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MP-1a: A CERTIFIED REFERENCE ORE

H.F. STEGER AND W.S. BOWMAN



MINERALS RESEARCH PROGRAM
MINERAL SCIENCES LABORATORIES



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MP-la: A Certified Reference Ore

bу

H.F. Steger* and W.S. Bowman**

SYNOPSIS

A 345-kg sample of a base metal ore from Mount Pleasant, New Brunswick, has been prepared as a compositional reference material to replace the similar certified ore, MP-1, of which the stock had been exhausted. MP-1a was ground to minus 74 µm and mixed in one lot. Approximately one half of this ore was bottled in 200-g units and tested for homogeneity with respect to its zinc and bismuth contents by chemical methods. The remaining material is being stored in bulk under periodic purging with nitrogen gas.

In a "free choice" analytical program, 21 laboratories contributed results for one or more of zinc, lead, copper, tin, arsenic, bismuth, indium, molybdenum, tungsten and silver in one bottle of MP-la. Based on a statistical analysis of the data, the following recommended values were assigned: Zn, 19.02%; Pb, 4.33%; Cu, 1.44%; Sn, 1.28%; As, 0.84%; In, 0.033%; Bi, 0.032%; Mo, 0.029%; and Ag, 69.7 µg/g.

^{*}Research Scientist and **Technologist, Mineral Sciences Laboratories, CANMET, Energy, Mines and Resources Canada, Ottawa.

Note: Major contributions were also made by other staff members of the Mineral Sciences Laboratories.

MP-la: Minerai de Référence

par

H.F. Steger* et W.S. Bowman**

SYNOPSIS

Un échantillon de 345-kg de minerai de métaux communs provenant de Mount Pleasant au Nouveau-Brunswick a été préparé comme matériau de référence de composition pour remplacer le minerai certifié analogue, MP-1, dont l'inventaire avait été épuisé. Le MP-la a été broyé à une granulométrie de moins 74 µm et mélangé en lot de minerai. Approximativement une moitié de ce minerai a été embouteillée en unités de 200-g et soumise à des essais d'homogénéité quant au zinc et bismuth par des méthodes chimiques. Le reste du matériau se met en réserve en gros avec une purge périodique avec du gaz d'azote.

En vertu d'un programme analytique de "libre choix", 21 laboratoires ont soumis les résultats pour un ou plusieurs des éléments suivants: zinc, plomb, cuivre, étain, arsenic, bismuth, indium, molybdène, tungstène et argent sur une bouteille du MP-la. Suite à analyse statistique des données, les valeurs recommandées suivantes ont été assignées: Zn, 19,02%; Pb, 4,33%; Cu, 1,44%; Sn, 1,28%; As, 0,84%; In, 0,033%; Bi, 0,032%; Mo, 0,029%; et Ag, 69,7 µg/g.

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Nota: D'autres membres du personnel des Laboratoires des sciences minérales ont également apporté une grande contribution à ce projet.

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INTRODUCTION

The preparation, characterization and certification of base metal ore MP-la is a further contribution of the Canadian Certified Reference Materials Project (CCRMP) in its endeavour to provide compositional reference ores, concentrates and related products typical of Canadian deposits and generally unavailable from other sources for use in analytical laboratories associated with mining, metallurgy and the earth sciences. Other certified reference materials are described in a catalogue available from CANMET, Energy, Mines and Resources, Ottawa, Canada (1).

MP-la is intended to replace MP-1, the supply of which was exhausted (2,3). MP-1 was certified in 1972 for zinc, lead, copper, tin, arsenic, bismuth, molybdenum, indium and silver. Only a provisional value for tungsten was given. MP-la was prepared with a higher tungsten content to provide a better chance for certification for this element. MP-1 had been a popular reference material because of the large number of certified elements and because of its mineralogical complexity.

An interlaboratory program was conducted to obtain results for 10 elements from 21 commercial, industrial and government laboratories using analytical methods of their choice. The results should therefore be indicative of the state-of-the-art of the analysis for these elements.

NATURE AND PREPARATION

The two raw materials which were blended to make MP-la were donated to CCRMP in October 1978 by Billitone Exploration Company Limited and are representative of the sulphide and of the wolframite-molybdenite-bismuth mineralization deposits at Mount Pleasant, New Brunswick. The materials were hand-picked at the mine site by CANMET geologists (4).

Both samples were dry-ground in March 1980 to pass a 74-µm screen. The powdered ore, consisting of 330.5 kg of the high sulphide-bearing material and of 14.5 kg of the wolframite-

molybdenite-bismuth-bearing material, was tumbled in a 570-L conical blender for 8 h. Approximately one half of the blended material was bottled in 200-g units which were heat-sealed in polyesteraluminium foil-polyethylene pouches to prevent oxidation while in storage at CANMET. The remainder of the material is being stored in bulk under periodic purging with nitrogen gas.

The analysis of 15 randomly-selected bottles of MP-la for both zinc and bismuth demonstrated the material to be sufficiently homogeneous for use as a reference material. The results of the evaluation of the homogeneity of MP-la are reported in Appendix A.

The approximate mineralogical and chemical composition and the particle size analysis are given in Tables 1 to 3.

Table 1 - Approximate mineralogical composition

Mineral	wt %
Quartz	36.5
Sphalerite	31.6
Chlorite	9
Galena	4.9
Chalcopyrite	4.1
Arsenopyrite	4.0
Topaz	4
Fluorite	2
Cassiterite	1.5
Kaolinite	1
Pyrite	0.7
Stannite	0.5
Rutile	0.5
Lepidocrocite	0.2
Molybdenite	0.04
Wolframite	0.04
Loellingite	0.04
Calcite	0.04
Bismuthinite	0.03
Bismuth	0.01
Tennantite	0.01
Monazite	0.004
Zircon	0.004

Table 2 - Approximate chemical composition

Element	wt %*
Si	19.4
Zn	19.02
S	12.8
Fe	6.2
Al	5.2
Pb	4.33
F	2.2
Ca	2.1
Cu	1.44
Sn	1.28
As	0.84
W	0.040
Мо	0.029
In	0.033
Bi	0.032
C (total)	0.03
Mg	0.02
Ag	69.7 µg/g
H ₂ O (105°)	0.01

^{*}Mean of minimum of two determinations or certified value.

Table 3 - Particle size analysis (wet screen)

Size of fraction (µm)	wt %*
bize of fraction (bin)	W C P
-104 + 74	0.2
-74 + 55	12.2
- 55 + 46	3.4
- 46 + 37	9.4
- 37	74.8

^{*}Mean of duplicate determinations

INTERLABORATORY PROGRAM FOR CERTIFICATION

The laboratories that participated in the certification program are listed in Appendix B. Each was assigned a code number which bears no relation to its alphabetical order. The results from CANMET are reported as Laboratories 30, 51, 52 and 53.

To prevent the situation wherein nearly all laboratories would submit results for elements such as zinc, lead, copper, etc., but where only a few would submit results for indium, bismuth, tungsten, etc., CCRMP divided the elements into three groups:

- (1) zinc, lead, copper, tin
- (2) arsenic, tin and
- (3) bismuth, indium, molybdenum, tungsten

and requested each participating laboratory to state which elements in each group it preferred to analyze for. An attempt was made to assign to as many laboratories as possible their two preferred elements from groups (1) and (3) and one preferred element from group (2) and still ensure a sufficient number of results for each element to permit certification by consensus.

Each laboratory was requested to contribute five replicate results for the assigned elements for one bottle of MP-la by methods of its own choice and to report the results on an "as is" basis. Some laboratories however deviated from the request for five results for an element or contributed results for more than five elements. When a laboratory submitted results by more than one method for an element, each set was considered statistically independent.

The recommended values for MP-la are presented in Table 4. Methodological and analytical information is presented in Tables 5 and 6.

STATISTICAL TREATMENT OF ANALYTICAL RESULTS

DETECTION OF OUTLIERS

Any sets of results obviously suspect for methodological reasons were rejected. Also, the sets of results whose means differed by more than twice the overall standard deviation from the initially calculated mean value were not used in subsequent computations to avoid biasing of the statistics. All results that were rejected are identified in Table 6.

