REGIONAL LAKE SEDIMENT AND WATER GEOCHEMICAL DATA, MANITOBA 1986, GSC OF1358, NGR 94-1986, NTS 53L, PARTS OF 53E, 53F, 53K, AND 53M

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GEOLOGICAL SURVEY OF CANADA OPEN FILE 1358. REGIONAL LAKE SEDIMENT AND WATER GEOCHEMICAL RECONNAISSANCE DATA, EAST-CENTRAL MANITOBA NTS, 53L AND PARTS OF 53E, 53F, 53K, AND 53M.

THE RECONNAISSANCE SURVEY WAS UNDERTAKEN BY THE GEOLOGICAL SURVEY OF CANADA IN CONJUNCTION WITH THE MANITOBA DEPARTMENT OF ENERGY AND MINES, MINERALS DIVISION UNDER THE CANADA-MANITOBA MINERAL DEVELOPMENT AGREEMENT (1984-1989).

E.H.W. HORNBROOK DIRECTED THE SURVEY PROGRAM.

P.W.B. FRISKE COORDINATED THE OPERATIONAL ACTIVITIES OF THE CONTRACTING AND GEOLOGICAL SURVEY OF CANADA STAFF THROUGHOUT THE SURVEY.

CONTRACTS LET FOR COLLECTION, SAMPLE PREPARATION AND ANALYSIS WERE THE RESPONSIBILITY OF, AND WERE SUPERVISED AND/OR MONITORED BY THE STAFF OF THE EXPLORATION GEOCHEMISTRY SUBDIVISION AS FOLLOWS:

COLLECTION: - SIAL COMPAGNIE INTERNATIONALE DE GEOPHYSIQUE INC., DORVAL, QUEBEC.

- E.H.W. HORNBROOK, H.R. SCHMITT

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ANALYSIS: - BONDAR CLEGG AND COMPANY LTD., OTTAWA

- BARRINGER MAGENTA LABORATORIES (ALBERTA) LTD.,

CALGARY, ALBERTA (WATERS)

- CHEMEX LABS. LTD, NORTH VANCOUVER, B.C. (GOLD)

- J.J. LYNCH

H.R. SCHMITT COORDINATED OPEN FILE PRODUCTION.

A.C GALLETTA MANAGED THE DIGITAL GEOCHEMICAL DATA AND PROVIDED COMPUTER PROCESSING SUPPORT.

D.J. ELLWOOD DEVELOPED SOFTWARE TO RASTER PLOT OPEN FILE VALUE, SYMBOL AND REGIONAL TREND MAPS. THE PLOTTING WAS DONE BY CANADA LANDS DATA SYSTEMS STAFF AT ENVIRONMENT CANADA, HULL QUEBEC.

M. MCCURDY AND S. COOK PROCESSED INCOMING AND OUTGOING MATERIALS, SUPPLIES AND SAMPLES.

COMPUTING, PLOTTING AND OPEN FILE TEXT LASER PRINTING SERVICES, WERE PROVIDED BY THE COMPUTER SCIENCE CENTER, E.M.R.

J. YELLE AND F. WILLIAMS OF THE GEOLOGICAL INFORMATION DIVISION SUPERVISED THE PREPARATION OF OPEN FILE MAPS BY CARTOGRAPHY UNIT A-2. HELICOPTER SUPPORTED SAMPLE COLLECTION WAS CARRIED OUT DURING THE SUMMER OF 1986.

LAKE SEDIMENT AND WATER SAMPLES WERE COLLECTED AT AN AVERAGE DENSITY OF ONE SAMPLE PER 11.4 SQUARE KILOMETERS THROUGHOUT THE 39,000 SQUARE KILOMETERS OF THE EAST-CENTRAL MANITOBA SURVEY AREA.

SAMPLE SITE DUPLICATE SAMPLES WERE ROUTINELY COLLECTED IN EACH ANALYTICAL BLOCK OF TWENTY SAMPLES.

IN OTTAWA, FIELD DRIED SAMPLES WERE AIR-DRIED, CRUSHED, BALL MILLED AND SIEVED. THE MINUS 80 MESH (177 MICRONS) FRACTION WAS USED FOR SUBSEQUENT ANALYSES. AT THIS TIME, CONTROL REFERENCE AND BLIND DUPLICATE SAMPLES WERE INSERTED INTO EACH BLOCK OF TWENTY SEDIMENT SAMPLES. FOR THE WATER SAMPLES, ONLY CONTROL REFERENCE SAMPLES WERE INSERTED INTO THE BLOCK. THERE WERE NO BLIND DUPLICATE WATER SAMPLES.

ON RECEIPT, FIELD AND ANALYTICAL DATA WERE PROCESSED WITH THE AID OF COMPUTERS THE FIELD DATA WERE RECORDED BY THE FIELD CONTRACT STAFF ON STANDARD LAKE SEDIMENT FIELD CARDS (REV. 74) USED BY THE GEOLOGICAL SURVEY OF CANADA (GARRETT, 1974).

