

Energy, Mines and Énergie, Mines et Resources Canada Ressources Canada Geological Survey of Canada Commission gélogique du Canada Resource Geophysics and Geochemistry Division



Province of British Columbia Ministry of Energy, Mines and Petroleum Resources Geological Division Mineral Resources Branch

NATIONAL GEOCHEMICAL RECONNAISSANCE 1:250 000 MAP SERIES

Bonaparte Lake, British Columbia

(N.T.S. 92P)

GSC Open File 775

BC RGS 4

REGIONAL GEOCHEMICAL SURVEY, BRITISH COLUMBIA, 1979 NTS 92P, BC RGS-4, 1979

BC RGS-4, 1979 GSC OPEN FILE 775

Page

INTRODUCTION	
DATA LEGEND	1
DATA LIST	
SUMMARY STATISTICS	20

AVAILABLE FOR REFERENCE AT:

- ALL LIBRARIES OF THE GEOLOGICAL SURVEY OF CANADA
- BRITISH COLUMBIA MINISTRY OF ENERGY, MINES AND PETROLEUM RESOURCES, VICTORIA, BRITISH COLUMBIA

AVAILABLE FOR SALE AT:

- BRITISH COLUMBIA MINISTRY OF ENERGY, MINES AND PETROLEUM RESOURCES, VICTORIA, BRITISH COLUMBIA AS BC RGS-4

REGIONAL GEOCHEMICAL SURVEYS 3 AND 4 WERE MANAGED AND FUNDED BY THE BRITISH COLUMBIA MINISTRY OF ENERGY, MINES AND PETROLEUM RESOURCES; DATA MANAGEMENT BY THE GEOLOGICAL SURVEY OF CANADA.

REGIONAL STREAM SEDIMENT AND WATER GEOCHEMICAL RECONNAISSANCE DATA, BRITISH COLUMBIA, 1979

BC RGS-4, GEOLOGICAL SURVEY OF CANADA OPEN FILE 775, NTS 92P

THE 1979 RECONNAISSANCE SURVEY WAS UNDERTAKEN BY THE BRITISH COLUMBIA MINISTRY OF ENERGY, MINES AND PETROLEUM RESOURCES USING THE METHODS OF SAMPLE COLLECTION AND ANALYSES ESTABLISHED BY THE GEOLOGICAL SURVEY OF CANADA DURING THE FEDERAL -PROVINCIAL JOINT URANIUM RECONNAISSANCE PROGRAM.

CONTRACTS LET FOR SAMPLE COLLECTION, PREPARATION, AND ANALYSES WERE SUPERVISED AND/OR MONITORED BY STAFF OF THE MINISTRY AS FOLLOWS:

- COLLECTION STAFF AND EQUIPMENT PROVIDED BY BEMA INDUSTRIES LTD., LANGLEY.
- SUPERVISION T.E. KALNINS, N.C. CARTER, W.M. JOHNSON.
- PREPARATION KAMLOOPS RESEARCH ASSAY AND LABORATORY LTD.
- SUPERVISION W.M. JOHNSON.
- ANALYTICAL CHEMEX LABS. LTD., NORTH VANCOUVER (STREAM SEDIMENTS). MINEN LABORATORIES LTD., NORTH VANCOUVER (STREAM WATERS). NOVATRACK ANALYSTS LIMITED, VANCOUVER (URANIUM IN STREAM SEDIMENTS).

SUPERVISION - W.M. JOHNSON.

THE GEOLOGICAL SURVEY OF CANADA WAS RESPONSIBLE FOR DATA MANAGEMENT.

FEDERAL LIAISON WAS WITH E.H.W. HORNBROOK AND S.B. BALLANTYNE, RESOURCE GEOCHEMISTRY SUBSECTION, GEOLOGICAL SURVEY OF CANADA.

