NMISA-PT104

Proficiency Test Description

Elements, Proximates and Amino Acids in Fortified Milk Powder

Issue date: 14 April 2023
1 FOREWORD

This is the call for participation in, and description of the NMISA proficiency test (PT) for the determination of selected nutritional labelling parameters in fortified milk powder material. Participants will be required to report on all parameters which form part of their routine or developing laboratory services. A confidential report will be issued to all participants after completion of the PT. Information on the material, parameters included for potential performance evaluation, as well as dates for the registration, distribution and reporting is listed in Table 1.

This forms part of a range of ISO 17043 accredited PT services offered by NMISA. Please consult our website www.nmisa.org for information on PT schemes on offer. NMISA can also assist with the preparation of traceable gravimetrically prepared spike solutions for benchmarking ad-hoc analyses for which commercial PT schemes are not available.

2 PT AIMS

This PT will assist laboratories that routinely analyse nutritional labelling parameters in food matrices, to monitor their laboratory performance. The PT allows laboratories to evaluate their accuracy and comparability of measurement results produced; the continued competency of analytical staff; and the maintenance and effectiveness of the current quality assurance systems within the laboratory. In addition, this information may also be used to provide accreditation bodies or clients with objective evidence of laboratory performance.

3 PARTICIPATION FEES AND ADDITIONAL CHARGES

The cost of participation in the PT is R 5 000. This fee includes the material and a confidential report upon completion, but excludes costs associated with delivery (0% VAT, please note that we are not a VAT registered company).

Since many of the South African participants are located within close proximity to NMISA, the option of collecting the PT scheme samples from NMISA premises is permitted. International laboratories will have test samples sent by courier and appropriately packaged to maintain sample integrity. International participants must provide NMISA with any import or quarantine permits that might be required to complete sample delivery well in advance of the shipment date and are liable for any customs or import duties charged.

Upon registration for participation an official quotation will be provided. Participation is confirmed following receipt of a purchase order and/or proof of payment.

4 PT DESCRIPTION

The timeline for the PT is presented in Table 1. Laboratories are requested to report results for as many of the parameters specified as possible, to allow for maximum benefit from the participation. This study is designed to support laboratories routinely performing nutritional labelling measurements. The levels of the analytes should be easily achievable using analytical methods typically applied, however, some of the trace elements can be expected to represent a measurement challenge. Instructions for proper handling and storage of the samples prior to sample preparation will accompany the PT samples. Participants should adhere to these instructions to ensure sample integrity and comparability of the results.
### Parameters

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Sample format</th>
<th>Distribution/ Dispatch</th>
<th>Result reporting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minerals (0.2 – 1200 mg/100 g): Iron, copper, zinc, sodium, magnesium, calcium, potassium, phosphorus, manganese</td>
<td>2 x 20 g fortified milk powder</td>
<td>Jan 2024</td>
<td>Feb 2024</td>
</tr>
<tr>
<td>*Proximates (2 – 20 g/100 g): Protein (N = 6.25)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Amino acids: Alanine, arginine, aspartic acid, cystine, glutamic acid, glycine, histidine, isoleucine, leucine, lysine, methionine, phenylalanine, proline, serine, threonine, tryptophan, tyrosine, valine</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Result Reporting

Participants will be required to perform the analysis using their normal laboratory procedures and are required to report two results for each parameter measured in each of the samples provided.

Participants are encouraged to include an uncertainty estimate for each result obtained. The result reporting form will be distributed to participants and will request additional information on the measurement technique and parameters, any recovery correction application, calibration standards used etc.

### Assigned value

- The assigned value for the elements will be the reference values obtained through ICP analysis at the NMISA Inorganic Analysis Laboratory.
- *The assigned values for the proximates and the amino acids will be the consensus values determined from participant results (and/or expert laboratories) in accordance with ISO 13528:2017 statistical principles.*

### Laboratory performance

- Laboratory performance will be evaluated using the z-score

### Standard deviation of proficiency assessment

- NMISA typically employs the modified Horwitz model to estimate the reproducibility standard deviation that can typically be expected.
- Where and if applicable, the standard deviation for proficiency assessment will be in accordance with the tolerances stipulated in section 3 of Guideline 5 referred to in the South African regulations related to food labelling (R146).
- The standard deviation of participant results will also be included in the final PT report for reference and the robust standard deviation of participant results may be considered as the standard deviation of proficiency assessment.

### PT report

- The PTS report will be distributed within 2 weeks following the result submission deadline. Reports will be provided in electronic format only (Adobe Acrobat-pdf) files.
- The scheme is fully confidential. Each participant will be issued with a unique identification number. For multiple participants within the same laboratory the participating laboratory is required to identify its analysts by a code known only to the laboratory.

*The assigned value and subsequent performance evaluation can only be determined on parameters where at least 10 results have been received.*