standard curve. Alternatively the 5% IRMM's CRM can be used and further diluted without target DNA. It has to be pointed out that matrix effects can impact the results as sample to be analyzed and reference material used for calibration might be different matrices and are extracted separately.

In cases where the  $\Delta$ CT-method is employed by a laboratory instead of a calibration based quantitative method, it will be the responsibility of the analyst to ensure that the overall amount of DNA is well within the range for which the assay was validated. The contrary is of course valuable.

*Recommendation: Typical LOQ are in the range of 0.1% biotechnology-derived material by molecules if the material is not highly processed.*

### **Specificity**

The specificity of GMO detection and reference genes should be demonstrated by providing experimental evidences from testing the method with non-target transgenic events and non-transgenic plants. This testing should include transformation events and preferably cases where the limits of the detection are truly tested. As the GMO detection method should be event-specific it should only be functional with the GMO or GM based product considered and ought not to be functional if applied to other events already authorized or not. In addition, if a reference gene system is a part of the method this should not recognize any sequence corresponding to even phylogenetically related species, and should give similar Ct-values, not statistically different, when amplifying equal amounts of DNA from different cultivars of very different origins of the same taxon.

The adequacy of the testing needs to be analysed on a method-by-method basis. It is necessary to provide Codex with information about the specificity testing in case of stacked genes at some stage.

*Recommendation: Specificity is the starting point for a method and needs to be considered during primer and probes design. Primers should be checked against the known sequence of the event insert and pertinent sequences databases for possible homologies. After such a theoretical specificity assessment, specificity must then be demonstrated experimentally. The following suggests a reasonable approach of experiments which should be performed during pre-validation of an assay.*

#### For event-specific assays:

- Analyse of at least a ten non-target transgenic events and any non-transgenic plants that may commonly be found as contaminants in the commodity.
- Test on sample from each source (total of at least 10 DNA samples).
- Analyse two replicates for each DNA sample which shall give identical results.

*Test results shall clearly indicate that no significant instrument reading or chemistry effects are observed.*

#### For assays on plant endogenous (reference) genes:

- Analyse of at least ten different plant samples comprising different varieties of very different genetic origins of the same plant species as well as other plants species important for food production (such as wheat, rice, corn, potato, and soybean) and that may commonly be found as contaminants in the commodity.
- Test one sample from each source (total of at least 10 DNA samples).
- Analyse two replicates for each DNA sample which shall give identical results .

*Test results shall clearly indicate that no significant instrument reading or chemistry effects are observed.*

## **Trueness**

*Recommendation: The trueness should be within  $\pm$  [30%] of the accepted reference value over the whole dynamic range. This refers to the PCR-step provided that the modular approach has been applied.*

## **ANALYTICAL CONTROL ACCEPTANCE CRITERIA AND INTERPRETATION OF RESULTS**

A validated method also includes values of criteria on which the validity of an observed measurement result can be assumed. It is important to follow these criteria and to observe the decision support system for data analysis and interpretation. In the case that it may be desired to deviate from said criteria and rules a new method validation study would be needed in order to demonstrate the validity of the new decision support system and procedures.

At a minimum, the following acceptance criteria are common to all quantitative PCR methods and applicable to each PCR run:

- The result of the positive DNA target control, with, for example 0.9% transgenic DNA, the mean of the replicates deviates less than 3 standard deviations from the assigned value. When applicable, a target DNA control is defined as reference DNA or DNA extracted from a certified reference material or known to be a positive sample representative of the sequence or organism under study. The control is intended to demonstrate what the result of analyses of test samples containing the target sequence should be.
- The amplification reagent control is  $\leq$ LOD. The amplification reagent control is defined as control containing all the reagents, except extracted test sample template DNA. Instead of the template DNA, a corresponding volume of nucleic acid free water is added to the reaction.
- The % of residual for each of the standards should be  $\leq$ 30%

To accept the result of an unknown sample, the relative standard deviation of the sample replicates should be  $\leq$ [35]%.

## **REFERENCES FOR ANNEX III**

1. Huebner P, Waiblinger H U, Pietsch K, Bordmann P (2001) Validation of PCR methods for quantitation of genetically modified plants in food. Journal of AOAC International 84(6) 1855-1864;
2. Kay S, Van den Eede G, The limits of GMO detection, Nature Biotech. 19(5) 504 (2001)).
3. Residue Chemistry Test Guidelines OPPTS 860.1340 "Residue Analytical Method" United States Environmental Protection Agency, August 1996, (Mihaliak & Berberich, 1995).

## ANNEX IV: VALIDATION OF A QUALITATIVE PCR METHOD

### Introduction

A qualitative PCR must be validated as much as possible in the same way as it is intended to be used for routine analyses – that means the sensitivity of the method must be shown to be such that it can reliably detect one positive particle I.E.? (food) in a test portion, and does not give rise to a significant number of false positives. A concept of using false-positive and false-negative rates to describe the accuracy and precision of a qualitative assay has been developed for microbial assays<sup>1</sup>. This concept can be applied to qualitative PCR assays. A critical issue in the validation of this type of method is the availability of test materials that are known to be either positive or negative. The provision of negative reference materials is particularly important and critical in the case of a qualitative method. However, it is recognized that in some cases, no reference material might be available, for instance in case of unapproved GMO. Any impurities must be present only at levels so low that they become negligible.