REGIONAL STREAM SEDIMENT AND WATER GEOCHEMICAL RECONNAISSANCE DATA, BRITISH COLUMBIA, 1979

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STREAM SEDIMENT AND WATER SAMPLES WERE COLLECTED AT AN AVERAGE DENSITY OF ONE SAMPLE PER 14 SQUARE KILOMETRES (5 SQUARE MILES) THROUGHOUT THE 30 000-SQUARE-KILOMETRE (11,600-SQUARE-MILE) TOTAL SURVEY AREA. THE HELICOPTER AND 4-WHEEL-DRIVE TRUCK-SUPPORTED SAMPLE COLLECTION WAS CARRIED OUT DURING THE SUMMER OF 1979.

ONE FIELD DUPLICATE SAMPLE WAS ROUTINELY COLLECTED IN EACH ANALYTICAL BLOCK OF 20 SAMPLES.

FIELD DRIED SAMPLES WERE AIR DRIED AND THE MINUS 80 MESH (177 MICRONS) FRACTION WAS OBTAINED FOR SUBSEQUENT ANALYSES.

FOR CONTROL, A SAMPLE BLOCK OF 20 SILT SAMPLES INCLUDES THE FIELD DUPLICATE AND ONE CONTROL REFERENCE AND ONE BLIND DUPLICATE SAMPLE. IN EACH ANALYTICAL BLOCK OF 20 WATER SAMPLES THE CONTROL REFERENCE AND BLIND DUPLICATE SAMPLE POSITIONS WERE FILLED WITH CONTROL REFERENCE WATER SAMPLES. THE SAMPLE PREPARATION LABORATORY IS RESPONSIBLE FOR INSERTION OF CONTROL SAMPLES.

THE DETERMINATION OF ZN, CU, PB, NI, CO, AG, MN, FE, MO, AS, W, AND HG IN STREAM SEDIMENTS WAS CARRIED OUT BY CHEMEX LABS LTD.

THE DETERMINATION OF U IN STREAM SEDIMENTS WAS CARRIED OUT BY NOVATRACK ANALYSTS LIMITED.

THE DETERMINATION OF U, F, AND pH IN STREAM WATERS WAS CARRIED OUT BY MINEN LABORATORIES LTD.

FOR THE DETERMINATION OF ZN, CU, PB, NI, CO, AG, MN, AND FE, A 1-GRAM SAMPLE WAS REACTED WITH 3 ML OF CONCENTRATED HNO₃ IN A TEST TUBE FOR 30 MINUTES AT 90° C. AT THIS POINT, 1 ML CONCENTRATED HCL WAS ADDED AND THE DIGESTION WAS CONTINUED AT 90° C FOR AN ADDITIONAL 90 MINUTES. THE SAMPLE SOLUTION WAS THEN DILUTED TO 20 ML WITH METAL-FREE WATER AND MIXED. ZN, CU, PB, NI, CO, AG, MN, AND FE WERE DETERMINED BY ATOMIC ABSORPTION SPECTROSCOPY USING AN AIR-ACETYLENE FLAME. BACKGROUND CORRECTIONS WERE MADE FOR PB, NI, CO, AND AG.

MOLYBDENUM WAS DETERMINED BY ATOMIC ABSORPTION SPECTROSCOPY USING A NITROUS OXIDE-ACETYLENE FLAME. A 0.5-GRAM SAMPLE WAS REACTED WITH 1.5 ML CONCENTRATED HNO₃ AT 90° C FOR 30 MINUTES. AT THIS POINT 0.5 ML CONCENTRATED HCL WAS ADDED AND THE DIGESTION WAS CONTINUED AT 90° C FOR AN ADDITIONAL 90 MINUTES. AFTER COOLING, 8 ML OF 1 250 PPM AL SOLUTION WERE ADDED AND THE SAMPLE SOLUTION WAS DILUTED TO 10 ML BEFORE ASPIRATION. MERCURY WAS DETERMINED BY THE HATCH AND OTT PROCEDURE WITH SOME MODIFICATIONS. THE METHOD IS DESCRIBED BY JONASSON, ET AL. (1973).