ESTIMATION OF CONSENSUS VALUES AND 95% CONFIDENCE LIMITS

A one-way analysis of variance technique was used to estimate the consensus value and vari-

Table 4 - Recommended values and statistical parameters (outliers excluded)

				95%	CL	σ *
Element	No. of	No. of results	Overall mean	Low	High	
	laboratories			wt %		
Zn	15	95	19.02	18.91	19.12	0.10
Pb	16	105	4.33	4.30	4.36	0.02
Cu	18	125	1.44	1.43	1.44	0.01
Sn	10	80	1,28	1.25	1.32	0.02
As	14	84	0.84	0.82	0.86	0.01
In	6	50	0.033	0.032	0.034	8000.0
Bi	10	65	0.032	0.030	0.034	0.0008
Мо	14	80	0.029	0.028	0.030	0.0006
Ag	14	90	69.7	67.9	71.4	1.1

^{*}Average within-set standard deviation.

ance. This approach considers the results of the described certification program to be only one sampling out of a universal set of results. The analytical data were assumed to fit the model (5).

$$x_{ij} = \mu + y_i + e_{ij}$$

where $x_{ij} = the j^{th}$ result in set i,

 μ = the true consensus value,

 y_i = the discrepancy between the mean of the results in the set i (\bar{x}_i^{\bullet}) and μ ,

 e_{ij} = the discrepancy between x_{ij} and \bar{x}_{i} .

It is assumed that both y_i and e_{ij} are normally distributed with means of zero and variances of ω^2 and σ^2 , respectively. The significance of ω^2 is detected by comparing the ratio of between-set mean squares to within-set mean squares with the F statistic at the 95% confidence level and with the appropriate degrees of freedom.

The consensus value of the assumed model is estimated by the overall mean \bar{x} .. by:

$$\bar{x} \dots = \frac{k}{\sum_{i}} n_{i} x_{i,j} / \sum_{i}^{k} n_{i}$$

where n_i = the number of results in set i, and k = the number of sets.

The value of σ^2 is estimated by $s_1^{\ 2}$ which is given by

$$s_1^2 = \sum_{i=j}^{k} \sum_{j=1}^{n_i} (x_{i,j} - \bar{x}_{i,j})^2 / \sum_{j=1}^{k} n_j - k.$$

The value of ω^2 is estimated by

$$\omega^{2} = (s_{2}^{2} - s_{1}^{2}) / \frac{1}{k-1} \begin{pmatrix} k & k & 2/k \\ \sum n_{i} - \sum n_{i}^{2}/\sum n_{i} \end{pmatrix}$$

where

$$s_2^2 = \sum_{i=1}^{k} n_i (\bar{x}_i - \bar{x}..)^2 / k-1.$$

The variance of the overall mean is given by

$$V[\vec{x}..] = \begin{pmatrix} k & n_1^2 / (\sum_i n_i)^2 \end{pmatrix} \omega^2 + \begin{pmatrix} k & n_i \\ 1 / \sum_i n_i \end{pmatrix} \sigma^2$$

and the 95% confidence limits for \bar{x} .. are

$$\bar{x}.. \pm t_{0.975, (k-1)} \sqrt{\bar{x}..}$$

To avoid giving an unduly heavy weighting to the contribution for zinc and bismuth from CANMET, only five results for each were selected at random out of the 45 available from the evaluation of the homogeneity of MP-la.

Table 5a - Summary of analytical methods for zinc (outliers excluded)

Method	Decomposition, separation, etc.	Lab No∙	n	x̄ (wt ¾)
Atomic absorption	HNO ₃ + one or more of HCl, HF, HClO ₄ , H ₂ SO ₄ ; 1-3M HCl	5, 11a, 12a, 19, 26a	25	18.98
	HF to dryness; taken up in HNO_3 + $HC1$ + $HC1O_4$	15	5	19.16
	HNO3; 5% HNO3 in final solution	7	5	19.08
	HClO ₄ ; 5% HCl in final solution	18, 28	10	19.31
Titrimetry	$^{\mathrm{HNO}}_3$ + one or more of HCl, HF, HClO $_{\!\mu}$; dissolved in	CANMET (2), 11b	30	19.05
	HCl + ${\rm H_2SO_{ll}}$; filtered off ${\rm PbSO_{ll}}$; separated Zn as thiocyanate by MIBK extraction; titrated with EDTA	12b, 14b, 23		
	$\mathrm{HNO_3}$ + HCl + HF + $\mathrm{H_2SO_4}$; separated $\mathrm{Re_2O_3}$; reduced with lead shot; titrated with ferrocyanide using ammonium molybdate as indicator	9, 26b	10	19.06
ICP-emission spectrometry	$\mathrm{HNO_3}$ + HF + $\mathrm{HClO_4}$; residue fused with $\mathrm{Na_2O_2}$; dissolved in 50% HCl	14a	5	18.56
X-ray fluorescence	Fused with $K_2S_2O_7$; ground and pressed into pellet	4	5	18.6

Table 5b - Summary of analytical methods for lead (outliers excluded)

				x
Method	Decomposition, separation, etc.	Lab No.	n	(wt %)
Atomic absorption	HNO_3 + one or more of HCl, HBr, HF, $HClO_{ij}$, H_2SO_{ij} ;	11, 12a, 15, 17, 19	35	4.30
	final solution was dilute $ ext{HNO}_3$ or $ ext{HCl}$	24,26a		
	${\rm HNO}_{3}$ + HCl + HF + ${\rm HClO}_{ij}$; tin and arsenic volatilized	CANMET (2)	10	4.30
	by HBr; PbCO ₃ co-precipitated with hydrous iron oxide			
	by (NH ₄) ₂ CO ₃ + NH ₄ OH; PbCO ₃ filtered and dissolved in			
	dilute HNO ₃			
	HNO ₃ + HF; residue fused with Na ₂ O ₂ + NaOH; final	2	5	4.41
	solution was 5% HCl	-	,	
	UNO a maridua formal with Li Do a filmal analytica mar	12	_	h 00
	HNO_3 ; residue fused with LiBO_2 ; final solution was 5% HNO_3	13	5	4.29
	J			
	Fused with Na ₂ 0 ₂ ; taken up in 5% HCl	3	5	4.35
	HNO3; final solution was 5% HNO3	7	5	4.33
	HClO ₄ ; final solution was 5% HCl	18, 28	10	4.39
Titrimetry	HNO_3 + $HC1$ + HF + Br_2 + H_2SO_4 ; $PbSO_4$ filtered and	12b, 14b	10	4.32
	dissolved in acetate medium; lead precipitated as			
	chromate; filtered and dissolved in dilute HCl; KI			
	added and I_2 titrated with $Na_2S_2O_3$			
	HNO_3 + HCl + HF + $HClO_{\mu}$; lead precipitated as $PbSO_{\mu}$;	CANMET	5	4.50
	filtered and dissolved in HCl + NaCl and complexed			
	with hexamethylene tetramine and titrated with EDTA			
	HNO_3 + HCl + HF + H_2SO_{ij} ; PbSO_{ij} filtered and	26b	5	4.31
	dissolved in ammonium acetate; titrated with ammonium			
	molybdate using tannic acid as indicator			
ICP-emission	$HNO_3 + HF + HClO_4$; residue fused with Na_2O_2 ; final	14a	5	4.37
spectrometry	solution was 50% HCl			-
X-ray fluorescence	Fused with K ₂ S ₂ O ₇ ; ground and pelletized	4	5	4.25

Table 5c - Summary of analytical methods for copper (outliers excluded)