By their very nature, qualitative test results refer to the identification above/below a detection limit. The measures of precision and accuracy are the frequencies of false negative and/or false positive results at the detection limit. False negative results indicate the absence of a given analyte when in fact the analyte is present in the sample, while false positive results indicate the presence of an analyte that is not present in the sample. Due to the inherent nature of the analytical technique, an increase in false negative results will be observed when the amount of analyte approaches the LOD of the method. Like the limit of detection for quantitative methods, the limit of detection for a qualitative method can be defined as the concentration at which a positive sample yields a positive result at least 95% of the time. This results in a rate of false negative results of 5% or less. During validation of a qualitative PCR assay, it is also important to determine the number of false positive results (a positive result obtained using a sample that is known to be negative). This is also expressed as a ratio or percentage.

### False Positive Rate

This is the probability that a known negative test sample has been classified as positive by the method. The false positive rate is the number of misclassified known negatives divided by the total number of negative test samples (misclassified positives plus the number of correctly classified known negatives) obtained with the method:

For convenience this rate can be expressed as percentage:

$$\% \text{ false positive results} = \frac{\text{number of misclassified known negative samples}}{\text{total number of negative test results [incl. misclassified]}}$$

### False Negative Rate

This is the probability that a known positive test sample has been classified as negative by the method. The false negative rate is the number of misclassified known positives divided by the total number of positive test samples (misclassified positives plus the number of correctly classified known positives) obtained with the method.

For convenience this rate can be expressed as percentage:

$$\% \text{ false negative results} = \frac{\text{number of misclassified known positive samples}}{\text{total number of positive test results [incl. misclassified]}}$$

Note: different sectors use different definitions here.

In order to demonstrate the false negative rate for qualitative assay, a series of samples (e.g. grain/seed pools) with a constant, known concentration of positive material in a pool of negative material (e.g., 1 positive kernel in 199 conventional corn kernels) have to be analysed and the results evaluated. It is important to

note that the concept of confidence intervals and statistical uncertainty needs to be applied to the risk of false positive and/or false negative results as well. The desired level of confidence determines the size and number of pools that need to be tested. For example, 100 positive test results obtained from 100 independent measurements on truly positive samples lead to the conclusion that the level of false negative results is below 4.5% at a confidence level of 99% for the tested concentration of positive kernels (expressed as the number of positive kernels in a pool of negative kernels).

### **Ruggedness**

As with any validated method, reasonable efforts must be made to demonstrate the ruggedness of the assay. This involves careful optimisation and investigation of the impact of small modifications that could occur for technical reasons.

### **Acceptance Criteria Values and Interpretation of Results**

A validated method includes performance criteria values on which the validity of an observed measurement result can be accepted as valid. It is important to follow these criteria and to observe the decision support systems for data analyses and interpretation. It is therefore important to make sure that the result of the positive DNA target control, when available, is positive. Similarly the amplification reagent control must be negative. In addition to these controls, it is desirable to carry out a parallel reaction on the same DNA sample using a primers set which detects an endogenous single copy sequence to determine the impact of PCR inhibitors putatively present. This reaction shall be carried out on every DNA sample on which no amplification has been observed, and can either be carried out in the same reaction (multiplexed) or as a separate reaction on the same DNA extract. In the case of multiplexed reactions, it is important that the endogenous reaction does not out compete the event specific reaction for reagents, as the endogenous sequence is likely to be present at up to 1000 fold the amount of the target sequence. Optimization of PCR conditions shall have taken into consideration these conditions. The inhibition control reaction with the endogenous sequence gives an indication of the quality of the DNA as a template for the PCR reaction. Table 1 sets out the acceptance/rejection criteria for the PCR reactions on a per lane basis, using the results of the PCR reaction with a endogenous sequence.

Alternatively, such a inhibition control reaction can be carried out with spiked DNA and the appropriate, not interfering, PCR test.

Table 1: Criteria for scoring Qualitative PCR analyses

PCR result (GM analyte)	PCR result (endogenous)	Scoring of test
+	+	+
-	+	-
+	-	[+][-]
-	-	Reject

A further complication is however introduced by the fact that qualitative PCR reactions are typically carried out at least in duplicate. Thus it can occur that the duplicates do not provide identical results. It is a common practice to repeat PCR reactions at least once on DNA samples that are rejected for discrepancy of results among replicates. A repeated indeterminate result is indicative that the analyte cannot be reliably detected. (Table 2), and that the assay is operating below the limit of detection as, by definition, a 95% or better detection rate would be achieved at the limit of detection. The sample shall therefore be scored negative. Similar criteria apply if more replicates are carried out on each DNA sample.

Table 2: Criteria for scoring duplicate qualitative PCR analyses

Lane 1	Lane 2	Scoring of test
+	+	Positive
-	+	Repeat/Indeterminate
+	-	Repeat/Indeterminate
-	-	Negative

#### REFERENCES FOR ANNEX IV

1. AOAC® Official Methods<sup>SM</sup> Program Manual, Appendix X p14f, May 2002, AOAC International; [http: www.aoac.org/vmeth/omamannual/htm](http://www.aoac.org/vmeth/omamannual/htm).

## **ANNEX V: VALIDATION OF A PROTEIN-BASED METHOD<sup>1</sup>**

### **QUANTITATIVE TESTING**

Quantitative immunoassays are used to determine levels of the protein analyte in specific parts of the plant (e.g. seed, leaf, root, stalk etc) of a cultivar or a mix of cultivars. Typical applications are given in Table 1. In order to perform any immunological detection method such as a microplate ELISA for quantitative determination of a protein analyte in plant tissue(s), it is first necessary to obtain a representative sample of the plant material. The sample amount and procedure to prepare test portions will influence the detection limit or sensitivity of the assay. The analyte is then extracted from the plant material by adding the appropriate liquid and blending, agitating, or applying sheering or sonic forces. Typical liquids used are water or buffered salt solutions. Sometimes detergents or surfactants are added according to the validated test and matrices. Some proteins require more rigorous procedures like homogenization or boiling in solvents, detergents, salts etc.