A 0.5-GRAM SAMPLE WAS REACTED WITH 20 ML CONCENTRATED HNO_3 AND 1 ML CONCENTRATED HCL IN A TEST TUBE FOR 10 MINUTES AT ROOM TEMPERATURE PRIOR TO 2 HOURS OF DIGESTION WITH MIXING AT 90° C IN A HOT WATER BATH. AFTER DIGESTION, THE SAMPLE SOLUTIONS WERE COOLED AND DILUTED TO 100 ML WITH METAL-FREE WATER. THE HG PRESENT WAS REDUCED TO THE ELEMENTAL STATE BY THE ADDITION OF 10 ML W/V SNSO₄ IN M H₂SO₄. THE HG VAPOUR WAS THEN FLUSHED BY A STREAM OF AIR INTO AN ABSORPTION CELL MOUNTED IN THE LIGHT PATH OF AN ATOMIC ABSORPTION SPECTROPHOTOMETER. ABSORPTION MEASUREMENTS WERE MADE AT 253.7 NM.

ARSENIC WAS DETERMINED BY HYDRIDE GENERATION/ATOMIC ABSORPTION SPECTROSCOPY ON AN ALIQUOT TAKEN FROM THE SAMPLE PREPARED FOR THE BASE METAL ANALYSES.

TUNGSTEN WAS DETERMINED COLOURIMETRICALLY SUBSEQUENT TO A PYROSULFATE FUSION AND A DITHIOLCARBONATE COMPLEXING FOR THE GENERATION OF THE COLOUR. URANIUM WAS DETERMINED USING A NEUTRON ACTIVATION METHOD WITH DELAYED NEUTRON COUNTING.

A DETAILED DESCRIPTION OF THE METHOD IS PROVIDED BY BOULANGER, ET AL. (1975). IN BRIEF, A 1-GRAM SAMPLE IS WEIGHED INTO A 7-DRAM POLYETHYLENE VIAL, CAPPED AND SEALED.

THE IRRADIATION IS PROVIDED BY THE TRIUMF CYCLOTRON WITH AN OPERATING FLUX OF 10^{11} NEUTRONS PER SQUARE CENTIMETRE PER SECOND.

THE SAMPLES ARE PNEUMATICALLY TRANSFERRED FROM AN AUTOMATIC LOADER TO THE REACTOR, WHERE EACH SAMPLE IS IRRADIATED FOR 60 SECONDS.

AFTER IRRADIATION, THE SAMPLE IS AGAIN TRANSFERRED PNEUMATICALLY TO THE COUNTING FACILITY WHERE AFTER A 20-SECOND DELAY THE SAMPLE IS COUNTED FOR 60 SECONDS WITH SIX BF3 DETECTOR TUBES EMBEDDED IN PARAFFIN.

FOLLOWING COUNTING, THE SAMPLES ARE AUTOMATICALLY EJECTED INTO A SHIELDED STORAGE CONTAINER.

CALIBRATION IS CARRIED OUT EVERY 40 SAMPLES AS A MINIMUM USING NATURAL MATERIALS OF KNOWN URANIUM CONCENTRATION.

URANIUM WAS DETERMINED IN THE WATER SAMPLES BY A FLUOROMETRIC METHOD. THE URANIUM WAS INITIALLY PRECONCENTRATED BY EVAPORATION. THE RESIDUE AFTER EVAPORATION WAS FUSED WITH A MIXTURE OF NA_2CO_3 , K_2CO_3 , AND NAF IN A PLATINUM DISH. AFTER COOLING THE FLUORESCENCE OF THE FUSED PELLET WAS MEASURED USING A TURNER FLUOROMETER MODEL 111.