				- x
Method	Decomposition, separation, etc.	Lab No.	n	(wt %)
Titrimetry	HNO_3 + $HC1$ + HF + Br_2 + H_2SO_{ij} ; $PbSO_{ij}$ filtered and	3, 12a, 13b,	10	4.32
	was dilute HCl or HNO3	15, 16, 19,		
	-	23, 24, 26a	,	
	${\rm HNO_3}$ + HCl + HF + HClO $_{\rm H}$; HBr to volatilize tin and arsenic, residue fused with ${\rm Na_2CO_3}$; final solution was dilute HCl	CANMET (2)	10	1.44
	HNO3; residue fused with LiBO2	13a	. 5	1.43
	HNO ₃ + HBr	lla	5	1.46
	HNO_3 + HF; residue fused with NaOH + Na_2O_2	2	5	1.42
	HC1 + HF + KNO ₃	20	5	1.44
	HNO3; final solution was 5% KNO3	7	5	1.45
	HClO ₄ ; final solution was 5% HCl	28	5	1.46
Titrimetry	${\rm HNO_3}$ + HCl + HF + ${\rm H_2SO_4}$ and ${\rm Br_2}$; fluoride, acetate and iodide added; titrated with ${\rm Na_2S_2O_3}$	12b , 26b	10	1.44
	${ m HClO_4}$; copper precipitated with ${ m Na_2S_2O_3}$; CuS digested and taken to dryness with ${ m HNO_3}$ + ${ m HClO_4}$; taken up with HCl; acetate added and titrated with ${ m Na_2S_2O_3}$	9, 11b	· 10	1.43
Colorimetry	${\rm HNO_3}$ + HCl + HF + ${\rm H_2SO_4}$ to dryness; taken up in dilute HCl; copper determined as cuproine complex in n-amyl alcohol	CANMET	5	1.44
Electrolysis	$\mathrm{HNO_3}$ + $\mathrm{HC1}$ + $\mathrm{Br_2}$ + $\mathrm{H_2SO_4}$; gravimetric electrolysis at 300 mV from tartrate and hydrazine solution	17	5	1.46
	HClO_{4} ; gravimetric electrolysis from dilute HNO_{3} + $\mathrm{H_{2}^{SO}}_{4}$	28	5	1.46
X-ray fluorescence	Fused with K ₂ S ₂ O ₇ ; ground and pelletized	4	5	1.30

Table 5d - Summary of analytical methods for tin (outliers excluded)

				x
Method	Decomposition, separation etc.	Lab No.	n	(wt %)
Atomic absorption	Fused with $Na_2O_2 + Na_2CO_3$; taken up in HCl	12a, 17	10	1.36
	Fused with Na ₂ O ₂ + Na ₂ CO ₃ ; iron and tin separated	CANMET (2),	21	1.28
	as hydrous oxides; dissolved in dilute HCl	206, 26		
	Fused with Na ₂ O ₂ + Na ₂ CO ₃ ; taken up in dilute H ₂ SO ₄ ;	CANMET	5	1.27
	SiF ₄ volatilized; tin extracted from 3M H ₂ SO ₄ - 1.5M KI			
	into toluene; back-extracted with 50% HNO 3 - 16% HCl;			
	determined in 10% HCl - 0.5% tartaric acid			
	HNO ₃ + HF; residue fused with NaOH + Na ₂ O ₂ ; determined in 5% HCl	2	5	1.28
	-1. 3, 1.0 <u>-</u>			
Titrimetry	Fused with Na ₂ O ₂ + Na ₂ CO ₃ ; iron and tin hydrous oxides	CANMET	5	1.25
	separated; reduced with iron powder and titrated with			
	potassium iodate			
	Fused with Na ₂ O ₂ ; taken up in dilute HCl; reduced with	14a	5	1.43
	aluminum powder and titrated with iodate			
X-ray fluorescence	l g sample + 9 g of 1:1 KCl:SiO ₂ ; pelletized	7	5	1.18
	No dobaile	5 00.	3.0	1 00
	No details	5, 20a	10	1.20
	Binder + internal standard pressed into pellets	4	4	1.24
ICP-emission	$\mathrm{HNO_3}$ + HF + $\mathrm{HClO_4}$; residue fused with $\mathrm{Na_2O_2}$ and leached	146	5	1.28
spectrometry	with 50% HCl			
Colorimetry	Fused with $Na_2O_2 + Na_2CO_3$; taken up in dilute H_2SO_4 ;	1 2b	5	1.35
	SnI ₁₁ extracted and stripped with 2% NaOH; tin(IV) -			
	gallein complex in n-amyl alcohol			

Table 5e - Summary of analytical results for arsenic (outliers excluded)

				x
Method	Decomposition, separation, etc.	Lab No.	n	(wt %)
Atomic absorption	$HNO_3 + H_2SO_{ll}$; taken up in 25% HCl	12a	5	0.88
·	Fused with KOH-MgO; taken up in 10% HCl; AsH $_3$ generated with NaBH $_4$	19	5	ა.88
	Br ₂ + HNO ₃	16	5	0.894
Titrimetry	Acid digestion; AsCl distillation; titrated with iodine	9, 26	10	0.806
	${\rm HNO_3} + {\rm H_2SO_4} + {\rm Br_2}; {\rm AsBr_3}$ distilled; titrated with iodine	28	5	0.83
	HNO3; KBr + hydrazine sulphate to dryness; taken up in HCl; AsCl3 distilled and titrated with KBrO3	12b	. 5	0.83
	${\rm HNO}_3$ + HCl; NaBr and hydrazine sulphate added; ${\rm AsBr}_3$ distilled and titrated with ${\rm KBrO}_3$ using Eriochrome SE as indicator	13 .	. 5	0.804
	Fused with ${\rm K_2S_2O_7}$ + ${\rm H_2SO_4}$; AsCl $_3$ distilled and titrated with ${\rm KBrO_3}$	CANMET	5	0.83
Colorimetry	${\rm HNO_3}$ + ${\rm HCl}$ + ${\rm H_2SO_4}$ + ${\rm Br_2}$; As(V) co-precipitated with hydrous iron oxide; dissolved in 11M HCl and reduced to As(III) with Fe(II); arsenic extracted as xanthate into chloroform; oxidized by ${\rm Br_2}$ and stripped into water; molybdenum blue complex	CANMET	5	0.775
	HNO_3 + $\mathrm{H}_2\mathrm{SO}_4$; arsenic extracted into chloroform - butanol as molybdenum blue complex	CANMET	5	0.85

Table 5e (cont'd)

				x
Method	Decomposition, separation, etc.	Lab No.	n	(wt %)
	$HNO_3 + HF + NaClO_3 + H_2SO_4$; arsenic extracted as iodide;	2	5	0.865
	determined as molybdenum blue complex			
	$\mathrm{HNO_3} + \mathrm{H_2SO_4}$; distilled as $\mathrm{AsCl_3}$; molybdenum blue complex	17	5	0.80
	${ m NaOH-Na}_2{ m O}_2$ fusion; AsH $_3$ generation; colour development using silver diethyldithiocarbamate in pyridine	24	5	0.90
ICP-AES	${\rm HNO_3}$ + HCl + HClO $_{\rm 4}$; taken up in 50% HCl; Te and HPO $_{\rm 2}$ added to precipitate arsenic; dissolved in HNO $_{\rm 3}$ + HClO $_{\rm 4}$	15	5	0.839
X-ray fluorescence	Sample + binder + internal standard are pelletized	Ц	Ц	0.83
Neutron activation analysis	Counting X-ray at 559 KeV	23	5	0.836

Table 5f - Summary of analytical methods for indium (outliers excluded)

				x () a l
Method	Decomposition, separation, etc.	Lab No.	n	(wt %)
Atomic absorption	$^{\mathrm{HNO}}_{3}$ + HCl + HF + HClO $_{\mathrm{4}}$; dissolved in 5% HCl	CANMET (2)	10	0.0320
	HNO_3 + HCl + HF + HClO $_{\mu}$; tin and arsenic volatilized	CANMET (2)	9	0.0337
1	with HBr; indium extracted into n-butyl acetate from			
	5 M HBr and back-extracted into H ₂ O; final solution			
	was 2-5% HCl			
	HNO ₃ + HF; residue fused with NaOH + Na ₂ O ₂ ; final	2	5	0.0343
	solution was 5% HCl			
	Indium concentrated by ion-exchange procedure - no other	14	5	0.0339
	details		-	0.000
X-ray fluorescence	No details	5, 24	8	0.0319
ray rraor obcence	NO GODALLO	J, 27		0.00127
	Loose powder technique	4	4	0.030
			*	
Polarography	See above for acid decomposition and indium separation	CANMET	4	0.0330
	by extraction from 5 M HBr; final solution was 5% HCl			
Emission	Sample mixed with 4 times its weight of 17.5:2.5	CANMET	5	0.0363
spectrometry	graphite:Na ₂ CO ₃			

Table 5g - Summary of analytical methods for bismuth (outliers excluded)