The following description of the procedure is only one of several possibilities to carry out an immunological detection assay for proteins expressed in GMOs.

After more or less specific parts of the detection tests such as the capture antibody's immobilization on the microplate well surface, a precise volume of the standard or sample extract solution is added to each well. The analyte in the test solution binds to the capture antibody. The enzyme-labelled second antibody is then added and also binds to the analyte, forming a sandwich. At this point, the well is washed to remove unbound analyte and antibodies, leaving only the antibody-analyte-antibody complex bound to the well surface. A colorimetric substrate is added which is processed by the enzyme and produces a coloured product. The reaction is stopped after a set period of time and the colour absorbance at a given wavelength is measured by a photometer. The standard curve is generated by plotting the optical density (OD) on the *y*-axis (linear scale) against the concentration on the *x*-axis (log scale) which produces a sigmoidal dose response curve Figure 4 according a calibration curve set up with reference material.

To obtain an accurate and precise quantitative value, the OD for the sample solutions must pertain to the linear portion of the calibration curve. If the OD is too high, the sample solution must be diluted until the OD falls within the quantification range of the assay. The concentration of the protein analyte in the original sample of plant material is calculated by correcting for any dilution factor that was introduced in preparing the sample for application to the microplate. The initial weight of the sample and the volume of extraction liquid, as well as any subsequent dilutions are used to calculate the dilution factor.

Various assay controls can be employed to demonstrate the performance of the assay. A blank sample such as an empty well or buffered solution can be run in the assay to determine any background response which shall be subtracted from sample and calibration responses if desired. A negative control sample (i.e. matrix extract solution known to contain no analyte) shall be used to demonstrate any non-specific response or matrix effect occurring in the assay. A positive control or matrix extract spiked with a known amount of the analyte can be run to demonstrate the accuracy of the test. Standards and samples can be run in an appropriate number of replicates to appreciate the precision of the test. Blanks, negative controls, positive controls, fortified sample extracts when necessary, reference materials, reference material extracts, and replicates are typically run on each microplate to control for plate-plate variation.

### **REFERENCE MATERIALS**

The reference material consists of the same matrix as the actual agricultural commodity to be tested. For example, if the matrix to be tested is soybean seed, the standardized reference material would be soybean seed containing a known proportion of transgenic seed. Alternatively, a pure sample or extract of the protein of interest may be used, providing the use of such protein reference materials has been validated against the matrix in question. In some cases the reference matrix, e.g. starch, may be unavailable. Access to reference materials is important during the development, validation, and use of immunoassays for analysis of introduced proteins in transgenic agricultural commodities. The best available reference material should be used in order to comply with regulations and testing requirements.

In the case of commodities such as grain or seed, where the commodity consists of discrete units, it is fairly straightforward to prepare a reference sample with a known proportion of transgenic material. In other cases, generating reference samples for certain matrices and analytes can be difficult. Stability and uniformity are important considerations. For example, if the matrix to be tested consists of a mixture of materials, it will be difficult to combine transgenic and non-transgenic material in such a way as to achieve a homogeneous reference sample with a known proportion of transgenic material. The stability of these materials would need to be evaluated under storage and test conditions. In any case, it is useful to have non-transgenic and transgenic material available to use as negative and positive controls.

In case of lack of continuity of standards furniture, such as the withdrawal of the market of a GMO, appropriate solutions have to be found to get positive controls.

During assay development, the reference material is used to help select assay parameters which would minimize any interfering effects of the matrix (e.g. non-specific binding of sample components to the antibodies or enzyme inhibitors). During validation and use of the assay, the reference materials can be extracted and analysed alongside the test samples so that the results can be directly compared.

## **VALIDATION OF A QUANTITATIVE PROTEIN-BASED METHOD**

The principles of method validation described in appendices III and IV for PCR methods also apply to protein methods. For commercially available immunoassay kits, assay performance is generally validated by the manufacturer in single laboratory or collaborative trials validation conditions and is documented in the product user's guide.

Validation should be conducted according to the harmonized ISO/IUPAC/AOAC protocol for chemical analytical methods. This document defines the procedures necessary to validate a method<sup>2</sup>.

**Accuracy:** Accuracy is demonstrated by measuring the recovery of analyte from fortified samples and is reported as the mean recovery at several levels across the quantitative range. Ideally, quantitative methods will have demonstrated recoveries between 70 and 120% and a coefficient of variation (CV) of less than 20% for measured recoveries at each fortification level (Mihaliak & Berberich, 1995).

**Extraction efficiency:** Extraction efficiency is a measure of how efficient a given extraction method is at separating the protein analyte from the matrix. It is expressed as percent analyte recovered from the sample. Since the introduced protein expressed is endogenous to the plant, it can be difficult to truly demonstrate efficiency of the extraction procedure. There may not be an alternate detection method against which to compare the immunoassay results. One approach to addressing extraction efficiency is to demonstrate the recovery of each type of introduced protein analyte from each type of food fraction by exhaustive extraction, i.e. repeatedly extracting the sample until no more of the protein is detected (Stave, 1999).

**Precision:** Intra-assay precision describes how much variation occurs within an assay. It can be evaluated by determining the variation (% CV) between replicates assayed at various concentrations on the standard curve and on the pooled variation (% CV) derived from absorbance values in standards from independent assays performed on different days. Inter-assay precision describes how much variation occurs between separate assays and can be measured by analysis of quality control samples on every microplate. The quality control samples required would consist of two pools of extracts, one extract from transgenic plant tissue and one from conventional plant tissue. These extracts would be stored frozen and a portion would be thawed and assayed on every microplate. Inter-assay precision can be evaluated over time and expressed as % CV (Rogan *et al*, 1999). The precision of protein-based quantitative methods is in general higher than PCR-based methods.

