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Correlation of test results and influence of a mass balance constraint on risks in conformity assessment of a substance or material

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# Measurement

## Correlation of test results and influence of a mass balance constraint on risks in conformity assessment of a substance or material

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<b>Abstract:</b>	<p>When components of a substance or material are subject to a mass balance constraint, test results of the components' contents are intrinsically correlated because of the constraint. That is in addition to possible metrologically-related correlation of test results, and natural and/or technological correlation of the components' contents. Such correlations may influence understanding of compositional data and evaluation of risks in conformity assessment of the substance or material due to measurement uncertainty. A Bayesian multivariate approach to evaluate the conformance probability of multicomponent materials or objects and corresponding risks of false decisions, able to take into account all observed correlations including spurious, is discussed for different scenarios of compositional data. A Monte Carlo method, which includes the mass balance constraint, written in the R programming environment is provided for the necessary calculations. A technique for separation of spurious correlations from experimental (natural and/or technological) correlations is proposed.</p>

## Cover letter

Professor Paolo Carbone,

Editor-in Chief, Measurement

09 April, 2020

Dear Prof. Carbone,

Please find attached the manuscript by Francesca R. Pennecchi, Aglaia Di Rocco, Ilya Kuselman, D. Brynn Hibbert and Michela Sega, titled "Correlation of test results and influence of a mass balance constraint on risks in conformity assessment of a substance or material", which we would like to publish in Measurement.

### **Novelty Statement:**

Three models of prior and likelihood describing composition of a multicomponent material or substance, which consider requirements of a mass balance and different kinds of correlation among the components' contents, are proposed in the framework of a Bayesian multivariate approach.

A Monte Carlo method, coded in the R programming environment, is reported for calculations of the prior coverage probability and the total global risks of false decisions on conformity of the material or substance, taking into account the mass balance constraint.

**Declaration of interest:** There is no any actual or potential conflict of interest of any author.

**Submission declaration and verification:** The work described has not been published previously, it is not also under consideration for publication elsewhere. The publication in Measurement is approved by all the authors.

**The manuscript was previously submitted** to Talanta (on 08.03.20) and rejected on due to outside the scope of the journal: the editor, Dr. Joaquim Nobrega, has recommended to submit it to a metrological journal.

Best regards,

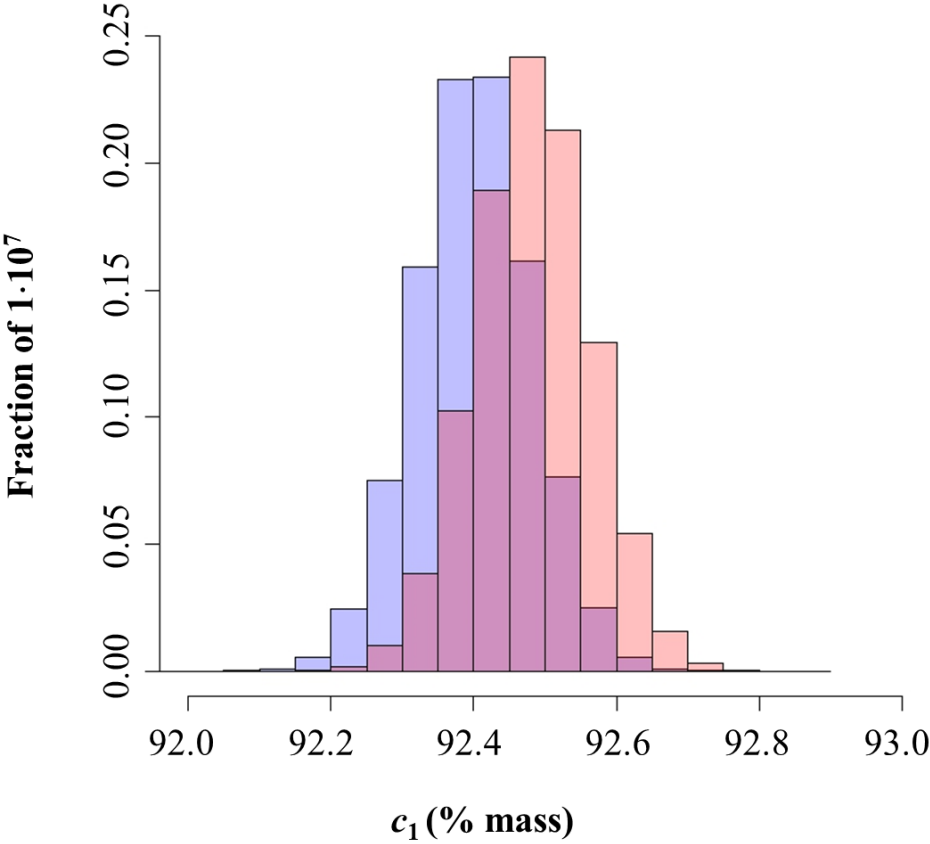


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## HIGHLIGHTS

- Three kinds of correlation of component contents involved in a mass balance are discussed
- Three Bayesian multivariate models describing compositional data are proposed
- A Monte Carlo method for conformity assessment at a mass balance constraint is developed



Histogram of the marginal pdf of the Pt content,  $c_1$ . Pink values represent the distribution of Pt content before closure operation, whereas blue values - after this operation taking into account the mass balance constraint.

## **Correlation of test results and influence of a mass balance constraint on risks in conformity assessment of a substance or material**

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## ABSTRACT

When components of a substance or material are subject to a mass balance constraint, test results of the components' contents are intrinsically correlated because of the constraint. That is in addition to possible metrologically-related correlation of test results, and natural and/or technological correlation of the components' contents. Such correlations may influence understanding of compositional data and evaluation of risks in conformity assessment of the substance or material due to measurement uncertainty. A Bayesian multivariate approach to evaluate the conformance probability of multicomponent materials or objects and corresponding risks of false decisions, able to take into account all observed correlations including spurious, is discussed for different scenarios of compositional data. A Monte Carlo method, which includes the mass balance constraint, written in the R programming environment is provided for the necessary calculations. A technique for separation of spurious correlations from experimental (natural and/or technological) correlations is proposed.

### *Keywords:*

Conformity assessment; Mass balance constraint; Compositional data; Bayesian multivariate modelling; Monte Carlo method; Risk of false decisions



## 1. Introduction

A number of techniques are used during development of a chemical analytical method to overcome possible correlations between test/measurement results of concentrations or contents of different components (main as well as impurities) of a substance or material. Some of these techniques include extraction of target components (analytes) from a sample and chromatographic separation of an analyte from other components of the sample [1]. Chemometrics software is applied for separation of spectral signals and multivariate calibrations of spectrometers [2]. Sample digestion [3] and standard additions of an analyte to a sample [4] are used for calibration of a measuring system to overcome multiplicative matrix effects, and so on. There are validation requirements for “analytical selectivity” of a standard operating procedure (SOP), as its performance characteristic, to prove the procedure’s fitness for purpose. IUPAC Recommendations [5] and the Eurachem Guide [6] define that “analytical selectivity relates to the extent to which the method can be used to determine particular analytes in mixtures or matrices without interferences from other components of similar behavior”. This definition is consistent with “selectivity of a measuring system” in JCGM 200 [7]. The corresponding procedure validation parameter in the pharmaceutical industry is termed in ICH Guideline [8] “specificity”. A procedure not able to answer the requirements of selectivity or specificity is not ‘fit-for-purpose’ and cannot be applied to the given task. Validation of the SOP, training analysts and proficiency testing, supervision and quality control are elements of the quality system of an analytical laboratory that should prevent human error causing metrologically-related correlation, unless, as shown in Fig. 1, the Swiss-cheese model [9] lines up an error in each element. Therefore, correlations that have arisen in the routine measurement process should be in general negligible and test results for two or more components of the same item (sample) are expected to be metrologically independent. In practice, if statistically significant correlation between measured values of concentrations or contents of components of the same sample is detected, analysis cannot be continued without a thorough chemical analytical inspection of the reason for the correlation in the laboratory. For example, when a medication is tested routinely with a pharmacopeial HPLC procedure, correlations might be related to the resolution of the chromatography column used, not able to separate the analytes completely, and the column must be replaced by another one. In the IUPAC/CITAC Guide [9] such an event is rated as a skill-based mistake or omission error (lapse).

Fig. 1

However, test results of chemical compositions of different batches, lots or objects are inevitably correlated when their actual (“true”) values are correlated. Correlations of contents of sample components can be caused by natural physicochemical properties of substances, such as stoichiometry [10-12], and by technological reasons in a material production [13, 14]. These correlations are taken into account using “conventional” multivariate statistical methods.