				\bar{x}
Method	Decomposition, separation, etc.	Lab No.	n	(wt %)
Atomic absorption	$HNO_3 + HF + H_2SO_4 + HBr$; taken up in dilute HCl;	CANMET (4)	20	0.0323
	bismuth and iron hydrous oxides co-precipitated;			
	filtered and dissolved in 20% HCl			
	HNO_3 + HF; residue fused with Na_2O_2 + $NaOH$; bismuth +	2	5	0.029
	iron + lanthanium hydrous oxides co-precipitated,			
	filtered and dissolved in HCl			
	${\rm HNO_3}$ + HCl + ${\rm (NH_4)_2SO_4}$ + ${\rm H_2SO_4}$; dissolved in HCl; boiled with ${\rm MnSO_4}$ and ${\rm KMnO_4}$ to precipitate bismuth hydrous oxide;	15	5	0.0267
	filtered and dissolved in 35% HC1			
	HNO ₃ + HCl; taken up in 5% HCl	17	5	0.033
	${\rm HNO_3}$ + ${\rm HF}$ + ${\rm HClO_4}$; taken up in 20% HCl; ${\rm BiH_3}$ generation	19	5	0.0370
	HNO3; no other details	20	5	0.0315
Atomic absorption- graphite furnace	${\rm HNO_3}$ + HF + NaClO $_3$ + HClO $_4$; bismuth precipitated with NH $_4$ OH + NH $_4$ Cl; dissolved in HCl + HNO $_3$	16	5	0.036
Emission spectrometry	Sample mixed with equal weight of KCl	7	5	0.0338
X-ray fluorescence	No details	5	5	0.0301
	Loose powder technique	4	5	0.026

Table 5h - Summary of analytical methods for molybdenum (outliers excluded)

		•		z
Method	Decomposition, separation, etc.	Lab No.	n	(wt %)
Atomic absorption	${\rm HNO_3}$ + one or more of HCl, HF, HClO $_{\rm H}$, HBr, H $_{\rm 2}{\rm SO_4}$; final solution was 5-10% HCl containing 1000-2000 mg/L AlCl $_{\rm 3}$	CANMET, 11, 17, 19, 23, 24, 26	35	0.029
		24, 20		
	${\rm HNO_3}$ + HCl + HF + Br ₂ + ${\rm H_2SO_4}$; tin and arsenic volatilized with HBr; dissolved in dilute tartaric	CANMET	5	0.0288
	acid; molybdenum and tungsten extracted as			
	$\alpha\text{benzoinoxine}$ complex into chloroform from 1.8 M			
	HCl; back-extracted with $\rm HNO_3$ + $\rm HClO_4$ + $\rm H_2SO_4$; taken to dryness and dissolved in 15% HCl + 1000 mg/L AlCl 3			
	$^{\rm HNO}_3$ + HF + HC1 + HC10 $_{\rm H}$; final solution was 10% HN0 $_3$ + 129 g/L KN0 $_3$	15	5	0.0282
	$\mathrm{HNO_3};$ residue treated with HF to remove $\mathrm{SiO_2};$ fused with $\mathrm{LiBO_2}$	13	5	0.0302
	$\mathrm{HNO_3}$ + HCl + HF + $\mathrm{HClO_4}$; final solution was 20% HCl + $\mathrm{Na_2^{SO_4}}$	3	5	0.0312
	Fused with NaOH + Na ₂ 0 ₂ ; taken up in dilute HNO ₃	9	5	0.032
Colorimetry	${ m HN0_3}$ + HCl + HF + HClO $_{\rm H}$; residue fused with ${ m Na_2CO_3}$; iron + copper removed by double precipitation with NaOH; molybdenum thiocyanate complex	CANMET	5	0.027
Emission spectrometry	Sample mixed with equal weight of carbon	7	5	0.0318
X-ray fluorescence	Loose powder technique	4	5	0.024
	Binder + internal standard pressed into pellets	5	5	0.029

Table 5i - Summary of analytical methods for silver (outliers excluded)

				z
Method	Decomposition, separation, etc.	Lab No.	n	(µg/g)
Fire assay- gravimetry		26b	5	62.7
Fire assay - atomic absorption	Lead button collection; scorification; parted in HNO3; complexation with diethylenetriamine	CANMET (2, 3)	15	68.2
Atomic absorption	$\text{HNO}_3^{}$ + one or more of HCl, HF, HClO $_{\!$	5, 11, 16, 17 20, 26a, 28	35	70.4
	${\rm HNO}_3$ + HCl + HF + HClO $_{\rm H}$; complexation with diethylenetriamine; 10% HCl	CANMET	5	68.3
	${\rm HNO}_3$ + ${\rm HF}$ + ${\rm HClO}_{\rm p};$ dissolved in ${\rm HNO}_3$ containing 10 g/L tartaric acid	15	5	68.5
	$HNO_3 + HF$; taken up in 3% HNO_3	19	5	65.7
	HClO ₄ ; taken up in 25% HCl	18	5	74.6
	${\rm HNO}_3$ + ${\rm Br}_2$ + ${\rm HF}$ + ${\rm H}_2{\rm SO}_4$; evaporated to fumes of ${\rm SO}_3$; silver extracted as bromide complex into chloroform + tribenzylamine from 2 M ${\rm H}_2{\rm SO}_4$ + KBr; stripped with HBr and taken to dryness; taken up in 10% HCl - 1% diethylenetriamine	CANMET	5	67.0
Emission spectrometry	Sample mixed with equal weight of carbon	7	5	76.0
DCP-Emission spectrometry	но 3	ц	5	73.6

Table 5j - Summary of analytical methods for tungsten

			,	z
Method	Decomposition, separation, etc.	Lab No.	n	(wt/%)
Atomic absorption	HNO ₃ + HF; dissolved in 0.5 M KOH	13	5	0.0382
Colorimetry	Fused with ${\rm NaHSO}_{\rm ll}$; dissolved in tartaric acid; tungsten reduced with ${\rm SnCl}_{\rm 2}$; determined as W(V)-thiocyanate-diantipyrylmethane complex in chloroform	CANMET	5	0.0440
	HCl + HF + H ₃ PO ₄ ; iron removed as chloride with isopropylether; tungsten reduced with SnCl ₂ ; determined as thiocyanate complex	CANMET	5	0.032
	${\rm HNO_3}$ + HCl + HF + ${\rm H_2SO_4}$; silicon volatilized as ${\rm SiF_4}$; determined as thiocyanate	9	5	0.042
	${\rm HNO_3}$ + HCl + ${\rm H_3PO_4}$ + ${\rm H_2SO_4}$ + HClO $_4$; residue fused with ${\rm Na_2O_2}$ + ${\rm Na_2CO_3}$; tungsten reduced with SnCl $_2$ and determined as toluene 3,4-dithiol complex in n-butyl acetate	14	5	0.049
	${ m Na_2^{0}}_2$ + ${ m Na_2^{C0}}_3$ fusion; tungsten reduced with ${ m SnCl}_2$ and determined as thiocyanate	15, 26	10	0.035
	${ m Na_2O_2}$ fusion; arsenic removed; tungsten determined as thiocyanate	20	5	0.0448
	HNO_3 + HCl + HF + HClO _{μ} ; tungsten extracted as tetraphenylarsonium chloride - Anal Chem Acta <u>30</u> , 501 (1969)	16	5	0.036
X-ray fluorescence	No details	5	5	0.0459
Emission spectrometry	Sample mixed with equal weight of carbon	7	5	0.0294
Neutron activation	Gamma ray count	4	['] 5	0.047

Table 6a - Laboratory results, means and standard deviations for zinc

						ME AN	S.D.
140 6 / VDC3	18.2	18.8	18.6	18.7	18.7	18.6000	. 2345
LA8- 4 (XRF)	19.09	18.72	19.50	19.23	18.95	19.0980	.2932
LA8- 5 (AA)		19.16	19.05	19.14	19.02	19.0760	.0695
LA8- 7 (AA)	19.01	19.10	19.33	19.33	19.28	19.3020	.0311
LA8- 9 (TITR)	19.26	18.9	18.9	19.0	18.9	18.9400	.0548
LA8-11 (AA)	19.0	19.16	19.12	19.09	19.20	19.1460	.0422
LA8-11 (TITR)	19.16		19.24	19.27	19.06	19.0940	.1667
LA8-12 (AA)	18.86	19.04	18.97	19.11	19.13	19.0840	.0882
LA8-12 (TITR)	19.19	19.02	18.99	18.94	18.97	18.9880	.0370
LA8-14 (TITR)	19.04	19.00		18.7	18.6	18.5600	.2074
LA8-14 (ICP)	18.4	18.3	18.8	19.25	19.15	19.1600	.0894
LAB-15 (AA)	19.05	19.25	19.10		19.40	19.4200	.1037
LA8-18 (AA)	19.25	19.50	19.45	19.50		18.9200	.1304
LA8-19 (AA)	19.1	18.8	19.0	18.9	18.8	19.1600	.0187
LA8-23 (TITR)	19.18	19.15	19.18	19.15	19.14	18.8400	.0894
LA8-26 (AA)	18.80	18.8	18.8	18.8	19.0		.0671
LAB-26 (TITR)	18.74	18.79	18.89	18.79	18.89	18.8200	
LA8-28 (AA)	19.2	19.1	19.3	19.2	19.2	19.2000	.0707
LA8-52 (TITR)	19.01	19.01	18.99	18.98	18.98	18.9940	.0152
LA8-53 (TITR)	18.88	18.86	18.92	18.97	18.87	18.9000	.0453