**Recommendation:** *The accuracy should be within [ $\pm 25\%$ ] of the accepted reference value over the whole dynamic range.*

**Sensitivity:** The sensitivity of the assay could be defined as the amount of analyte that can be measured by an absorbance reading of two standard deviations above background absorbance (Rogan *et al*, 1992). The detection limit could be expressed as the lowest dilution of transgenic crop that could be detected when transgenic and non-transgenic crop are combined (Rogan *et al*, 1999).

### **Dynamic Range - Range Of Quantification**

The scope of the methods defines the concentration range over which the analyte will be accurately determined. In most cases the analytical range for a GM product will range from a tenth of a percent up to a few percent. This desired concentration range defines the standard curves and a sufficient number of standards must be used to adequately define the relationship between concentration and test's response. The relationship between response and concentration should be demonstrated to be continuous, reproducible and should be linear after suitable transformation.

Interpretation of the percentage values (e.g. dynamic range from 10% to 500% the target value) can be difficult when using quantitative methods. Quantitative protein methods generally give an estimate of the concentration of the GM protein in the matrix, due to variations in the expression of the amount of protein in different tissues of plants or among cultivars, and within the same tissue at different locations. The use of qualitative protein-based methods is thus much more prevalent. In addition, care must be taken to employ a method which can detect the protein in the analyzed matrix. For example, it is believed that proteins undergo modification or degradation due to processing to a greater degree than DNA, and thus loss of signal due to food processing effects must be considered.

It is worth noting that a determination of an LOD or LOQ is not necessarily needed to establish the validity of a method for a given application. For example, it does not add much value if an LOD is determined to be 1ng/kg, while the scope of the method validation extends only for concentrations ranging in g/kg. In this and similar cases the reliability of the method will be proven by the other parameters and no efforts are included in the method validation to assess the LOD. However, the LOQ shall always be established and included in the validation study.

### **Limit of Detection (LOD)**

LOD is defined in annex II. Proteins are present in GM foods at higher concentrations than the target DNA for PCR methods. Thus stochastic effects have less influence on the determination of the LOD than when using PCR.

It is common practice when estimating the LOD to assume that it is the signal strength of a blank increased by three times the standard deviation of the blank. This method gives at best an estimate, and relies on normal Gaussian distribution of the blank measurements around zero. This can generally be assumed for methods such as ELISA, but the LOD is best determined experimentally. Alternatively the LOD is commonly defined as a concentration equal to the lowest standard used in the assay, should a positive value be consistently obtained with that standard.

### **Limit of Quantification (LOQ)**

For a quantitative method, it is important to know whether the LOQ for a particular matrix is close to the values to be measured. Using the traditional approach, the LOQ can be expressed as the signal strength of a blank equal to the LOD increased by 6-10 times the standard deviation of the blank, unless it is known from other sources that the measured values range so high above the LOQ that its knowledge becomes irrelevant. However, this method to determine the LOQ leads only to an estimate of the true LOQ that may be an artificially high or low approximation.

In practice, two procedures have been employed to determine the LOQ. The first approach is to assay a number of negative samples that have been supplemented (spiked) with known amounts of analyte. The LOQ is then the level at which the variability of the result and percent recovery of the analyte meet certain preset criteria. For small molecules, these criteria have typically been a CV of  $\leq 20\%$  and 70-120%

recovery<sup>3</sup>. Protein recovery, however, may be difficult from some matrices, e.g. starches or oils, and lower recovery efficiencies may have to be accepted. When recovery efficiencies are low, this must be stated in the validation data and in the analytical report. A more complete approach is to test the method using a number of samples that contain known amounts of the GM material. This is more complicated as it requires access to significant quantities of reference materials that contain a known range of concentrations of the GM event of interest. Procedures for assessing LOD and LOQ during the validation of quantitative PCR methods are also discussed in annexes III and IV.

### **Specificity**

The specificity is the degree to which analogs or other molecules bind to the antibodies and should be characterized and described in the method. Specificity should be demonstrated by showing experimental results from testing the method with non-target transgenic events and non-transgenic plants. This testing should include closely related events and cases where the limits of the detection are truly tested. As the method should be protein-specific it should only be functional with the GMO or GM based products considered and ought not to be functional if applied to events which do not express the protein in question. Interferences: the potential for interferences from reagents and labware can be evaluated by assaying extracts from non-transgenic plant material.

Matrix effects: if the response of the method is affected by a substance in the final extract other than the specific protein analyte, the non-specific response is referred to as a matrix effect. One way to manage matrix effects is to demonstrate that the analytical method gives identical results with or without sample matrix present in the extract. In this approach, freedom from matrix effects would have to be demonstrated in all matrices for which the assay is to be used. Another approach (although less desirable) to managing matrix effects would be to prepare the standard solutions in extracts from non-transgenic matrix, i.e. matrix-matched standards. This would ensure that any matrix effects would be consistent between the standards and the samples.

### **Ruggedness (Robustness)**

The evaluation of ruggedness (robustness) demonstrates the reliability of a method with respect to inadvertent variation in assay parameters. Variations that may be included are reaction volumes incubation temperature (e.g., plus and minus 5-10°C) and/or other relevant variations. The experiments need to be performed at least in triplicates and the recovery needs to be calculated. The response of an assay with respect to these small changes should not deviate more than  $\pm 30\%$  from the response obtained under the original conditions. Experiments which may be performed to establish ruggedness include repeated analysis of a sample or samples on several days and measurement of accuracy and precision in fortified samples using control material from several sources.