When components  $i = 1, 2, \dots, n$  of a substance or material are subject to a mass balance constraint (sum of their contents  $c_i$  - mass fractions, mole fractions or any other positive quantity ratios - is 100 % or 1), test results of the components’ contents are called “compositional data”. These data are intrinsically correlated because of the constraint, and the relevant correlation was named by Karl Pearson in 1897 as “spurious” [15-17]. It is a kind of mathematical property of the data, not related to physicochemical interdependence of true values of contents of a material component. Compositional data may be depicted in a multi-dimensional simplex, as in Fig. 2, in which, in general, Euclidean geometry cannot be blindly applied. Compositional Data Analysis (CoDA) based on an isometric logratio transformation of the original test results was developed in the 1980’s by John Aitchison [18-20]. Logratios of amalgamations of test results for pairs of components (parts of the data) – pairwise logratios – were recently used for practical simplification of CoDA [21-24]. There are a number of examples of CoDA applications in geochemistry [25, 26], agriculture and environmental analysis [27-30], bioinformatics [31, 32], forensic science [33, 34], materials [35], and other fields. More publication references, software, slides and lecture notes can be found at website CODAWEB [36].

Fig. 2

There is also a strong message in the literature (e.g., in refs. [25, 26, 37]) stressing how traditional statistical techniques may produce inadequate results if applied to raw compositional data without suitable transformation. However, the relevant techniques of CoDA are still not implemented widely, neither in metrology in chemistry nor in conformity assessment. Spurious correlations may influence evaluation of measurement uncertainty of compositional data (e.g. of standard gas mixtures [38, 39]) and quantification of risks of false decisions due to measurement uncertainty in the conformity assessment of a substance or material. A special case is characterization of materials based on a mass balance [40-43], for example matrix reference materials, as well as evaluation of purity of substances and corresponding (pure) certified reference materials.

A multivariate Bayesian model was recently elaborated for evaluation of risks of false decisions in conformity assessment of multicomponent materials or objects, also taking into account possible correlations between measured concentrations or contents of an item's components [13, 14, 44-47]. This “conventional” approach applies integration of the relevant posterior multivariate probability density function on the tolerance/specification multi-domain of the material (or object) compositions in order to obtain total specific risks, or integration of the joint probability density function of true and measured values to give total global risks. All observed correlations are embedded in the corresponding experimental covariance matrix. This matrix influences the subsequent multivariate results.

A problem of application of CoDA in this field is that a subset of the components may not be related directly to the mass balance, i.e. be outside of the multidimensional simplex; for example, when a sum of impurities' contents is a member of the mass balance, but individual impurities or a subset of those impurities also have to undergo conformity assessment. Since each component participating in conformity assessment has content tolerance/specification limits, the mass balance constraint forms a multivariate sub-domain of the feasible material or object compositions in the domain of specification limits. Moreover, spurious correlation coefficients may be comparable or smaller (in absolute values) than coefficients of correlations caused by natural properties of the materials and technological reasons [14], and in general be not easily distinguishable from them. In addition, using isometric logratio transformation of data, it might be difficult to interpret relevant estimates for conformity assessment purposes in the original variable space.

In this regard, a new IUPAC project [48] was started with the aim of developing guidelines for treatment of influence of a mass balance constraint on measurement uncertainty of test results of a substance or material and risks in its conformity assessment. In the present position paper of the project, a Bayesian multivariate approach for evaluating the probability of conformance of a substance or material and the risks of false decisions is discussed for different scenarios of compositional data. A Monte Carlo method, which includes the mass balance constraint, written in the R programming environment, is developed for the necessary calculations. A technique for separation of spurious correlations from experimental (natural and/or technological) correlations is also proposed.

## 2. A Bayesian multivariate approach to conformity assessment of compositional data

The chemical composition of a material or substance is considered to conform when the actual (“true”) value of the content  $c_i$  of each  $i$ -th component under control,  $i = 1, 2, \dots, n$ , is within its tolerance (specification) interval  $T_i = [T_{Li}, T_{Ui}]$ , where  $T_{Li}$  and  $T_{Ui}$  are lower and upper tolerance limits of the interval, respectively. The term “component” relates to a main component or an impurity or a group of impurities in the item (sample, batch or lot). To decide whether the material or substance conforms or not, the measured content value  $c_{im}$  is compared with the limits of the acceptance interval  $A_i = [A_{Li}, A_{Ui}]$ , where  $A_{Li}$  and  $A_{Ui}$  are lower and upper limits of the interval, respectively, taking into account the standard measurement uncertainty  $u_i$  associated with  $c_{im}$ . This uncertainty causes risks of a false decision on conformity [49].

### 2.1. Compositional data properties

The vector of measured values  $\mathbf{c}_m = [c_{1m}, c_{2m}, \dots, c_{nm}]$  describes the  $n$ -component ( $n$ -part) composition of a material or substance. The compositional data space is the simplex

$$S^n = \{\mathbf{c}_m = [c_{1m}, c_{2m}, \dots, c_{nm}] \mid c_{im} > 0, i = 1, 2, \dots, n; \sum_{i=1}^n c_{im} = k\}, \quad (1)$$

where  $k$  is usually equal to 1 or 100 %, but might be any other positive constant value. As  $c_{im}$  are positive quantity ratios, a vector of measured values  $\mathbf{c}_m$  multiplied by any positive constant contains the same information as the original one, i.e. represents the same composition and can be considered as an equivalence class. This property is termed “scale invariance”. In other words, if  $\mathbf{c}_m$  is scaled by a constant, e.g. measured content values  $c_{im}$  changing from parts-per-unit to percentages, the information which  $\mathbf{c}_m$  conveys is completely equivalent. Therefore, it is natural to select a representative of the equivalence class to facilitate data analysis and interpretation of corresponding results. This selection is formalized by the closure operation:

$$clo(\mathbf{c}_m) = \left[ \frac{k \cdot c_{1m}}{\sum_{i=1}^n c_{im}}, \dots, \frac{k \cdot c_{nm}}{\sum_{i=1}^n c_{im}} \right]. \quad (2)$$

Compositional data do not depend on the order of measured contents  $c_{im}$  within vector  $\mathbf{c}_m$ . The order does not influence conclusions of any analysis on  $\mathbf{c}_m$ . This is known as the “permutation invariance” property. Another property is “subcompositional coherence”, which means that analyses concerning a subset of the component contents should not depend on the remaining part. If a metric is used to compare two compositions on the simplex, the distance between the two should be greater than or equal to that obtained comparing any couple of corresponding subcompositions. That is called “subcompositional dominance”. This property is used to measure distances between compositions and subcompositions following the rule of a projection: distances become smaller in a projection.

Note, ordinary correlations (natural and/or technological) depend on the subcomposition considered and violate subcompositional coherence. The ordinary Euclidean distance between the vectors cannot be evaluated here, as both scale invariance and subcompositional dominance are violated [18].

## 2.2. Modelling composition of a material or substance

Vector  $\mathbf{c} = [c_1, c_2, \dots, c_n]$  represents the actual (“true”)  $n$ -component composition of a material or substance. Component contents  $c_i, i = 1, 2, \dots, n$ , are the measurands in conformity assessment, and  $\mathbf{c}$  is the vector of the measurands. Using a multivariate Bayesian approach [13, 14, 44-47], knowledge about a composition  $\mathbf{c}$  can be modelled by a random multivariate posterior variable and expressed in terms of its probability density function (pdf). Such a pdf combines prior knowledge about the measurands and new information acquired during the measurements:

$$g(\mathbf{c} | \mathbf{c}_m) = C g_0(\mathbf{c}) h(\mathbf{c}_m | \mathbf{c}), \quad (3)$$

where  $g(\mathbf{c} | \mathbf{c}_m)$  is the posterior pdf;  $C$  is a normalizing constant;  $g_0(\mathbf{c})$  is the multivariate prior pdf; and  $h(\mathbf{c}_m | \mathbf{c})$  is the multivariate likelihood function taking into account the measurement uncertainties and possible covariance terms.

A large enough dataset of results of testing items of the same material produced at the same factory, or results of monitoring the same environmental compartment, can be used for modelling the prior pdf  $g_0(\mathbf{c})$ . The assumption is that the actual content values are approximated by the test/measurement results adequately, since measurement uncertainty is negligible in comparison with item-to-item (batch-to-batch) variations caused by changes of conditions of the material production, environmental conditions, etc. Based on this assumption, normal and gamma distributions in JCGM 106 [49], as well as normal, lognormal and Weibull distributions in IUPAC/CITAC Guide [50], were considered as prior pdfs in univariate conformity assessment. Multivariate normal distributions in refs. [13, 14], normal and lognormal distributions in refs. [44, 45] were used in conformity assessment of materials and an environmental object.

If there is no detailed prior knowledge about distribution of the component content in the tested items, the prior pdf is vague. According to the principle of maximum entropy, a multivariate normal distribution is usually considered as the joint prior pdf for the vector of actual component contents  $\mathbf{c}$  when the mean  $\boldsymbol{\mu}$  and covariance matrix  $\mathbf{V}$  constitute the only available information about the vector quantity [51]. In case of compositional data  $\mathbf{c}$ , however, their properties need to be taken into account when assigning a corresponding pdf. Since the vector  $\mathbf{c}$  is non-negative, and the first two moments of the distribution (the expected value and variance) are known, the maximum entropy distribution is a truncated multivariate normal distribution, not a logarithmic one [52]. This is also the case for an elliptical truncation region [53], when the pdf is supposed to be nonzero-valued just for  $(\mathbf{c} - \boldsymbol{\mu})^T \mathbf{V}^{-1}(\mathbf{c} - \boldsymbol{\mu}) \leq k$ .