Table 6b - Laboratory results, means and standard deviations for lead

						ME AN	S.D.
LA8- 2 (AA)	4.43	4.42	4.41	4.42	4.38	4.4120	.0192
LAB- 3 (AA)	4.36	4.35	4.35	4.32	4.36	4.3480	. 0164
LAB- 4 (XRF)	4.26	4.30	4.22	4.24	4.24	4.2520	.0303
LA8- 5 (XRF)*	3.83	3.57	3.83	3.57	3.83	3.7260	.1424
LAS- 7 (AA)	4.32	4.35	4.36	4.30	4.34	4.3340	.0241
LA8-11 (AA)	4.21	4.22	4.22	4.19	4.20	4.2080	.0130
LA8-12 (AA)	4.27	4.28	4.28	4.26	4.29	4.2760	.0114
LA8-12 (TITR)	4.34	4.37	4.42	4.41	4.39	4.3860	.0321
LA8-13 (AA)	4.27	4.27	4.30	4.30	4.30	4.2880	.0164
LA8-14 (TITR)	4.24	4.19	4.27	4.22	4.30	4.2440	.0428
LA8-14 (ICP)	4.38	4.38	4.41	4.36	4.34	4.3740	.0261
LA8-15 (AA)	4.416	4.401	4.427	4.413	4.404	4.4122	.0103
LAB-17 (AA)	4.28	4.28	4.29	4.29	4.28	4.2840	.0055
LA8-18 (AA)	4.42	4.40	4.45	4.41	4.39	4.4140	.0230
LA8-19 (AA)	4.30	4.30	4.30	4.30	4.30	4.3000	.0000
LA8-24 (AA)	4.32	4.28	4.28	4.24	4.28	4.2800	.0283
LA8-26 (AA)	4.40	4.38	4.38	4.35	4.35	4.3720	.0217
LA8-26 (TITR)	4.33	4.38	4.28	4.28	4.28	4.3100	.0447
	4.35	4.36	4.35	4.36	4.36	4.3560	•0055
LA8-28 (AA) LA8-52 (AA)	4.283	4.300	4.263	4.257	4.264	4.2734	.0178
	4.35	4.31	4.37	4.32	4.33	4.3360	.0241
LA8-53 (AA) LA8-53 (TITR)	4.52	4.54	4.53	4.42	4.50	4.5020	.0482

^{*}Outlying set.

Table 6c - Laboratory results, means and standard deviations for copper

						ME AN	S.D.
LAB- 2 (AA)	1.44	1.42	1.42	1.42	1.42	1.4240	.0089
LAB- 3 (AA)	1.43	1.43	1.44	1.43	1.44	1.4340	•0055
LAB- 4 (XRF)	1.42	1.36	1.40	1.36	1.37	1.3820	.0268
LAB- 5 (AA)*	1.65	1.65	1.60	1.65	1.65	1.6400	.0224
LAB- 7 (AA)	1.42	1.46	1.48	1.44	1.47	1.4540	.0241
LAB- 9 (TITR)	1.42	1.43	1.42	1.43	1.42	1.4240	• 0055
LAB-11 (TITR)	1.43	1.44	1.43	1.43	1.43	1.4320	.0045
LAB-li (AA)	1.45	1.46	1.46	1.46	1.45	1.4560	•0055
LAB-12 (AA)	1.44	1.46	1.47	1.47	1.48	1.4640	•0152
LAB-12 (TITR)	1.43	1.45	1.45	1.47	1.45	1.4500	•0141
LAB-13 (AA)	1.44	1.43	1.43	1.43	1.44	1.4340	.0055
LAB-13 (AA)	1.43	1.42	1.40	1.40	1.41	1.4120	•0130
LAB-15 (AA)	1.458	1.462	1.461	1.460	1.433	1.4548	.0123
LAB-16 (AA)	1.415	1.403	1.406	1.408	1.413	1.4090	.0049
LAB-17 (ELECTR)	1.46	1.47	1.44	1.49	1.44	1.4600	.0212
LAB-18 (AA)*	1.35	1.38	1.35	1.36	1.35	1.3580	.0130
LAB-19 (AA)	1.45	1.44	1.44	1.45	1.44	1.4440	.0055
LAB-20 (AA)	1.444	1.441	1.443	1.443	1.446	1.4434	.0018
LAB-23 (AA)	1.431	1.439	1.442	1.440	1.438	1.4380	.0042
LAB-24 (AA)	1.42	1.42	1.40	1.42	1.42	1.4160	.0089
LAB-26 (AA)	1.44	1.43	1.43	1.43	1.44	1.4340	.0055
LAB-26 (TITR)	1.42	1.42	1.42	1.45	1.42	1.4260	.0134
LAB-28 (ELECTR)	1.45	1.46	1.45	1.46	1.46	1.4560	.0055
LAB-28 (AA)	1.45	1.46	1.44	1.45	1.45	1.4500	.0071
LAB-52 (AA)	1.431	1.437	1.424	1.421	1.426	1.4278	.0063
LAB-53 (AA)	1.47	1.47	1.43	1.45	1.44	1.4520	.0179
LAB-53 (COLOR)	1.44	1.45	1.42	1.45	1.44	1.4400	.0122

^{*}Outlying set.

Table 6d - Laboratory results, means and standard deviations for tin

						ME AN	S.D.
LAB- 2 (AA)	1.27	1.26	1.30	1.27	1.28	1.2760	.0152
LAB- 4 (XRF)	1.24	1.24	1.26	1.22		1.2400	.0163
LAB- 5 (XRF)	1.16	1.15	1.14	1.18	1.14	1.1540	.0167
LAB- 7 (XRF)	1.13	1.17	1.21	1.20	1.18	1.1780	.0311
LAB-12 (AA)	1.34	1.40	1.44	1.33	1.39	1.3800	•0453
LAB-12 (COLOR)	1.30	1.37	1.37	1.40	1.33	1.3540	•0391
LAB-14 (TITR)	1.46	1.33	1.47	1.46	1.44	1.4320	.0581
LAB-14 (ICP)	1.27	1.31	1.29	1.26	1.29	1.2840	.0195
LAB-17 (AA)	1.37	1.34	1.31	1.33	1.34	1.3380	.0217
LAB-20 (XRF)	1.253	1.253	1.246	1.261	1.243	1.2512	.0070
LAB-20 (AA)	1.259	1.257	1.262	1.255	1.254	1.2574	.0032
LAB-26 (AA)	1.28	1.30	1.30	1.27	1.29	1.2880	
LAB-30 (AA)	1.33	1.32	1.30	1.33	1.28	1.3117	.0130
	1.31	,		1033	1.4.20	1.3111.	.0194
LAB-52 (AA)	1.282	1.274	1.277	1.251	1.262	1 5/02	0124
LAB-52 (TITR)	1.264	1.246	1.255	1.246	1.255	1.2692	•0126
LAB-53 (AA)	1.289	1.279	1.222	1.269	1.238	1.2532	• 0075
LAB-53 (TITR)*	1.02	1.10	1.06	1.02	1.11	1.2694 1.0620	•0277 •0427

^{*}Outlying set.