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FLUORIDE IN STREAM WATER SAMPLES WAS DETERMINED USING A SPECIFIC ION ELECTRODE. AN ALIQUOT OF THE SAMPLE WAS MIXED WITH AN EQUAL VOLUME OF A TISAB SOLUTION (TOTAL IONIC STRENGTH ADJUSTMENT BUFFER). THE FLUORIDE WAS MEASURED USING A CORNING 101 ELECTROMETER WITH AN ORION FLUORIDE ELECTRODE.

FOR THE DETERMINATION OF pH AN ALIQUOT OF THE WATER SAMPLE WAS TRANSFERRED TO A CLEAN DRY BEAKER. THE pH WAS MEASURED USING A FISHER ACCUMET pH METER.

ON RECEIPT, FIELD AND ANALYTICAL DATA WERE PUNCHED ONTO 80-COLUMN CARDS AND ALL SUBSEQUENT PROCESSING WAS CARRIED OUT WITH THE AID OF COMPUTERS. THE FIELD DATA WERE RECORDED BY THE FIELD CONTRACT STAFF ONTO STANDARD GEOCHEMICAL STREAM WATER AND SEDIMENT SAMPLE FIELD CARDS BASED ON FORM REV. 77 USED BY THE GEOLOGICAL SURVEY OF CANADA. THE SAMPLE SITE POSITIONS WERE SELECTED ON 1:50 000 BASE MAPS AND TRANSFERRED TO APPROPRIATE 1:250 000-SCALE NTS MAPS IN THE FIELD.

THESE MAPS WERE DIGITIZED AT THE GEOLOGICAL SURVEY IN OTTAWA TO OBTAIN THE SAMPLE SITE UTM COORDINATES.

THE DOMINANT ROCK TYPES IN THE STREAM CATCHMENT BASINS WERE IDENTIFIED BY T.E. KALNINS ON A GEOLOGICAL MAP WITH A MODIFIED LEGEND COMPILED BY T.E. KALNINS AND S.B. BALLANTYNE FOR THE NGS GEOCHEMICAL MAPS AND GEOLOGICAL SOURCES ARE GIVEN IN THE REFERENCES.

THE ANALYTICAL DATA WERE RECORDED AS FOLLOWS (SEE GARRETT, 1974, FOR DETAILS) AND ALSO FOR CONVENIENCE THE DETECTION LIMITS OF THE ANALYTICAL METHODS USED ARE ALSO GIVEN:

	ELEMENT	DETECTION LIMIT (D.L.)	VALUE REPORTED IF LESS THAN D.L.
SEDIMENT			
	ZN	2	1
	CU	2	1
	PB	2	1
	NI	2	1
	CO	2	1
	AG	0.2	0.1
	MN	5	2
	FE	0.02	0.01
	AS	1	1
	MO	2	1
	W	2	1
	HG PPB	10	5
	U	0.2	0.1
WATER			
	U PPB	0.05	0.02
	F PPB	20	10

UNLESS OTHERWISE NOTED THE UNITS OF MEASUREMENT FOR THE ANALYSES ARE PPM. THE FIGURES TO WHICH VALUES WERE SET IF THEY FELL BELOW THE DETECTION LIMIT WERE CHOSEN ARBITRARILY.

DETAILED INSPECTIONS OF THE FIELD AND ANALYTICAL DATA WERE MADE TO CHECK FOR ANY MISSING INFORMATION AND/OR GROSS ERRORS.

THE SAMPLE SITE COORDINATES WERE CHECKED BY PLOTTING SAMPLING LOCATION MAPS ON A FLAT BED PLOTTER FROM THE DIGITIZED COORDINATES AND THEN OVERLYING THESE OVER THE FIELD CONTRACTOR'S SAMPLE LOCATION BASE MAPS.

QUALITY CONTROL AND MONITORING OF THE ANALYTICAL DATA WAS BASED ON CONTROL REFERENCE AND BLIND DUPLICATE SAMPLES AND CRITERIA STIPULATED IN THE ANALYTICAL CONTRACTS.