Therefore, in the present work, a truncated multivariate normal (TMN) distribution on the  $n$ D region  $[0, k]^n$  is employed in modelling the prior pdf of  $\mathbf{c}$ . It is written  $\text{TMN}(\boldsymbol{\mu}, \mathbf{V})$ ,  $\boldsymbol{\mu}$  and  $\mathbf{V}$  being the location and scale parameter, respectively, of the original normal pdf from which the TMN distribution arises by truncation on  $[0, k]^n$ . For the univariate case, the TMN pdf for  $c_i$  on the interval  $[0, k]$  is

$$f_i = \frac{\phi\left(\frac{c_i - \mu_i}{\sigma_i}\right)}{\sigma_i \left( \Phi\left(\frac{k - \mu_i}{\sigma_i}\right) - \Phi\left(\frac{-\mu_i}{\sigma_i}\right) \right)}, \quad (4)$$

where  $\mu_i$  and  $\sigma_i$  are the mean and standard deviation (the location and scale parameters) of the normal pdf from which this TMN distribution arises;  $\phi(\cdot)$  is the pdf of the standard normal distribution and  $\Phi(\cdot)$  is its cumulative distribution function. Outside the interval  $[0, k]$ , the pdf  $f_i$  is equal to zero.

Accordingly, the likelihood function  $h(\mathbf{c}_m|\mathbf{c})$  can be also modelled on the base of a TMN of the measured values, having location parameter equal to the vector  $\mathbf{c}$  of actual/“true” values, i.e.  $\text{TMN}(\mathbf{c}, \mathbf{U})$ , where  $\mathbf{U}$  is the covariance matrix of measurement uncertainties  $u_i$  and covariance terms  $u_{ij}$  whose corresponding correlation coefficients are the same as for  $\mathbf{V}$ . Correlation between measured content values  $c_{im}$  and  $c_{jm}$  is the same as between the corresponding actual ones  $c_i$  and  $c_j$ , when no other (metrologically-related) correlation arose in the measurement process.

The multivariate truncated distribution accommodates the constraint of compositional data to lie on the region  $[0, k]^n$  and is promising for overcoming the above-mentioned problems of application of CoDA for evaluation of risks of false decisions in conformity assessment.

### 2.3. Models for the prior distribution

Let us consider the case of  $c_1 + c_2 + c_3 = 100\%$ , or without loss of generality  $c_1 = 100\% - c_2 - c_3$ . As an example, the composition of a PtRh alloy described in paper [14] is discussed below. Here,  $c_1$  is the Pt content in the PtRh alloy,  $c_2$  is the Rh content, and  $c_3$  is the content of eight impurities (sum of their mass fractions). Prior means  $\mu_i$  and standard deviations  $\sigma_i$ , as well as observed correlation coefficients  $r_{ij}$ ,  $i \neq j$ , for these three components are shown in Table 1. The content of an additional component of the alloy that is under control in conformity assessment – the sum of mass fractions of the three precious impurities (a part of  $c_3$ ) – is not considered here as it is not related directly to the mass balance.

Table 1

The following scenarios have been considered for modelling the multivariate prior probability distribution of the actual (“true”) three-component composition described by the vector  $\mathbf{c}$ :

- 1) Modelling all the actual values of the components’ contents: the closure operation  $clo(\mathbf{c})$  is applied to  $\mathbf{c}$  following a  $\text{TMN}(\boldsymbol{\mu}, \mathbf{V})$  distribution on the 3D region  $[0, 100]^3$ .
- 2) Modelling actual values of two components’ contents and deriving the third:  $[c_2, c_3]$  follows a bivariate  $\text{TMN}(\boldsymbol{\mu}_{23}, \mathbf{V}_{23})$  distribution on the 2D region  $[0, 100]^2$ , where subscript

“23” indicates that the applied operation refers just to components 2 and 3. The scale parameter  $V_{23}$  involves the correlation coefficient  $r_{23} = 0.228$ , and  $c_1$  is deterministically calculated as  $c_1 = 100 \% - c_2 - c_3$ , disregarding possible negative values.

- 3) Sequential modelling:  $c_2$  follows a  $TN(\mu_2, \sigma_2)$ , a univariate normal distribution truncated on the interval  $[0, 100]$ ;  $c_3|c_2$  has a conditional probability distribution following a  $TN(\mu_3, \sigma_3)$  on the interval  $[0, 100 - c_2]$ ; and, again deterministically,  $c_1|(c_2, c_3) = 100 \% - c_2 - c_3$ .

In order to assess the suitability of the models, two parameters are considered: 1) the correlation matrix encompassing correlation coefficients  $r_{ij}$  of the multivariate prior pdf, and 2) the pdf coverage probability  $p$ , calculated as its integral over the multivariate tolerance region, which is  $[92.2 \%, 92.8 \%) \times [7.3 \%, 7.7 \%) \times [0 \%, 0.18 \%)$  for the present example. By definition,  $p$  is the probability that the prior values of all the three variables lie within corresponding tolerance intervals. The calculation of  $p$  using the Monte Carlo (MC) method is described in Appendix 1. Equivalence or dissimilarity of this parameter's values from model to model indicate that the multivariate pdfs generated by the various models are actually the same or different, respectively. Results of calculating  $r_{ij}$  and  $p$  obtained by means of  $10^7$  MC simulations for each model, are reported in Table 2. For comparison,  $p = 0.979$  was obtained when the ordinary multivariate normal distribution was applied in ref. [14] and the original/experimental correlation coefficients shown in Table 1 were used.

Table 2

Correlation coefficients in Table 2 for the prior pdf obtained by models 1 and 2 are equal, within the precision of the MC simulation, to the experimental ones in Table 1, considering that two decimal digits in the coefficient estimates can be taken as reliable when  $10^7$  MC simulations are performed. For both models 1 and 2, coverage probabilities slightly greater than  $p = 0.979$  are obtained, meaning that the prior pdfs generated by these models are narrower than the ordinary multivariate normal pdf.

To show the influence of the closure operation on a TMN pdf, boxplots of the marginal pdfs of the three variables by model 1, before the closure operation (boxplot 1) and after it (boxplot 2), are depicted in Fig. 3. The closure operation shifts the marginal pdf location towards smaller values, especially for  $c_1$ , i.e. the Pt content. More details of this pdf are shown in the histogram of  $c_1$  in Fig. 4.

Fig. 3

Fig. 4



By model 3, the contents of Rh ( $c_2$ ) and the impurities ( $c_3$ ) are each drawn from a univariate truncated normal distribution, one of which depends on the other just in its truncation limits. Pt content values are deterministically generated to satisfy the mass balance constraint. In this model, there is no place for a covariance matrix encompassing the correlation coefficients which reflect different kinds of correlation between the contents. The correlation matrix by model 3 can reveal only the spurious correlations arising from the truncation effect and the mass balance constraint. In the present example, as the sum of Rh and impurities contents is always far away from 100 %, correlation between the two due to the truncation effect is negligible. Thus, the correlation coefficient arising from spurious correlations between Rh and the impurities contents is zero, despite the fact that the experimental correlation coefficient is 0.288. Also, the spurious correlation coefficient for Pt and the impurities contents is different from the observed correlation coefficient, due to both the mass balance constraint and the nature of the raw materials. Nonetheless, the coverage probability obtained for model 3 is very close or equal to those by models 1 and 2.

In general, modelling 1 is suitable for matrix reference materials, standard gas mixtures, food products, etc., i.e. for materials whose composition is tested completely and the test results should correspond to a mass balance. Modelling 2 can be applied for pure substances or materials in which the main component content is not tested using physicochemical measurement methods but calculated based on a mass balance constraint. Sequential modelling 3 is more complicated, since the sequence of  $c_2$  and  $c_3$  can be exchanged and that may lead to different results as the conditional probabilities are different. Moreover, the number of such models increases significantly with the number of components  $n > 3$ . This modelling does not allow consideration of types of correlation other than spurious and cannot be implemented adequately when natural (stoichiometric) and/or technological correlations are significant. On the other hand, modelling 3 could be a helpful tool for understanding sources of the experimental correlation, disentangling its spurious part from the rest.

#### *2.4. Likelihood modelling for a component content calculated based on a mass balance constraint*

According to the propagation of uncertainty prescribed by JCGM 100 [55] for correlated input quantities, when contents of a main component ( $i = 1$ ) are calculated based on a mass balance

constraint  $c_1 = 100 \% - \sum_{i=2}^n c_i$ , the squared measurement uncertainty associated with the calculated  $c_{1m}$  is

$$u_1^2 = \sum_{i=2}^n u_i^2 + 2 \sum_{i=2}^n \sum_{j=i+1}^n u_i u_j r_{ij}. \quad (5)$$

In the example of the three components of the PtRh alloy participating in the mass balance, the standard measurement uncertainty  $u_1$  associated with the calculated Pt content  $c_{1m}$  is evaluated as following:

$$u_1 = (u_2^2 + u_3^2 + 2u_2 u_3 r_{23})^{0.5} = (0.040^2 + 0.011^2 + 2 \cdot 0.040 \cdot 0.011 \cdot 0.228)^{0.5} = 0.044 \text{ (\%)},$$

where  $u_2 = 0.040 \%$  is the standard uncertainty associated with the measured Rh content  $c_{2m}$  in its tolerance (specification) interval, and  $u_3 = 0.011 \%$  is the standard measurement uncertainty associated with the measured mean of the impurities content  $c_{3m} = 0.059 \%$  (the relative standard uncertainty being  $u_3/c_{3m} = 0.18$ ). When correlation between contents of components 2 and 3 is ignored, e.g. in model 3 where  $r_{23} = 0$ , the measurement uncertainty associated with the calculated  $c_{1m}$  is  $u_1 = 0.041 \%$ .

