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Certificate of Analyses

B.C.S. No. 301/1 EURO-CRM 651-LINCOLNSHIRE IRON ORE

Prepared under rigorous laboratory conditions and, AFTER STANDARDIZATION BY ANALYSTS IN GREAT BRITAIN, issued by the Bureau of Analysed Samples, Ltd.

The material for this standard was kindly donated by the British Steel Corporation, Scunthorpe Group. It was crushed to pass a 180 micrometre (85 mesh) sieve and the sieved material passed through a magnetic separator.

CO-OPERATING ANALYSTS AND FIRMS

INDEPENDENT ANALYSTS

- 1. Coppins, W. C., M.Sc., F.R.I.C., Ridsdale & Co. Ltd., Middlesbrough.
- 2. WHITEHEAD, T. C. G., B.Sc., A.R.I.C., Pattinson & Stead, Middlesbrough.

GOVERNMENT DEPARTMENT

3. Moore, P. J., B.Sc., A.R.I.C., and Hutchison D., Institute of Geological Sciences, London.

ANALYSTS representing MAKERS and USERS

- 4. Borrowdale, J., B.Sc., B.S.C., Redbourn Works, Scunthorpe.
- 5. HARRISON, T. S., B.Sc., Ph.D., B.Met., F.R.I.C., B.S.C., Group Chemical Laboratories, Scunthorpe.
- 6. McArdle, A. J., A.I.M., B.S.C., Lackenby Works, Middlesbrough.
- 7. Muir, S.,

B.S.C., Corby Works, Corby.

ANALYSES

ALL RESULTS RELATE TO THE DRIED (105°C SAMPLE)

			1.5								,			Loss on
Analyst No.	Fe %	SiO ₂	Al ₂ O ₃	TiO ₂	MnO %	CaO %	MgO %	Na ₂ O %	K₂O %	S %	P %	°CO ₂	H ₂ O %	Ignition %
1 2 3 4 5 6 7	23·83 23·89 23·89 23·85 23·85 23·86 23·79	7·40 7·40 7·43 7·42 7·41 7·38 7·37	$\begin{array}{c} 4 \cdot 21 \\ 4 \cdot 22 \\ 4 \cdot 31 \\ 4 \cdot 22 \\ 4 \cdot 31 \\ 4 \cdot 26 \\ 4 \cdot 27 \end{array}$	0·16 0·18 0·16 0·17 0·16 0·16 0·16	1·21 1·24 1·29 1·27 1·28 1·22 1·24	22.7 22.4 22.8 22.6 22.5 22.5 22.6	1·69 1·74 1·67 1·74 1·75 1·75 1·76	0.06 0.05 0.06 0.07 0.08 0.08	0·31 0·29 0·31 0·34 0·34 0·31	0·39 0·41 0·40 0·41 0·40 0·40	0·36 0·35 0·37 0·35 0·34 0·35	22·2 22·2 22·9 21·8 22·0 21·9 22·0	5.5 5.5 5.4 4.8 5.1 5.4	25.5 25.9 25.6 26.0 25.5 25.9 26.0
Average	23.85	7.40	4.26	0.16	1.25	22.6	1.73	0.07	0.32	0.40	0.35	••	••	

EQUIVALENT % ELEMENT CONTENT OF OXIDE CONSTITUENTS

Si	A1	Ti	Mn	Ca	Mg	Na	K
3.46	2.25	0.10	0.97	16.2	1.04	0.05	0.27

The above figures are those which each analyst has decided upon after careful verification Figures in bold type standardized, figures in small italic type only approximate

B.C.S. No. 301/1 LINCOLNSHIRE IRON ORE

NOTES ON METHODS USED

IRON

Ali analysts determined iron by titration of the reduced solution with standard dichromate solution. Nos. 1, 5 and 7 used the Standard Method B.S. 4158: Part 1: 1967 which depends on reduction of the ferric iron with hydrogen sulphide. No. 3 used a similar method. Nos. 2, 4 and 6 reduced with stannous chloride.

Analyst No. 5 also used a stannous chloride reduction method and found 24.00%. No. 6 also used the British Standard Method and found 23.64%.

Analyst No. 4 also determined ferrous iron and found 7.4%; this will result in an ignition gain of 1.1%.

SILICA

All analysts determined silica gravimetrically after double dehydration with perchloric acid. Nos. 1, 4, 5 and 6 followed the procedure of the Standard Method B.S. 4158: Part 3: 1972.