REFERENCES

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CAMPBELL, R.B. AND TIPPER, H.W. (1971) GEOLOGY OF BONAPARTE LAKE MAP-AREA, BRITISH COLUMBIA, GEOL. SURV., CANADA, MEM. 363, MAP 1278A (92P).

- GARRETT, ROBERT G. (1974): FIELD DATA ACQUISITION METHODS FOR APPLIED GEOCHEMISTRY SURVEYS AT THE GEOLOGICAL SURVEY OF CANADA, GEOL. SURV., CANADA, PAPER 74-52.
- JONASSON, I.R., LYNCH, J.J., AND TRIP, L.J. (1973): FIELD AND LABORATORY METHODS USED BY THE GEOLOGICAL SURVEY OF CANADA IN GEOCHEMICAL SURVEYS, NO. 12, MERCURY IN ORES, ROCKS, SOILS, SEDIMENTS, AND WATER, GEOL. SURV., CANADA, PAPER 73-21.
- TIPPER, H.W. (1978): GEOLOGY OF TASEKO LAKES MAP-AREA, BRITISH COLUMBIA, GEOL. SURV., CANADA, OPEN FILE 534, MAP NO. 1-125000 (920).

DATA LIST LEGEND

MAP-	NATIONAL TOPOGRAPHIC SYSTEM (NTS) - LETTERED SIXTEENTH (SCALE 1:50000). PART OF SAMPLE NUMBER
SAMPLE-	REMAINDER OF SAMPLE NUMBER - YEAR (2), FIELD CREW (1), SAMPLE SEQUENCE NUMBER (3)
UTM COORDINATES-	UNIVERSAL TRANSVERSE MERCATOR (UTM) COORDINATE SYSTEM - SAMPLE COORDINATES
ZN- EAST- NORTH-	ZONE EASTING (METERS) NORTHING (METERS)
ROCK TYPE- Age-	MAJOR ROCK TYPE OF CATCHMENT AREA STRATIGRAPHIC AGE OF ROCK TYPE
WD- DT-	WIDTH OF STREAM (DECIMETER) AT THE SAMPLE SITE DEPTH OF STREAM SAMPLED TO NEAREST DECIMETER
SAMP-	TYPE OF MATERIAL SAMPLED
RP ST-	REPLICATE STATUS - RELATIONSHIP OF SAMPLE WITH RESPECT TO OTHERS WITHIN THE SURVEY
CONT-	CONTAMINATION
BANK-	BANK TYPE
WCOL-	WATER COLOUR AND SUSPENDED LOAD
RATE-	WATER FLOW RATE
SCOL-	PREDOMINANT SEDIMENT COLOUR
SMP CMP-	SAMPLE COMPOSITION - BULK MECHANICAL COMPOSITION OF SAND, FINES, ORGANICS RESPECTIVELY
PPPS-	PRECIPITATE OR STAIN ON SEDIMENTS AT SAMPLE SITE
PRPB-	DISTINCTIVE PRECIPITATE, STAIN, WEATHERING, BLOOMS ON ROCKS IN THE IMMEDIATE CATCHMENT AREA
PHYS-	GENERAL PHSYIOGRAPHY
PATT-	DRAINAGE PATTERN
TYPE-	STREAM TYPE
CLSS-	STREAM CLASS
SRCE-	SOURCE OF WATER

DATA LIST LEGEND (CONT'D)