The modelling of the likelihood function for measured content values  $\mathbf{c}_m$  is based on the idea that an appropriate pdf with zero expectation is chosen for an error  $\mathbf{e}_m$  and then translated to the vector of actual (“true”) content values  $\mathbf{c}$  generated for the prior. Therefore,  $\mathbf{c}_m$  is recovered as  $\mathbf{c}_m = \mathbf{c} + \mathbf{e}_m$ . The covariance matrix  $\mathbf{U}$  associated with  $\mathbf{c}_m$  contains the squared above-mentioned uncertainties  $u_i$  and the covariance terms  $u_{ij}$  whose corresponding correlation coefficients are the same as for  $\mathbf{V}$ . The error  $\mathbf{e}_m$  pdf, having zero mean, is the same as that for the likelihood when translated into the vector  $\mathbf{c}$ . The truncation limits are chosen such that each component of the vector  $\mathbf{c}_m$  has to be between zero and 100 %:  $0 \% < \mathbf{c}_m = \mathbf{c} + \mathbf{e}_m < 100 \%$ . Hence, a vector of errors is extracted between  $-\mathbf{c} \%$  and  $(100 - \mathbf{c}) \%$ , where  $\mathbf{c}$  is drawn from the prior pdf. For ease of computation, the  $\mathbf{c}$  values in these truncation limits are approximated with corresponding mean values  $\boldsymbol{\mu}$ . Thus, the modelling of the likelihood follows that of the prior:

- 1) Modelling the likelihood of all the measured component contents:  $\mathbf{e}_m|\mathbf{c}$  is used as a TMN( $\mathbf{0}, \mathbf{U}$ ) distribution on the region  $[-\mu_1, 100 - \mu_1] \% \times [-\mu_2, 100 - \mu_2] \% \times [-\mu_3,$

$100 - \mu_3$  % for the  $c_m$  generation, where  $c$  is approximated by  $\mu$ , and then applying the closure operation  $clo(c_m)$ .

- 2) Modelling the likelihood of two measured component contents and deriving the third:  $[e_{2m}, e_{3m}]|c_{23}$  is taken as a bivariate TMN( $\mathbf{0}, \mathbf{U}_{23}$ ) distribution on the region  $[-\mu_2, 100 - \mu_2] \% \times [-\mu_3, 100 - \mu_3] \%$  for the  $c_{2m}$  and  $c_{3m}$  generation, where  $c_{23}$  is approximated by  $\mu_{23}$ , and  $\mathbf{U}_{23}$  involves the correlation coefficient  $r_{23} = 0.228$ . Then,  $c_{1m} = 100 \% - c_{2m} - c_{3m}$  is calculated directly, disregarding its possible negative values.
- 3) Sequential modelling the likelihood:  $e_{2m}|c_2$  is represented by a univariate TN( $0, u_2$ ) distribution on  $[-\mu_2, 100 - \mu_2] \%$  interval for the  $c_{2m}$  generation, where  $c_2$  is approximated by  $\mu_2$ , and  $e_{3m}|c_{23}$  by a univariate TN( $0, u_3$ ) on the interval  $[-\mu_3, 100 - \mu_3 - c_{2m}] \%$  for the  $c_{3m}$  generation, where  $c_3$  is approximated by  $\mu_3$ . Then,  $c_{1m} = 100 \% - c_{2m} - c_{3m}$  is calculated directly, as in the previous modelling.

When the prior pdf and likelihood function are modelled, the joint pdf for true and measured values given by the product  $g_0(c)h(c_m | c)$ , and the posterior pdf for the measurand  $c$  may be obtained by eqn. (3). Thus, evaluation of risks in conformity assessment of a material or substance, using compositional data, is possible.

### 3. Risks in conformity assessment of compositional data

#### 3.1. Kinds of risks

Several kinds of risk of a false decision on conformity of an item due to measurement uncertainty may be evaluated. The probability of accepting a batch of the material, when it should be rejected, is the ‘consumer’s risk’, whereas the probability of falsely rejecting the batch is the ‘producer’s risk’. For a specified batch, they are referred to as the ‘specific consumer’s risk’  $R_{ci}^*$  and the ‘specific producer’s risk’  $R_{pi}^*$ , respectively, for the  $i$ -th particular component of the material under control [49]:

$$R_{ci}^* = \int_{T_i^c} g(c_i | c_{im}) dc_i \quad \text{and} \quad R_{pi}^* = \int_{T_i} g(c_i | c_{im}) dc_i, \quad (6)$$

where  $T_i^c$  is the interval complementary to the tolerance (specification) interval  $T_i$ , and measured value  $c_{im}$  is inside of the acceptance interval in the case of  $R_{ci}^*$  and outside – in the case of  $R_{pi}^*$ . Thus,  $R_{ci}^*$  relates to the situation when a batch is accepted at the value  $c_i$  out of  $T_i$ , while  $R_{pi}^*$  – when a batch is rejected at  $c_i$  in  $T_i$ .

The risks of incorrect conformity assessment of a batch randomly drawn from a statistical population of such batches are the ‘global consumer’s risk’  $R_{ci}$  and the ‘global producer’s risk’  $R_{pi}$ , respectively, as they characterize the material production globally [49]:

$$R_{ci} = \int_{T_i^c} \int_{A_i} g_0(c_i) h(c_{im} | c_i) dc_{im} dc_i \quad \text{and} \quad R_{pi} = \int_{T_i} \int_{A_i^c} g_0(c_i) h(c_{im} | c_i) dc_{im} dc_i, \quad (7)$$

where  $A_i^c$  is the interval complementary to the acceptance interval  $A_i$ .

When conformity assessment for each  $i$ -th component of a material is successful (i.e. the particular specific or global risks are small enough), the total probability of a false decision concerning the material as a whole (the *total* specific  $R^*$  or *total* global  $R$  risk) might still be significant. The total risks are also consumer’s and producer’s, as shown below with subscripts “c” and “p”, respectively:

$$R_c^* = \int_{T^c} g(\mathbf{c} | \mathbf{c}_m) d\mathbf{c} \quad (\text{for } \mathbf{c}_m \text{ in } A) \quad \text{and} \quad R_p^* = \int_T g(\mathbf{c} | \mathbf{c}_m) d\mathbf{c} \quad (\text{for } \mathbf{c}_m \text{ outside } A), \quad (8)$$

$$R_c = \int_{T^c} \int_A g_0(\mathbf{c}) h(\mathbf{c}_m | \mathbf{c}) d\mathbf{c}_m d\mathbf{c} \quad \text{and} \quad R_p = \int_T \int_{A^c} g_0(\mathbf{c}) h(\mathbf{c}_m | \mathbf{c}) d\mathbf{c}_m d\mathbf{c}, \quad (9)$$

where  $T$  is the tolerance region  $T_1 \times T_2 \times \dots \times T_n$ ;  $A$  is the acceptance region  $A_1 \times A_2 \times \dots \times A_n$ ; and superscript “c” means “complementary”.

Altogether, for  $n > 1$  components under control, one can distinguish  $4(n+1)$  kinds of risks of false decisions [56]. Evaluation of the total global risks of false decisions on the conformity of compositional data (constrained by a mass balance) using a Monte Carlo method in the R programming environment is discussed below. Quantification of total specific risks will be addressed in the framework of the project [48] at a subsequent stage.

### 3.2. Evaluation of the total global risks

To evaluate the global risks in the case of three variables related to a mass balance, prior pdf and likelihood function are modelled as in sections 2.3 and 2.4, respectively. Then,  $10^7$  MC simulations of the joint pdf  $g_0(\mathbf{c})h(\mathbf{c}_m | \mathbf{c})$  are generated simulating prior values from  $g_0(\mathbf{c})$  and corresponding likelihood values from  $h(\mathbf{c}_m | \mathbf{c})$ . The three models for the prior and likelihood discussed in the previous section give three ways to obtain the joint pdf. The function “rtmvnorm” [57] is applied in the first and the second model, and the function “rtnorm” [58] in the third model. Finally, the joint pdf is represented by values of the vector  $[c_1, c_2, c_3, c_{1m}, c_{2m}, c_{3m}]$ .

The total producer's risk  $R_p$  is evaluated considering the fraction of the number of cases (from the total number of  $10^7$  MC simulations) when all the actual values  $c_1, c_2, c_3$  are within the corresponding tolerance region  $T$ , while at least one of the measured values  $c_{1m}, c_{2m}, c_{3m}$  is out of its acceptance interval  $A_i, i = 1, 2, 3$ , respectively. The total consumer's risk  $R_c$  is evaluated as the fraction of the number of cases when all the measured values  $c_{1m}, c_{2m}, c_{3m}$  are within the acceptance region  $A$ , while at least one of the actual values  $c_1, c_2, c_3$  is out of its tolerance interval  $T_i, i = 1, 2, 3$ , respectively. Core calculations of the developed R code are presented in Appendix B.

