

Bureau of Mines Information Circular/1985

Bulk Mineralogy and Geochemistry of Selected Alaskan Chromian Spinel Samples

By William S. Roberts



UNITED STATES DEPARTMENT OF THE INTERIOR



Information Circular 9023

Bulk Mineralogy and Geochemistry of Selected Alaskan Chromian Spinel Samples

By William S. Roberts



UNITED STATES DEPARTMENT OF THE INTERIOR Donald Paul Hodel, Secretary

BUREAU OF MINES Robert C. Horton, Director

Library of Congress Cataloging in Publication Data:

Roberts, William S

Bulk mineralogy and geochemistry of selected Alaskan chromian spinel samples.

.

(Information circular / United States Department of the Interior, Bureau of Mines ; 9023).

Bibliography: p.9.

Supt. of Docs. no.: I 28,27:9023.

1. Spinel-Alaska-Composition. 2. Chromite-Alaska-Composition. I. Title. II. Series: Information circular (United States. Bureau of Mines); 9023.

 $TN295.U4 \quad \left[\mbox{QE391.S68} \right] \quad 622s \ \left[549 \ .526 \right] \quad 84\text{-}600362$

CONTENTS

Abstract	1
Introduction	2
Acknowledgments	2
Samples	2
Procedures	2
Nomenclature and mineralogy	4
Results	5
X-ray diffraction	5
X-ray fluorescence	5
Conclusions	8
References	9
Appendix AAnalytical results	10
Appendix BNumber of cations per 32 oxygens	12
Appendix CMineral terms and sample key	13

ILLUSTRATIONS

1.	Locations of ultramafic rocks hosting chromian spinel occurrences charac-
	terized in this report
2.	Schematic diagram of sample preparation procedures
3.	Nomenclature of chromium-bearing spinels based on molecular proportion plot of divalent and trivalent cations
4.	Unit cell dimensions of chromian spinel based on (113) reflection
5.	Molecular proportion plot of purified chromian spinel samples
6.	Trace Ni versus Cr
7.	Trace Zn versus Cr
8.	Trace Ti versus Cr

TABLES

1.	Range of oxide concentrations in chromian spinel samples	7
2.	Statistical comparison of oxides and lattice data for the interior Alaska	
	and Chugach group of samples	7

Page

Ĩ

	UNIT OF	MEASURE	ABBREVIATIONS	USED	IN THIS	REPORT		
Á	angstrom				wt %	weight	percent	
ppm	n part per	million						

BULK MINERALOGY AND GEOCHEMISTRY OF SELECTED ALASKAN CHROMIAN SPINEL SAMPLES

By William S. Roberts¹

ABSTRACT

As part of the Bureau of Mines critical and strategic minerals program in Alaska, 34 high-purity chromian spinel samples were analyzed for trace and major element geochemistry and mineralogy. The samples were analyzed by X-ray diffraction, X-ray fluorescence, and chemical techniques.

Systematic differences in chemistry are evident when samples collected from the Chugach Mountains and their related extension are compared with chromian spinel samples collected from interior Alaska. Compared with the Chugach samples, the interior Alaska samples have higher average amounts of Al_2O_3 , Fe_2O_3 , TiO_2 , and NiO, have lower average amounts of FeO and Cr_2O_3 , and exhibit greater dispersion in values.

Considerable mineralogical variability of chromian spinel exists between and within individual Alaskan chromite-bearing ultramafic complexes. Of 34 samples analyzed, 25 samples are magnesiochromite, 5 are chromite, 3 are spinel, and 1 is hercynite. As a group the interior Alaska samples have a smaller average unit cell dimension than the Chugach group of samples.

The chemical and mineralogical differences between the two groups of samples can be attributed to different genetic histories and may have implications for locating chromian spinel deposits amenable to standard beneficiating techniques.

¹Physical scientist, Alaska Field Operations Center, Bureau of Mines, Juneau, AK.

This report summarizes mineralogical and geochemical work done on samples collected during field investigations of Alaskan chromite occurrences. Field work assessing chromite deposits, as part of a critical and strategic minerals program, began in 1981 by the Bureau of Mines Alaska Field Operation Center (AFOC), and several deposits have recently been described (6-7, 11).² Beneficiation studies of chromian spinel samples have been done by the Bureau's Albany Research Center (ALRC) (3-4).

The purpose of this study is to characterize and compare the geochemical determinations for 34 purified chromian spinel samples. The samples, plus 18 other chromian spinel samples, were analyzed by X-ray diffraction. Chromite mineralogy is sufficiently complex to require a combination of X-ray diffraction, X-ray fluorescence, and chemical analyses for characterization.