Table 6e - Laboratory results, means and standard deviations for arsenic

						MEAN	S.D.
LAB- 2 (COLOR)	.863	.860	. 869	. 869	•865	.8652	.0039
LAB- 4 (XRF)	•82	. 82	a 84	.82		.8250	.0100
LAB- 5 (XRF)*	1.16	1.16	1.16	1.16	1.17	1.1620	.0045
LAB- 9 (TITR)	0.804	0.804	0.798	0.804	0.798	.8016	.0033
LAB-12 (AA)	0.92	0.90	0.86	0.86	0.86	.8800	.0283
LAB-12 (TITR)	0.83	0.83	0.82	0.83	0.83	.8280	.0045
LAB-13 (TITR)	0.80	0.80	0.81	0.80	0.81	.8040	•0055
LAB-15 (ICP)	0.850	0.828	0.838	0.842	0.836	.8388	.0081
LAB-16 (AA)	0.906	0.897	0.899	0.884	0.882	.8936	.0103
LAB-17 (COLOR)	0.82	0.80	0.79	0.78	0.81	.8000	.0158
LAB-17 (COLUR)	0.87	0.86	0.92	0.88	0.88	.8820	.0228
	0.840	0.833	0.837	0.835	0.837	.8364	.0026
LAB-23 (NAA)	0.94	0.88	0.88	0.88	0.90	.8960	.0261
LAB-24 (COLOR)	0.815	0.815	0.800	0.808	0.815	.8106	.0067
LAB-26 (TITR)	0.82	0.82	0.82	0.84	0.83	. 8 260	.0089
LAB-28 (TITR)	0.02	0.775	0.776	0.775	0.772	.7754	.0025
LAB-52 (COLOR)		0.81	0.84	0.83	0.80	.8260	.0207
LAB-53 (TITR) LAB-53 (COLOR)	0.85 0.86	0.89	0.85	0.85	0.79	.8480	• 0363

^{*}Outlying set.

Table 6f - Laboratory results, means and standard deviations for indium

						MEAN	S.D.
LAB- 2 (AA) LAB- 4 (XRF) LAB- 5 (XRF) LAB- 7 (ES)* LAB-14 (AA) LAB-24 (XRF) LAB-30 (AA) LAB-30 (AA) LAB-30 (ES) LAB-30 (ES) LAB-52 (AA) LAB-53 (AA)	.0345 .030 0.0303 .0390 .0340 0.035 0.0313 .0335 0.0326 .0350 0.0344	.0340 .029 0.0299 .0400 .0336 0.036 0.0319 .0333 0.0329 .0380 0.0337	.0340 .030 0.0297 .0385 .0340 0.037 0.0319 .0337 0.0340 .0365 0.0341	.0340 .031 0.0282 .0405 .0336 0.0330 .0333 0.0324 .0345 0.0335 0.031	.0350 0.0296 .0390 .0344 0.0324 .0375 0.0336 0.033	.0343 .0300 .0295 .0394 .0339 .0360 .0321 .0335 .0330 .0363	.0004 .0008 .0008 .0003 .0010 .0006 .0002 .0007 .0015 .0004

^{*}Outlying set.

Table 6g - Laboratory results, means and standard deviations for bismuth

						ME AN	S.D.
LAB- 2 (AA)	• 028	•028	•030	•030	•029	• 0290	.0010
LAB- 4 (XRF)	. 028	• 026	•027	.024	.027	.0264	.0015
LAB- 5 (XRF)	0.0296	0.0303	0.0295	0.0300	0.0309	•0301	.0006
LAB- 7 (ES)	.0340	•0330	•0335	.0340	. 0345	•0338	.0006
LAB-15 (AA)	0.0244	0.0274	0.0278	0.0254	0.0287	.0267	.0018
LAB-16 (AA)	0.035	0.036	0.036	0.036	0.037	.0360	.0007
LAB-17 (AA)	0.032	0.033	0.034	0.033	0.032	.0328	.0008
LAB-19 (AA)	•0384	.0372	• 0366	.0357	•0369	.0370	.0010
LAB-20 (AA)	0.0314	0.0313	0.0314	0.0318	0.0316	.0315	.0002
LAB-30 (AA)	• 0335	•0325	.0318	.0329	•0333	.0328	.0007
LAB-52 (AA)	•0318	.0324	.0321	.0320	.0320	.0321	.0002
LAB-52 (AA)	.0314	• 03 20	.0319	• 0327	.0316	.0319	.0005
LAB-53 (AA)	.031	.032	•032	.034	.033	.0324	.0011

Table 6h - Laboratory results, means and standard deviations for molybdenum

•						MEAN	S.D.
LAB- 3 (AA)	0.031	0 021	0.022	0.021	0.001		
LAB- 4 (XRF)		0.031	0.032	0.031	0.031	•0312	• 0004
	•024	.025	.024	. 023	•024	•0240	• 0007
LAB- 5 (XRF)	0.0294	0.0293	0.0283	0.0284	0.0289	•0289	• 0005
LAB- 7 (ES)	•0316	•0318	•0316	•0320	•0318	•0318	.0002
LAB- 9 (AA)	0.032	0.031	0.033	0.033	0.033	.0324	.0009
LAB-11 (AA)	0.032	0.032	0.031	0.032	0.030	.0314	.0009
LAB-13 (AA)	0.0301	0.0302	0.0301	0.0302	0.0304	•0302	.0001
LAB-15 (AA)	0.0282	0.0276	0.0285	0.0281	0.0285	.0282	.0004
LAB-17 (AA)	0.029	0.028	0.029	0.029	0.029	•0288	.0004
LAB-18 (AA)*	.038	.044	.035	.040	.033	.0380	•0043
LAB-19 (AA)	• 0285	.0280	•0279	.0277	.0285	.0281	.0004
LAB-23 (AA)	0.0278	0.0277	0.0285	0.0286	0.0298	.0285	• 0008
LAB-24 (AA)	0.028	0.028	0.026	0.028	0.028	.0276	.0009
LAB-26 (AA)	0.032	0.032	0.031	0.032	0.032	.0318	•0004
LAB-52 (AA)	0.0298	0.0304	0.0297	0.0301	0.032	•0299	
LAB-53 (AA)	0.030	0.028	0.030	0.029			.0003
LAB-53 (COLOR)	0.027	0.027	0.037	0.027	0.027 0.026	•0288 •0268	.0013

^{*}Outlying set.

Table 6i - Laboratory results, means and standard deviations for silver

						MEAN S.	. D .
						· · · · · · · · · · · · · · · · · ·	
LAB- 3 (FA-AA)	72.7	71.3	71.7	72.7	71.0	71.8800 .78	
LAB- 4 (ES)	69.	72.	75.	75.	77.	73.6000 3.13	
LAB- 5 (AA)	66.	66.	66.	69.	69•	67.2000 1.64	
LAB- 7 (ES)	76.	78.	76.	74.	76.	76.0000 1.41	
LAB-11 (AA)	69.2	67.6	71.8	69.7	68.6	69.3800 1.56	
LAB-15 (AA)	69.0	68.5	68.0	68.5	68.5	68.5000 .35	
LAB-16 (AA)	71.5	71.5	71.4	71.6	71.5	71.5000 .07	707
LAB-17 (AA)	69.	69.	68.	69.	68.	68.6000 .54	477
LAB-18 (AA)	75.	76.	74.	73.	75.	74.6000 1.14	402
LAB-19 (AA)	66.	65.	65.5	66.	66.	65.7000 .44	472
LAB-20 (AA)	72.95	73.15	72.80	73.40	73.45	73.1500 .28	30 t
LAB-26 (AA)	72.0	72.0	71.0	71.0	72.0	71.6000 .54	477
LAB-26 (FA-G)	63.8	62.7	67.2	59.7	60.3	62.7400 3.00	380
LAB-28 (AA)	72.1	71.3	71.2	72.1	71.3	71.6000 -45	583
LAB-30 (FA-AA)	64.1	67.0	66.3	64.3	67.1	65.7600 1.45	586
LAB-30 (AA)	69.2	68.1	67.6	69.1	67.5	68.3000 .80	093
LAB-50 (AA7	66.9	56.5	67.9	67.3	66.5	67.0200 .59	933
LAB-52 (AA)	66.7	67.0	66.7	69.6	65.0	67.0000 1.65	538

Table 6j - Laboratory results, means and standard deviations for tungsten

						ME AN	S.D.
LAB- 4 (NAA) LAB- 5 (XRF) LAB- 7 (ES) LAB- 9 (COLOR) LAB-13 (AA) LAB-14 (COLOR) LAB-15 (COLOR)	.047 0.0477 .0290 0.044 0.036 0.053	.044 0.0446 .0294 0.040 0.036 0.049	.049 0.0431 .0298 0.040 0.041 0.048 0.038	.046 0.0459 .0294 0.040 0.038 0.047	.049 0.0482 .0296 0.044 0.040 0.049	.0470 .0459 .0294 .0416 .0382 .0492	.0021 .0021 .0003 .0022 .0023 .0023
LAB-16 (COLOR) LAB-20 (COLOR) LAB-26 (COLOR) LAB-52 (COLOR) LAB-53 (COLOR)	0.035 0.045 0.035 0.0436 0.031	0.036 0.045 0.032 0.0467 0.032	0.036 0.040 0.032 0.0422 0.032	0.037 0.047 0.029 0.0417 0.032	0.038 0.047 0.035 0.0456 0.033	.0364 .0448 .0326 .0440 .0320	.0011 .0029 .0025 .0022

It should be noted that 95% confidence limits denote that if the certification program were performed 100 times, the overall mean in 95 would fall within the prescribed limits.