THE SAMPLE SITE POSITIONS WERE MARKED ON 1/100,000 SCALE PROVINCIAL MAPS IN THE FIELD AND TRANSFERRED TO APPROPRIATE 1/250,000 SCALE NTS MAPS IN OTTAWA THESE MAPS WERE DIGITIZED AT THE GEOLOGICAL SURVEY IN OTTAWA TO OBTAIN THE SAMPLE SITE UTM COORDINATES, WHICH WERE VERIFIED BY D. SCHOLTZ.

THE SAMPLE SITE COORDINATES WERE CHECKED AS FOLLOWS: A SAMPLE LOCATION MAP WAS PRODUCED ON A CALCOMP 1051 DRUM PLOTTER USING THE DIGITIZED COORDINATES; THE FIELD CONTRACTORS'S SAMPLE LOCATION MAP WAS THEN OVERLAYED WITH THE CALCOMP MAP; THE TWO SETS OF POINTS WERE CHECKED FOR COINCIDENCE. THE DOMINANT ROCK TYPES IN THE LAKE CATCHMENT BASINS WERE IDENTIFIED ON APPROPRIATE GEOLOGICAL MAPS USED AS THE BEDROCK GEOLOGICAL BASE ON RGR MAPS.

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

QUALITY CONTROL AND MONITORING OF THE GEOCHEMICAL DATA WAS UNDERTAKEN BY A STANDARD METHOD USED BY THE EXPLORATION GEOCHEMISTRY SUBDIVISION AT THE GEOLOGICAL SURVEY OF CANADA.

FOR THE DETERMINATION OF ZN, CU, PB, NI, CO, AG, MN, FE, CD, AND AS A 1 GRAM SAMPLE WAS REACTED WITH 6 ML OF A MIXTURE OF 4M HNO3 AND M HCL IN A TEST-TUBE OVERNIGHT AT ROOM TEMPERATURE.

AFTER DIGESTION, THE TEST-TUBE WAS IMMERSED IN A HOT WATER BATH AT ROOM TEMPERATURE AND BROUGHT UP TO 90C AND HELD AT THIS TEMPERATURE FOR 2 HOURS WITH PERIODIC SHAKING.

THE SAMPLE SOLUTION WAS THEN DILUTED TO 20 ML WITH METAL FREE WATER AND MIXED. ZN, CU, PB, NI, CO, AG, MN, FE AND CD WERE DETERMINED BY ATOMIC ABSORPTION SPECTROSCOPY USING AN AIR-ACETYLENE FLAME.

BACKGROUND CORRECTIONS WERE MADE FOR PB, NI, CO, AG AND CD.
AS WAS DETERMINED BY ATOMIC ABSORPTION USING A HYDRIDE EVOLUTION METHOD WHEREIN THE HYDRIDE (ASH3) IS EVOLVED, PASSED THROUGH A HEATED QUARTZ TUBE IN THE LIGHT PATH OF AN ATOMIC ABSORPTION SPECTROPHOTOMETER.
THE METHOD IS DESCRIBED BY ASLIN (1976).

MOLYBDENUM AND VANADIUM WERE DETERMINED BY ATOMIC ABSORPTION SPECTROSCOPY USING A NITROUS OXIDE ACETYLENE FLAME.

A 0.5 GRAM SAMPLE WAS REACTED WITH 1.5 ML CONCENTRATED HN03 AT 90C FOR 30 MINUTES.

AT THIS POINT 0.5 ML CONCENTRATED HCL WAS ADDED AND THE DIGESTION WAS CONTINUED AT 90C FOR AN ADDITIONAL 90 MINUTES.

AFTER COOLING, 8 ML OF 1250 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 HNO3 AND 1 ML CONCENTRATED HCL IN A TEST-TUBE FOR 10 MINUTES AT ROOM TEMPERATURE PRIOR TO 2 HOURS OF DIGESTION WITH MIXING AT 90C 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 10% 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.

LOSS ON IGNITION WAS DETERMINED USING A 500 MG SAMPLE.

THE SAMPLE, WEIGHED INTO 30 ML BEAKER, WAS PLACED IN A COLD MUFFLE FURNACE AND BROUGHT UP TO 500C OVER A PERIOD OF 2-3 HOURS.

THE SAMPLE WAS LEFT AT THIS TEMPERATURE FOR 4 HOURS, THEN ALLOWED TO COOL TO ROOM TEMPERATURE FOR WEIGHING.

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 WAS WEIGHED INTO A 7 DRAM POLYETHYLENE VIAL, CAPPED AND SEALED.

THE IRRADIATION WAS PROVIDED BY THE SLOWPOKE REACTOR WITH AN OPERATING FLUX OF 5** 10 NEUTRONS/SO.CM./SEC.

THE SAMPLES WERE PNEUMATICALLY TRANSFERRED FROM AN AUTOMATIC LOADER TO THE REACTOR, WHERE EACH SAMPLE WAS IRRADIATED FOR 20 SECONDS. AFTER IRRADIATION, THE SAMPLE WAS AGAIN TRANSFERRED PNEUMATICALLY TO THE COUNTING FACILITY WHERE AFTER A 10 SECOND DELAY THE SAMPLE WAS COUNTED FOR 20 SECONDS WITH SIX HELIUM DETECTOR TUBES EMBEDDED IN PARAFFIN.

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

CALIBRATION WAS CARRIED OUT ONCE A DAY AS A MINIMUM, USING NATURAL MATERIALS OF KNOWN URANIUM CONCENTRATION.

FLUORINE WAS DETERMINED IN LAKE SEDIMENTS AS DESCRIBED BY FICKLIN (1970). A 250 MG SAMPLE IS SINTERED WITH 1 GRAM OF A FLUX CONSISTING OF TWO PARTS BY WEIGHT SODIUM CARBONATE AND 1 PART BY WEIGHT POTASSIUM NITRATE. THE RESIDUE IS THEN LEACHED WITH WATER, THE SODIUM CARBONATE IS NEUTRALIZED WITH 10 ML 10% (W/V) CITRIC ACID AND THE RESULTING SOLUTION IS DILUTED TO 100 ML WITH WATER.

THE PH OF THE RESULTING SOLUTION SHOULD BE FROM 5.5 TO 6.5.
THE FLUORIDE CONTENT OF THE TEST SOLUTION IS THEN MEASURED USING A FLUORIDE ION ELECTRODE.

STANDARD SOLUTIONS CONTAIN SODIUM CARBONATE AND CITRIC ACID IN THE SAME QUANTITIES AS THE SAMPLE SOLUTION.
A DETECTION LIMIT OF 40 PPM IS ACHIEVED.

ANTIMONY WAS DETERMINED IN LAKE SEDIMENTS AS DESCRIBED BY (ASLIN,1976). A 500 MG SAMPLE IS PLACED IN A TEST TUBE; 3 ML CONCENTRATED HNO3 AND 9 ML CONCENTRATED HCL ARE ADDED AND THE MIXTURE IS ALLOWED TO STAND OVERNIGHT AT ROOM TEMPERATURE. THE MIXTURE IS HEATED SLOWLY TO 90C AND MAINTAINED AT THIS TEMPERATURE FOR AT LEAST 90 MINUTES. THE SOLUTION IS COOLED AND DILUTED TO 10 ML. A 400 MICRO L ALIQUOT OF THIS TEST SOLUTION IS REMOVED AND DILUTED TO 10 ML WITH 1.8M HCL.THE ANTIMONY IN AN ALIQUOT OF THIS DILUTE SOLUTION IS THEN DETERMINED BY HYDRIDE EVOLUTION—ATOMIC ABSORPTION SPECTROMETRY.

GOLD WAS USUALLY DETERMINED ON A 10 GRAM LAKE SEDIMENT SAMPLE;
DEPENDING ON THE AMOUNT OF SAMPLE AVAILABLE, LESSER
WEIGHTS WERE SOMETIMES USED. THIS RESULTED IN A VARIABLE DETECTION
LIMIT: 2 PPB FOR A 5 GRAM SAMPLE, 1 PPB FOR A 10 GRAM SAMPLE... THE SAMPLE WAS
FUSED TO PRODUCE A LEAD BUTTON, COLLECTING ANY GOLD IN THE SAMPLE, WHICH
WAS CUPELLED IN A MUFFLE FURNACE TO PRODUCE A SILVER (DORE) BEAD. THE
SILVER BEADS WERE IRRIATED IN A NEUTRON FLUX FOR 1 HOUR, COOLED FOR 4
HOURS, AND COUNTED BY GAMMA RAY SPECTROMETRY.CALIBRATION WAS CARRIED OUT
USING STANDARD AND BLANK BEADS.

FLUORIDE IN LAKE WATER SAMPLES WAS DETERMINED USING A FLUORIDE ELECTRODE. PRIOR TO MEASUREMENT AN ALIQUOT OF THE SAMPLE WAS MIXED WITH AN EQUAL VOLUME OF TISAB II SOLUTION (TOTAL IONIC STRENGTH ADJUSTMENT BUFFER).

THE TISAB II BUFFER SOLUTION WAS PREPARED AS FOLLOWS: 58 GM NACL AND 5 GM CDTA (CYCLOHEXYLENE DINITRILO ACETIC ACID) WERE DISSOLVED IN A MIXTURE OF 50 ML METAL FREE WATER AND 57 ML GLACIAL ACETIC ACID. THE SOLUTION WAS COOLED TO ROOM TEMPERATURE AND THE PH ADJUSTED TO BETWEEN 5.0 AND 5.5 BY NTHE SLOW ADDITION OF 5M NAOH SOLUTION. THE SOLUTION WAS COOLED AND DILUTED TO 1 LITER IN A VOLUMETRIC FLASK.

HYDROGEN ION ACTIVITY (PH) WAS MEASURED WITH A COMBINATION GLASS-CALOMEL ELECTRODE AND A PH METER.

URANIUM IN WATERS WAS DETERMINED BY A LASER-INDUCED FLUOROMETRIC METHOD USING A SCINTREX UA-3 URANIUM ANALYSER.

A COMPLEXING AGENT, KNOWN COMMERCIALLY AS FLURAN AND COMPOSED OF SODIUM PYROPHOSPHATE AND SODIUM MONOPHOSPHATE, (HALL, G.E.M., 1979) WAS ADDED TO PRODUCE THE URANYL PYROPHOSATE SPECIES WHICH FLUORESCES WHEN EXPOSED TO THE LASER.

SINCE ORGANIC MATTER IN THE SAMPLE CAN CAUSE UNPREDICTABLE BEHAVIOUR, A STANDARD ADDITION METHOD WAS USED.

FURTHER, THERE HAVE BEEN INSTANCES AT THE G.S.C. WHERE THE REACTION OF URANIUM WITH FLURAN IS EITHER DELAYED OR SLUGGISH; FOR THIS REASON AN ARBITRARY 24 HOUR TIME DELAY BETWEEN THE ADDITION OF THE FLURAN AND THE ACTUAL READING WAS INCORPORATED INTO THIS METHOD.

IN PRACTICE, 500 UL OF FLURAN SOLUTION WERE ADDED TO A 5 ML SAMPLE AND ALLOWED TO STAND FOR 24 HOURS. AT THE END OF THIS PERIOD FLUORESCENCE READINGS WERE MADE WITH THE ADDITION OF 0.0, 0.2 AND 0.4 PPB U.

FOR HIGH SAMPLES THE ADDITIONS WERE 0.0, 2.0 AND 4.0 (20 UL ALIQUOTS OF EITHER 55 OR 550 PPB U WERE USED).

ALL READINGS WERE TAKEN AGAINST A SAMPLE BLANK.

ALKALINITY IN WATERS WAS DETERMINED BY TITRATING A 25 ML ALIQUOT OF THE SAMPLE WITH 0.02N H2SO4 USING A CORNING COMBINATION ELECTRODE AND A CORNING MODEL 135 PH METER. THE END POINT WAS PH 4.5

CALCIUM AND MAGNESIUM IN WATERS WERE DETERMINED BY INDUCTIVELY COUPLED PLASMA EMISSION SPECTROSCOPY (ICP). AN ALIQUOT FROM THE SAMPLE BOTTLE WAS TRANSFERRED TO A SEPARATE CONTAINER AND ASPIRATED DIRECTLY INTO THE ICP SPECTROMETER (INSTRUMENTATION LABORATORY MODEL 200). MEASUREMENTS WERE MADE AT 317.9MM FOR CA AND 279.8NM FOR MG. THE INSTRUMENT WAS CALIBRATED WITH AQUEOUS STANDARDS.

TABLE -1 DISPLAYS THE DATA RECORD FORMAT SPECIFICATIONS AND TABLE -2 THE DETECTION LIMITS OF THE ANALYTICAL METHODS.

THE SECOND FIGURE UNDER THE DETECTION LIMIT HEADING CORRESPONDS TO AN ARBITRARILY SET VALUE IF THE RESULTS FALL BELOW THE DETECTION LIMIT (USUALLY 1/2 THE DETECTION LIMIT) AND ARE USED IN SOME OF THE STATISTICAL CALCULATIONS.

TABLE -1

ELEMENT		CARD	COLUMNS
FIELD			
MAP		1	01-06
ID		1	07-12
UTM ZONE		1	13-14
UTM EAST	(METER)	1	15-20
UTM NORTH	(METER)	1	21-27
ROCK TYPE		1	28-31
LAKE AREA		1	32-35
SAMPLE DEPTH	(METER)	1	36-38
REPLICATE ST.	AUS	1	39-40
RELIEF		1	41-43
CONTAMINATION		1	48-51
SAMPLE COLOUR		1	52-57
SUSPENDED MA	TTER	1	58-59
AGE		1	60-61

TABLE -2

THE ANALYTICAL DATA WERE RECORDED AS FOLLOWS:

	ELEM		UNITS	CARD	COLUMNS	DETECTION LIMIT
	SEDIMENT	_				
		ZN	PPM	2	16-20	<2=1
		CU	PPM	2	21-25	<2=1
		PB	PPM	2	26-30	<2=1
		NI	PPM	2	31-35	<2=1
		CO	PPM	2	36-40	<2=1
		AG	PPM	2	41-47	<0.2=0.1
		MN	PPM	2	48-52	<5=3
		AS	PPM	2	53-59	<1=.5
		MO	PPM	2	60-64	<2=1
		FE	PCT	2	65-69	<0.02=0.01
		HG	PPB	2	70-74	<10=5
		LOI	PCT	2	75-79	<1.0=.5
		U	PPM	3	16-22	<0.5=0.3
		F	PPM	3	23-27	<40=20
		V	PPM	3	28-32	<5=3
		CD	PPM	3	33-39	<0.2=0.1
		SB	PPM	3	40-46	<0.2=0.1
		AU	PPB	4	31-35	VARIABLE
	REPEAT	AU	PPB	4	36-40	VARIABLE
	AU WEIGH	T	GRAMS	4	41-44	
REPEAT AU WEIGHT GRAMS			IGHT GRAM	S 4	45-48	
	WATER					
		F	PPB	4	26-30	<20=10
		PH	LOG	4	31-35	
		U	PPB	4	36-40	<0.05=0.03
		CA	PPM	5	26-30	<0.2=0.1
		MG	PPM	5	31-35	<0.02=0.01
	T-A	ALK	PPM	5	36-40	<2=1

PRESENTATION OF GOLD DATA AND COMMENTS REGARDING

INTERPRETATION OF RESULTS

THE FOLLOWING DISCUSSION REVIEWS THE FORMAT USED TO PRESENT THE AU GEOCHEMICAL DATA AND OUTLINES SOME IMPORTANT POINTS TO CONSIDER WHEN INTERPRETING THIS DATA. THIS DISCUSSION IS INCLUDED IN RECOGNITION OF THE SPECIAL GEOCHEMICAL BEHAVIOUR AND MODE OF OCCURRENCE OF AU IN NATURE AND THE RESULTANT DIFFICULTIES IN OBTAINING AND ANALYZING SAMPLES WHICH REFLECT THE ACTUAL CONCENTRATION LEVEL AT A GIVEN SITE.

UNDERSTANDING AU GEOCHEMICAL DATA FROM REGIONAL STREAM SEDIMENT OR LAKE SEDIMENT SURVEYS REQUIRES AN APPRECIATION OF THE UNIQUE CHEMICAL AND PHYSICAL CHARACTERISTICS OF AU AND ITS MOBILITY IN THE SURFICIAL ENVIRONMENT. KEY PROPERTIES OF AU THAT DISTINGUISH ITS GEOCHEMICAL BEHAVIOUR FROM MOST OTHER ELEMENTS INCLUDE :

- 1) AU OCCURS MOST COMMONLY IN THE NATIVE FORM WHICH IS CHEMICALLY AND PHYSICALLY RESISTANT. A HIGH PROPORTION OF THE METAL IS DISPERSED IN MICRON-SIZED PARTICULATE FORM. GOLD'S HIGH SPECIFIC GRAVITY ENSURES HETEROGENEOUS DISTRIBUTION ESPECIALLY IN STREAM SEDIMENT AND CLASTIC-RICH (LOW LOI) LAKE SEDIMENT ENVIRONMENTS. AU DISTRIBUTION APPEARS TO BE MORE HOMOGENEOUS IN ORGANIC-RICH FLUVIATILE AND LAKE SEDIMENT ENVIRONMENTS.
- 2) AU TYPICALLY OCCURS AT LOW CONCENTRATIONS IN THE PPB RANGE. AU CONCENTRATIONS OF A FEW PPM MAY REPRESENT ECONOMIC DEPOSITS. BACKGROUND LEVELS ENCOUNTERED FOR STREAM AND CENTRE-LAKE SEDIMENTS SELDOM EXCEED 10 PPB, AND COMMONLY ARE NEAR THE DETECTION LIMIT OF 1 PPB.

THE MANY FOREGOING FACTORS CAN RESULT IN A PARTICLE SPARSITY EFFECT WHEREIN VERY LOW CONCENTRATIONS OF AU ARE HETEROGENEOUSLY DISTRIBUTED IN THE SURFICIAL ENVIRONMENT. HENCE, A MAJOR PROBLEM FACING THE GEOCHEMIST IS OBTAINING A REPRESENTATIVE SAMPLE. IN GENERAL THE LOWER THE ACTUAL CONCENTRATION OF AU, THE LARGER THE SAMPLE SIZE, OR THE SMALLER THE GRAIN SIZE REQUIRED TO REDUCE UNCERTAINTY OVER WHETHER SUBSAMPLE ANALYTICAL VALUES TRULY REPRESENT ACTUAL VALUES.CONVERSELY, AS ACTUAL AU CONCENTRATIONS INCREASE OR GRAIN SIZE DECREASES, THE NUMBER OF AU PARTICLES TO BE SHARED IN RANDOM SUBSAMPLES INCREASES AND THE VARIABILITY OF RESULTS DECREASES (CLIFTON ET AL., 1969; HARRIS, 1982). THE LIMITED AMOUNT OF MATERIAL COLLECTED DURING THE RAPID, RECONNAISSANCE-STYLE REGIONAL SURVEYS AND THE NEED TO ANALYZE FOR A BROAD SPECTRUM OF ELEMENTS, PRECLUDES THE USE OF A SIGNIFICANTLY LARGE SAMPLE WEIGHT FOR THE AU ANALYSES. THEREFORE, TO THE EXTENT THAT SAMPLE REPRESENTIVITY CAN BE INCREASED, SAMPLE GRAIN SIZE IS REDUCED BY SIEVING AND BALL MILLING OF ALL SAMPLES. IF THE AU IS PRESENT IN A METALLIC STATE, BALL MILLING MAY NOT REDUCE

IT'S PARTICLE SIZE SIGNIFICANTLY BECAUSE OF ITS MALLEABILITY.

GOLD DATA DISCUSSION CONTINUED

THE FOLLOWING CONTROL METHODS ARE CURRENTLY EMPLOYED TO EVALUATE AND MONITOR THE SAMPLING AND ANALYTICAL VARIABILITY WHICH ARE INHERENT IN THE ANALYSIS OF AU IN GEOCHEMICAL MEDIUMS:

- 1) FOR EACH BLOCK OF TWENTY SAMPLES:
 - A) RANDOM INSERTION OF A STANDARD REFERENCE SAMPLE TO MONITOR
 AND CONTROL ANALYTICAL ACCURACY AND LONG-TERM PRECISION,
 - B) COLLECTION OF A FIELD DUPLICATE (TWO SAMPLES SEPARATELY COLLECTED FROM ONE SITE) TO MONITOR SAMPLING VARIANCE,
 - C) ANALYSIS OF A SECOND SUBSAMPLE (BLIND DUPLICATE) FROM ONE SAMPLE TO MONITOR AND CONTROL SHORT-TERM PRECISION;
- 2) FOR BOTH STREAM SEDIMENTS AND LAKE SEDIMENTS, REPEAT ANALYSES ON A SECOND SUBSAMPLE ARE PERFORMED FOR ALL SAMPLES HAVING VALUES THAT ARE STATISTICALLY ABOVE APPROXIMATELY THE 90TH PERCENTILE OF THE TOTAL DATA SET; WITHIN THE SURVEY AREA
- 3) FOR LAKE SEDIMENTS ONLY, REPEAT ANALYSIS ON A SECOND SUBSAMPLE WAS PERFORMED ON THOSE SAMPLES WITH LOI VALUES BELOW 10%, INDICATING A LARGE CLASTIC COMPONENT. ON-GOING STUDIES SUGGEST THAT THE AU DISTRIBUTION IN THESE SAMPLES IS MORE LIKELY TO BE HIGHLY VARIABLE THAN IN SAMPLES WITH A HIGHER LOI CONTENT.

AU DATA PRESENTATION, STATISTICAL TREATMENT AND THE VALUE MAP FORMAT ARE SOMEWHAT DIFFERENT THAN FOR OTHER ELEMENTS. AU DATA LISTED IN THIS OPEN FILE INCLUDES INITIAL ANALYTICAL RESULTS, VALUES DETERMINED FROM REPEAT ANALYSES, TOGETHER WITH SAMPLE WEIGHTS AND CORRESPONDING DETECTION LIMITS FOR ALL ANALYZED SAMPLES.

THE GOLD HISTOGRAM, STATISTICAL PARAMETERS, AND REGIONAL TREND MAP ARE DETERMINED USING THE FOLLOWING DATA POPULATION SELECTION CRITERIA:

- 1) ONLY THE FIRST VALUE OF A REPEAT ANALYSIS IS UTILIZED;
- AU VALUES DETERMINED FROM SAMPLE WEIGHTS LESS THAN 10 G ARE EXCLUDED.
- 3) AU VALUES LESS THAN THE DETECTION LIMIT(<1PPB) FOR 10 G SAMPLES ARE SET TO 0.5 PPB.

GOLD DATA DISCUSSION CONTINUED

ON THE VALUE MAPS, REPEAT ANALYSIS VALUES (NOT FIELD DUPLICATES) ARE PLACED IN BRACKETS FOLLOWING THE INITIAL VALUE DETERMINATION. ALL VALUES DETERMINED ON A SAMPLE LESS THAN 10 G ARE DENOTED BY AN ASTERISK.

ACTUAL SAMPLE WEIGHT USED CAN BE DETERMINED FROM THE TEXT. FOLLOWING ARE POSSIBLE VARIATIONS IN DATA PRESENTATION ON A VALUE MAP:

*	NO DATA
+27	SINGLE ANALYSIS,10 G SAMPLE WEIGHT
+27*	SINGLE ANALYSIS,<10 G SAMPLE WEIGHT
+27 (14)	REPEAT ANALYSIS, BOTH SAMPLES 10 G
+27 (14*)	REPEAT ANALYSIS, FIRST SAMPLE 10 G, REPEAT <10 G
+<1	SINGLE ANALYSIS, 10 G SAMPLE, LESS THAN DETECTION LIMIT OF 1 PPB

IN SUMMARY, GEOCHEMICAL FOLLOW-UP INVESTIGATIONS FOR AU SHOULD BE BASED ON A CAREFUL CONSIDERATION OF ALL GEOLOGICAL AND GEOCHEMICAL INFORMATION, AND ESPECIALLY A CAREFUL APPRAISAL OF GOLD GEOCHEMICAL DATA AND ITS VARIABILITY. IN SOME INSTANCES, PROSPECTIVE FOLLOW-UP AREAS MAY BE INDIRECTLY IDENTIFIED BY PATHFINDER ELEMENT ASSOCIATIONS IN FAVOURABLE GEOLOGY, ALTHOUGH A COMPLEMENTARY AU RESPONSE DUE TO NATURAL VARIABILITY MAY BE LACKING. ONCE AN ANOMALOUS AREA HAS BEEN IDENTIFIED, FIELD INVESTIGATIONS SHOULD BE DESIGNED TO INCLUDE DETAILED GEOCHEMICAL FOLLOW-UP SURVEYS AND COLLECTION OF LARGE REPRESENTATIVE SAMPLES. SUBSEQUENT REPEAT SUBSAMPLE ANALYSES WILL INCREASE THE RELIABILITY OF RESULTS AND PERMIT A BETTER UNDERSTANDING OF NATURAL VARIABILITY WHICH CAN THEN BE USED TO IMPROVE SAMPLING METHODOLOGY AND INTERPRETATION.

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DATA LIST LEGEND

MAP- NATIONAL TOPOGRAPHIC SYSTEM(NTS) - LETTERED QUADRANGLE

(SCALE 1:250000). PART OF SAMPLE NUMBER

ID- REMAINDER OF SAMPLE NUMBER- YEAR(2), FIELD CREW(1),

SAMPLE SEQUENCE NUMBER (3)

UTM COORDINATS- UNIVERSAL TRANSVERSE MERCATOR (UTM) COORDINATE

SYSTEM- SAMPLE COORDINATES

ZN- ZONE

EAST- EASTING (METERS)

NORTH- NORTHING (METERS)

ROCK TYPE- MAJOR ROCK TYPE OF LAKE CATCHMENT AREA

AGE- STRATIGRAPHIC AGE OF ROCK TYPE

LAKE AREA- AREA OF LAKE SAMPLED

SMP DTH- LAKE DEPTH AT SAMPLE SITE MEASURED TO THE NEAREST METER

RP ST- REPLICATE STATUS- RELATIONSHIP OF SAMPLE TO OTHERS

WITHIN THE BLOCK OF TWENTY

RELF- RELIEF OF THE SURROUNDING LAKE CATCHMENT BASIN

CONT- CONTAMINATION- HUMAN OR NATURAL (WORK-DRILL/TRENCH,

CAMP, FUEL OR GOSSAN)

SMPL COLOR- SEDIMENT COLOUR

SUSP- SUSPENDED MATTER

LAKE AREA: POND- POND

LT 1- 1/4 TO 1 SQ KM

1-5- 1 TO 5 SQ KM

GT 5- GREATER THAN 5 SQ KM

RP ST: 00- ROUTINE REGIONAL SAMPLE

10- FIRST OF FIELD DUPLICATE

20- SECOND OF FIELD DUPLICATE

RELF: L- LOW

M- MEDIUM

H- HIGH

CONT: BLANK- NONE

1- PRESENT

SMPL COLOR: TN- TAN GY- GREY

YL- YELLOW BR- BROWN

GN- GREEN BK- BLACK

SUSP: BLANK- NONE

L- LIGHT

H- HEAVY

ROCK TYPES:

QUATERNARY

(QU 64) - AREAS OF DRIFT COVER; MAY BE SPARSE OUTCROP.

PRECAMBRIAN

DYKE SUITE

- (ADA 04) MOLSON DYKES; GABBRO, DIABASE, LAMPROPHYRE.
- (ADC 04) AMPHIBOLITE, PLAGIOCLASE PORPHYRITIC, ULTRAMAFIC AMPHIBOLITE.
- (ADD 04) DIORITE AND SYENITE, GABBRO PLUGS, LOCALLY ZONED AND BRECCIATED.

GODS LAKE DOMAIN

POST OXFORD LAKE GROUP PLUTONIC ROCKS

- (AGB 02) GABBRO
- (AGG 02) GRANITE, PEGMATITIC GRANITE, GRANODIORITE,
 PORPHYRITIC GRANODIORITE.

OXFORD LAKE GROUP

- (AOW 02) FELDSPATHIC GREYWACKE, ARKOSIC WACKE,

 MUDSTONE, AMPHIBOLE-BIOTITE GNEISS, GARNET
 BIOTITE GNEISS, POLYMICTIC CONGLOMERATE.
- (AOB 02) GABBRO, PORPHYRITIC GABBRO, DIABASE.
- (AOV 02) MAFIC TO FELSIC VOLCANIC ROCKS, INCLUDING FRAGMENTALS, FELSIC TO INTERMEDIATE TUFF, AND RELATED SEDIMENTARY ROCKS.

BAYLY LAKE COMPLEX

- (ABG 02) GRANITE, PORPHYRITIC, MICROCLINE-PHYRIC, OR GNEISSOID GRANODIORITE.
- (ABT 02) TONALITE, HORNBLENDE GRANODIORITE PORPHYITIC,
 MYLONITIC AND GNEISSIC TONALITE, SUBORDINATE
 AMPHIBOLITE INTERLAYERS MAY INCLUDE TONALITE
 AND GRANODIORITE NOT PART OF BAYLY LAKE COMPLEX.
- (ABRZ 02) PLAGIOCLASE PORPHYRY, QUARTZ PORPHYRY,
 PROTOMYLONITIC QUARTZ MONZONITE, SYENITE,
 PEGMATITIC SYENITE.
- (ABHU 02) GABBRO, DIABASE, HORNBLENDE, CLINOPYROXENITE.

ISLAND LAKE GROUP

- (AIV 02) ANDESITE FLOWS, TUFF AND AGGLOMERATE, MINOR BASALT, SILTSTONE, CONGLOMERATE.
- (AIS 02) CONGLOMERATE, QUARTZ WACKE (REGOLITH),
 SILTSTONE, TURBIDITIC GREYWACKE AND ARGILLITE.

ROCK TYPES (CONTINUED):

HAYES RIVER GROUP

- (AHS 02) VOLCANOGENIC SEDIMENTARY ROCKS; FELDSPATHIC GREYWACKE, GREYWACKE, SILTSTONE, CONGLOMERATE AND CORRELATIVE SCHIST AND PARAGNEISS; MINOR CHERT, IRON FORMATION, CARBONATE AND TUFF.
- (AHF 02) FELSIC AND INTERMEDIATE VOLCANIC ROCKS; MAINLY FLOWS AND RELATED SILLS, APHYRIC FELSITE, PORPHYRITIC DACITE AND RHYOLITE, TUFF AND TUFF BRECCIA, ALBITIC SCHISTS AND GNEISS DERIVED FROM FELSIC VOLCANICS.
- (AHV 02) MAFIC AND INTERMEDIATE VOLCANIC ROCKS; MAINLY
 APHYRIC FLOWS, BASALT, ANDESITE, VOLCANIC BRECCIA,
 CORRELATIVE AMPHIBOLITIC SCHIST AND GNEISS.
- (AHBU 02) GABBRO, PLAGIOCLASE-PHYRIC GABBRO, SERPENTINIZED PERIDOTITE, RELATED ULTRAMAFIC ROCKS.

UNNAMED VOLCANIC AND SEDIMENTARY ROCKS

- (AS 02) ARKOSE TO FELDSPATHIC GREYWACKE, POLYMICTIC AND VOLCANOGENIC CONGLOMERATE.
- (AF 02) FELSIC AND INTERMEDIATE VOLCANIC ROCKS; FLOWS, SILLS, FRAGMENTALS, TUFF AND RELATED SEDIMENTS.
- (AV 02) MAFIC AND INTERMEDIATE VOLCANIC ROCKS; MAINLY FLOWS, PILLOWED FLOWS, TUFF AND RELATED SEDIMENTS, AMPHIBOLITE.
- (ABU 02) GABBRO, PYROXENE PHYRIC GABBRO, PLAGIOCLASE-PHYRIC GABBRO, SERPENTINIZED PERIDOTITE.

MOLSON/KALLICAHOOLIE DOMAIN

- (AMG 02) GRANITE, PEGMATITIC GRANITE, PYROXENE-BEARING MONZODIORITE.
- (AMN 02) HYBRID GRANODIORITIC TO GRANITIC GNEISS.
- (AMGD 02) GRANODIORITE, INCLUDES MEGACRYSTIC AND HORNBLENDE-BEARING PHASES.
- (AMB 02) PYROXENE-PHYRIC GABBRO AND PLAGIOCLASE-PHYRIC GABBRO.
- (AMT 02) LEUCOTONALITE, TONALITE WITH GRANODIORITE
 INTRUSIONS, TONALITE TO TONALITIC GNEISS WITH
 AMPHIBOLITIC INTERLAYERS, BIOTITE GRANODIORITE,
 MINOR BIOTITE-MUSCOVITE GRANITE; PLUTONS MAY
 SHOW MARKED POSITIVE MAGNETIC ANOMALIES.
- (AMNG 02) GARNET-BEARING QUARTZOFELDSPATHIC GNEISS.
- (AM 02) AMPHIBOLITE.

AGES:

- 64 OUATERNARY
 - 04 PRECAMBRIAN
 - 02 ARCHEAN

- 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)
- AS- ARSENIC BY HYDRIDE EVOLUTION-ATOMIC ABSORPTION SPECTROSCOPY (PPM)
- MO- MOLYBDENUM BY ATOMIC ABSORPTION SPECTROSCOPY (PPM)
- FE- IRON BY ATOMIC ABSORPTION SPECTROSCOPY (%)
- HG- MERCURY BY FLAMELESS SPECTROSCOPY (PPB)
- LOI- LOSS ON IGNITION BY WEIGHT DIFFERENCE (%)
- U- URANIUM BY NEUTRON ACTIVATION DELAYED
 - NEUTRON COUNTING (PPM)
- F- FLUORINE BY SPECIFIC ION ELECTRODE (PPM)
- V- VANADIUM BY ATOMIC ABSORPTION SPECTROSCOPY (PPM)
- CD- CADMIUM BY ATOMIC ABSORPTION SPECTROSCOPY (PPM)
- SB- ANTIMONY BY HYDRIDE EVOLUTION-ATOMIC
 ABSORPTION SPECTROSCOPY (PPM)
- AU- GOLD BY FIRE ASSAY PRECONCENTRATION-NEUTRON
 ACTIVATION (PPB)
- AU-R- GOLD REPEAT ANALYSIS BY FIRE ASSAY PRECONCENTRATION-NEUTRON ACTIVATION(PPB)
- AU WT1- WEIGHT IN GRAMS OF ORIGINAL GOLD SAMPLE ANALYZED
- AU WT2- WEIGHT IN GRAMS OF REPEAT GOLD SAMPLE ANALYZED
 - DL1- GOLD DETECTION LIMIT BASED ON ANALYSIS SAMPLE WEIGHT FOR INITIAL GOLD ANALYSIS
 - DL2- GOLD DETECTION LIMIT BASED ON ANALYSIS SAMPLE WEIGHT FOR REPEAT GOLD ANALYSIS
 - F-W- FLUORIDE IN WATERS BY SPECIFIC ION ELECTRODE (PPB)
 - PH- PH BY COMBINATION GLASS-CALOMEL ELECTRODE
 - U-W- URANIUM IN WATERS BY LASER INDUCED FLOURESCENCE (PPB)
 - T-ALK- ALKALINITY (AS PPM CACO3) BY TITRATION
 - CA-W- CALCIUM IN WATERS BY INDUCTIVELY COUPLED ARGON PLASMA
 - EMISSION SPECTROMETRY (PPB)
 - MG-W- MAGNESIUM IN WATERS BY INDUCTIVELY COUPLED ARGON PLASMA EMISSION SPECTROMETRY (PPB)