## **QUALITATIVE (THRESHOLD) TESTING**

Lateral flow devices are useful tools for on-site or field threshold testing. This type of testing requires a quick, accurate and cost-effective approach. In order to ensure reliable results, the manufacturers of the lateral flow devices must conduct a method validation and provide a description of the performance characteristics of the product in the package insert. If this has been completed there is generally no need for validation studies to be performed by users for implementation of the technique within their laboratory. Each lateral flow device is an individual stand-alone unit, capable of performing to the standards described in the product package insert according to the quality assurance scheme of the provider.

In order to establish an on-site procedure for threshold testing, the threshold level must first be established. To establish that the lateral flow device is able to differentiate between samples containing transgenic protein above or below the threshold, both a negative reference and a threshold reference containing a known proportion of transgenic grain should be assayed concurrently. The negative reference is a sample of the test matrix known to contain none of the protein analyte and is assayed to demonstrate that the method can distinguish between zero and the threshold level. A sufficient number of these samples are run to ensure that assay sensitivity is adequate to determine whether the level in the test sample is greater or less than the

threshold level. During routine testing of bulk commodity samples, the lateral flow devices would typically be used without running the concurrent negative and threshold reference samples.

### **VALIDATION OF A QUALITATIVE PROTEIN-BASED METHOD**

The same principles apply to qualitative protein-based testing as to qualitative PCR testing. These approaches, including calculation of false positive and false negative rates, can therefore be applied to protein-based methods. In general, due to the more reliable nature of protein-based lateral flow strip methods, they are not performed in duplicate on each sample. However, if ELISA testing is performed, duplicate wells should be used.

The same types of control samples, and criteria for acceptance/rejection of the result can be used as for qualitative PCR methods. The LOD is expressed as the amount of analyte at which the analytical method detects the presence of the analyte at least 95% of the time (<5% false negative results). However, lateral flow strip tests are generally applied at test concentrations that are at least two fold (or more) above the LOD.

### **REFERENCES FOR ANNEX V**

1. Guidelines for the Validation and Use of Immunoassays for Determination of Introduced Proteins in Biotechnology Enhanced Crops and Derived Food Ingredients. Lipton et al., Food and Agricultural Immunology, 2000, 12, 153-164.
2. Horwitz W; Protocol for the design, conduct and interpretation of method-performance studies. Pure and Applied Chemistry, 67, 331 (1995).
3. Residue Chemistry Test Guidelines OPPTS 860.1340 "Residue Analytical Method" United States Environmental Protection Agency, August 1996, (Mihaliak & Berberich, 1995).

## **ANNEX VI: GMO METHODS OF ANALYSIS VALIDATED THROUGH COLLABORATIVE TRIALS**

The results from validation studies may be used by analysts to aid their measurement uncertainty estimations. This Annex gives brief descriptions of collaborative trials which have been carried out by various institutions/groups in order to provide information about the capacity and limits of GMO detection methods. The list includes only those collaborative studies in which at least eight analysts participated. Thirty-one of the 32 reported validated PCR-based methods; only one method using the protein-based ELISA approach has been validated.

Eleven out of 32 studies carried out were intended to validate a qualitative detection of the GMO under investigation. The majority of validation studies have been designed in order to quantify the GM material present in a given matrix. However, one of the major concerns regarding some of the listed validation studies is the use of the same source of material for the establishment of the standard curves necessary to determine the genome copy number present in the sample and for the unknown samples, respectively. Therefore, the results obtained with these samples cannot be used for the evaluation of the trueness of the methods applied. Four of the described studies used only DNA as unknown sample material distributed to the participants. The measurement uncertainty of the DNA extraction module needs to be determined independently. A collaborative trial aimed to validate the DNA extraction module individually has not yet been carried out.

The results do indicate that the PCR technique (as well as protein-based methods) does have the capability to be applied for the GMO detection and quantification. The overall reproducibility of real time PCR methods is in the range of 10 to 45%. Moreover there is a significant improvement in the accuracy comparing the results of the first studies carried out to validate GMO methods and those carried out more. Thus, it seems to be realistic to end-up with real time PCR methods for GMO quantification with an average of  $RSD_R$  value of about 20 to 25%.

No	Purpose (GMO)	Qualitative/ quantitative	Co-ordinator	Year	Results	Comments
1	GTS 40-3-2 (Roundup® Ready soya) <sup>1</sup>	Qualitative	Federal Institute for Risk Assessment (BfR), Berlin, Germany	1998	No false-positive and false- negative	Construct-specific PCR method; only soya beans as samples.
2	Genetically modified tomato (Zeneca) <sup>1</sup>	Qualitative	BfR	1999	No false-positive and false- negative	Construct-specific PCR method; only tomato pieces as samples.
3	Bt 11 maize <sup>1</sup>	Qualitative	BfR	2000	False-positive (%): 5 False-negative (%): 10	Construct-specific PCR method; only maize kernels as samples.