For the three discussed models, the total global consumer's risk at the acceptance limits equal to the tolerance limits ( $A = T$ ) was the same  $R_c = 4.7 \times 10^{-3}$ . A similar result  $R_c = 5.1 \times 10^{-3}$  was reported in the paper [14] for a scenario when Pt, Rh and the eight impurities contents only were taken into account: the slight difference in the risk values is due to different numerical implementations of the risk calculation. For more details, the  $R_c$  dependence on  $A_2$  and  $A_3$  (at  $A_1 = T_1$ ) was also studied, as shown in Fig. 5a for model 1. The arrows on both the plots in Fig. 5 show the acceptance intervals from  $A_2 = T_2$  and  $A_3 = T_3$ , at the beginning of the axes, to the acceptance intervals with limits differing from the tolerance limits by three times the relevant measurement uncertainty ( $3u_i$ ), at the axes' ends. Specifically, in Fig. 5a, these are the lower acceptance limit  $A_{L2} = T_{L2} + 3u_2 = 7.3 + 3 \times 0.04 = 7.42$  (%) and the upper acceptance limit  $A_{U2} = T_{U2} - 3u_2 = 7.7 - 3 \times 0.04 = 7.58$  (%). Since the upper only tolerance limit  $T_{U3}$  is set for the content of the impurities, at the end of the  $A_3$  axis  $A_{L3} = 0$  and  $A_{U3} = T_{U3} - 3u_3 = 0.18 - 3 \times (0.059 \times 0.18) = 0.15$  (%), where 0.059 % is the value of the content prior mean  $\mu_3$  of the impurities (Table 1). The color column bar in the plot gives indication of the risk  $R_c$  values between the minimum and the maximum on the surface, from  $3 \times 10^{-6}$  to  $4.7 \times 10^{-3}$ . One can see on the plot that the influence of the impurities'

Fig. 5

content on the risk is negligible under these conditions. That is because the content prior mean  $\mu_3 = 0.059\%$  is far enough from the tolerance limit  $T_{U3} = 0.18\%$ .

The dependences of  $R_c$  on the acceptance intervals for the second and the third models are very similar ( $R_c$  varying from  $4 \times 10^{-6}$  to  $4.7 \times 10^{-3}$  and  $R_c$  from  $6 \times 10^{-6}$  to  $4.6 \times 10^{-3}$ , respectively). These results show that the mass balance influence on  $R_c$  in the small sub-domain of feasible alloy compositions is practically negligible.

The total global producer's risk at  $A = T$  was  $R_p = 2.4 \times 10^{-2}$  for models 1 and 2, and a bit smaller  $R_p = 2.0 \times 10^{-2}$  for model 3. The producer's risk values were not discussed in ref. [14] but have been calculated, for sake of comparison, according to the approach proposed there, leading to the same value  $R_p = 2.4 \times 10^{-2}$ . The  $R_p$  dependence on  $A_2$  and  $A_3$  (at  $A_1 = T_1$ ) is presented for model 1 in Fig. 5b, where, at the end of the axes,  $A_{L2} = T_{L2} - 3u_2 = 7.3 - 3 \times 0.04 = 7.18\%$  and  $A_{U2} = T_{U2} + 3u_2 = 7.7 + 3 \times 0.04 = 7.82\%$ , and  $A_{L3} = 0$  and  $A_{U3} = T_{U3} + 3u_3 = 0.18 + 3 \times (0.059 \times 0.18) = 0.21\%$ , respectively. The color column bar gives indication of  $R_p$  values from  $4.9 \times 10^{-3}$  to  $2.4 \times 10^{-2}$ . The influence of the impurities' content on the producer's risk is minor at these conditions, as it is for the consumer's risk for the same reason.

The dependence of  $R_p$  on the acceptance intervals for the second model is very similar as well, its values being practically the same:  $R_p$  from  $4.8 \times 10^{-3}$  to  $2.4 \times 10^{-3}$ . However, the third model produces smaller values of  $R_p$  from  $3.7 \times 10^{-4}$  to  $2.0 \times 10^{-3}$ .

The obtained results show that the mass balance influence on the consumer's and producer's risks, evaluated with models 1 and 2, is not visible in the small sub-domain of feasible alloy compositions. The plots by these two models, like in Fig. 5, allow choice of acceptance limits corresponding to the suitable risks of the consumer(s) and the producer. The departure of results by model 3, with respect to the results obtained with models 1 and 2, is caused by the fact that model 3 does not take into account correlations other than spurious (natural and technological). Therefore, this model is helpful not only to separate spurious correlation from the observed/experimental (cumulative) correlation, but also to indicate how significant the difference between them in a material conformity assessment is.

## 4. Conclusions

Three kinds of correlation can be considered between components' contents of a substance or material which meet requirements of a mass balance: 1) metrologically-related, 2) natural and technological, and 3) spurious. All these kinds of correlation have been taken into account in the Bayesian multivariate approach modified for evaluating the conformance probability of a substance or material and the risks of false decisions in its conformity assessment when the material components' contents satisfy a mass balance constraint.

Three models of prior pdfs and likelihood functions describing the compositional data in the framework of the Bayesian multivariate approach can be applied. Model 1 is suitable for materials whose composition is tested completely and for which the test results should correspond to a mass balance. Model 2 can be applied for testing pure substances or materials in which the main component content is not measured using physicochemical measurement methods, but calculated based on a mass balance constraint. Model 3 is a helpful tool for understanding sources of experimental correlation, disentangling the spurious part from natural and/or technological correlations.

A Monte Carlo method, coded in the R programming environment, was developed for calculations of the coverage probability of the prior pdf and the total global risks of false decisions on conformity of a material or substance, taking into account the mass balance constraint of the data.

The approach to conformity assessment of compositional data is demonstrated using an example of a PtRh alloy, for which spurious correlation effects were minor.

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## **Appendix A. Evaluation of coverage probability**

Evaluation of the coverage probabilities for three variables according to the models of prior pdfs described in sec. 2.3 was performed in the R programming environment using the library “compositions” [59] and a Monte Carlo method with  $10^7$  simulations.

For the first model, where  $\mathbf{c}$  follows a TMN( $\boldsymbol{\mu}, \mathbf{V}$ ) distribution on the region  $[0, 100]^3$ , a prior pdf as a trivariate truncated normal distribution was generated using the function “rtmvnorm” with truncation limits equal to  $[0, 0, 0]$  and  $[100, 100, 100]$ . Note, a non-singular matrix is inserted as an approximation of matrix  $\mathbf{V}$ . Then, the function “clo” was applied for performing the closure operation with the constraint equal to 100 %.

For the second model the function “rtmvnorm” was applied to generate a bivariate truncated normal distribution with truncation limits  $[0, 0]$  and  $[100, 100]$ . Then, the third variable was computed as  $c_1|(c_2, c_3) = 100 \% - c_2 - c_3$ . Rows  $[c_1, c_2, c_3]$  of the matrix, containing negative generated values of the third component content, were removed.

In the third model,  $c_2$  is generated using the function “rtnorm” with the truncation lower limit 0 % and the upper limit 100 %. Then, variable  $c_3|c_2$  was given by the same function, “rtnorm”, but with the truncation limits 0 % and  $100 \% - c_2$ , i.e. for each value of  $c_2$  the corresponding value of  $c_3$  was in the interval  $[0, 100 - c_2]$ . In this way  $0 < c_2 + c_3|c_2 < 100$ . Finally, the third variable was computed as  $c_1|(c_2, c_3) = 100 \% - c_2 - c_3$ .

When the prior pdf is generated in the form of a matrix containing rows of values  $[c_1, c_2, c_3]$ , the coverage probability can be evaluated using function “length” to count cases where  $c_1, c_2$  and  $c_3$  values are simultaneously within their limits. For each row corresponding to this condition, the function gives the number 3, because three variables were treated. Thus, to obtain the coverage probability, the result of the “length” function should be divided by  $3 \times 10^7$ . One can also compute marginal coverage probability, taking into account the variables one by one. Core calculations of the developed R code are presented in Appendix B.