Previous studies have focused on the description of chromian spinel occurrences and standard beneficiation tests of bulk samples. Mineralogical determinations have generally been restricted to the identification of gangue and trace minerals associated with chromian spinel. This report provides information on the mineralogy of Alaskan chromian spinel samples and reports trace and major oxide chemical signatures of samples collected statewide.

ACKNOWLEDGMENTS

K. Broadhead, metallurgical engineer, Reno Research Center, provided the FeO determinations, and D. Dahlin, metallurgist, Albany Research Center, provided splits of chromian spinel samples.

SAMPLES

Fifty-two samples were collected by Bureau personnel. The samples represent splits taken from grab or bulk samples that were high-graded in the field for mineralogical and beneficiation tests. The results of this study represent reconnaissance-type information since a few samples from each occurrence do not constitute a thorough characterization for any single deposit, area, or region.

The samples were obtained from widely scattered ultramafic occurrences across

PROCEDURES

The procedures used for this study are not standard concentration techniques. High-purity chromian spinel samples are required for mineralogical characterization using molecular proportion plots, so modified laboratory techniques were used.

the State; however, all chromian spinel occurrences in Alaska are not represented in this report (fig. 1). For comparative purposes the samples are lumped into two logical groups that are based on geographic location or extension of geographic or geologic features. The two groups include the interior Alaska and General localities and Chugach groups. grouping of the samples are indicated in appendix C.

Two different sources for samples resulted in slight variations in the concentration procedures. Samples archived at AFOC were prepared in the manner outlined in figure 2. Samples received from ALRC were table concentrates of minus 28-plus 65-mesh sized fraction, so crushing and hand picking were not required.

²Underlined numbers in parentheses refer to items in the list of references preceding the appendixes.



FIGURE 1. - Locations of ultramafic rocks hosting chromian spinel occurrences characterized in this report. *A*, Avan Hills; *B*, Caribou Mountain, Kanuti River; *C*, Yuki River; *D*, Mount Hurst; *E*, Tonsina-Bernard, Sheep Hill; *F*, Wolverine Complex, Eklutna; *G*, Claim Point, Red Mountain; H, Miners Bay, Grant Lagoon, Halibut Bay; *I*, Red Bluff Bay.

The samples were run through a laboratory model isodynamic magnetic separator to remove all magnetite, and a hydrofluoric acid (HF) bath effectively removed unwanted silicate minerals. The HF bath altered the magnesium silicates to chondrodite, a relatively low density, fragile fluorohydroxide which could be washed out by agitation and wet screening. There is no evidence the HF bath altered the chemistry or mineralogy of the chromian spinels.

High-purity chromian spinel concentrates free of magnetite and silicates were analyzed by energy-dispersive X-ray fluorescence spectrometry (XRF), with the exception of FeO, which was analyzed by wet chemical methods. Ferric oxide (Fe_2O_3) was calculated from total Fe.

The analytical precision is less than a few percent relative error, while errors in accuracy may range as high as $\pm 20\%$ in the case of Mg or trace elements because of low counting rates. Comparison of Fe and Cr determinations by ALRC and AFOC during this study generally indicates a relative error in accuracy of $\pm 5\%$ to 10% or less. National Bureau of Standards NBS103A was the primary reference standard used in the XRF determinations.



FIGURE 2. - Schematic diagram of sample preparation procedures. Samples received from Albany were concentrated using tabing methods, and splits consisted of minus 28-plus 65-mesh fractions. These samples were treated beginning with the magnetic separator.

After analysis by XRF the samples were analyzed by X-ray diffraction using two modes: a scanning speed of $1/4^{\circ}$ two-theta per minute from 31° to 37° two-theta, and a scanning speed of 2°



FIGURE 3. - Nomenclature of chromium-bearing spinels based on molecular proportion plot of divalent and trivalent cations. Note: Changes along horizontal axis reflect different proportions of divalent cations (Fe, Mg), and changes on the vertical axis represent different proportions of trivalent cations (Cr, Al). Unit cell dimensions, in angstroms, are indicated in parentheses and are for end member minerals only.

two-theta per minute from 2 to 62° twotheta. Measured precision of d-spacings in the former mode is ± 0.0005 Å. An internal standard (NaCl) was used to standardize the d-spacing measurements using a full-width-half-maximum technique of marking peak centroids.

The (113) d-spacing, the strongest spinel reflection, was used to calculate unit cell dimensions. Since multiple reflections from different crystallographic planes were not used to calculate the unit cell dimensions, the values reported must be considered provisional.