The average within-set standard deviation, σ_A , is a measure of the average within-bottle precision as determined by the analytical methods used. The implication exists therefore that a laboratory using a method of average or better reproducibility should obtain individual results for a given certified element with a precision that is at least comparable to the reported value of σ_A .

CRITERION FOR CERTIFICATION

The ratio of the between-laboratory to the within-laboratory standard deviation, σ_B/σ_A , where

$$\sigma_{\rm B} = \sqrt{\left[\begin{smallmatrix} k \\ \Sigma \\ i \end{smallmatrix} \left(\begin{smallmatrix} \bar{x} \\ i \end{smallmatrix} - (\begin{smallmatrix} k \\ \Sigma \\ i \end{smallmatrix} \bar{x}_i .)/k \right) \right]^2 / k-1}$$

is a measure of the quality of the certification data for the reference materials of CCRMP (7). The acceptable upper limit for σ_B/σ_A is 3 for all elements except uranium for which an upper limit of 2 is more realistic.

The criterion for the certification of an element in a reference material is RP, the percentage of sets of results that must be rejected to give a value of σ_B/σ_A equal to or less than the acceptable upper limit. RP should not exceed 15%.

The values of σ_B/σ_A and RP for MP-la are reported in Table 7. It appears that zinc, lead, copper, tin, arsenic, tungsten, indium and silver but not bismuth and molybdenum can be certified according to the described criteria. For reasons stated below bismuth and molybdenum are being certified in MP-la. To the contrary, tungsten will not be certified.

DISCUSSION

Table 5 is a summary of a methodological classification of accepted analytical results

Table 7 - Values of $\sigma_{\rm B}/\sigma_{\rm A}$ and RP for MP-la

	Number of sets	$\sigma_{\rm B}/\sigma_{\rm A}$		RP
Element	of results	All results	Final	%
Zņ	19	2.21	2.21	0.0
Pb	21	3.22	2.93	4.8
Cu	27	4.30	2.35	3.7
Sn	16	3.16	2.58	12.5
As	17	2.79	2.79	0.0
W	12	3.42	2.87	8.3
In	12	3.49	2.76	8.3
Bi	13	3.79	2.99	23.1
Мо	17	3.78	2.83	17.6
Ag	18	3.19	2.96	11.1

where there is a clear-cut distinction between types of methods in decomposition, separations and determination steps. No attempt was made for any element to detect a statistically significant difference between the overall means of the more popular methods because there was generally not a sufficient number to warrant the test.

Table 8 summarizes the values of the between-laboratories spread given by

$$2(t_{0.975,(k-1)} \sqrt{v[\bar{x}..]}) \times 100\% \bar{x}$$
.

and the mean within-laboratory coefficient of variation given by $\sigma_A \propto 100\%/\bar{x}$. for the interlaboratory program for the 10 elements for both MP-1 and MP-1a. An examination of the between-laboratories spread readily illustrates an appreciable improvement in MP-1a over MP-1 for all elements except indium. In particular, the large decrease in the between-laboratories spread for MP-1a compared with MP-1 noted for tungsten is the main reason this element has a relatively low σ_B/σ_A ratio in MP-1a. The availability of a reference material such as MP-1 is likely one reason for the improved between-laboratories agreement noted for MP-1a.

A comparison of the values of the average within-laboratory coefficients of variation for MP-1 and MP-1a indicates that the elements fall

Table 8 - Between-laboratories spread and within-laboratory coefficient of variation for MP-l and MP-la

	Between-lab	spread (%)	Average within-lab coef	ff of variation (%)
Element	MP-1	MP-la	MP-1	MP-la
Zn	1.53	1.09	0.46	0.52
Pb	3.10	1.49	1.19	0.51
Cu	2.99	1.10	0.82	0.71
Sn	8.90	5 . 79	2.16	1.69
As	5.71	4.38	1.47	1.50
W*	57.09	20.61	7.87	4.65
In	8.86	8.80	2.41	2.39
Bi	16.02	11.86	5.56	2.70
Мо	13.16	8.00	7.62	1.98
Ag	7.30	5.03	1.50	1.60

^{*}W not certified in MP-1

into two groups - one for which the within-laboratory precision has remained essentially the same, and the other for which improvement has occurred. Of the latter, bismuth and molybdenum show the greatest improvement in the within-laboratory precision for MP-la compared with MP-l. Indeed, the magnitude of this improvement is sufficiently large that in spite of a significant improvement in the between-laboratories agreement, the ratio of σ_{R}/σ_{A} is increased to such an extent that RP of < 15% cannot be attained. Similar calculations readily illustrate that bismuth and molybdenum meet the criterion of RP <15% for MP-1 since $\sigma_{\rm B}/\sigma_{\rm A}$ \leq 3 initially. Therefore, a paradoxical situation exists wherein bismuth and molybdenum were certified for MP-1 but do not satisfy the certification criterion for its replacement, MP-la, even though the quality of the results from the interlaboratory program is superior with respect to both between-laboratories agreement and within-laboratory precision. Consequently, CCRMP considers it justifiable to waive the certification criterion for these two elements.

Figures la-lj, which show the histograms for the elements in MP-la after removal of outliers, illustrate that a calculation of a consensus value is warranted for all elements except tungsten. The histogram for tungsten clearly

shows the lack of any clustering of the analytical results, a fact which leads to the relatively large value of 20.61% for the between-laboratories spread. Indeed, it is only the relatively poor within-laboratory precision that allows tungsten to meet the certification criterion. Therefore, it is the opinion of CCRMP that using the certification criterion for tungsten would lead to an erroneous result and the lack of consensus precludes its certification at this time. The provisional value for tungsten is given in Table 9.

Indium presents a special example where there appears to have been no change in the between-laboratories agreement and the within-laboratory precision for MP-1 and MP-1a. Since the determination of this element is known not to be especially troublesome, it must be concluded

Table 9 - Provisional value for tungsten

Number of laboratories		11
Number of results		60
Overall mean		0.040%
95% confidence limits,	low	0.036%
	high	0.044%
σд		0.002%

that it is of minor interest in the routine operation of analytical laboratories. Supporting this is the fact that only six laboratories besides CANMET submitted results for indium. It may be concluded therefore that indium is of only minor commercial importance.