## Appendix B. Core calculations of the R code

### *B.1. Calculation of the coverage probability of the prior pdf*

```
library(mvtnorm)
library(msm)
library(tmvtnorm)
library(compositions)
library(nortest)

rm(list=ls())
```



```

# Tolerance limits
lrl1 = 92.2
url1 = 92.8
lrl2 = 7.3
url2 = 7.7
lrl3 = 0
url3 = 0.18

#####
# MODEL 1

n = 10000000
mu1 = 92.483
mu2 = 7.547
mu3 = 0.059
uc1 = 0.081
uc2 = 0.073
uc3 = 0.021
mean = c(mu1,mu2,mu3)
sigma = matrix(c(uc1^2,-0.967*uc1*uc2,-0.467*uc1*uc3,-
0.967*uc1*uc2,uc2^2,0.228*uc2*uc3,-0.467*uc1*uc3,0.228*uc2*uc3,uc3^2), nrow = 3)
lower = rep(0,3)
upper = rep(100, 3)

priorTMN = rtmvnorm(n, mean = mean, sigma = sigma, lower, upper)
prior = clo(priorTMN, total=100)

prior1 = prior[,1]
prior2 = prior[,2]
prior3 = prior[,3]

# Correlation matrix of the modelled multivariate prior PDF
cor(cbind(prior1, prior2, prior3))

# Joint coverage probability
Pcov123 = length(prior[(prior1>=lrl1 & prior1<=url1) & (prior2>=lrl2 & prior2<=url2) &
(prior3>=lrl3 & prior3<=url3)])/(3*n)
Pcov123

# X Marginal coverage probability
Pcov1 = length(prior1[prior1>=lrl1 & prior1<=url1])/n
Pcov1

# Y Marginal coverage probability
Pcov2 = length(prior2[prior2>=lrl2 & prior2<=url2])/n

```

Pcov2

# Z Marginal coverage probability

Pcov3 = length(prior3[prior3>=lrl3 & prior3<=url3])/n

Pcov3

#####

# MODEL 2

n = 10000000

mu2 = 7.547

mu3 = 0.059

uc2 = 0.073

uc3 = 0.021

mean23 = c(mu2,mu3)

sigma3 = matrix(c(uc2^2,0.228\*uc2\*uc3,0.228\*uc2\*uc3,uc3^2), nrow = 2)

lower = rep(0,2)

upper = rep(100,2)

prior23 = rtmvnorm(n,mean23,sigma3,lower,upper)

prior2 = prior23[,1]

prior3 = prior23[,2]

prior1 = c(rep(100,n)) - prior2 - prior3

prior123 = cbind(prior1,prior2,prior3)

prior = prior123[prior123[,1]>=0,] # Only generated vectors in which X values are non-negative are retained

prior1 = prior[,1]

prior2 = prior[,2]

prior3 = prior[,3]

# Correlation matrix of the modelled multivariate prior PDF

cor(prior)

# Joint coverage probability

Pcov123 = length(prior[(prior1>=lrl1 & prior1<=url1) & (prior2>=lrl2 & prior2<=url2) & (prior3>=lrl3 & prior3<=url3)])/(3\*n)

# X Marginal coverage probability

Pcov1 = length(prior1[prior1>=lrl1 & prior1<=url1])/length(prior1)

# Y Marginal coverage probability

Pcov2 = length(prior2[prior2>=lrl2 & prior2<=url2])/length(prior1)

# Z Marginal coverage probability

Pcov3 = length(prior3[prior3>=lrl3 & prior3<=url3])/length(prior1)

```
#####
```

```
# MODEL 3
```

```
n = 10000000
```

```
mu2 = 7.547
```

```
mu3 = 0.059
```

```
uc2 = 0.073
```

```
uc3 = 0.021
```

```
prior2 = rtnorm(n,mu2,uc2,0,100)
```

```
prior3 = rtnorm(n,mu3,uc3,0,100-prior2)
```

```
prior1 = c(rep(100,n)) - prior2 - prior3
```

```
prior = cbind(prior1,prior2,prior3)
```

```
# Correlation matrix of the modelled multivariate prior PDF
```

```
cor(prior)
```

```
# Joint coverage probability
```

```
Pcov123 = length(prior[(prior1>=lrl1 & prior1<=url1) & (prior2>=lrl2 & prior2<=url2) &
(prior3>=lrl3 & prior3<=url3)])/(3*n)
```

```
# X Marginal coverage probability
```

```
Pcov1 = length(prior1[prior1>=lrl1 & prior1<=url1])/n
```

```
# Y Marginal coverage probability
```

```
Pcov2 = length(prior2[prior2>=lrl2 & prior2<=url2])/n
```

```
# Z Marginal coverage probability
```

```
Pcov3 = length(prior3[prior3>=lrl3 & prior3<=url3])/n
```

## *B.2. Calculation of the total global risks*

```
# Inizializations and setting
```

```
library(mvtnorm)
```

```
library(msm)
```

```
library(tmvtnorm)
```

```
library(compositions)
```

```
library(nortest)
```

```
rm(list=ls())
```

```
lrl1 = 92.2
```

```
# Lower regulation limit
```

```
url1 = 92.8
```

```
# Upper regulation limit
```

```

lrl2 = 7.3          # Lower regulation limit
url2 = 7.7          # Upper regulation limit
lrl3 = 0            # Lower regulation limit
url3 = 0.18         # Upper regulation limit
lal1 = lrl1         # Lower acceptance limit
lal2 = lrl2         # Lower acceptance limit
lal3 = lrl3         # Lower acceptance limit
ual1 = url1         # Upper acceptance limit
ual2 = url2         # Upper acceptance limit
ual3 = url3         # Upper acceptance limit
lrl = c(lrl1,lrl2,lrl3)
url = c(url1,url2,url3)
lal = c(lal1,lal2,lal3)
ual = c(ual1,ual2,ual3)

mu1 = 92.483        # Prior mean value of the true results
mu2 = 7.457         # Prior mean value of the true results
mu3 = 0.059         # Prior mean value of the true results
uc1 = 0.081         # Prior sd
uc2 = 0.073         # Prior sd
uc3 = 0.021         # Prior sd

rucm2 = 0.04/mu2*100 # RELATIVE (%) measurement uncertainty
rucm3 = 0.18*100     # RELATIVE (%) measurement uncertainty
rucm1 = sqrt(0.04^2+0.18^2*mu3^2+2*0.228*0.04*0.18*mu3)/mu1*100 # RELATIVE (%)
measurement uncertainty
ucm1 = rucm1*mu1/100 # ABSOLUTE (%) measurement uncertainty
ucm2 = rucm2*mu2/100 # ABSOLUTE (%) measurement uncertainty
ucm3 = rucm3*mu3/100 # ABSOLUTE (%) measurement uncertainty

#####
# Model 1
mu = c(mu1,mu2,mu3)
sigma = matrix(c(uc1^2, -0.967*uc1*uc2, -0.467*uc1*uc3, -0.967*uc1*uc2, uc2^2,
0.228*uc2*uc3, -0.467*uc1*uc3, 0.228*uc2*uc3,uc3^2), nrow = 3)
sigma_lik = matrix(c(ucm1^2, -0.967*ucm1*ucm2, -0.467*ucm1*ucm3, -0.967*ucm1*ucm2,
ucm2^2, 0.228*ucm2*ucm3, -0.467*ucm1*ucm3, 0.228*ucm2*ucm3, ucm3^2), nrow = 3)

# PRIOR
n = 10000000
lowerT = rep(0,3)
upperT = rep(100,3)
prior = rtmvnorm(n,mu,sigma,lowerT,upperT)
prior = clo(prior, total = 100)

# LIKELIHOOD

```

```
like = prior + rtmvnorm(n, c(rep(0,3)), sigma_lik, lower = -mu, upper = 100 - mu)
like = clo(like, total = 100)
```

#### # PRODUCER'S RISK

```
m = 3 # Number of components
z=0
for(i in 1:n) {
  if(all(prior[i,]>=lrl[1:m] & prior[i,]<=url[1:m]) & any(like[i,]>ual[1:m] | like[i,]<lal[1:m])) {
    z=z+1 }
}
RtotglobP = z/n
```

#### # CONSUMER'S RISK

```
m = 3 # Number of components
q=0
for(i in 1:n) {
  if(all(like[i,]>=lal[1:m] & like[i,]<=ual[1:m]) & any(prior[i,]>url[1:m] | prior[i,]<lrl[1:m])) {
    q=q+1 }
}
RtotglobC = q/n
```

```
#####
```

#### # Model 2

```
mu = c(mu2,mu3)
sigma = matrix(c(uc2^2, 0.228*uc2*uc3, 0.228*uc2*uc3, uc3^2), nrow = 2)
sigma_lik = matrix(c(ucm2^2, 0.228*ucm2*ucm3, 0.228*ucm2*ucm3, ucm3^2), nrow = 2)
lowerT = rep(0,2)
upperT = rep(100,2)
```

#### # PRIOR

```
n = 10000000
prior = rtmvnorm(n,mu,sigma,lowerT,upperT)
prior2 = prior[,1]
prior3 = prior[,2]
prior1 = c(rep(100,n)) - prior2 - prior3
prior123 = cbind(prior1,prior2,prior3)
prior = prior123[prior123[,1]>=0,] # Only generated vectors in which X values are non-negative
are retained
```

#### # LIKELIHOOD

```
like23 = prior[,2:3] + rtmvnorm(n, c(rep(0,2)), sigma_lik, lower = -mu, upper = 100 - mu)
like2 = like23[,1]
like3 = like23[,2]
like1 = 100 - like2 - like3
like123 = cbind(like1,like2,like3)
```

```
like = like123[like123[,1]>=0,]
prior = prior[like123[,1]>=0,]      # Updated prior, disregarding values corresponding to
negative likelihood values for the first quantity
```

#### # PRODUCER'S RISK

```
m = 3 # Number of components
```

```
z=0
```

```
for(i in 1:dim(prior)[1]) {
  if(all(prior[i,]>=lrl[1:m] & prior[i,]<=url[1:m]) & any(like[i,]>ual[1:m] | like[i,]<lal[1:m])) {
    z=z+1 }
  }
```

```
RtotglobP = z/n
```