NOMENCLATURE AND MINERALOGY

Mineral nomenclature is adapted from Palache (10) and conforms to terminology used by the Joint Committee Powder Diffraction Society (1). Chromite from podiform and stratiform ultramafics is mineralogically part of the spinel group, which consists of the magnetite, spinel, and chromite series. Spinel-group minerals may be represented by the general formula AB_2X_4 , which may be expressed as $R^{2+}R_2^{3+}O_4$ (<u>13</u>). Extensive ionic substitution of divalent cations (Fe, Mg, and Ti) and trivalent cations (Cr, Al, Fe, and Mn) makes characterization by Xray diffraction difficult since crystal lattice dimensions are variably affected by substitution at the octahedral and tetrahedral positions. Although there are natural limits on ionic substitution, characterization is best achieved by combining X-ray diffraction and quantitative chemistry.

Low-titanium chromium-bearing spinels consist of six idealized end members: chromite (FeCr $_2O_4$), magnesiochromite $(MgCr_2O_4)$, hercynite (FeAl₂O₄), spinel $(MgAl_2O_4)$, magnesioferrite $(MgFe_2O_4)$, and magnetite (Fe₃ O_4) (9). A multicomponent prism described by Haggerty (8) best characterizes chromium spinels, but for this study only the base of the prism is used. Figure 3 outlines, on a molecular proportion plot, the fields represented by four of the end-member minerals Magnesioferrite and maglisted above. netite are not represented on the plot, although there is an appreciable ferrian $(Fe+^3)$ component represented by the analyses.

RESULTS

The results of the chromian spinel chemical analyses are presented after a brief discussion of X-ray diffraction results. Diffraction data for 52 samples are listed in appendix A, along with chemical results for 34 samples. Molecular proportions of the principal spinel cations are listed in appendix B, and mineral terms and general localities are tabulated in appendix C.

X-RAY DIFFRACTION

High-resolution X-ray diffraction indicates that average d-spacings of the (113) reflection vary from 2.4835 to 2.5260 Å. These measurements indicate unit cell dimensions of 8.237 to 8.378, respectively. Figure 4 represents a plot of calculated cell dimensions with summary envelopes of the Chugach and interior Alaska groups of samples. There is a clear difference in average cell dimensions, with the Chugach group averaging 8.312 Å and the interior Alaska group averaging 8.279 Å. The disparity in unit cell dimensions is apparently due to a difference in average amounts of Al, Mg, Fe, and Cr.

Eleven samples have multiple (113) reflections ("doublet"), indicating more than one spinel mineral is present. Nine of the 11 samples with doublets are from the Chugach group of samples. Dahlin (3) and Bliss (2) describe zoned chromian spinel minerals, which are the probable cause for the doublets.

X-RAY FLUORESCENCE

Plots comparing molecular proportions of $Fe^{2+}/Fe^{2+} + Mg^{2+}$ and $Cr^{3+}/Cr^{3+} + Al^{3+}$ concentrations are presented in figure 5. The plots define R^{2+} and R^{3+} cation proportions and the mineralogy of the high-purity bulk chromian spinel splits. Fe^{+3} is not represented on the plot even though the chemical analyses indicate an appreciable ferrian component.

Mineral terms for 34 samples are based mainly on molecular proportions. Analysis by X-ray diffraction was used to check purity of the samples and to confirm the presence of a spinel. A total of 25 samples are characterized as magnesiochromite, 5 are chromite, 3 are spinel, and 1 is hercynite.



FIGURE 4. - Unit cell dimensions of chromian spinel based on (113) reflection. Note: Envelopes are used to diagrammatically illustrate differences between the interior Alaska and Chugach groups of samples. The range of cell dimensions shows the variation in chromian spinel compositions within a related area or complex.

The concentrations of major oxides in the chromian spinel samples exhibit considerable variability (table 1). This variability exists not only between regions, but between samples taken within the same ultramafic complex. This emphasizes the importance of understanding the chemistry and mineralogy of chromian spinel occurrences since the



FIGURE 5. - Molecular proportion plot of purified chromian spinel samples. Note: Refer to figure 3 for mineral nomenclature.

beneficiating procedures and market depend heavily on ore chemistry.

Systematic chemical and unit cell differences exist between the Chugach and interior Alaska samples when the groups, rather than specific samples, are compared. Compared with the Chugach group of samples, the interior Alaska group

TABLE 1. - Range of oxide concentrations in chromian spinel samples, weight percent

Oxide ¹	Minimum value	Maximum value	Mean value
MgO	4.5	17.7	11.4
AI 20 3	7.0	26.7	13.9
$\operatorname{Cr}_{2}\overline{\operatorname{O}}_{3}$	25.8	59.8	49.5
Fe ₂ 0 ₃	0	15.6	8.2
FeÖ.	4.2	22.0	14.5

¹Oxides based on X-ray fluorescence analyses except FeO which was determined chemically by the Bureau of Mines, Reno Research Center. The Fe_2O_3 values were calculated from total iron. samples have higher average amounts of $Al_{2}O_3$, Fe_2O_3 , TiO_2 , and NiO and exhibit greater dispersion in values (table 2). Figures 6, 7, and 8 graphically summarize the differences between the two groups

when Ni, Zn, and Ti concentrations are compared to Cr content. The significance of the chemical differences between the groups can be attributed to different genetic histories.