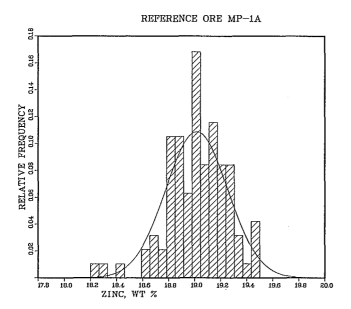


Fig. la - Histogram for zinc

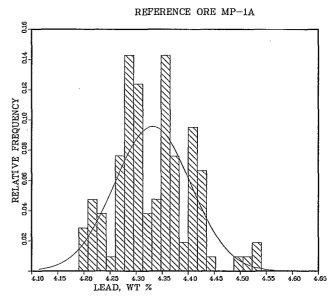


Fig. 1b - Histogram for lead

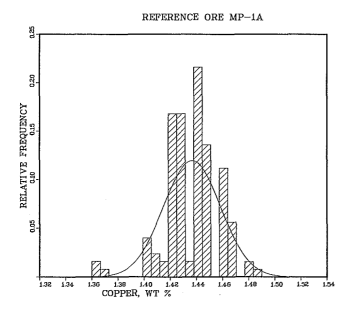


Fig. lc - Histogram for copper

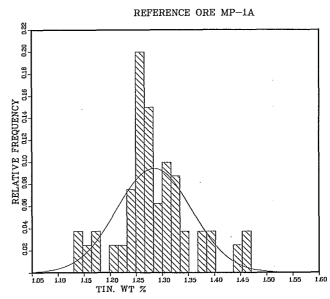
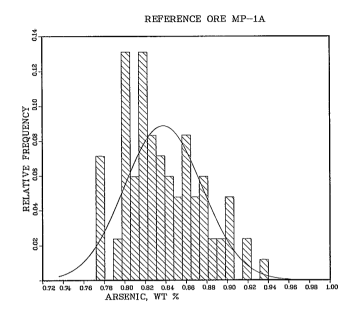


Fig. 1d - Histogram for tin



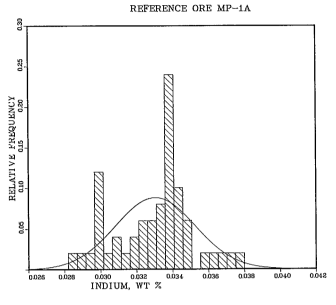
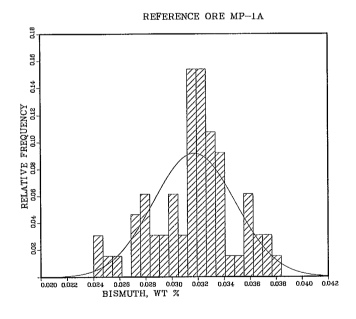


Fig. le - Histogram for arsenic

Fig. 1f - Histogram for indium



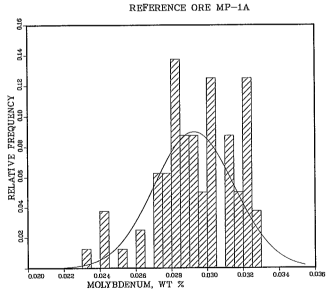
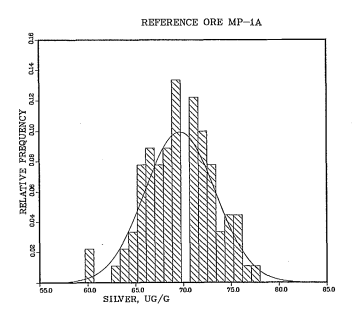


Fig. 1g - Histogram for bismuth

Fig. lh - Histogram for molybdenum



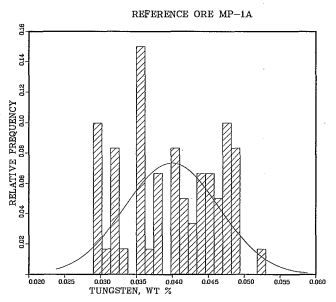


Fig. 1i - Histogram for silver

Fig. 1j - Histogram for tungsten

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APPENDIX A

CONFIRMATION OF HOMOGENEITY

CONFIRMATION OF HOMOGENEITY

The homogeneity of MP-la was confirmed at CANMET by analyzing in triplicate 15 bottles selected from a stock of 840 for zinc and bismuth. The stock was divided into 15 lots of 56 bottles. The code number of the first bottle was selected at random out of the first lot. The code numbers of the remaining bottles selected were given by

the code number of the preceding bottle plus 56. The results are shown in Tables 10 and 11.

A one-way analysis of variance technique was used to assess the homogeneity (5). Herein, the ratio of the between-bottle to within-bottle mean square is compared with the F statistic at the 95% level of probability. No evidence of bottle-to-bottle inhomogeneity was found for either zinc or bismuth.

Table 10 - Confirmation of homogeneity of MP-la for zinc

	···-	Zn (w	t %)	··
Bottle No	0.	Individual		Mean
44	19.08	19.01	19.01	19.03
100	18.96	19.04	18.94	18.98
156	19.00	18.93	18.99	18.97
212	19.06	19.00	18.97	19.01
268	19.02	19.01	19.05	19.03
324	18.97	18.49	18.99	18.98
380	19.01	18.96	18.98	18.98
436	18.97	18.96	18.96	18.96
492	18.98	19.04	18.96	18.99
548	18.98	19.00	18.99	18.99
604	19.00	18.96	19.04	19.00
660	18.96	19.03	18.95	18.98
716	18.99	19.02	18.97	18.99
772	18.97	18.97	18.95	18.96
828	18.97	19.03 _	18.99	19.00

Overall mean = 18.99

Table 11 - Confirmation of homogeneity of MP-la for bismuth

		Zn (1	wt %)	
Bottle 1	No.	Individual		Mean
44	0.0330	0.0313	0.0316	0.0320
100	0.0320	0.0325	0.0322	0.0322
156	0.0314	0.0318	0.0312	0.0315
212	0.0320	0.0317	0.0320	0.0319
268	0.0315	0.0314	0.0309	0.0313
324	0.0327	0.0309	0.0319	0.0318
380	0.0316	0.0316	0.0317	0.0316
436	0.0318	0.0322	0.0324	0.0321
492	0.0324	0.0314	0.0318	0.0319
548	0.0319	0.0318	0.0319	0.0319
604	0.0314	0.0317	0.0315	0.0315
660	0.0327	0.0327	0.0320	0.0325
716	0.0318	0.0312	0.0321	0.0317
772	0.0320	0.0316	0.0318	0.0318
828	0.0318	0.0329	0.0315	0.0321

Overall mean = 0.0318

Analysis of variance table for zinc

Source of	Degrees of	Mean
<u>variation</u>	freedom	square
Between bottles	14	1.232 x 10 ⁻³
Within bottles	30	1.082×10^{-3}
Total	44	
Calculated F statistic	= 1.139	
F.95(14,30)	= 2.037	
Null hypothesis of no	difference	between bottles
is accepted for zinc		

Analysis of variance table for bismuth

Source of	Degrees of	Mean
variation	freedom	square
Between bottles	14	2.852 x 10 ⁻⁷
Within bottles	30	2.173×10^{-7}
Total	44	
Calculated F statistic	= 1.312	
F.95(14,30)	= 2.037	
Null hypothesis of no	difference	between bottles
is accepted for bismuth	I	

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APPENDIX B

PARTICIPATING LABORATORIES

PARTICIPATING LABORATORIES

Bondar-Clegg and Company Ltd. North Vancouver, British Columbia R.K. Rogers

Bondar-Clegg and Company Ltd. Ottawa, Ontario P. Haulena

Brenda Mines Limited Peachland, British Columbia D. Perkins

CANMET, Energy, Mines and Resources Canada Mineral Sciences Laboratories Ottawa, Ontario (6 independent analysts)

Chemex Labs. Ltd.
North Vancouver, British Columbia
B.L. Twaites

Falconbridge Nickel Mines Ltd. Metallurgical Laboratories Thornhill, Ontario W.L. Ott

Falconbridge Nickel Mines Ltd. Sudbury Division Falconbridge, Ontario R.J. Wiseman

Geological Survey of India Central Chemical Laboratory Calcutta, India A.N. Chowdbury

Geological Survey of West Malaysia, Ipoh, Perak, Malaysia Tong Yik Lum

Hudson Bay Mining and Smelting Company Ltd. Flin Flon, Manitoba W.W. Henderson

Inco Ltd., Analytical Services Process Technology Copper Cliff, Ontario J. Bozic Inco Ltd.
J. Roy Gordon Research Laboratory
Sheridan Park, Ontario
St. J.H. Blakely

Lakefield Research of Canada Ltd. Lakefield, Ontario D.M. Wyslouzil

National Institute for Metallurgy Randburg, South Africa E.J. Ring

Noranda Research Centre Pointe Claire, Quebec J.D. Kerbyson

Ontario Ministry of Natural Resources Geoscience Laboratories Toronto, Ontario C. Riddle

Sherritt-Gordon Mines Ltd. Research and Development Division Fort Saskatchewan, Alberta D.J. Whitehead

Sherritt-Gordon Mines Ltd. Mining Division Lynn Lake, Manitoba R. Klassen

Surinam Government Geological and Mining Services Paramaribo, Surinam K.E. Burke

United Keno Hill Mines Elsa, Yukon V. Rafuse

X-ray Assay Laboratories Ltd. Don Mills, Ontario E.J. Brooker