4	Event 176 maize <sup>1</sup>	Qualitative	BfR	2000	No false-positive and false-negative	Construct-specific PCR method; only maize kernels as samples.
5	T 25 maize (“LibertyLink”) <sup>1</sup>	Qualitative	BfR	2001	False-positive (%): 0 False-negative (%): 12	Construct-specific PCR method; ground seed samples e.g. containing 0.1 and 1% (w/w) T 25
6	MON 810 maize <sup>1</sup>	Qualitative	BfR	2001	No false-positive and false-negative	Event-specific PCR method; ground seed samples e.g. containing 0.1 and 1% (w/w) MON 810
7	35S promoter from Cauliflower mosaic virus (CaMV) <sup>1</sup>	Qualitative	BfR	1999	No false-positive and false-negative	Screening method; only soya beans (RRS and non-GM soya beans) as samples.
8	35S promoter from CaMV <sup>1</sup>	Qualitative	Joint research Centre of the European Commission (EC-JRC), Ispra, Italy	1999	False-positive (%): 3.9 False-negative (%): 1.9	Screening method; various food matrices as samples containing 0, 2, 100% RRS and Event 176 maize (w/w), respectively
9	Nopaline synthase (nos) terminator of <i>Agrobacterium tumefaciens</i> <sup>1</sup>	Qualitative	EC-JRC	1999	False-positive (%): 1.8 False-negative (%): 2.1	Screening method; various food matrices as samples containing 0, 2, 100% RRS and Event 176 maize (w/w), respectively
10	neomycin phosphotransferase gene ( <i>nptII</i> ) <sup>1</sup>	Qualitative	BfR	1998	No false-positive and false-negative	Screening method; only tomato pieces as samples.
11	Genetically modified papaya	Qualitative	BfR	2005	False-positive (%): 3.3 False-negative (%): 3.3	Construct-specific PCR method; raw samples containing 0, 10 and 100% GM papaya (w/w), respectively
12	35S promoter from CaMV <sup>2</sup>	Quantitative	Kantonales Laboratorium Basel, Switzerland	1999	Reproducibility relative standard deviation (RSD <sub>R</sub> ) in the range of 17 to 34 %	Real time PCR screening method; samples containing between 0,7 % and 3 % (w/w) of RRS powder. Same material for unknowns and calibrants

13	Roundup <sup>®</sup> Ready soya <sup>2</sup>	Quantitative	Kantonales Laboratorium Basel, Switzerland	1999	RSD <sub>R</sub> in the range of 16 to 28 %	Real time PCR construct-specific method; samples containing between 0,7 % and 3 % (w/w) of RRS powder. Same material for unknowns and calibrants
14	Roundup <sup>®</sup> Ready soya <sup>2</sup>	Quantitative	BfR	2000	RSD <sub>R</sub> in the range of 27 to 44 %	Real time PCR construct-specific method; samples containing between 0.1 % and 5 % (w/w) of RRS powder. Same material for unknowns and calibrants
15	Event 176 maize <sup>2</sup>	Quantitative	BfR	2000	RSD <sub>R</sub> in the range of 33 to 42 %	Real time PCR construct-specific method; samples containing between 0.1 % and 5 % (w/w) of Event 176 powder. Same material for unknowns and calibrants
16	Roundup <sup>®</sup> Ready soya <sup>2</sup>	Quantitative	National Food Research Institute (NFRI), Tsukuba, Japan	2001/2002	RSD <sub>R</sub> in the range of 11 to 16 %	Real time PCR construct-specific method; samples containing between 0.1 % and 10 % (w/w) of RRS powder. For calibration plasmid standards containing the target DNA were used
17	MON 810 maize <sup>2</sup>	Quantitative	NFRI	2001/2002	RSD <sub>R</sub> in the range of 12 to 32 %	Real time PCR construct-specific method; samples containing between 0.1 % and 10 % (w/w) of MON 810 maize powders. For calibration plasmid standards containing the target DNA were used
18	Event 176 maize <sup>2</sup>	Quantitative	NFRI	2001/2002	RSD <sub>R</sub> in the range of 9 to 21 %	Real time PCR construct-specific method; samples containing between 0.1 % and 10 % (w/w) of Event 176 maize powders. For calibration plasmid standards containing the target DNA were used

19	Bt 11 maize <sup>2</sup>	Quantitative	NFRI	2001/2002	RSD <sub>R</sub> in the range of 11 to 24 %	Real time PCR construct-specific method; samples containing between 0.1 % and 10 % (w/w) of Bt 11 maize powders. For calibration plasmid standards containing the target DNA were used
20	GA 21 maize <sup>2</sup>	Quantitative	NFRI	2001/2002	RSD <sub>R</sub> in the range of 14 to 22 %	Real time PCR construct-specific method; samples containing between 0.1 % and 10 % (w/w) of GA 21 maize powders. For calibration plasmid standards containing the target DNA were used
21	T 25 maize <sup>2</sup>	Quantitative	NFRI	2001/2002	RSD <sub>R</sub> in the range of 11 to 28 %	Real time PCR construct-specific method; samples containing between 0.1 % and 10 % (w/w) of T 25 maize powders. For calibration plasmid standards containing the target DNA were used
22	Bt 11 maize <sup>2</sup>	Quantitative	EC-JRC	2003	RSD <sub>R</sub> in the range of 13 to 33 %	Real time PCR event-specific method; samples consist of Bt 11 DNA mixtures corresponding to 0.1 % to 2 % (w/w) of Bt 11 maize. Same source of DNA material for unknowns and calibrants
23	MON810 maize <sup>2</sup>	Quantitative	BfR, EC-JRC, AACC, Genescan Europe	2003/2004	RSD <sub>R</sub> in the range of 32 to 45 %	Real time PCR event-specific method; samples containing between 0.1 % and 5 % (w/w) of MON 810 maize powders. Same material for unknowns and calibrants
24	NK 603 maize <sup>3</sup>	Quantitative	EC-JRC	2004	RSD <sub>R</sub> in the range of 25 to 37 %	Real time PCR event-specific method; samples containing between 0.1 % and 5 % (w/w) of NK 603 maize powders. Same material for unknowns and calibrants

25	GA 21 maize <sup>3</sup>	Quantitative	EC-JRC	2004	RSD <sub>R</sub> in the range of 29 to 44 %	Real time PCR event-specific method; samples containing between 0.1 % and 4.26 % (w/w) of GA 21 maize powders. Same material for unknowns and calibrants
26	MON 863 maize <sup>3</sup>	Quantitative	EC-JRC	2004	RSD <sub>R</sub> in the range of 18 to 35 %	Real time PCR event-specific method; samples containing between 0.1 % and 10 % (w/w) of MON 863 maize powders. Same material for unknowns and calibrants
27	TC 1507 maize (Herculex™ I) <sup>3</sup>	Quantitative	EC-JRC	2004	RSD <sub>R</sub> in the range of 18 to 35 %	Real time PCR event-specific method; samples consist of TC 1507 DNA mixtures corresponding to 0.1 % to 5 % of TC 1507 maize. Same source of DNA material for unknowns and calibrants
28	T 25 maize <sup>3</sup>	Quantitative	EC-JRC	2005	RSD <sub>R</sub> in the range of 18 to 26 %	Real time PCR event-specific method; samples consist of T 25 DNA mixtures corresponding to 0.15 % to 3.3 % of T 25 maize. Same source of DNA material for unknowns and calibrants
29	DAS-59122-7 maize <sup>3</sup>	Quantitative	EC-JRC	2005	RSD <sub>R</sub> in the range of 13 to 25 %	Real time PCR event-specific method; samples consist of DAS-59122-7 DNA mixtures corresponding to 0.1 % to 4.5 % of DAS-59122-7 maize. Same source of DNA material for unknowns and calibrants
30	Sugar beet line H7-1 <sup>3</sup>	Quantitative	EC-JRC	2005	RSD <sub>R</sub> in the range of 13 to 20 %	Real time PCR event-specific method; samples consist of H7-1 DNA mixtures corresponding to 0.1 % to 5 % of sugar beet line H7-1. Same source of DNA material for unknowns and calibrants

31	Roundup <sup>®</sup> Ready soya <sup>4</sup>	Quantitative	Central Science laboratory (CSL), York, United Kingdom	2002	RSD <sub>R</sub> in the range of 20 to 34 %	Real time PCR construct-specific method; samples containing between 0.5 % and 3.9 % (w/w) of RRS powder. Same material for unknowns and calibrants
32	Roundup <sup>®</sup> Ready soya <sup>5</sup>	Quantitative	EC-JRC	1999	RSD <sub>R</sub> in the range of 9 to 17 %	ELISA method; samples containing between 0.5 % and 2 % (w/w) of RRS powder.

<sup>1</sup> A detailed description is given in ISO/FDIS 21569 “Methods of analysis for the detection of genetically modified organisms and derived products — Qualitative nucleic acid based methods”.

<sup>2</sup> A detailed description is given in ISO/FDIS 21570 “Methods of analysis for the detection of genetically modified organisms and derived products — Quantitative nucleic acid based methods”.

<sup>3</sup> A detailed description of the validation study is given by the Community Reference Laboratory (CRL), Ispra, Italy (<http://gmo-crl.jrc.it/statusofdoss.htm>).

<sup>4</sup> Hird H, Powell J, Johnson ML, Oehlschlager S. Determination of percentage of RoundUp Ready soya in soya flour using real-time polymerase chain reaction: interlaboratory study. J AOAC Int. (2003): **86** (1):66-71.

<sup>5</sup> A detailed description is given in ISO/FDIS 21572 “Methods of analysis for the detection of genetically modified organisms and derived products — Protein based methods”.

In some proficiency tests concerned with measuring the proportion of genetically modified organism (GMO) in food the results produced are log-transformed (converted into logarithms) before z-scores are calculated [1]. The transformation can be justified both theoretically and practically. However, the transformation gives rise to z-scores that are not on the same type of scale as the original data, and are therefore less readily interpreted. A certain amount of background in logarithmic transformation may be helpful.

### What is a lognormal distribution?

Figure 1 shows the density of a lognormally distributed variable. It is asymmetric, with a positive skew and all values of  $x$  necessarily greater than zero. If alternatively we plot the density against the logarithm of  $x$ , we see the familiar shape of the normal distribution (Figure 2). (Note that logarithms base ten are implied throughout this Brief.)

*Definition: a variable  $x$  is lognormally distributed if  $\log x$  is normally distributed.*

While all normal distributions are essentially the same shape, the shape of a lognormal distribution depends on its RSD (relative standard deviation, here expressed as a fraction). For example, the highly-skewed distribution in figure 1 has an RSD of 0.3, while figure 3, also a lognormal but with an RSD of 0.1, shows only a slight skew. (For reference, results from a round of a GMO proficiency test commonly have an RSD of about 0.7).

### Data From GMO Proficiency Testing

At present, nearly all quantitative measurements of a genetically modified species in a food are based on the polymerase chain reaction (PCR). In interlaboratory studies such as the proficiency test, the results almost invariably show a strongly skewed distribution of results, Figure 4 for example. There are *a priori* reasons for expecting this outcome. Firstly the procedure may start with a small number of copies of the gene, so that there is a binomial distribution of copies in the sample taken for PCR. Binomial distributions are positively skewed for small number of copies. If the DNA is associated with a small number of particles, sampling these particles could give rise to a skewed result even if the number of copies of the gene is reasonably large. The calibration function in PCR is log-linear in form and this will tend to produce a lognormal distribution of results from a normal input. Finally there is the usual normal distribution of errors from the instrumental readout system.

As an outcome of all this, the distribution of errors is expected to be a complex convolution of distribution types, but with a tendency towards a positive skew. It is therefore tempting to suggest that log-transformation of participants' results may be appropriate before the formation of z-scores. A detailed study of proficiency test data has justified this action in practice [2]. But how are we to relate such z-scores to everyday practice and the performance of individual laboratories?

### Z-Scoring In GMO Proficiency Testing

What we have to bear in mind is that, in quantitative GMO testing, errors seem to be largely multiplicative, rather than additive as in most other analytical work. In that context, a very useful property of log-transformation is that various datasets with the same *relative* standard deviation in the original scale have the same *absolute* standard deviation in the log-scale. In the instance of GMO proficiency testing, this enables providers to set a single  $\sigma_p$  value for the scheme, regardless of the concentration of the analyte (except, of course, where the concentration is zero, or very close to it). A z-score can then be calculated from a result  $x$  and an assigned value  $x_a$  (both in the original scale) according to the equation

$$z = (\log x - \log x_a) / \sigma_p = \log(x/x_a) / \sigma_p .$$

$\log x_a$  will usually be the robust mean of the  $\log x$  values. The  $\sigma_p$  value (the standard deviation for proficiency, previously called the ‘target value’) should be a fitness-for-purpose criterion, if at all possible.

So if fitness for purpose demanded that a satisfactory result should be within limits of (say)  $0.5x_a$  and  $2.0x_a$ , we would need to set  $\sigma_p$  such that these limiting results produced z-scores of  $-2$  and  $+2$  respectively.

Substituting the corresponding values  $z = 2$  and  $x = 2.0x_a$  in the equation above gives us

$$2 = (\log 2x_a - \log x_a) / \sigma_p, \text{ or}$$

$$\sigma_p = \frac{1}{2} \log 2 = 0.1505.$$

(Using  $z = -2$  and  $x = 0.5x_a$  gives the same result: try it!)

Generalising, if limits given by  $x_a/q$  and  $qx_a$  are required, we need  $\sigma_p = \frac{1}{2} \log q$ . Furthermore, we can easily toggle between a z-score and the corresponding value of  $r = x/x_a$  (the factor by which a result differs from the assigned value) by using the equations  $r = 10^{z\sigma_p}$  and  $z = \log r / \sigma_p$ . For instance, if  $z = 3.5$  and  $\sigma_p = 0.1505$ , we have  $r = 10^{0.527} = 3.36$ : the result exceeds the assigned value by a factor of 3.36.

### **In conclusion**

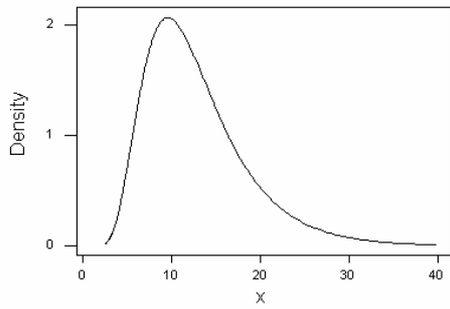
The essential point here is that the major errors seem to be multiplicative in quantitative GMO testing based on PCR. As a consequence, the uncertainty on the original measurement scale is not symmetrically disposed around the result. Regardless of this, z-scores based on log-transformed data can still be treated as symmetric: a z-score of  $-3.5$  has the same importance as one of  $+3.5$ . Similar considerations might apply to any measurement system (such as quantitative microbiology) based on a multiplicative procedure.

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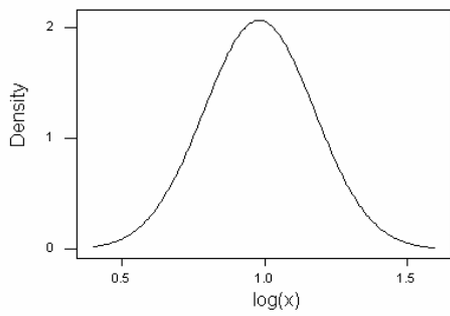
### **References**

1. J Powell and L Owen, *Accred Qual Assur*, 2002, **7**, 392-402.
2. Scoring in Genetically Modified Organism Proficiency Tests Based on Log-Transformed Results, Michael Thompson, Stephen I. R. Ellison, Linda Owen, Kenneth Mathieson, Joanne Powell, Roger Wood and Andrew P Damant, *Journal of AOAC International*, 2006, **89(1)**, 232- 239

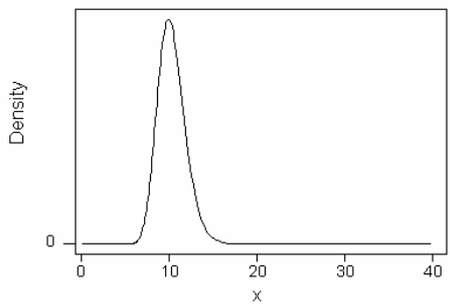
Figures:



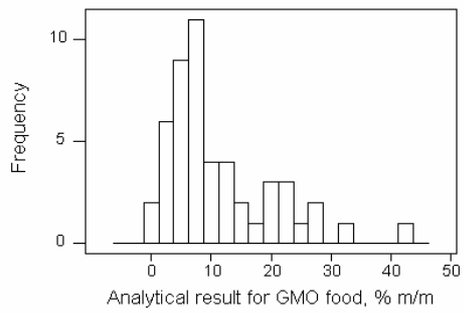
**Figure 1.** A lognormal distribution with an RSD of 0.3.



**Figure 2.** The same distribution as Figure 1, with the density plotted against  $\log x$ .



**Figure 3.** A lognormal distribution with an RSD of 0.1.



**Figure 4.** Results from a single round of a proficiency test involving measuring the concentration of GMO soya.