TABLE 2. - Statistical comparison of oxides and lattice data for the interior Alaska and Chugach group of samples

Statistical parameter	In	terior Alas	ka group	Chugach group				
	N 1	Mean wt %	SD, ² wt %	N ¹	Mean wt %	SD, ² wt %		
Mg0	17	11.2	2.1	17	11.5	3.9		
A1 20 3	17	17.4	8.9	17	10.4	3.8		
Fe0.	17	12.7	4.3	22	15.8	4.0		
Fe ₂ 0 ₃	17	9.5	4.1	17	6.9	3.5		
$\operatorname{Cr}_{2}\overline{\operatorname{O}}_{3}$	17	46.4	11.6	17	52.5	6.1		
\$i0 2	17	0.27	0.19	17	0.27	0.21		
P ₂ 0 ₅	17	0.12	0.04	17	0.11	0.04		
Ti0 ₂	17	0.50	0.22	17	0.30	0.15		
Nio	17	0.10	0.04	17	0.06	0.03		
Zn0	17	0.11	0.10	17	0.09	0.09		
d-spacingÅ.	27	2.4965	0.0075	27	2.5060	0.0081		
Unit cellÅ	27	8.279	0.024	27	8.312	0.027		
^{1}N = number of observations. ^{2}SD = standard deviation.								

45 40 nterior Alaska Group 35 Ø 30 Cr, w† % C 25 Chugach Grou 20 0 15 10 600 800 1,400 1,600 200 400 1,000 1,200 0 Ni, ppm KEY Red Mountain, Claim Point Avan Δ 0 Kanuti, Caribou ÷ Kodiak ø Yuki River Tonsing Mount Hurst æ Southeast ۵ Eklutna, Wolverine



FIGURE 6. - Trace Niversus Cr. Note: Envelopes are used to diagrammatically illustrate differences between the interior Alaska and Chugach groups of samples.

FIGURE 7. - Trace Zn versus Cr. Note: Envelopes are used to diagrammatically illustrate differences between the interior Alaska and Chugach groups of samples.



FIGURE 8. - Trace Ti versus Cr. Note: Envelopes are used to diagrammatically illustrate differences between the interior Alaska and Chugach groups of samples.

CONCLUSIONS

The following conclusions are based on X-ray diffraction, X-ray fluorescence, and chemical analyses of 34 high-purity chromian spinel samples:

1. Samples that are grouped by region or geologic extension have systematic chemical differences when average values are compared. The interior Alaska group of samples, when compared to the Chugach group of samples, have higher average amounts of Al₂O₃, Fe₂O₃, TiO₂, and NiO, exhibit greater dispersion in values, have lower average amounts of FeO and Cr_2O_3 , and have similar average Mg0 concentrations.

2. Considerable mineralogical variability exists between and within individual Alaskan chromian spinel-bearing

34 chromian ultramafic complexes. 0f spinel samples analyzed, 25 consist of ferroan or aluminian magnesiochromite, 5 are aluminian or magnesian chromite, 3 are chromian spinel, and 1 is chromian hercynite. The Chugach sample group has a larger average unit cell dimension when compared to the interior Alaska group of samples and has a higher incidence of doublet (113) diffraction reflections, indicating the presence of more than one spinel phase.

3. The chemical and mineralogical differences between the groups may be a significant factor in the location of chromian spinel deposits amenable to standard beneficiating techniques.

REFERENCES

1. Berry, L. G. (ed.). Selected Powder Diffraction Data for Minerals. Joint Committee on Powder Diffraction Standards, Swarthmore, PA, 1974, 833 pp.

2. Bliss, N. W., and W. H. MacLean. The Paragenesis of Zoned Chromite From Central Manitoba. Geochim. et Cosmochim. Acta, v. 39, 1975, pp. 973-990.

3. Dahlin, D. C., L. L. Brown, and J. J. Kinney. Podiform Chromite Occurrences in the Caribou Mountain and Lower Kanuti River Areas, Central Alaska. Part 2: Beneficiation. BuMines IC 8916, 1983, 15 pp.

4. Dahlin, D. C., D. E. Kirby, and L. L. Brown. Low-Grade Chromite Deposits Along the Border Ranges Fault, Southern Alaska. 2. Beneficiation. BuMines IC 8991, 1984.

5. Deer, W. A., R. A. Howie, and J. Zussman. An Introduction to the Rock-Forming Minerals. Longman Group Limited, 1966, pp. 424-433.

6. Foley, J. Y., and J. C. Barker. Low-Grade Chromite Deposits Along the Border Ranges Fault, Southern Alaska. 1. Field Investigations and Descriptions of Chromite Deposits. BuMines IC 8990, 1984.

7. Foley, J. Y., and M. M. McDermott. Podiform Chromite Occurrences in the Caribou Mountain and Lower Kanuti River Areas, Central Alaska. Part 1. Reconnaissance Investigations. BuMines IC 8915, 1983, 27 pp.

8. Haggerty, S. E. Opaque Mineral Oxides in Terrestrial Igneous Rocks. Sec. in Oxide Minerals. Mineralogical Society of America Short Course Notes, v. 3, Nov. 1976, pp. 101-150.

9. MacGregor, I. D., and C. H. Smith. The Use of Chrome Spinels in Petrographic Studies of Ultramafic Intrusions. Can. Mineral., 1962, pp. 403-412.

10. Palache, C., H. Berman, and C. Frondel. The System of Mineralogy of J. D. Dana and E. S. Dana. Wiley, v. 1, 7th ed., 1944, pp. 687-712.

11. Roberts, W. S. Economic Potential for Chromium, Platinum, and Palladium in the Mount Hurst Ultramafics, West-Central Area, Alaska. BuMines OFR 22-84, 1984, 52 pp.

12. Stevens, R. E. Composition of Some Chromites of the Western Hemisphere. Am. Mineral., v. 29, Nos. 1-2, 1944, pp. 1-34.

13. Thayer, T. P. Principal Features and Origin of Podiform Chromite Deposits, and Some Observations on the Guleman-Soridag District, Turkey. Econ. Geol., v. 59, 1964, pp. 1497-1524.

9

APPENDIX A. -- ANALYTICAL RESULTS¹

Cample No.	1				E	6	7	0	0	10
Sample No	¹	Z				0		0	7	10
Analysis, wt %:	NT A		0.0	NT A	NT A	NT A	NT A	0.1	NTA.	NT A
MgU	NA	9.8	9.3	NA		NA	NA	8.1	NA	NA NA
$AL_{2}O_{3}$	NA	11.5	26.7	NA	NA	NA	NA	29.8	NA	NA
reo	Na	15.0	15.0	NA	NA	NA	NA	18.0	NA.	NA
re ₂ 0 ₃	NA	2.5	/.5	NA	NA	NA	NA	14.3	NA	NA
$\operatorname{Cr}_2 \operatorname{O}_3 \ldots \ldots \ldots$	NA	58.7	38.3	NA	NA	NA	NA	27.4	NA	NA
Si0 ₂	NA	0.06	0.63	NA	NA	NA	NA	0.24	NA NA	NA
P ₂ 0 ₅	NA	0.15	0.08	NA	NA	NA	NA	0.10	NA	NA
Ti0 ₂	NA	0.56	0.76	NA	NA	NA	NA	0.89	NA	NA
NiO	NA	0.09	0.12	NA	NA	NA	NA	0.09	NA	NA
Zn0	NA	0.05	0.08	NA	NA	NA	NA	0.14	NA	NA
d-spacingÅ	2.5010	2.5015	2.4845	2.4970	2.5025	2.4970	2.4985	2.4870	2.4875	2.4950
Unit cellÅ	8.295	8.297	8.240	8.282	8.300	8.282	8.287	8.248	8.250	8.275
Sample No	11	12	13	14	15	16	17	18	19	20
Analysis, wt %:										
Mg0	NA	NA	11.0	12.3	11.3	11.4	11.7	9.9	8.6	12.9
A1 ₂ 0 ₃	NA	NA	15.2	11.7	22.9	31.9	8.5	34.6	15.0	12.9
Fe0	NA	NA	9.3	14.0	15.0	17.0	17.0	12.0	15.0	13.0
Fe ₂ 0 ₃	NA	NA	7.7	6.3	5.6	13.4	3.5	11.1	14.1	9.2
$Cr_{2}O_{3}$	NA	NA	54.2	52.7	43.1	25.8	53.9	28.2	46.1	49.8
Si0 ₂	NA	NA	0.16	0.12	0.11	0.30	0.19	0.12	0.17	0.09
P ₂ 0 ₅	NA	NA	0.12	0.19	0.07	0.09	0.07	0.08	0.19	0.20
Ti0,	NA	NA	0.50	0.45	0.53	1.05	0.22	0.48	0.32	0.45
Ni0	NA	NA	0.12	0.08	0.11	0.13	0.03	0.18	0.17	0.10
Zn0	NA	NA	0.05	0.10	0.07	0.09	0.47	0.112	0.14	0.06
d-spacingÂ	2.5065	2.5005	2.5040	2.5005	2.4890	2.4835	2.5045	2.4775	2.5045	2.5015
Unit cellÅ	8.313	8.293	8.305	8.293	8.255	8.237	8.306	8.217	8.306	8.296
Sample No	21	22	23	24	25	26	27	28	29	30
Analysis, wt %:										
Mg0	11.7	11.6	16.4	NA	13.4	8.4	12.0	13.5	NA	NA
A1 20 3	8.5	7.8	16.7	NA	10.1	22.4	9.8	3.6	NA	NA
Fe0	4.2	6.8	14.0	NA	9.1	16.0	4.6	14.0	20.0	19.0
Fe ₂ 03	15.6	9.6	7.4	NA	6.5	15.2	12.3	3.6	NA	NA
$\operatorname{Cr}_{2}O_{3}$	58.2	59.5	43.6	NA	57.4	36.0	56.4	59.8	NA	NA
Si02	0.23	0.39	0.22	NA	0.53	0.64	0.45	0.24	NA	NA
P ₂ O ₅	0.17	0.10	0.12	NA	0.11	0.09	0.09	0.14	NA	NA
Ti02	0.39	0.30	0.44	NA NA	0.31	0.44	0.32	0.03	NA	NA
NiO	0.07	0.12	0 08			0.08	0.02		NA	
ZnO	0.05		0.00		0.07	0.00	0.00	0.02	NIA NIA	
d-spacing 8	2 /025	2 /075	2 /000	2 5000	2 /000	2 /0/5	2 6040	2 5120	2 4055	2 /070
u spacingA	2.472J	2.47/3	0 100	2.5000	2.4990	2.4943	0 070	2. 0 221	4.4700	2.49/0
Contraction Area	0.207	0.203	0.200	0.291	0.200	0.2/3	0.218	0.331	0.211	0.202

See explanatory notes at end of table.

Sample No	31	32	33	34	35	36	37	38	39	40
Analysis, wt %:										
Mg0	14.6	17.7	NA	17.5	10.0	11.0	9.1	6.6	10.7	4.5
Al ₂ 03	11.8	7.2	NA	7.6	8.2	9.9	9.1	7.0	9.8	15.5
Fe0	11.0	8.5	10.0	16.0	13.0	16.0	13.0	16.0	22.0	21.0
Fe ₂ 03	5.8	7.4	NA	1.9	6.2	4.5	7.6	4.9	0.0	9.2
Cr 203	54.6	53.6	NA	51.6	58.8	57.1	57.8	59.7	60.3	46.9
Si02	.20	-28	NA	.49	.01	.18	.19	NA	.08	.70
P ₂ 0 ₅	.08	.10	NA	.11	.21	.10	.07	.04	.11	.08
Ti02	. 44	.30	NA	.29	.24	28	.29	.27	31	38
NiO	.09	.09	NA	.09	.04	.06	.07	.03	.05	.03
Zn0	.04	.05	NA	.05	.05	.05	.04	.05	.04	.14
d-spacingÅ.	2.5020	2.5045	2.5025	2.5040	2 5050	2 5065	2 5070	2 5100	2 5065	2 5030
Unit cellÅ.	8.298	8 306	8 300	8 305	8 308	8 313	8 315	8 325	8 313	8 302
Sample No	61	42	43	44	45	46	47	4.9	40	50
Analysis wt %		42				40		40	49	
Man	87	NA	NA	NTA	15.2	12 2	NIA	11.2	10.2	NT A
A1_0_	10.2				11.0	17.6	INA NA	11.2	10.5	IN A
Fa0	10.5				12.0	17.0	15 O	17.0	10.1	
For	19.0		INA NA		15.0	13.0			10.0	20.0
Cr 0	9.0 /0.0	INA NA			5.4	1.2	NA NA	0.2	14.0	INA NA
CL 203	49.2	NA NA	NA NA		51./	40./	NA NA	48.2	38.1	NA NA
B 0	•40	NA NA			.4/	•15	NA	.08	•21	NA
r ₂ 05••••••••	•11	NA NA	NA		•13	.12	NA	•14	•14	NA
110 ₂	.36	NA	NA	NA	.27	.25	NA	.23	./5	NA
N10	.10	NA	NA	NA	.09	•08	NA	•04	.09	NA
ZnO	•09	NA	NA	NA	.05	•08	NA	.10	.23	NA
d-spacingA	2.5120	2.5155	2.5255	2.5260	2.4995	2.4915	2.5090	2.5045	2.4985	2.5005
Unit cellA	8.331	8.343	8.375	8.380	8.290	8.263	8.321	8.306	8.286	8.293
Sample No	51	52								
Analysis, wt %:										
Mg0	15.7	6.2								
A1 ₂ 0 ₃	9.4	8.3								
Fe0	11.0	22.0								
Fe ₂ 0 ₃	8.7	12.8								
Cr ₂ 0 ₃	49.4	48.2								
Si0 ₂	•66	.15								
P ₂ 0 ₅	.08	.15								
Ti0 ₂	•22	.14								
Ni0	•09	.03								
Zn0	.06	.38								
d-spacingÅ	2.5075	2.5095								
Unit cellÅ	Unit cellA 8.316 8.323									
NA Not analyzed.										
¹ All analyses by X-ray flu-										
orescence except FeO which										
was done by H	Reno Re	esearch								
Center using	wet cl	nemical								
methods. X-ray fluorescence										

methods. X-ray fluorescence and X-ray diffraction results by Alaska Field Operations Center, Juneau, AK.

								1		
Sample No	2	3	8	13	14	15	16	1/	18	19
Cation:]]	j	ļ		}			
Fe ²⁺	3.70	3.80	4.44	2.57	3.12	3.42	3.65	3.59	3.24	3.96
$M\sigma^{2+}$	4.30	4.20	3.56	5.43	4,88	4,58	4.35	4.41	4.76	4.04
Fe ³⁺	0.49	1.34	2.55	1 39	1.26	1.03	2.37	0.76	1.87	2.62
c_{r}^{3+}	12 00	7 10	5 13	10 30	11 07	9.35	1. 70	12 34	4 00	0 00
01 ⁻ •••••••••••••	12.00		0.00	10.30	11.07	0.33	4./9	12.54	4.33	1 4 27
AL ⁹⁴	3.51	/.4/	8.32	4.31	3.6/	6.62	8.84	2.90	9.14	4.3/
$RO/R_2O_3^2$	0.88	0.78	0.80	0.73	1.00	0.90	0.92	1.15	0.69	0.78
Molecular ratio: ³)]	J	J	ļ			1
$Fe^{2+}/Fe^{2+}Mg^{2+}$	0.46	0.47	0.55	0.32	0.39	0.43	0.45	0.45	0.41	0.49
Cr^{3+}/Cr^{3+A1}^{3+}	0.77	0.49	0.38	0 71	0.75	0.56	0.35	0.81	0.35	0.67
Sample No	20	21	22	22	25	26	27	20	21	32
			<u> </u>		<u> </u>				<u> </u>	<u> </u>
-2+										
re ²	2.89	1.34	1.98	2.59	2.21	4.13	1.42	2.94	2.38	1.70
Mg ² + • • • • • • • • • • • • • • • • • •	5.11	6.65	6.02	5.41	5.79	3.87	6.58	5.06	5.62	6.30
Fe ³⁺	1.80	6.65	1.82	1.49	1.26	2.76	2.26	0.80	1.14	1.58
Cr ³⁺	10.24	3.67	11.86	9.23	11.68	6.87	10.91	13.95	11.24	12.01
A1 ³⁺	3.96	5.68	2.32	5 27	3 06	6 37	2 83	1 25	3 62	2 41
	3.70	5.00	2.52	5.21	5.00	0.57	2.05	1.25	5.02	2.41
$PO/P = 0^2$	0.00	1 / 0	0.70	1 01	0.00	0.70	0.00	1 17	1	1 10
M_{2}	0.90	1.40	0.72	1.41	0.09	0.78	0.00	1.1/	1.01	1.19
Molecular ratio:										
Fe ²⁺ /Fe ²⁺ Mg ²⁺	0.36	0.17	0.25	0.32	0.27	0.52	0.17	0.37	0.30	0.21
$Cr^{3+}/Cr^{3+}Al^{3+}$	0.72	0.82	0.84	0.64	0.79	0.52	0.79	0.92	0.76	0.83
Sample No	34	35	36	37	38	39	40	41	45	46
Cation:										<u> </u>
F_{0}^{2+}	2 71	2 27	2 60	2 5 6	1. 61	1, 20	5 70	6 4 1	2 50	2 05
M_{π}^{2+}	2./1	1. (2)	3.00	3.30	4.01	4.29	5.79	4.41	2.50	2.05
mg^{-1}	5.29	4.03	4.40	4.44	3.39	3./1	2.21	3.59	5.42	5.15
re	0.45	1.23	0.90	1.47	1.00	0.00	1.78	1.98	1.10	1.37
Cr^{J}	12.75	12.23	12.00	11.76	12.77	12.88	9.53	10.68	11.11	9.36
A1 ⁵⁺	2.80	2.54	3.10	2.76	2.23	3.12	4.69	3.33	3.78	5.26
										ĺ
$RO/R_{2}O_{2}^{2}$	1.54	0.85	0.99	0.77	0.79	1.16	0.78	0.99	1.14	0.97
Molecular ratio: ³				••••			0,,0	0.77	1.11	0.77
$F_0^{2+}/F_0^{2+}M_q^{2+}$	0.34	0 42	0 45	0.22	0 50	0 52	0.70	0 55	0.22	0.25
re / re re re	0.34	0.42	0.43	0.33	0.50	0.55	0.72	0.55	0.32	0.35
	0.82	0.83	0.79	0.81	0.85	0.81	0.67	0.76	0.75	0.64
Sample No	48	49	51	52						
Cation:										
Fe ²⁺	3.68	3.96	2.26	5.33						
Mg ²⁺	4.32	4.04	5.74	2.67						
Fe ³⁺	1.71	2.72	1.85	2.68						
Cr^{3+}	10 58	7 77	11 02	10 50						
λ1 3+	2 70	F F1	2 12	10.00						
AL	3.70	2.21	3.13	2.72						
DO (D. O. 2										
KU/R ₂ U ₃ ²	1.07	0.98	1.15	0.96						
Molecular ratio: ³										
Fe ²⁺ /Fe ²⁺ Mg ²⁺	0.46	0,50	0.28	0.66						
$Cr^{3+}/Cr^{3+}A1^{3+}$	0.74	0.58	0.78	0.80						
Cation proportion	calcul	ations	are ha	sed on						
Stevens (12) Unit a		taina O	p^{2+}	tiona						
Stevens (12). Unit cell contains 8 K ² cations,										

16 R³⁺ cations, and 32 oxygen atoms.
²Ratio based on molecular proportions calculated from data in appendix A.

³Molecular ratios are calculated from data in appendix A and are plotted on figure 5. Fe²⁺/ Fe²⁺ + Mg²⁺ represent values along the x-axis, and $Cr^{3+}/Cr^{3+} + Al^{3+}$ represent values along the y-axis.

Sample	Field	Locality	Mineral terms
	****	LOCALITIES IN INTERIOR ALASKA	
1	WB20438	Avan Hills	ND.
2	WB16537	do	Magnesiochromite, aluminian.
3	WB16538	do	Spinel, chromian.
4	WB20435	do	ND.
5	WB20436	do	ND.
6	WB20437	••••do••••••	ND.
7	WB16539	••••do••••••	ND.
8	WB16540	••••do••••••	Hercynite, chromian.
9	WB16541	do	ND.
10	WB20825	do	ND.
11	WB20826	do	ND.
12	WB20827	••••do••••••	ND.
13	WB16768	do	Magnesiochromite, aluminian.
14	PT16641	Caribou Mountain	Do.
15	PT16637	••••do••••••	Do.
16	PB15787	••••do••••••	Spinel, chromian.
1/	PT16635	Kanuti River	Magnesiochromite, ferroan.
18	PT16638	••••do••••••••••••••••••••••••	Spinel, chromian.
19	PT16639	••••do••••••••••••••••••••••••	Magnesiochromite, aluminian.
20	P110030	Valad Desar	Do.
21	KW20700	iuki kiver	Magnesiochromite, ferrian.
22	KW20702		Magnesiochromite, aluminian.
23	NW20779	Mount Ilungt	NO •
24	NW194/0		ND.
25	KW194//	do	Magnesiochromite, aluminian.
27	KW19000	do	Magnagiachromita aluminian
	LOCALITIE	S ALONG EXTENSION OF CHUCACH MO	HAGHESTOCHTOMICE, aldminian.
28	CM20261	Grant Lagoon	Magnesiochromite ferroan
29	СМ20268	Halibut Bay	ND.
30	СМ19277	••••do	ND.
31	СМ18624	do	Magnesiochromite, aluminian.
32	СМ19649	do	Do.
33	CM11168	do	ND.
34	СМ19641	Miners Bay	Magnesiochromite, ferroan.
35	СМ17679	Claim Point	Do.
36	СМ19680	••••do••••••	Do.
37	СМ17675	Red Mtd-Kenai	Do.
38	СМ17670	••••do••••••	Chromite, magnesian.
39	СМ17670	••••do••••••	Do.
40	СМ19312	Eklutna	Do.
41	СМ19322	Wolverine Complex	Do.
42	СМ19326	••••do•••••••	ND.
43	CM19371	••••do•••••••	ND.
44	СМ19373	do	ND.
45	СМ20488	Tonsina-Bernard	Magnesiochromite, aluminian.
46	CM20497	do	Do.
4/	CM186/8	do	ND.
Σö	CM20467	Sheep Hill	Magnesiochromite, ferroan.
49	UM20466	•••do•••••	Do.
51	CM2U443	Dust Mountain	ND.
52	15155	ked Blutt Bay	Magnesiochromite, aluminian.

APPENDIX C.--MINERAL TERMS AND SAMPLE KEY

52.....2S430.....Do.NDInsufficient data to designate mineral term (no chemical data).