#### # CONSUMER'S RISK

```
m = 3 # Number of components
```

```
q=0
```

```
for(i in 1:dim(prior)[1]) {
  if(all(like[i,]>=lal[1:m] & like[i,]<=ual[1:m]) & any(prior[i,]>url[1:m] | prior[i,]<lrl[1:m])) {
    q=q+1 }
  }
```

```
RtotglobC = q/n
```

```
#####
```

```
# Model 3
```

```
n = 10000000
```

```
prior2 = rtnorm(n,mu2,uc2,0,100)
```

```
like2 = prior2 + rtnorm(n, 0, ucm2, lower = -prior2, upper = 100 - prior2)
```

```
prior3 = rtnorm(n,mu3,uc3,0,100-prior2)
```

```
like3 = prior3 + rtnorm(n, 0, ucm3, lower = -prior3, upper = 100 - like2 - prior3)
```

```
prior1 = c(rep(100,n)) - prior2 - prior3
```

```
like1 = 100 - like2 - like3
```

```
joint123 = cbind(prior1,prior2,prior3,like1,like2,like3)
```

#### # PRODUCER'S RISK

```
RtotglobP = length(joint123[(prior1>=lrl1 & prior1<=url1) & (prior2>=lrl2 & prior2<=url2) &
(prior3>=lrl3 & prior3<=url3) & (like1<lal1 | like1>ual1 | like2<lal2 | like2>ual2 | like3<lal3 |
like3>ual3)])/(6*n)
```

#### # CONSUMER'S RISK

```
RtotglobC = length(joint123[(like1>=lal1 & like1<=ual1) & (like2>=lal2 & like2<=ual2) &
(like3>=lal3 & like3<=ual3) & (prior1<lrl1 | prior1>url1 | prior2<lrl2 | prior2>url2 | prior3<lrl3 |
prior3>url3)])/(6*n)
```

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## Figure captions

**Fig. 1. Metrologically-related correlation arising in a multicomponent chemical analysis - the Swiss-cheese model.**

**Fig. 2. Orthogonal coordinates (Euclidian space) and simplex.** The variables  $c_i$  are contents of material components  $i = 1, 2, 3$  and  $4$ , %; each simplex vertex is  $c_i = 100$  %;  $D$  is dimension. In each case mass balance constraint reduces the number of dimensions by one.

**Fig. 3. Boxplots of the marginal prior pdfs of the components' contents.** Boxplot 1 is before closure operation, and boxplot 2 after it; for a) Pt content,  $c_1$ ; b) Rh content,  $c_2$ ; and c) content of the eight impurities,  $c_3$ . The band near the middle of each box is the 50 % percentile (the median, equal to the mean when the distribution is symmetrical); the bottom and top of the box corresponds to 25 % and 75 % percentiles, respectively; the distance between upper and/or lower whiskers and the box is equal to the 1.5 box length [54].

**Fig. 4. Histogram of the marginal pdf of the Pt content,  $c_1$ .** Pink bars show frequencies of the Pt content (fractions of  $1 \times 10^7$  MC simulations) before the closure operation, while blue bars present the frequencies after this operation (i.e. after taking into account the mass balance constraint).

**Fig. 5. Surfaces of the total global consumer's risk  $R_c$  and producer's risk  $R_p$  vs. acceptance intervals for measured Rh content ( $A_2$ ) and measured content of the eight impurities ( $A_3$ ).** The arrows show the acceptance intervals from  $A_2 = T_2$  and  $A_3 = T_3$  at the axes beginning, to the acceptance intervals with limits differing from the tolerance limits by three times the relevant measurement uncertainties at the axes end, when for Pt content  $A_1 = T_1$ . The color column bar in plot 5a gives indication of the  $R_c$  values between the minimum and the maximum on the surface, i.e. from  $3 \times 10^{-6}$  to  $4.7 \times 10^{-3}$ ; in plot 5b the bar is for the  $R_p$  values between  $4.9 \times 10^{-3}$  and  $2.4 \times 10^{-2}$ .

Fig. 1

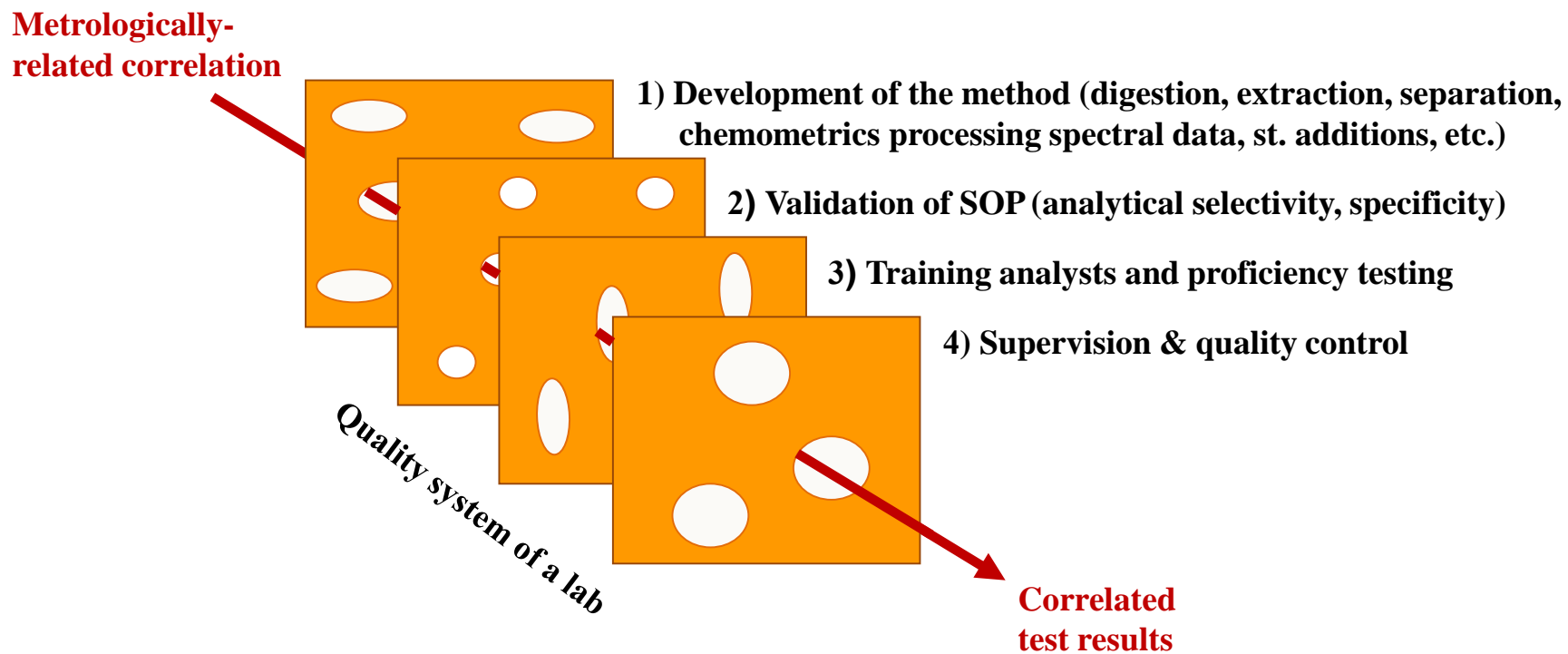
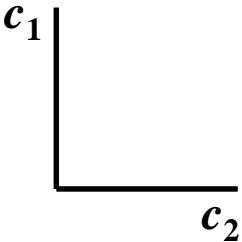