ALUMINA

Analysts Nos. 1, 2, 3, 4 and 7 separated aluminium together with other Group III elements by precipitation with ammonia solution; iron was separated by treatment with sodium hydroxide solution and the aluminium was precipitated as oxinate and determined by titration with bromate¹ (Nos. 1, 4 and 7) or gravimetrically (Nos. 2 and 3). Nos. 5 and 6 used atomic absorption spectroscopy.

Analyst No. 5 also determined aluminium as the oxinate by titration and found 4.26%.

TITANIA

Analysts Nos. 1, 2, 3, 4, 6 and 7 determined titania colorimetrically with hydrogen peroxide. Nos. 1, 2, 3 and 6 carried out a preliminary separation with cupterron, No. 4 first separated titanium, together with aluminium, as the phosphate. No. 5 determined titania colorimetrically with diantipyrylmethane.

Analyst No. 5 also used a cupferron/hydrogen peroxide method and found 0.17%.

MANGANESE OXIDE

Analysts Nos. 1, 2 and 4 determined manganese oxide by titration. No. 1 oxidized with persulphate/silver nitrate and titrated with arsenite/nitrite solution (Analoid Method No. 53), No. 2 oxidized with bismuthate and titrated with ammonium ferrous sulphate/permanganate. No. 4 oxidized with persulphate/silver nitrate and titrated with arsenite. Nos. 3, 5, 6 and 7 determined manganese oxide colorimetrically after oxidation with periodate.

LIME

Analysts Nos. 1 and 4 separated Group III elements by double precipitation with ammonia and separated manganese with bromine/ammonia; the lime was then precipitated as oxalate and titrated. No. 2 separated the bulk of the iron by extraction into amyl acetate, then separated any remaining Group III elements together with manganese by means of bromine/ammonia and finally precipitated lime as oxalate and completed gravimetrically by ignition to calcium oxide. No. 3 separated Group III elements by double precipitation with ammonia and determined lime gravimetrically by ignition of the oxalate; a correction was made for co-precipitated manganese which was determined colorimetrically. Nos. 5 and 6 carried out a basic acetate separation followed by a bromine/ammonia separation, then precipitated lime as oxalate and completed by titration. No. 7 separated lime by a basic acetate separation followed by bromine/ammonia separation but completed by titration with EDTA.

MAGNESIA

Analysts Nos. 1 and 6 determined magnesia by atomic absorption spectroscopy. Nos. 2, 4 and 5 determined magnesia in the filtrate from the lime determination by precipitation as magnesium ammonium phosphate and gravimetric determination as Mg₂P₂O₇. Nos. 3 and 7 used the filtrate from the lime determination but titrated with EDTA.

Analysts Nos. 2 and 5 also used atomic absorption spectroscopy and found 1.74% and 1.69% respectively. No. 6 also determined magnesia gravimetrically as the pyrophosphate and found 1.84%.

ALKALIS

Analysts Nos. 1, 2, 3 and 7 determined alkalis with a flame photometer. Nos. 5 and 6 used atomic absorption spectroscopy.

SULPHUR

Analysts Nos. 1, 2, 4, 5 and 6 determined sulphur gravimetrically as barium sulphate. Nos. 1, 2, 5 and 6 used a method involving reduction of the iron with hydroxylammonium chloride¹. No. 7 determined sulphur by combustion in the presence of tungstic oxide using nitrogen as carrier gas.²

Analyst No. 4 also used a combustion method and found 0.40%.

PHOSPHORUS

All analysts except No. 5 determined phosphorus by titration after precipitation as ammonium phosphomolybdate. Nos. 1, 6 and 7 oxidized the phosphorus by evaporation with perchloric acid and eliminated arsenic by treatment with hydrobromic acid before precipitating the phosphorus. No. 5 determined phosphorus colorimetrically as phosphovanadomolybdic acid according to the Standard method BS 4158: Part 2: 1970.

CARBON DIOXIDE

All analysts determined carbon dioxide gravimetrically by evolution with dilute acid and absorption in soda-asbestos1.

COMBINED WATER

All analysts determined combined water by heating the dried sample at 1000°C in a stream of dry air and collecting the evolved water in magnesium perchlorate.¹

LOSS ON IGNITION

All analysts except Nos. 3 and 7 determined loss on ignition by heating at 1000°C. for 1 hour. No. 3 heated at 1050°C to constant weight. No. 7 heated at 1000°C to constant weight.

- 1 British Iron and Steel Research Association: Report on Recommended Methods of Analysis for Iron Ores MG/D/56/59.
- 2 British Iron and Steel Research Association: Report on the Determination of Sulphur in Non-Metallic Materials, MG/D/412/70.

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