ROCK TYPE:	TILL- SLSN- BSLT- LNSN- DCIT-	ANDESITE TILLITE SILTSTONE BASALT LIMESTONE = CALCILUTITE DACITE GRANITE
AGE:	30- 32- 36- 41- 42-	SILURIAN-DEVONIAN CARBONIFEROUS CARBONIFEROUS MIDDLE PERMIAN MESOZOIC UNDIVIDED TRIASSIC TRIASSIC MIDDLE
SAMP:		STREAM BED SEDIMENT SIMULTANEOUS STREAM WATER AND SEDIMENT
RP ST:	10-	ROUTINE REGIONAL SAMPLE FIRST OF FIELD DUPLICATE SECOND OF FIELD DUPLICATE
CONT:	1- 2-	NONE POSSIBLE PROBABLE FOREST ACTIVITY
BANK:	1- 2- 3- 5- 6-	UNDEFINED UNCONSOLIDATED MATERIAL ALLUVIAL COLLUVIAL (RESIDUAL AND MOUNTAIN SOILS) GLACIAL TILL, TILLITE BARE ROCK TALUS, SCREE ORGANIC PREDOMINANT (DEBRIS, PEAT, MUSKEG, SWAMP)
WCOL:	1- 2-	COLOUR BROWN TRANSPARENT WHITE CLOUDY BROWN CLOUDY

	1- SLOW 2- MODERATE 3- FAST 4- TORRENTIAL
SCOL:	1- RED, BROWN 2- WHITE, BUFF 3- BLACK 6- GREY, BLUE-GREY
SMP CMP:	0- ABSENT 1- MINOR <33% 2- MEDIUM 33-67% 3- MAJOR >67%
PRPS:	0- NONE 1- RED, BROWN 2- WHITE
PRPB:	0- FEATURELESS 1- RED, BROWN 2- WHITE, BUFF 3- BLACK 4- YELLOW
PHYS:	 2- PENEPLAIN, PLATEAU 3- HILLY, UNDULATING 4- MOUNTAINOUS MATURE 5- MOUNTAINOUS YOUTHFUL (PRECIPITOUS)
PATT:	 0- POORLY DEFINED, HAPHAZARD 1- DENDRITIC 2- HERRING BONE 3- RECTANGULAR (JOINTED) 5- DISCONTINUOUS SHEILD TYPE (CHAINS OF LAKES, SWAMPS)
TYPE:	0- UNDEFINED 1- PERMANENT, CONTINUOUS 2- INTERMITTENT, SEASONAL 3- RE-EMERGENT, DISCONTINUOUS
CLSS:	2- SECONDARY 3- TERTIARY 4- QUATERNARY
SRCE:	0- UNKNOWN 1- GROUNDWATER

RATE: 0- ZERO

DATA LIST LEGEND (CONT'D)

ZN- ZINC BY ATOMIC ABSORPTION SPECTROSCOPY (PPM) CU- COPPER BY ATOMIC ABSORPTION SPECTROSCOPY (PPM) PB- LEAD BY ATOMIC ABSORPTION SPECTROSCOPY (PPM) NI- NICKEL BY ATOMIC ABSORPTION SPECTROSCOPY (PPM) CO- COBALT BY ATOMIC ABSORPTION SPECTROSCOPY (PPM) AG- SILVER BY ATOMIC ABSORPTION SPECTROSCOPY (PPM) MN- MANGANESE BY ATOMIC ABSORPTION SPECTROSCOPY (PPM) FE- IRON BY ATOMIC ABSORPTION SPECTROSCOPY (PCT) AS- ARSENIC BY FLAMELESS SPECTROSCOPY (PPM) MO- MOLYBDENUM BY ATOMIC ABSORPTION SPECTROSCOPY (PPM) W- TUNGSTEN BY COLORIMETRY USING DITHIOL (PPM) HG- MERCURY BY FLAMELESS SPECTROSCOPY (PPB) U- URANIUM BY DELAYED NEUTRON ACTIVATION (PPM) U-W- URANIUM IN WATERS FLUOROMETRICALLY (PPB) F-W- FLUORINE IN WATERS BY SPECIFIC ION ELECTRODE (PPB) PH- PH BY COMBINATION GLASS - CALOMEL ELECTRODE