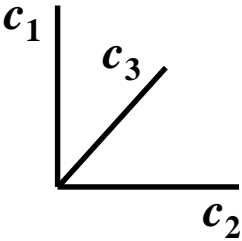
Fig. 2



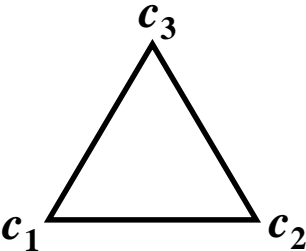
**2D → 1D**



**$c_1 + c_2 = 100 \%$**

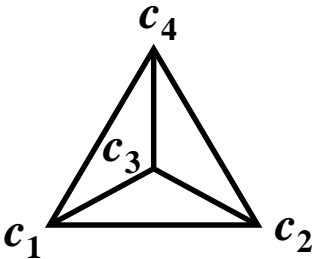


**3D → 2D**



**$c_1 + c_2 + c_3 = 100 \%$**

**Orthogonal/Euclidian 4D → 3D**



**$c_1 + c_2 + c_3 + c_4 = 100 \%$**

Fig. 3





Fig. 4

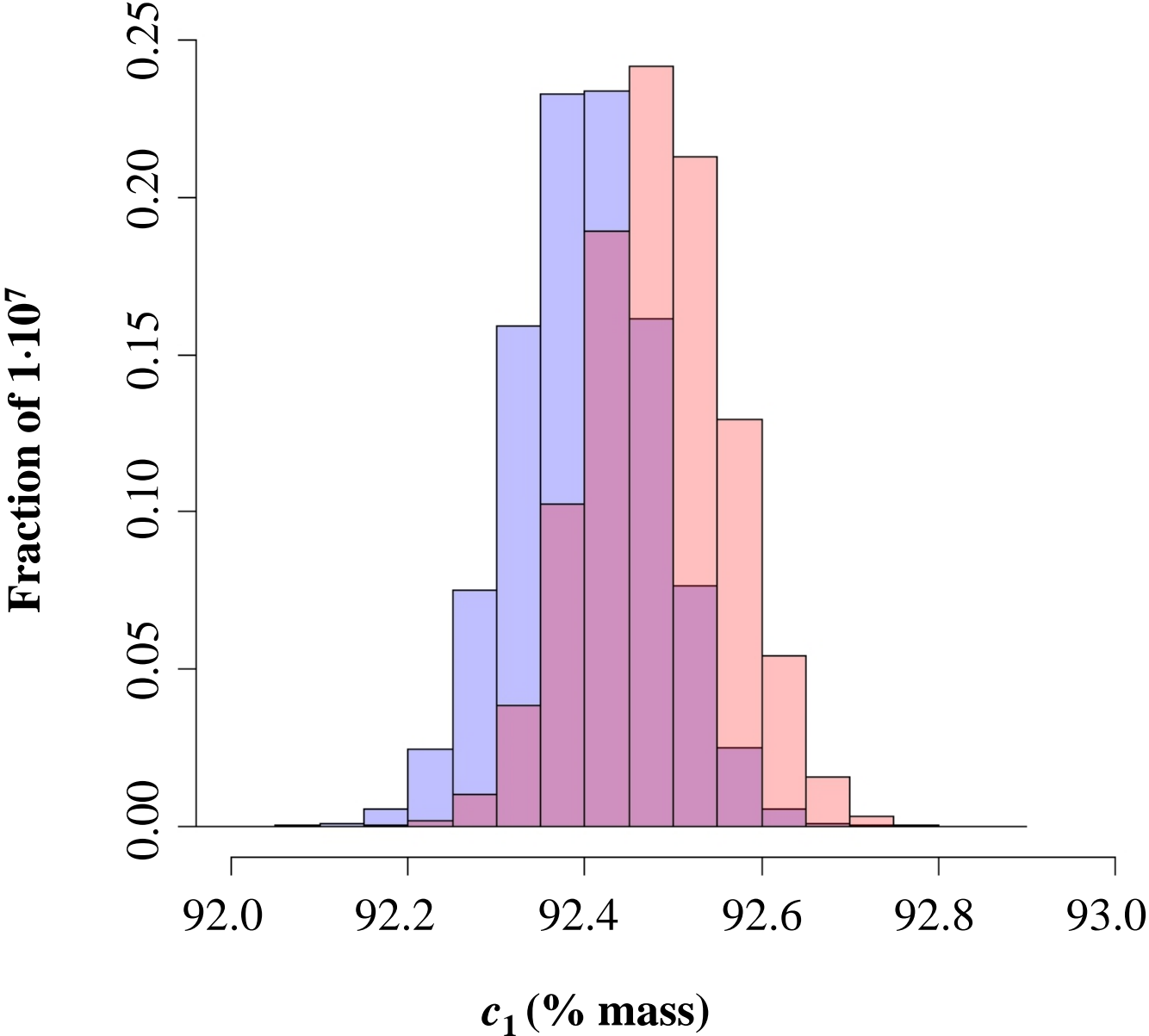
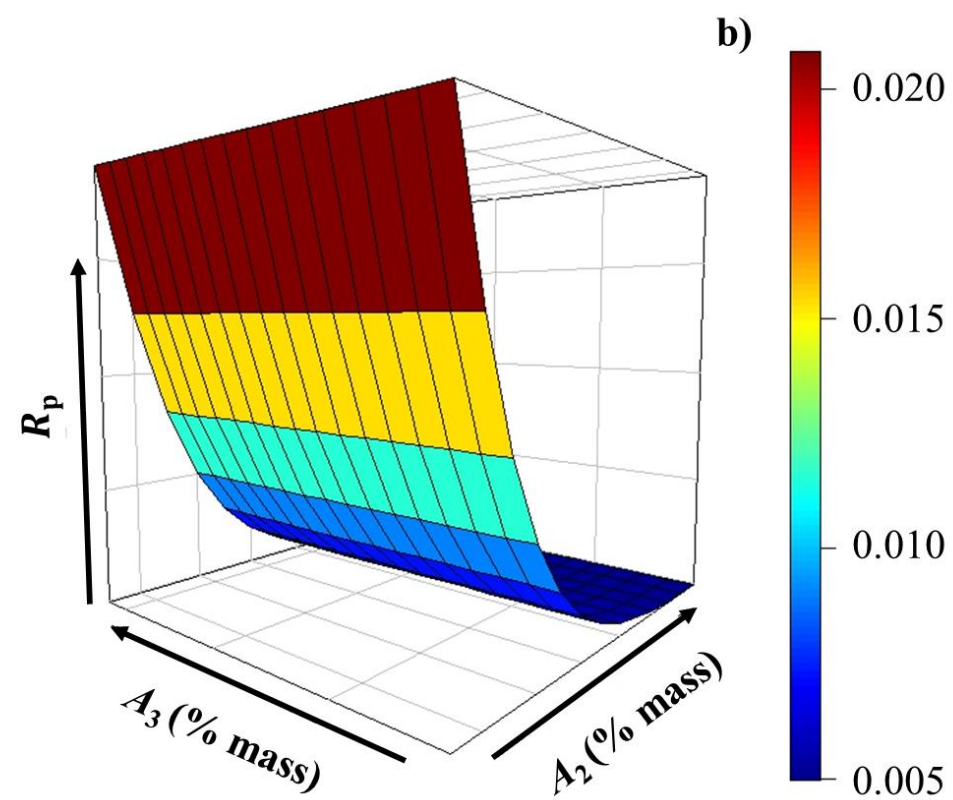
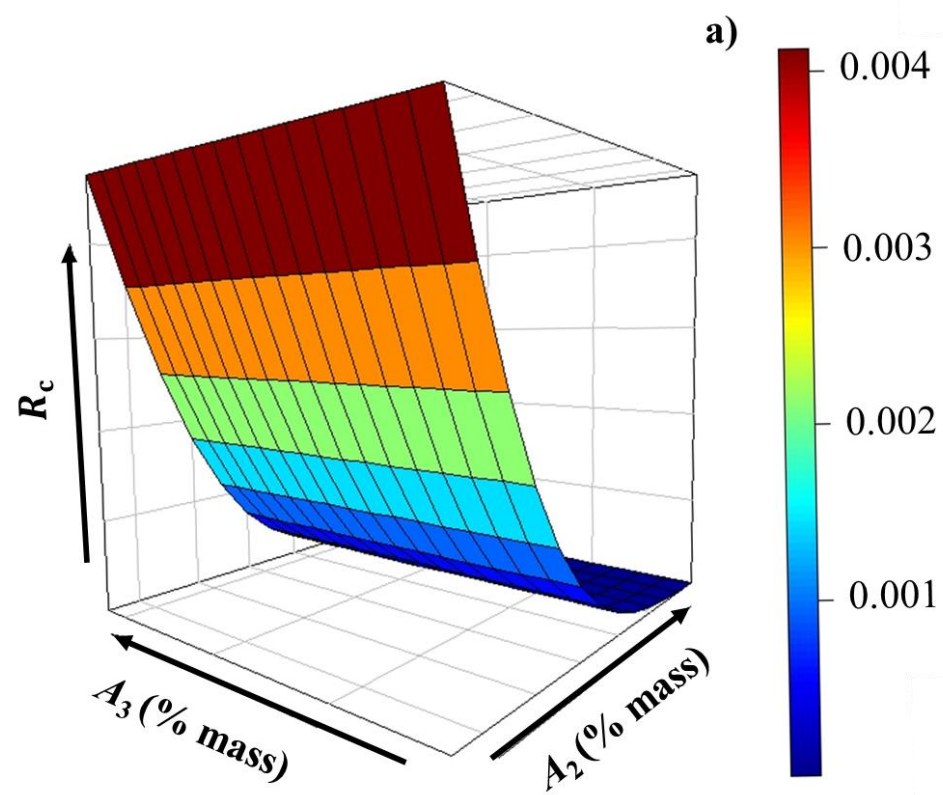


Fig. 5



**Table 1. Parameters of the prior distribution and measurement uncertainties for a PtRh alloy [14].**

component	$i$	$\mu_i$ , %	$\sigma_i$ , %	$ij$	$r_{ij}$
Pt	1	92.483	0.081	12	-0.967
Rh	2	7.457	0.073	13	-0.467
8 impurities	3	0.059	0.021	23	0.228

**Table 2. Correlation coefficients and coverage probabilities calculated from  $10^7$  Monte Carlo simulations for the PtRh alloy system**

Parameters	Model		
	1	2	3
$r_{12}$	-0.968	-0.968	-0.962
$r_{13}$	-0.464	-0.464	-0.274
$r_{23}$	0.226	0.226	0.000
$p$	0.985	0.981	0.981

**Declaration of interests**

☒ The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

☐The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: