Environment Canada
Environmental Protection Service
Pacific Region
Yukon Branch

RPR

ENVIRONMENTAL QUALITY

OF ROSE CREEK AS AFFECTED BY DISCHARGES

FROM THE CYPRUS ANVIL MINE,

YUKON TERRITORY (1983)

Regional Program Report No. 85-12

bу

Benoit Godin and Tim Osler

October, 1985

LIBRARY
ENVIRONMENT CANADA
CONSERVATION AND PROTECTION
PACIFIC REGION

ABSTRACT

An assessment of the water quality and biological conditions of Rose Creek, adjacent to Cyprus Anvil mine, was carried out during the summer of 1983. The water quality, sediment and bottom fauna characteristics were documented at 10 environmental quality monitoring stations. Water quality information from Cyprus Anvil Mining Corporation's 21 Surveillance Network Stations was evaluated.

Environmental quality monitoring program results have indicated an improvement in the water quality of Rose Creek since the CAMC's June 15, 1982 mine shutdown. Sediment analysis have revealed little change to previous studies. Benthic invertebrates have increased in abundance compared to previous studies and bioassays with Daphnia pulex showed no acute toxic effect from the tailings pond seepage. The surveillance network indicated incidents of non-compliance in 1982-83 with elevated levels of zinc at the tailings pond and in the winter months of 1983-84 at the north fork of Rose Creek.

RESUME

Une évaluation de la qualité et des conditions biologiques de l'eau de ruisseau Rose adjacent à la mine Cyprus Anvil, fut entrepris durant l'été 1983. La qualité de l'eau, les caractéristiques des sédiments et de la faune benthique ont été analysées at dix stations d'échantillonnage pour l'évaluation de la qualité environmentale. Vingt-une stations on été échantillonnés pour la qualité de l'eau du programme de surveillance a la Corporation Minière Cyprus Anvil.

Les résultats ont indiques une amélioration de la qualité de l'eau de ruisseau Rose durant de la période inopérante de la mine. L'analyse des sediments montre un accroissement important du niveau de métaux lourds aux stations intermediaires du ruisseau, qui doivent être reliés au déversement des bassins de décantation en 1975. Les invertebres benthiques augmentent en abondance et les bio-essais avec <u>Daphnia pulex</u> ne montrent aucune toxicité aigu des infiltrations des bassins de décantation. Le programme de surveillance indique par contre des incidents non conforme au permit en 1982-83, indicant de hautes concentrations de zinc aux bassins de décantation et aux mois d'hiver 1983-84 dans la branche nord du ruisseau Rose.

TABLE OF CONTENTS

		<u>Pag</u>
ABSTRACT		i
RESUME	·	ii
TABLE OF	CONTENTS	iii
	List of Figures	vii
	List of Tables	ix
1	INTRODUCTION	1
2	STUDY AREA	2
3	METHODS	9
3.1	Water Quality	9
3.2	Sediments	10
3.3	Bottom Fauna	11
3.3.1	Diversity/Evenness Indices	12
3.3.2	Community Index	12
3.4	Attached Algae	13
3.5	Bioassay	13
3.5.1	Control Water Details	13
3.5.2	Test Organism	13
3.5.3	Statistics	14
4	RESULTS AND DISCUSSION	15
4.1	Water Quality-Physical and Chemical Parameters	15
4.1.1	Flow	15
4.1.2	Dissolved Oxygen	15
4.1.3	рН	17
4.1.4	Conductivity and Filterable Residue	17
4.1.5	Color	19
4.1.6	Turbidity	19
4.1.7	Non-filterable Residue	21
4.1.8	Alkalinity and Hardness	21
4.1.9	Phosphates	21
4.1.10	Ammonia	23

TABLE OF CONTENTS (continued)

		Page
4.1.11	Nitrite and Nitrate	23
4.1.12	Sulfates	24
4.2	Water Quality-Metals	24
4.2.1	Extractable Metal Analysis	24
4.2.2	Copper	26
4.2.3	Zinc	29
4.3	Sediment	31
4.3.1	Particle Size	31
4.3.2	Sediment Metal Analysis	31
4.4	Bottom Fauna	32
4.4.1	Artifical Substrate Sampler	32
4.4.2	Taxonomic Features	35
4.4.3	Diversity Index	40
4.4.4	Community Index	42
4.5	Attached Algae	44
4.6	Bioassay	46
5	ENVIRONMENTAL COMPLIANCE MONITORING	48
5.1	Regulatory Requirements of the Cyprus Anvil Mine	48
5.2	Monitoring Data 1974-1983	48
5.3	Non-Compliance Incidents 1982-1984	53
CONCLUSI	ONS	55
REFERENC	ES	56
ACKNOWLEDGEMENTS		

TABLE OF CONTENTS (continued)

			<u>Page</u>
APPENDICES			61
APPENDIX I	COLLECTIO	ON, PRESERVATION, ANALYSIS OR	
	IDENTIFIC	CATION METHODS AND WATER QUALITY	
	CRITERIA		63
	TABLE 1	WATER SAMPLE COLLECTION,	
		PRESERVATION AND ANALYSIS METHODS	64
	TABLE 2	SEDIMENT COLLECTION,	
		PREPARATION AND ANALYSIS METHODS	69
	TABLE 3	BOTTOM FAUNA COLLECTION,	
		PRESERVATION AND IDENTIFICATION	
		METHODS	71
	TABLE 4	ATTACHED ALGAE COLLECTION, PRESER-	
		VATION AND IDENTIFICATION METHODS	72
	TABLE 5	WATER QUALITY CRITERIA FOR	
		DRINKING WATER AND AQUATIC LIFE	73
APPENDIX II	WATER QUA	ALITY DATA	77
	TABLE 1	WATER QUALITY - PHYSICAL AND	
		CHEMICAL PARAMETERS	78
	TABLE 2	WATER QUALITY - EXTRACTABLE	
		METALS (mg/1)	82
APPENDIX III	SEDIMENT	DATA	89
•	TABLE 1	SEDIMENT PARTICLE SIZE - MEAN	
		AND STANDARD DEVIATION	90
	TABLE 2	SEDIMENT - LEACHABLE METALS	91

TABLE OF CONTENTS (continued)

		<u>Page</u>
APPENDIX IV	BOTTOM FAUNA DATA	97
	TABLE 1 TAXONOMIC LIST OF THE BOTTOM	1
	FAUNA AT CYPRUS ANVIL -	
	SEPTEMBER, 1983	98
	TABLE 2 CYRPUS ANVIL MACRO-INVERTEBR	ATE
	DATA - SEPTEMBER, 1983	102
APPENDIX V	ATTACHED ALGAE DATA	105
	TABLE 1 ATTACHED ALGAE - EXTRACTABLE	•
	METALS - STATION 3	106
APPENDIX VI	ENVIRONMENTAL COMPLIANCE MONITORING DA	ATA 107
•	TABLE 1 CYPRUS ANVIL MINING CORPORAT	ION -
	HISTORIAL WATER QUALITY DATA	1
	(STATION X-3, X-14, X-15)	108

LIST OF FIGURES

Figure		Pag
1	GEOGRAPHIC LOCATION OF CYPRUS ANVIL MINING CORPORATION LTD.	3
2	ENVIRONMENTAL QUALITY MONITORING SAMPLE STATION LOCATIONS	4
3	CYRPUS ANVIL MINE SITE PLAN SHOWING EPS WATER QUALITY SAMPLE STATIONS AND CAMC COMPLIANCE MONITORING STATIONS	6
4	DISSOLVED OXYGEN - PERCENTAGE OF SATURATION - IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983	16
5	MEAN VALUES OF FILTERABLE RESIDUE (F.R.) AND CONDUCTIVITY IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983	18
6	MEAN VALUES OF TURBIDITY AND NON-FILTERABLE RESIDUE (NFR) IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983	20
7	MEAN VALUES OF ALKALINITY AND HARDNESS IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983	22
8	MEAN VALUES OF SULPHATES IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983	25
9	MEAN VALUES OF EXTRACTABLE COPPER IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983	27

LIST OF FIGURES

Figure		<u>Page</u>
10	MEAN VALUES OF EXTRACTABLE ZINC IN WATER	
	SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER	
	15, 1983	30
11	MEAN VALUES OF EXTRACTABLE CADMIUM IN SEDIMENT	
	SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER	
	15, 1983	33
12	MEAN VALUES OF EXTRACTABLE COPPER IN SEDIMENT	
	SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER	
	15, 1983	34
13	INVERTEBRATE TAXONOMIC GROUP DOMINANCE IN	
	CYPRUS ANVIL MINE STUDY AREA	41

LIST OF TABLES

<u>Table</u>		Page
1	STATION DESCRIPTION OF THE ENVIRONMENTAL QUALITY MONITORING PROGRAM	5
2	CYRPUS ANVIL MINING CORPORATION - SURVEILLANCE NETWORK PROGRAM	7
3	GENUS PRESENCE OF BENTHIC INVERTEBRATES AT THE VARIOUS STATIONS	38
4	PERCENTAGE OF DIFFERENT TAXONOMIC GROUPS REPRESENTED AT EACH STATION	39
5	COMPARISON OF DIVERSITY INDICES FOR SAMPLE STATIONS SINCE 1973	43
6	THE PERCENT SIMILARITY INDEX MATRIX FOR THE 1983 MONITORING PROGRAM	45
7	<u>Daphnia</u> <u>pulex</u> BIOASSAY RESULTS OF CYPRUS ANVIL TAILINGS POND SEEPAGE	50
8	WATER QUALITY, SELECTED PARAMETERS - JUNE, 1982 - DECEMBER, 1983	51
9	COMPLIANCE MONITORING - FOR Zn AND pH, 1982-1983	52

1 INTRODUCTION

The Environmental Protection Service (EPS) has monitored the biological communities, water quality and sediment characteristics of Rose and Anvil Creeks near the Cyprus Anvil mine since 1973 (Hoos 1973, Baker 1979, Weagle 1981). The purpose of these studies is to ensure that the receiving water quality downstream of the mine is not being seriously degraded. This 1983 study updates the environmental quality monitoring information and utilized the same station locations in most cases as previous surveys. These efforts compliment the biological monitoring program presently required as part of the water licence for the Cyprus Anvil Mine Corporation (CAMC).

Mining activity at the time of the survey consisted only of stripping overburden and waste rock covering the ore body to the east of the main pit. No processing of ore has occurred since June 1982 when the mill was shut down for an indefinite period.

The results of this survey are compared to results obtained in previous years to provide a relatively detailed picture of the environmental quality of Rose Creek and Anvil Creek.

Also included in the report is a section on water quality compliance based on the program as defined by the CAMC water licence.

2 STUDY AREA

The Cyprus Anvil Mine Corporation (CAMC) operates a lead-zinc-silver mine situated in the Anvil Range near Faro (62°20'N, 133°25'W) about 240 kilometres (150 miles) by air north of Whitehorse, Yukon (Figure 1). The mine area is drained by Rose Creek, Faro Creek and the north fork of Rose Creek (Figure 2). Rose Creek joins, 17km downstream of the mine, with Anvil Creek which in turn joins the Pelly River 64km from the mine. Figure 2 depicts and Table 1 describes sample stations for the environmental quality monitoring component of the study. Figure 3, an enlargement of the mine site area, shows station numbers and locations which are specified in the CAMC water licence for water quality compliance sampling. Table 2, describes the locations of the compliance monitoring network stations and details the activities regularly conducted by CAMC at each station.

There is some overlap of sampling stations between the environmental quality stations and the compliance monitoring network. These are identified in Figure 3 and described in Tables 1 and 2.

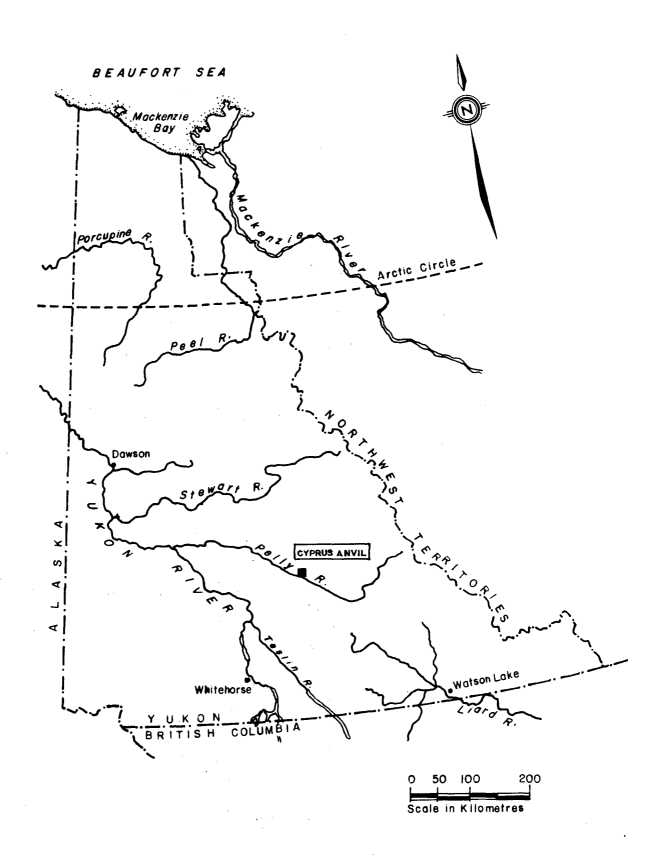
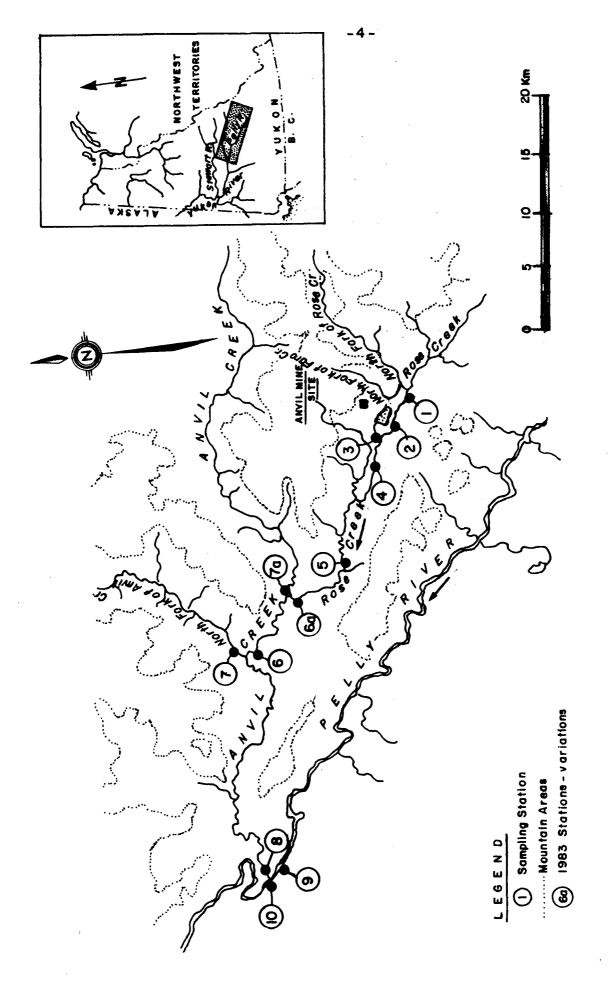


FIGURE I GEOGRAPHIC LOCATION OF CYPRUS ANVIL MINING CORPORATION LTD.



ENVIRONMENTAL QUALITY MONITORING SAMPLE STATION LOCATIONS FIGURE 2

TABLE 1 STATION DESCRIPTIONS OF THE ENVIRONMENTAL QUALITY MONITORING LOCATIONS

EBC	
EPS STATION	STATION LOCATION DESCRIPTION
SIATION	STATION LOCATION DESCRIPTION
1	On Rose Creek, above the influence of the mine and mill effluent, 250 meters upstream of the north fork of Rose Creek confluence and compliance monitoring network Station X-3. Stream bed consists of boulder, cobbles, coarse and fine gravel.
2	On the diversion canal, near the new tailings pond, 700 meters upstream of compliance monitoring network Station X-10. Canal bed consists of mainly big boulders, cobbles and minor fine sediments.
3	Combined seepage flows from X-11 and X-12 upstream of decant confluence. The same location as compliance monitoring network Station X-13. Stream bed consists of coarse and fine gravel.
4	On Rose Creek, 1 km downstream of the entry of decant stream. Streambed consists of cobbies and coarse gravel.
5	On Rose Creek, about 5km above the confluence with Anvil Creek. Streambed consists of cobbles, coarse and fine gravel.
6	On Anvil Creek, above confluence with the north fork of Anvil Creek.
6 a	On Rose Creek, above the confluence with Anvil Creek. The same location as compliance monitoring network Station X-15. Stream bed consists of boulders, cobbies, and coarse gravel.
7	On north fork of Anvil Creek, upstream of the confluence with Anvil Creek.
7a \	On Anvil Cr., upstream of the confluence with Rose Creek. Streambed consists of coarse and fine gravel and sand.
8	On Anvil Creek, upstream of the confluence with the Pelly River. Stream bed consists of coarse and fine gravel and sand.
9	On Pelly River, upstream of the confluence of Anvil Cr. Stream bed consists of cobbles, coarse and fine gravel.
10	On Pelly River, downstream of the confluence of Anvil Creek. Stream bed consists of cobbles, coarse and fine gravel.

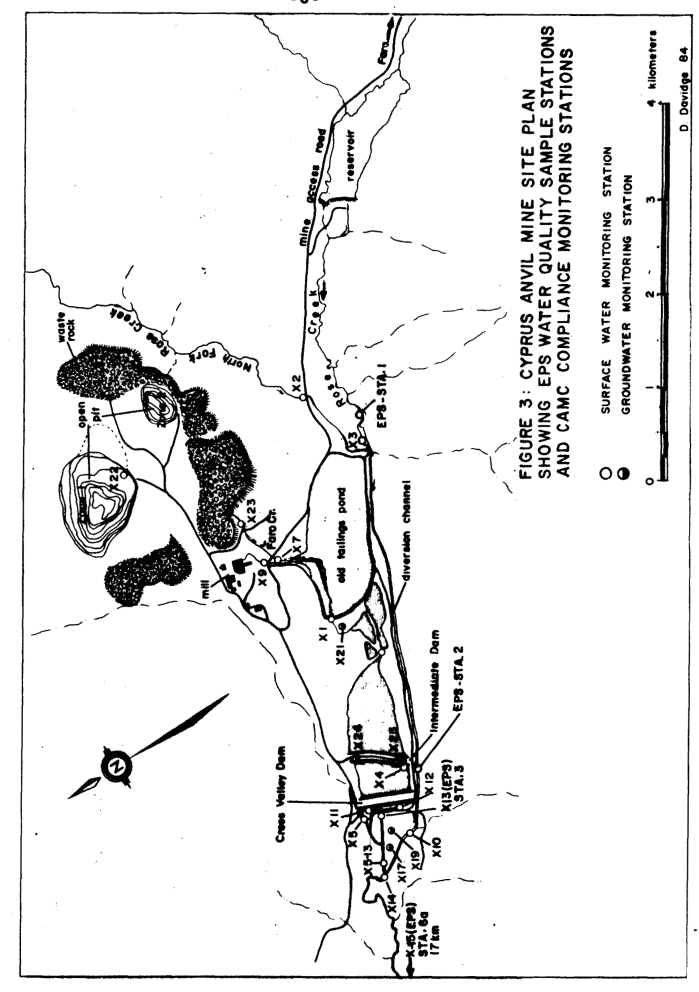


TABLE 2 CYPRUS ANVIL MINING CORPORATION - SURVEILLANCE NETWORK PROGRAM

Parameters Monitored:	pH, Temperature, Suspended Solids, Flow, Ammonia (NH ₃), Copper (Cu), Lead (Pb), Zinc (Zn), Cyanide (CN), Manganese (Mn) and Sodium (Na).
IDENTIFICATION NUMBER	DESCRIPTION OF SAME STATION AND PARAMETERS
X-2	North Fork Rose Creek, upstream of mine road bridge; sampled monthly for all parameters.
X-3	Rose Creek freshwater at pumphouse; sampled monthly for all parameters.
X-4	Intermediate Dam decant; sampled weekly for all parameters.
X-5	Cross Valley Dam decant; sampled weekly for all parameters.
X-7	Mine water at road crossing; sampled monthly for all parameters except Na.
X-10	Rose Creek Diversion Canal below wiers; sampled monthly for all parameters.
X-11	Seepage from north toe of the Cross Valley Dam; sampled weekly for all parameters.
X-12	Seepage from south toe of the Cross Valley Dam; sampled weekly for all parameters.
X-13	Combined seepage flows from X-11 and X-12 upstream of decant confluence; sampled weekly for all parameters.
X-14	Rose Creek downstream of diversion canal, seepage channel and decant channel confluences; Sampled monthly for all parameters.
X-15	Rose Creek at mouth. Sample site is the same as EPS Station 6(a).
X-16, X-17, X-18 X-19, X-20, X-21	Groundwater monitor wells K-1 to K-6 sampled every two months for all parameters except flow and CN.

TABLE 2 CYPRUS ANVIL MINING CORPORATION - SURVEILLANCE NETWORK PROGRAM (Continued)

Parameters Monitored:	pH, Temperature, Suspended Solids, Flow, Ammonia (NH ₃), Copper (Cu), Lead (Pb), Zinc (Zn), Cyanide (CN), Manganese (Mn) and Sodium (Na).		
IDENTIFICATION NUMBER	DESCRIPTION OF SAME STATION AND PARAMETERS		
X-22	Discharge from Faro #1 Pit pumps; sampled weekly for pH, suspended solids, NH ₃ , Cu, Pb, Zn and Mn.		
X-23	Pit drainage at toe of waste rock pile upstream of X-7; sampled weekly for pH, suspended solids, NH ₃ , Cu, Pb, Zn and Mn.		
X-24, X-25	Groundwater station monitor wells 1.0.2 and 1.0.6; sample every two months for all parameters except flow and CN.		
·			

3 METHODS

Environmental quality monitoring of the 10 different stations at CAMC was done on July 26 and September 15, 1983. All sample stations are in similar locations to past studies by Environmental Protection Service except stations 6 and 7. The 1983 sampling program altered Station 6 to be on Rose Creek upstream of its confluence with Anvil Creek instead of on Anvil Creek (Figure 2) and Station 7 to be on Anvil Creek upstream of the Rose Creek confluence instead of the north fork of Anvil Creek. Access to station 1 - 3 was by road and to stations 4 - 10 was by helicopter. The historic EPS monitoring stations will be referred to by number whereas the 1983 stations 6 and 7 are accompanied by a letter [6 (a) and 7 (a)]. Information was collected on water quality and quantity, sediment characteristics and benthic invertebrates at each station. In addition, bioassay testing and attached algal tissue analysis were performed on samples from specific stations.

3.1 Water Quality and Quantity

Water samples were collected in triplicate on July 26 and in duplicate on September 15, 1983. Sample collection, preservation, and analysis methods are described in Appendix 1, Table 1. Flow measurements were taken on July 26, 1983 using a Price type AA electronic current meter, except for Stations 9 and 10 on the Pelly The Pelly River was too large to sample with the equipment available. When taking flow measurements, an area with a uniform cross section was chosen for the metering. Ten readings were taken across the profile of the stream at each station whenever possible. discharge was calculated as the summation of these fractional discharges. Water quality field measurements included temperature. conductivity, pH and dissolved oxygen. Replicate values were not collected for these parameters. Most of the analyses were conducted at EPS Laboratory Services, 4195 Marine Drive, West Vancouver, B.C. These included pH, conductivity, color, turbidity, non-filterable residue,

total alkalinity, total hardness, total phosphates, nitrites, nitrates, ammonia, sulfate, chloride and the following extractable metals:

Aluminum (Al)	Cobalt (Co)	Selenium (Se)
Antimony (Sb)	Copper (Cu)	Silicon (Si)
Arsenic (As)	Iron (Fe)	Silver (Ag)
Boron (B)	Lead (Pb)	Sodium (Na)
Barium (Ba)	Magnesium (Mg)	Strontium (Sr)
Beryllium (Be)	Manganese (Mn)	Tin (Sn)
Cadmium (Cd)	Molybdenum (Mo)	Titanium (Ti)
Calcium (Ca)	Nickel (Ni)	Vanadium (V)
Chromium (Cr)	Potassium (K)	Zinc (Zn)

The percent dissolved oxygen saturation (% D0) was calculated by first determining the dissolved oxygen saturation concentration (S') from the formula:

$$S' = S P (APHA et al 1975)$$

- where S' = dissolved oxygen (DO) saturation concentration at the in situ temperature and atmospheric pressure
 - S = dissolved oxygen (DO) saturation concentration at sea level for in situ temperatur
 - P = atmospheric pressure in millimeter of mercury (mm Hg) at site elevation

The percent dissolved oxygen saturation was obtained by using the ratio of field dissolved oxygen and S' in the following formula:

$$\frac{\text{Field DO}}{\text{S'}} \times 100 = \% \text{ DO Saturation}$$

where Field DO = Dissolved Oxygen concentration measured in the field

3.2 Sediment

Sediment samples were collected at the same time as water samples. Sediments were collected near the bank below the water level, and selected for their representativeness of the streambed as much as

possible. Three sediment samples were collected at each site using an aluminum shovel to scoop the samples into pre-labelled geochemical sampling bags within Whirl-PakTM bags. A description of sediment collection, preparation and analysis methods is given in Appendix I, Table 2. All sediment samples were air shipped to Vancouver for analysis at Laboratory Services, Environmental Protection Service, 4195 Marine Drive, West Vancouver, B.C. The replicate sediment samples were each analysed for particle size and the following leachable metals.

Aluminum (Al)	Iron (Fe)	Silicon (Si)
Arsenic (As)	Lead (Pb)	Silver (Ag)
Barium (Ba)	Magnesium (Mg)	Sodium (Na)
Beryllium (Be)	Manganese (Mn)	Strontium (Sr)
Cadmium (Cd)	Mercury (Hg)	Tin (Sn)
Calcium (Ca)	Molybdenum (Mo)	Titanium (Ti)
Chromium (Cr)	Nickel (Ni)	Vanadium (V)
Cobalt (Co)	Phosphorus (P)	Zinc (Zn)
Copper (Cu)	Potassium (K)	

3.3 Bottom Fauna

Bottom fauna collection, preservation and identification methods are summarized in Appendix I, Table 3. The benthic invertebrates were sampled at each of 10 stations using an artificial substrate sampler. The samplers were placed in the stream on July 26, 1983 and left to colonize for 50 days before retrieving on September 15, 1983. Three replicate samples were collected at each station. The substrate used in the artificial substrate samplers was taken directly from the streambed at each stations and consisted of rock sizes ranging from 1cm to 8cm in size. The rocks were not washed or brushed in order to loosen debris, sediment and benthic invertebrates. The maximum volume of the artificial substrate sampler is 0.0057 m³.

The samplers were located in the creeks at various depths (30-60 cm) along a cross-section of the stream, deep pools and fast currents were avoided. A plankton net (mesh opening 0.76 mm) was used to capture any organisms lost from the sampler during retrieval.

3.3.1 <u>Diversity/Evenness Indices</u>. Diversity/evenness indices were calculated from the bottom fauna data using the Shannon-Weiner diversity index described by Pielou (1975) and modified as follows:

Species Diversity (H') =
$$-\Sigma (P_i \log_{10} P_i)$$

 $i=1$

where $P_i = n_i/N$

 n_i = total number of individuals in the i^{th} genus

N = total number of individuals identified to genus level and/or species taxonomic level in one sample

g = total number of genera in one sample

The use of individuals identified to genus level instead of to species level results in slightly lower and conservative diversity index (H') values (Hughes, 1978). Individuals that were not identified to genus or species level, were not included in the the Species Diversity calculation but are listed collectively with all individuals that were (Appendix IV, Tables 1 and 2).

The diversity of the benthic invertebrate community depends on the number of species and the evenness with which the individuals are apportioned among them. The method of measuring evenness used in this report is described by Pielou (1975) and is given by the formula:

Evenness
$$J' = \frac{H'}{\log g}$$

where $H' =$ the species diversity
 $g =$ the number of species

3.2.2 <u>Community Index</u>. The benthic communities were also analysed using the Percent similarity index (Psc) as the community index, comparing each station to other stations using the formula as described by Brock (1977).

k

Psc = $100 - 0.5 \Sigma | a-b |$

where a and b are, for a given genus, percentages of the total samples A and B which that genus represents. The absolute value of their difference is summed over all genera, k.

- 3.4 <u>Attached Algae</u>. Tissues of a filamentous algae (Class Bacillariophyceae) from Station 3 (seepage) were analysed for extractable metals. The analysis involved one sample on July 26 and three on September 15, 1983.
- 3.5 <u>Bioassay</u>. Duplicate bioassay samples were collected at Station 1 (Rose Creek) and a single sample at Station 3 (seepage) on July 25, 1983. The bioassays were performed at the Whitehorse EPS laboratory using <u>Daphnia pulex</u>. The results involved an acute toxicity test for 48 hours (48h-LC₅₀) and 96 hours (96h-LC₅₀).
- 3.5.1 <u>Control Water Details</u>. The tap water used for dilutions of bioassays and experimental controls originated from the City of Whitehorse water supply. The water was filtered through a carbon filter and aged to ensure dechlorination. The water was aerated to saturation with oxygen and adjusted to 20°C prior to use.
- Test Organism Culture. The Daphnia pulex culture originated from EPS Atlantic Region. The Daphnia pulex population was adapted to a light cycle of 16 hours of day and 8 hours of night. Culture temperatures were kept between 20-23°C, pH 7.5-8.2 and dissolved oxygen (DO) 5.0-8.0 ppm. Feeding occurred three times a week and included a composition of trout chow, yeast and cerophyllr. The test organisms used, were less than 48 hours old. The criteria for the acute toxicity test was the mortality of the organisms as defined by the lack of antenna, phyllopodes or abdominal movements. Two different exposures were performed, a 48 hour test for Station 1 (Rose Creek) and a 96 hour

test for Station 3 (seepage). The tests were conducted at 20° C with no aeration in the 200 ml test vessels. The loading density was one Daphnia pulex per 20 ml.

3.5.3 Statistics. Statistical treatment of the replicate data involved calculating the arithmetic mean (\bar{x}) and standard deviation (S.D.). Certain comparisons involved the use of confidence limits using the Student's t-distribution for small samples and correlations (r). The formula used for these calculations are according to Sokal and Rohlf (1981).

4 RESULTS AND DISCUSSION

4.1 Water Quality Physical and Chemical Parameters

Data of the replicate water chemistry collected is presented as an arithmetic mean (\bar{x}) and standard deviation (S.D.) for each station in Tables 1 and 2 of Appendix II. Figures 4 to 9 provide graphic representation of the trends of selected parameters, their distribution and mixing in the stream.

4.1.1 Flow. Flow measurements were taken only on July 26, 1983 and variations in the flow regime between the two sampling periods can be assessed only by water level observations on the river bank. September 15, 1983 water levels were equal to or higher than July 26, 1983 at all stations. The increase in flow between Stations 1 and 2 on July 26 is a result of increases from the north fork of Rose Creek and Faro Creek. The cross valley pond seepage, measured at Station 3 (compliance monitoring sampling Station X-13) remained unchanged during the sampling periods, with mixing proportions with Rose Creek being approximately 1:20 during July. The flow measured at Station 6(a) (Rose Creek) and Station 7(a) (Anvil Creek) were comparable, 4.50 and 4.32 m³/sec respectively.

The Water Survey of Canada data collected for Rose Creek below Faro Creek (upstream of Station 2 on Rose Creek) is available for 1968 and 1969. The monthly means indicate the peak flow for the year is expected to occur during spring breakup (April-June) with a secondary smaller peak occurring before freeze up (Sept.-Oct.)(Hoos, 1973).

4.1.2 <u>Dissolved Oxygen</u>. Percentage dissolved oxygen (% DO) levels recorded at Stations 1-10 are illustrated in Figure 4 and listed in Appendix II Table 1. Rose Creek, Anvil Creek and north fork of Anvil Creek had stable percent dissolved oxygen (% DO) levels ranging between 85 and 98% saturation. Percent dissolved oxygen (%DO) at Station 3 (seepage from the toe of the Cross Valley dam) was 38% during both sampling periods. Station 9 (Pelly River) had a mean for both sampling

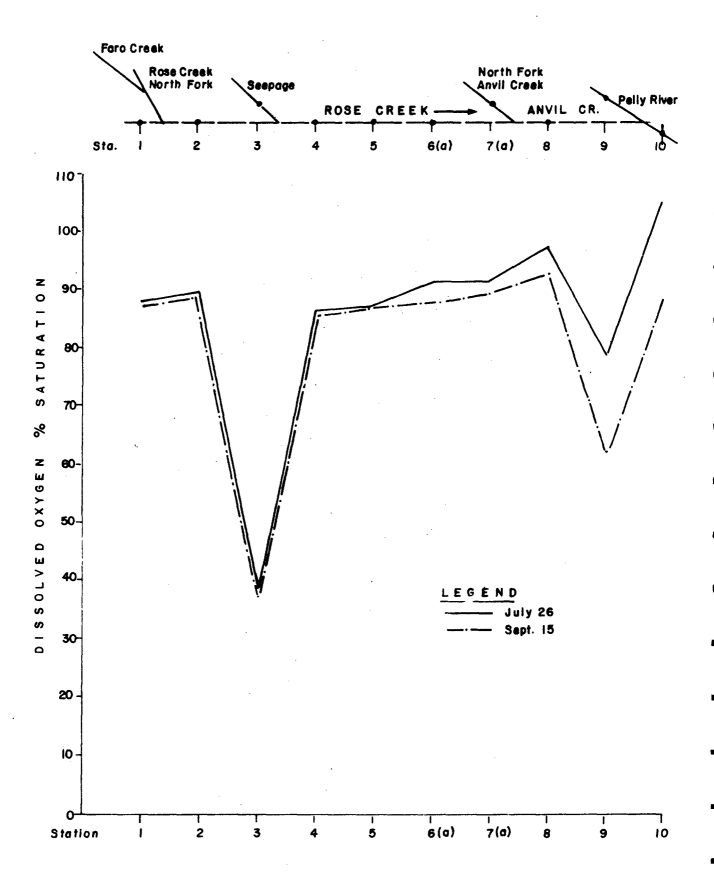


FIGURE 4 DISSOLVED OXYGEN-PERCENTAGE OF SATURATION-IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983

dates of 71% DO. The low % DO levels at Station 3 (seepage) indicate the source of flow may be partially from groundwater origin. The origin of the seepage (Station 3) is discussed to a greater extent later in the text. The results at Station 10 (Pelly River downstream of Anvil Creek confluence) suggest that Anvil Creek enhances the dissolved oxygen content in the river, at least along the north bank where the plume from Anvil Creek is located.

4.1.3 pH. The pH of the water in the study area tends to increase uniformly progressing downstream from Rose Creek Station 1 (pH 7.3) to Anvil Creek Station 8 of (pH 8.1). Laboratory pH varies less than 0.3 pH units between samplings on July 26, 1983 and September 15, 1983. Station 3 (seepage) averaged a pH of 7.6 whereas Station 2 (Rose Creek) averaged 7.8. McNeely (1979) indicates surface water generally tends to be alkaline, whereas ground waters are more acidic.

The data presented by Hoos (1973) and Baker (1979) present a similarity in the range and the spatial variation of pH as found in this study. Weagle (1981) found higher pH values ranging between pH 8 and pH 10 as far as Stations 4 and 5. The cause for these higher values was not discovered but suspected to be the reagent additives (lime) in the mill effluent.

d.1.4 Conductivity and Filterable Residue. Conductivity gives a measure of the ions in the water while filterable residue is a measure of the exact content of dissolved solids. As shown in Figure 5 for both parameters, Station 3 (seepage) adds a high input of mineral to the water. This impacts on the stream by increasing the level of minerals downstream relative to the upstream stations (Stations 1 and 2, Rose Creek). Once the seepage and Rose Creek are uniformly mixed, the level of mineral at downstream Station 5 (252-289 umhos/cm), is twice that of Rose Creek upstream of any mine influence (108-141 umhos/cm).

The conductivity values have varied temporally over the years Hoos (1973) reported spatial similarities in conductivity at most

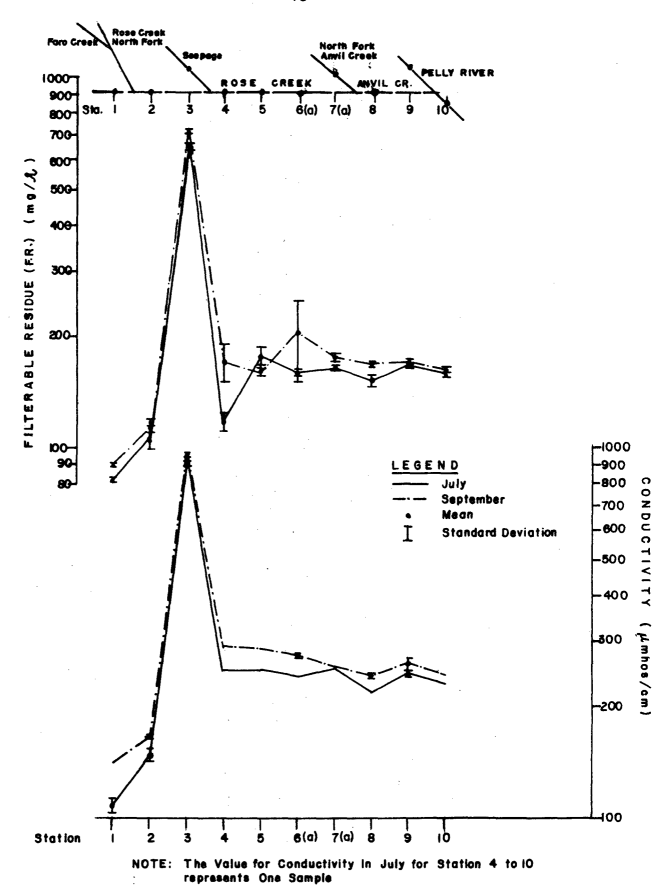


FIGURE 5 MEAN VALUES OF FILTERABLE RESIDUE (F.R.) AND CONDUCTIVITY IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983

stations. In 1974 and 1976 all EPS laboratory conductivity values of downstream stations (4-10) ranged between 150-200 umhos/cm and tended to be lower than results of this study (228-289 umhos/cm). In 1975, the readings were in the same range as this study except for Station 2 (Rose Creek) which was lower (Baker, 1979). Baker (1979) has indicated that for the years that seepage from the tailings pond (EPS Station 3, X-13) has been measured, the conductivity has fluctuated greatly, from 136 umhos/cm to 94010 umhos/cm. Conductivity of the seepage at Station 3 (X-13) during this study had lab conductivity of 929-955 umhos/cm.

- 4.1.5 <u>Color</u>. Color of the water is a result of metallic ions and soluble coloured organic compounds remaining after the turbidity is removed. In July, color was equal to or higher than the September samples at upstream Stations 1-4. Downstream stations (Stations 5-10) in July had values less than or equal to the September samples. This may reflect heavy rains a few days prior to sampling and the increase of organics leached from the land which contain various humic acids that contribute to color.
- 4.1.6 <u>Turbidity</u>. Turbidity is an expression of the optical property in water that causes light to be scattered and absorbed rather than be transmitted in straight lines through a sample. The turbidity measured during this study is generally low, in the range of 0.10-0.20 FTU, but did show an increase at Pelly River Stations 9 and 10 with values up to 4.4 FTU (Figure 6). Turbidity at Pelly River Station 10 was slightly lower than Station 9 Pelly River as a result of the influence of Anvil Creek outflow. The higher level of turbidity in the Pelly River on the July sample date may be attributed to an increased level of rain induced colloidals and organic acids.

Comparison of our results with the previous studies (Hoos 1973, Baker 1979, Weagle 1981) clearly shows that low turbidity levels at sampling stations existed in 1983. Turbidity level comparisons of 1973 to 1978 are generally 10 to 20 times greater and 30 times greater in 1974, than our results. For those years, the turbidity levels at Rose Creek control station (Station 1) varied between 0.55 and 2.50 FTU higher than observed in 1983.

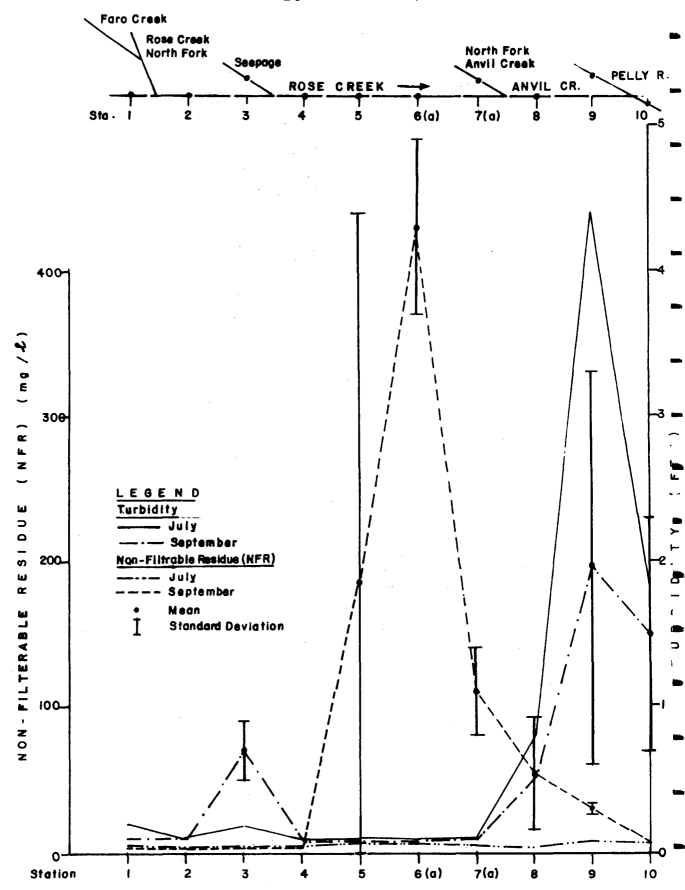


FIGURE 6 MEAN VALUES OF TURBIDITY AND NON-FILTERABLE RESIDUE (NFR) IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15,1983

- 4.1.7 <u>Non-filterable Residue (NFR)</u>. The NFR or suspended solids is that portion of a sample remaining on a filter after filtration. NFR values (Figure 6) for all stations in July and September were at the detection limit (5.0 mg/l) except Station 5-10 in September where an incident caused by increased precipitation and resulting erosion was recorded. Figure 6 indicates a lack of correlation between NFR and turbidity values, particularly at Stations 5 to 7 in September when increased NFR values were not accompanied by an expected increase in turbidity. Station 5 (Rose Creek) in September, shows very high NFR variability as reflected by the standard deviation, indicating this Station to be a mixing zone.
- 4.1.8 Alkalinity and Hardness. Alkalinity is a measurement of the power of a solution to neutralize hydrogen ions (EPS, 1976). Hardness is principally determined by the sum of calcium (Ca) and magnesium (Mg). Alkalinity and hardness values follow similar patterns (Figure 7) temporally and spatial. Hardness and alkalinity were elevated in the tailings seepage (Station 3) relative to creek and river stations.

Station 7(a) (Anvil Creek) shows slightly higher alkalinity values than do stations on Rose Creek or further downstream. The difference in alkalinity between Rose and Anvil Creeks may be due to the neutralization of acidity generated from the tailings ponds at the Cyprus Anvil mine site. Previous studies (Hoos, 1973, Baker 1979, Weagle 1981) have shown similar patterns of alkalinity and hardness.

4.1.9 Total Phosphate. Phosphate levels are low for both Rose and Anvil Creeks. They vary from below detection limit (0.0050 mg/l) to 0.0080 mg/l \pm .002. Station 9 (Pelly River) had a phosphate level on July 25 of 0.016 mg/l \pm 0.002. These are lowered at Station 10 (Pelly River) as a result of Anvil Creek input to 0.010 mg/l \pm 0.001. Phosphates are eliminated and recycled fairly quickly in streams and small quantities may actually occur in solution in the water (Hynes 1972).

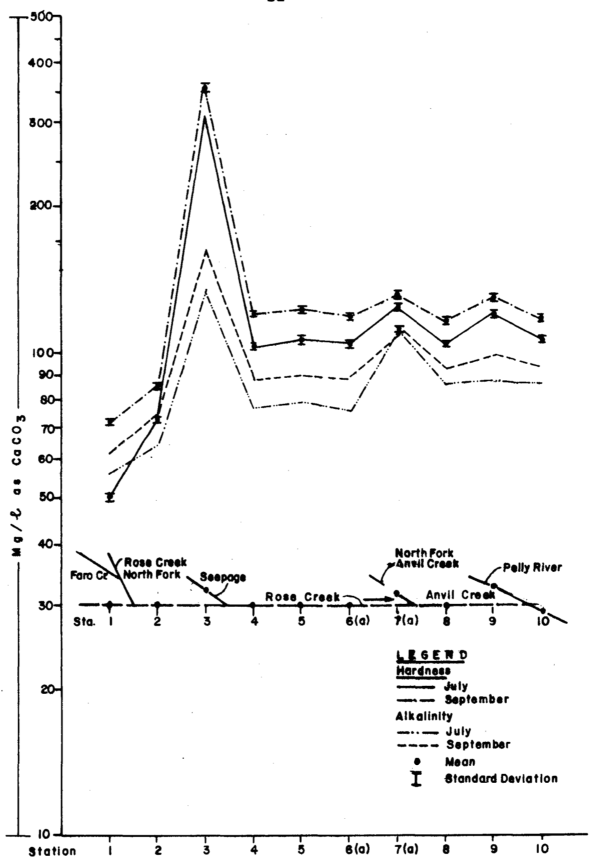


FIGURE 7 MEAN VALUES OF ALKALINITY AND HARDNESS IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983

4.1.10 The levels of ammonia (NH3) in the seepage water Ammonia. (Station 3) of 0.5 to 0.7 mg/l, are 2 orders of magnitude higher than in Rose Creek (Station 1)(Appendix II, Table 1). Ammonia (NH3) is present in groundwater, particularly at colder temperatures (EPS, 1972). The toxicity of ammonia decreases with temperature, attributable to a decrease in the concentration of the un-ionized ammonia fraction, but increases at low oxygen concentrations. Yake and James (1983) report that at a pH of 7.5 to 8.0 and a temperature of 5°C, the percentage of un-ionized ammonia in an ammonia water solution is 0.4 to 1.0. Ammonia (NH₃) toxicity at Station 3 (seepage) increased at low oxygen levels to a concentration of 0.0106 mg $NH_3/1$. Recommended toxicity of ammonia is not to exceed 0.020 mg NH₃/1 (Thurston et al, 1979). synergism occurs with cyanide, and ammonia toxicity is additive to the toxicity of zinc and copper (Thurston et al, 1979).

4.1.11 <u>Nitrite and Nitrate</u>. On July 25, 1983 nitrite levels at all stations were below the detection limit (<0.0050 mg/l) while in September the means of each station ranged from 0.0090 ± 0.0000 mg/l to 0.013 + 0.0007 mg/l (Appendix II, Table 1).

The mean nitrate values ranged from less than the detection limit of 0.010 mg/l to 0.065 ± 0.004 mg/l. The amount of nitrate was higher in September than in July. The levels of those two nitrogen compounds were very low at the upstream Rose Creek stations (Stations 1 and 2) but increased significantly in September at the seepage location (Station 3). Knowing that the ammonia level is higher in the seepage at Station 3 (Appendix II, Table 1) indicates the process of transformation to a more oxidized form may occur. This transformation may also contribute to a certain extent to deplete the percent dissolved oxygen (% DO). Another cause of the difference between the two sample periods may be attributable to the rainfall and surface runoff experienced, which may contribute to increases in the level of nitrate in the stream (Hynes 1979).

4.1.12 Sulphates. The level of sulphates (Figure 8) at Station 3 (seepage) was 30-40 times greater than upstream Rose Creek Stations 1 and 2 (Appendix II, Table 1) on both sample dates. The mean amount varies in September from 468 + 10.6 mg/l (Station 3 seepage) to 10.4 + 0.07 mg/l (Rose Creek Station 1). This chemical is one of the major compounds responsible for the increase of conductivity, hardness, alkalinity and filterable residue at Station 3 (seepage). potential hazard associated with a high concentration of sulphates, is that in acidic deoxygenated water, the sulfates could be reduced to the form of H₂S which is toxic to fish in its undissociated form to a level of 2 ug/l (Thurston et al 1979). In neutral and alkaline deoxygenated water, the sulfate will be reduced to the form of FeS which precipitates but when the iron is all utilized, the reduction will be to the form of H2S. At Station 3 (seepage), ionized iron is still available and the alkalinity pH indicates further buffering available therefore there is little concern for the transformation of sulphate to sulphite.

The effect of sulphur from CAMC is highlighted when comparing levels at Station 7(a) (Anvil Creek) and Station 6(a) (Rose Creek) with mean values of 19 mg/l and 46 mg/l respectively. Cprus Anvil's ores are massive sulfides consisting primarily of pyrite, pyrrhotite, galena and sphalerite. The tailings from this ore body are high in sulfide minerals and have been studied by others (Davidge, 1984).

4.2 Water Quality - Metals

4.2.1 Extractable Metal Analysis. The extractable metals in the waters of Rose and Anvil Creeks, the settling pond seepage and the Pelly River are given in Appendix II, Table 2. The mean (\bar{x}) values of the data from the July 26 survey is calculated from three grab samples while the September 15 survey is calculated on two grab samples.

For all the stations and on both sampling dates, silver (Ag), beryllium (Be), cadmium (Cd), cobalt (Co), chromium (Cr),

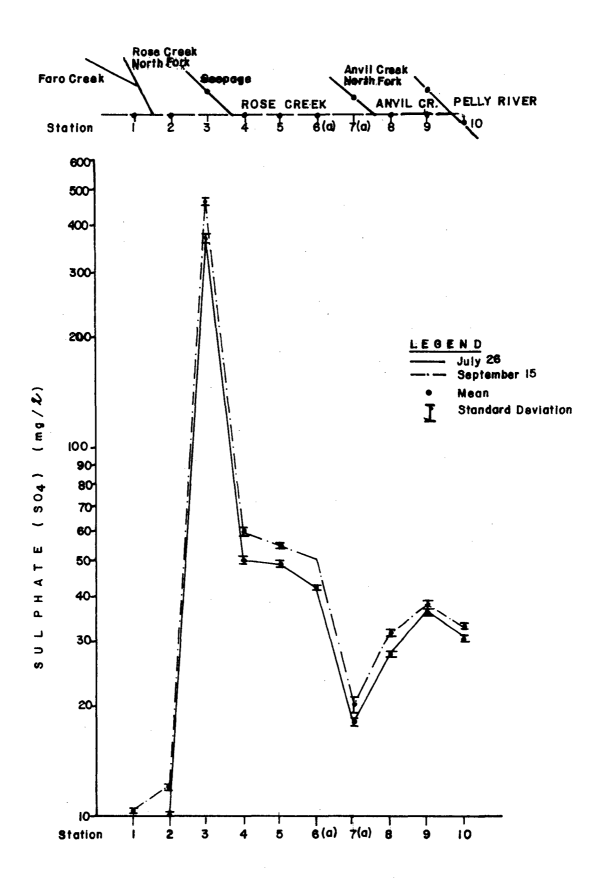


FIGURE 8 MEAN VALUES OF SULPHATES IN WATER SAMPLES
COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983

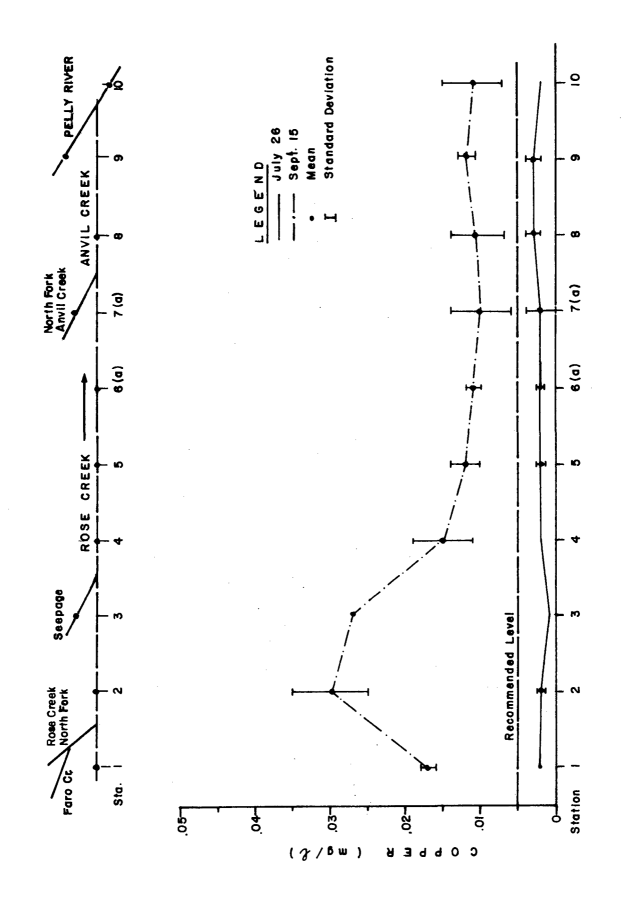
molydenum (Mo), antimony (Sb), selenium (Se), tin (Sn), and vanadium (V) values were below the EPS Laboratory detection limit for that element. Arsenic (As), boron (B), barium (Ba), nickel (Ni), lead (Pb), silica (Si) and titanium (Ti) levels were over the detection limit but below acceptable levels for drinking water and for aquatic life. These metals showed little fluctuation spatially and temporally along the creek, suggesting no impact from the mine.

Calcium (Ca), potassium (K), magnesium (Mg) and strontium (Sr) values were all less than the acceptable level for the protection of aquatic life. Seepage at Station 3 contained levels of these metals which exceeded Rose Creek (Station 1) by 3 to 6 times. Levels of these metals at Station 4 (Rose Creek) and further downstream, were lower than the seepage (Station 3) values but remained elevated over Station 1, indicating the influence of the seepage. This increase may be due to the input of those elements by erosion because comparable levels of metal exist in Anvil Creek at Station 7(a).

Manganese (Mn) and sodium (Na) at Station 3(seepage) were from 40 times to 150 times higher than the background levels measured at Stations 1 and 2 (Rose Creek). Associated dilution decreased Station 3 seepage level effect in Rose Creek (Stati n 6) to a value which remained marginally higher than Station 7(a) Anvil Creek.

Iron (Fe) and aluminum (Al) showed variation without pattern. Iron (Fe) levels increased 8 times at Station 3 (seepage) between sample dates but this is not seen at any other location. The aluminum (Al) at Station 7(a) (Anvil Creek) exceeds the recommended level for aquatic life as outlined by Health and Welfare Canada, on September 15, 1983, but was the only sample to do so.

4.2.2 <u>Copper</u>. The level of copper differs notably between the sampling dates (Figure 9). All samples collected on July 26, 1983 had copper levels below the recommended level for aquatic life while on September 15, all samples exceeded this level of 0.0050 mg/l. Demayo and Taylor (1981) suggest a limit of copper of 0.0020 mg/l total in water but also mention that existing levels and effects of local water



COPPER IN WATER SAMPLES COLLECTED 15, 1983. MEAN VALUES OF EXTRACTABLE ON JULY 26 AND SEPTEMBER თ FIGURE

quality must be taken into account when setting an objective for copper for specific bodies of water. The Yukon Territory Water Board require Cyprus Anvil to meet a level of 0.20 mg/l (extractable) for copper in the effluent discharge. Station 3 (seepage) exceeded this requirement on September 26, 1983.

According to Nriague (Demayo and Taylor 1981), approximately 25% of the copper is from natural sources (windblown dusts, vegetation exudates, volcanic emissions, sea salt spray) and the rest is from anthropogenic sources. In an intensive study of Susquehanna River by Troup and Bricker (Demayo and Taylor, 1981) it was found that the total metal concentrations (Mn, Fe, Cd, Cr, Co, Cu, Ni and Zn) were correlated with the discharge of the river and with the total suspended solids, thus in a period of high discharge, most of the metals were carried in the suspended solids phase. Another study by Jennett et al (Demayo and Taylor, 1981) monitoring the water quality of a region of mining and milling, found that during peak runoff periods following a storm, dissolved metal concentrations increased abruptly and suspended solids increased signficantly. The concentration of copper in the water may also increase by the formation of copper complexes. increase of organic acid concentration derived from complexing also results in increases of pH (Bolter and Butz, 1975, quote in Demayo and Taylor, 1981).

The analysis performed on our water quality is for extractable metal which include the dissolved metals, the absorbed metal on small inorganic particles and the organic complex which are digested by the acid. The highest peak recorded in September is at Station 2 (Rose Creek) which drains the northeast side of the mine property. Significant increases in Copper (Cu) levels at Station 1 (Rose Creek Control Station) may be attributable to the high flows in September.

Previous data on Rose Creek (Hoos 1973, Baker 1979, Weagle 1981) for extractable copper shows levels only at the detection limit of 0.01 mg/l, which is still twice the recommended level for safe aquatic life (Thurston et al, 1979). Generally, downstream stations in September had levels twice the detection limit of earlier studies. In

1978, significantly higher (6-7 times) copper (Cu) levels were recorded in Rose Creek (Baker, 1979). In July, copper (Cu) levels are below acceptable for drinking water (1.0 mg/l) at all stations.

4.2.3 Zinc. The levels of extractable zinc in the water were all close to or higher than the EPA recommended level for aquatic life of 0.030 mg/l (Thurston et al, 1979), except Station 3 (Seepage) on July 26 which was 0.0020 mg/l (Figure 10). DeMayo and Taylor (1981) suggest different acceptable levels of zinc based hardness on the classifications given in the Hydrological Atlas of Canada 1978. For a hardness as mg/l CaCO₃ O to 120, the maximum acceptable level of total zinc is 0.050 mg/l; hardness 120-180, the level is 0.10 mg/l of zinc; hardness 180-300, 0.2 mg/l zinc and over 300 the zinc level may be up to 0.30 mg/l.

The distribution of zinc in the water throughout Rose Creek, follows similar patterns as copper (Cu). September levels of zinc exceeded both recommended EPA and Health and Welfare Canada levels except Station 3 (seepage) which, because of hardness (300 mg/l), did not exceed the Canadian guideline. Zinc levels of upstreams stations are high in the fall and it may be caused by high flows increasing erosion into the stream. The values observed at Station 2 (Rose Creek) which were the highest value recorded in this study, are attributed to the drainage of the east part of the mine by the the north fork of Rose Creek. The water licence issued by the Yukon Territory Water Board, identifies the maximum level of zinc permitted in the effluent to be 0.50 mg/l zinc (extractable). All stations fell within this requirement.

Demayo and Taylor (1981) suggested that in case of high natural levels, the objectives for safe aquatic life should meet the natural level.

The background levels of zinc (Station 1) have increased since the baseline study was completed in 1973. At that time zinc (Zn) showed less than 0.010 mg/l as total zinc (Hoos, 1973). In 1974, background levels increased to a mean of 0.275 mg/l + 0.5

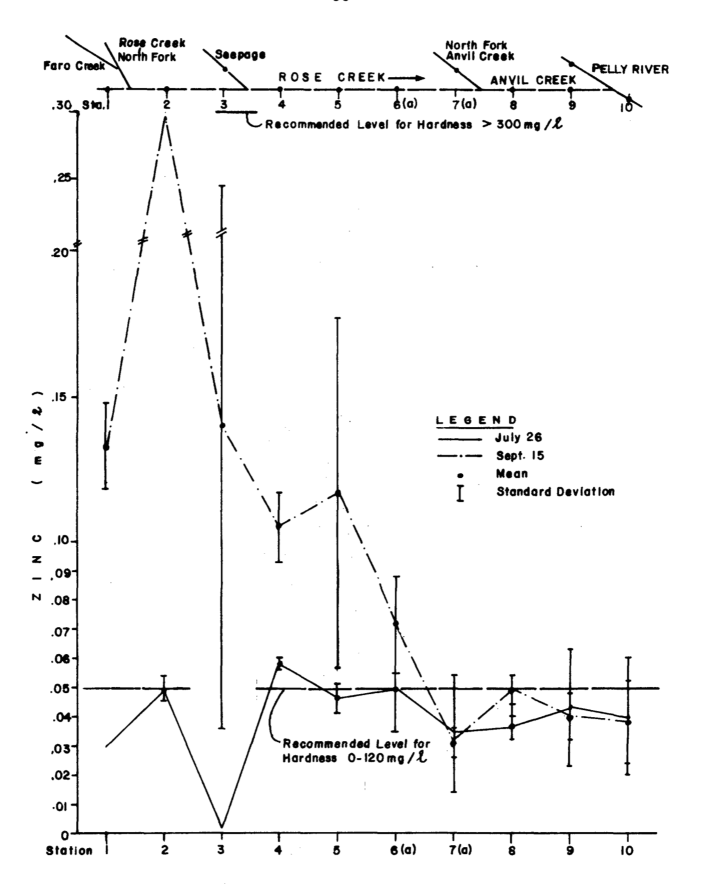


FIGURE 10 MEAN VALUES OF EXTRACTABLE ZINC IN WATER SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983

giving an idea of the variability of the zinc content at that station. During the following years (Baker 1979, Weagle 1981), zinc (Zn) content stabilized at 0.10 mg/l. Zinc levels decreased from Station 4 (Rose Creek) highs as further dilutions occured towards downstream stations. The downstream level in previous years, are quite similar to what we see in our present study (Baker 1979, Weagel 1981). Zinc (Zn) levels at downstream stations from this study are similar to previous studies (Baker 1979, Weagle 1981).

4.3 Sediment

4.3.1 <u>Particle Size Analysis</u>. Results of the particle size analysis are shown in Appendix III, Table 1. When analyzing and discussing the results of sediment analysis, it is necessary to be aware of the variability that can occur among replicate samples due to natural variation in the streambed composition.

Results show a high percentage of gravel except at Station 7(a) (Anvil Creek) and 10 (Pelly River), at which fine sand and silt are predominant. When sediment samples were collected, all material larger than 32mm (pebbles) was selectively removed by hand and discarded. Therefore, the particle size distribution data must be handled with caution.

Particle size affects leachable metals results and the benthic invertebrate distribution and abundance (to be discussed in following sections). Only that portion of sediment smaller than 149 um was analysed for leachable metals since these particles can adsorb more metals on the surface which affect the aquatic environment.

4.3.2. <u>Sediment Metal Analysis</u>. The results of analysis for metals levels in the sediments are presented in Appendix III Table 2.

The metals analysis shows variability in the results. Only cadmium (Cd), molybdenum (Mo), silicon (Si), and titanium (Ti) show significant changes between sample dates (p<0.05) using Student's

t-distribution. Cadmium (Cd) and silicon (Si) are higher in July while molybdenum (Mo) and titanium (Ti) are higher in September at Station 4 (Rose Creek). Other significant differences occur at Stations 3 (seepage) and 8 (Anvil Creek) but cannot be tied into an obvious pattern (Figure 11).

An overall view of the leachable metal concentrations in the sediments exhibit a general trend of increase from lows at Station 1 to highs at Stations 4, 5 and 6(a) (Rose Creek) followed by a decrease to Station 10 (Pelly River). This particular increase in heavy metals (Ag, As, Ba, Cu, Fe, Hg, Mo, Pb, Zn) at middle stations on Rose Creek (Figure 12), may be explained by the remaining effects of a tailings pond spill in March 1975, when 2.45 x $10^5 \, \mathrm{m}^3$ of tailings slurry contaminated Rose Creek. In 1975, the slurrys were seen up to the confluence of Rose Creek and Anvil Creek. The cadmium (Cd) results for September did not follow this general increase/decrease trend.

The present study results show copper (Cu), zinc (Zn), arsenic (As) and Barium (Ba) to have higher levels present at middle stations (Stations 4 and 5) than existed in 1973 (Hoos, 1973). The 1973 detection limits were too high for comparative assessment of cadmium (Cd) with results from this study.

4.4 Bottom Fauna

The use of the benthic invertebrates as indicators of stream quality is guided by some biological considerations. Invertebrates have relatively low mobility which reflect conditions of that particular location. In stream pollution, the variability of contaminant discharges can be integrated by the benthic community and give an idea of pollution over an extended period, which is not the case with water grab samples.

4.4.1 <u>Artificial Substrate Sampler</u>. The artificial substrate sampler allows collection of data from locations that cannot be sampled effectively by other methods. It eliminates subjectivity of choice and

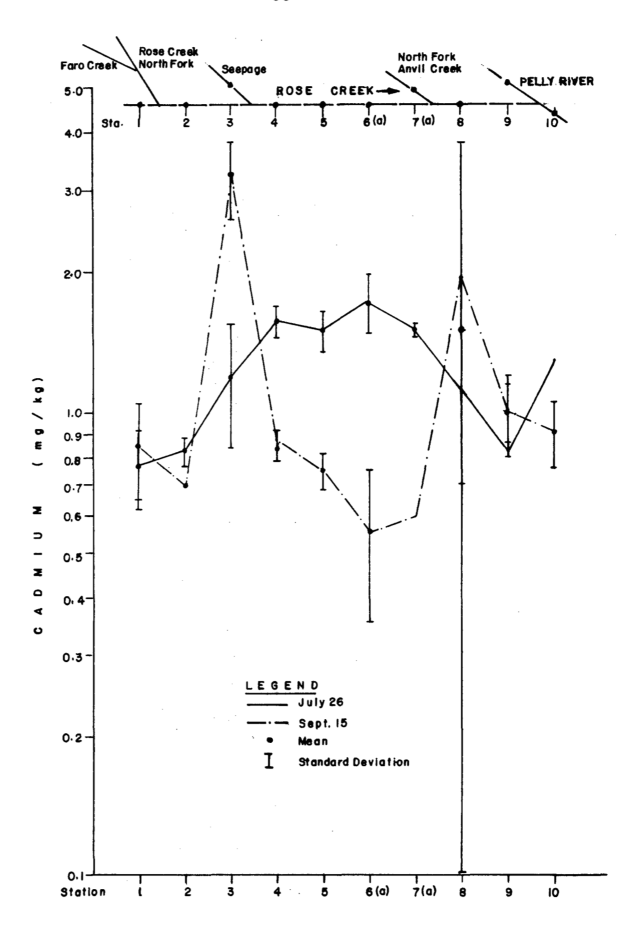


FIGURE II MEAN VALUES OF EXTRACTABLE CADMIUM IN SEDIMENT SAMPLES COLLECTED ON JULY 26 AND SEPT. 15, 1983

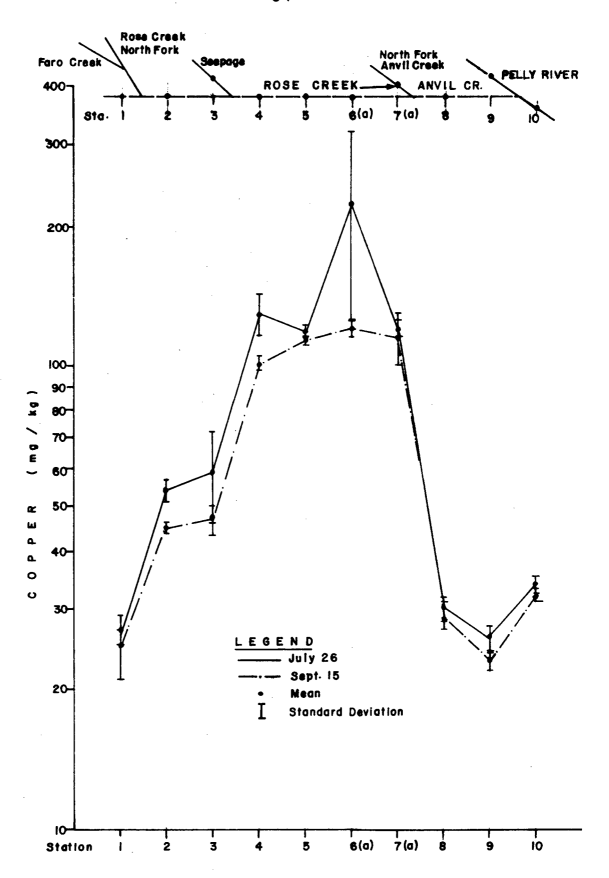


FIGURE 12 MEAN VALUES OF EXTRACTABLE COPPER IN SEDIMENT SAMPLES COLLECTED ON JULY 26 AND SEPTEMBER 15, 1983

sampling at a number of different stations, reducing confounding effects of differences in habitat substrate, depth and time (Rosenberg and Resh 1982). The variability in replicates using the artificial substrate sampler is reduced compared with other types of sampling methods (Khalaf and Tachet 1978, Rosenberg and Resh 1982).

EPS (unpublished data) has experimented with the artificial substrate sampler to determine the sampling effectiveness and the number of replicates required to obtain an adequate sample. The results of this work with this type of sampler indicates that 3 replicates are required to obtain an adequate sampling to represent the community.

The samplers were retrieved after 50 days colonization time. The samplers at Station 8 could not be retrieved on September 15, 1983 due to the high water conditions and fast current. Station 3, due to low water levels could not meet the requirement of depth, current and texture of substrate particles. All substrate particles were completely covered with ferrous (Fe) precipitate at Station 3 (seepage) and for this reason, Station 3 can be considered as atypical in the sampling program.

4.4.2. <u>Taxonomic Features</u>. The taxonomic levels involved in the discussion may refer to the class, order, family or genus. Organisms which are rare and are not well represented are referred to at the class level. Class Insecta had the greatest variety of organisms present. Separation of Class Insecta into orders is usually based on functional morphology structures. The Diptera were too diverse to specifically comment on at that taxonomic level and so were analysed at the family level. Comparisons between specific organisms was dealt with at the genus level.

A total of 32,994 organisms were collected from the 9 stations (substrate samplers were lost at Station 8). They comprised 71 different taxa, of which 51 are at the genus level. A taxonomic list of the benthic invertebrates collected during the survey on September 15, 1983 is given in Appendix IV, Table 1. The complete

description of the numbers and percentage abundance of organisms sampled at each station and their relative density per cubic meter is found in Appendix IV, Table 2. Appendix IV, Table 3 provides a summary comparison of diversity indices for benthic invertebrate data from previous studies, including monitoring programs sponsored by CAMC.

The greatest numbers of genera are found at Station 1 with 41 genera, followed by Stations 10, 5 and 4 with 37, 32 and 31 genera respectively. The lowest number of genera, 17, occurred at Station 3.

The abundance varies from station to station. The greatest abundance can be found at Station 10 with 7407 organisms (433,158 organisms per metre³). Stations 4 and 5 had 5187 (303,333 organisms per m^3) and 5436 (317,895 organisms per m^3) organisms respectively. The lowest number of organisms is found at Station 7a (Anvil Creek) with only 2094 individuals (122,456 organisms per m^3). The low abundance of individuals at Station 7a partially caused by a loss of rock substrate from one of the 3 replicate artificial substrate sampler which broke open during the survey. Only a quarter of the basket was filled at time of retrieval.

The two dominant genera are <u>Capnia</u> sp (Plecoptera) and <u>Baetis</u> sp (Ephemeroptera). Both are found at all static is and represent 18.5% and 17.6% respectively of the total organisms encountered at the genus level.

Seven genera were present at all stations of which three are representatives of the order Plecoptera, one in the order Ephemeroptera and three in the Family Chironomidae. Seven other genera, three in the order Plecoptera, two in the order Ephemeroptera, one in the order Trichoptera and one in the family Simuliidae were present (Table 3) at all stations except Station 3 (seepage). Table 4 lists the percentage composition of major taxa at each station.

The percentage dissolved oxygen (%DO) present at Station 3 (seepage) during the two surveys is below 40% saturation. The nymphal stage of the Plecoptera are typically found in water with a relatively high oxygen content (Wetzel, 1975), and would account for a very low representation of Plecoptera in the seepage from the Cross Valley Dam

(Station 3). The high percentage composition of Plecoptera at Stations 9 and 10 cannot be explained only by the oxygen levels. The use of an artificial substrate sampler may provide a better habitat for the Plecoptera than for other groups therefore their density relative to other groups increases.

The Ephemeroptera (mayflies) also require high levels of dissolved oxygen for optimum development and their percentage composition is also very low at Station 3. Sprague et al (Winner, 1980) indicates that mayflies are sensitive to copper and zinc. Cairns and Dickson (1971) consider the Ephemeroptera as a pollution sensitive group. The results of the present study indicate that the distribution of the Ephemeroptera does not seem to be influenced by the levels of extractable zinc and copper present in the water. Station 2, had the highest zinc and copper levels and was also the station at which all the genera of Ephemeroptera found in the survey were represented. Wetzel (1972) indicates that substrate type and specific sediment particle size may restrict species distribution. Khalaf and Tachet (1980) have shown that the Ephemeroptera is the only order of aquatic insects that show differences in numbers recovered using the artificial substrate sampler. The texture of the rocks and substrate, as well as the low dissolved oxygen levels at Station 3 (seepage) may partially explain the poor representation of Ephemeroptera.

The Trichoptera sampled at all stations were few, with a mean percentage composition of 2.07%. Station 6(a) (Rose Creek) shows the highest percentage (10.94%) of order Trichoptera and station 3 (seepage) the lowest (0.04%). The percentage at Station 6(a) is due mainly to the presence of an Hydropsychidae, Arctopsyche sp., which constitutes 9.53% of the population at that station.

Although the Chironomids (Family Chronomidae) represent 87% of the population at Station 3 (seepage), no common chironomid genera were particularly dominant. The two major genera involved in the population are <u>Brillia</u> sp. (48%) and <u>Phaenopsectra</u> sp. (29%), both of which are well represented only at Station 3 (seepage). This and other data supports the diversity of tolerances of species within the family.

TABLE 3 GENUS PRESENCE OF BENTHIC INVERTEBRATES AT THE VARIOUS STATIONS

Α.	Present	at	all	the	static	ns

GENUS

ORDER OR FAMILY

Capnia sp
Zapada sp
Brachyptera sp
Baetis sp
Cardiociadius sp
Cricotopus sp
Heterotrissociadius sp

Plecoptera
Plecoptera
Plecoptera
Ephemeroptera
Chironomidae
Chrionomidae
Chironomidae

B. Present at all stations except Station 3

GENUS

ORDER OR FAMILY

Amphinemura sp
Prostaia sp
Isoperia sp
Ephemerella sp
Cinygmula sp
Brachycentrus sp
Prosimulium sp

Plecoptera
Plecoptera
Plecoptera
Ephemeroptera
Ephemeroptera
Trichoptera
Simulidae

C. Unique to Station 1

GENUS

CLASS OR ORDER OR FAMILY

Hydra sp
Paranais sp
Corynoneura sp
Lymnacea sp
Valvata sp
Pisidium sp

Hydrolda Oligochueta Chironomidae Gastropoda Gastropoda Pelecypoda

D. Unique to Station 10

GENUS

ORDER OR FAMILY

Lepidostoma sp Onocosmoecus sp Paralauterborniella sp Trichoptera Trichoptera Chironomidae

TABLE 4 PERCENTAGE OF DIFFERENT TAXONOMIC GROUPS REPRESENTED AT EACH STATION

	STATION								
	1	2	3	4	5	6(a)	7(a)	9	10
Plecoptera	16.28	39.98	0.31	41.43	24.27	42.44	21.59	52.17	62.87
Ephemeroptera	25.86	25.26	0.18	36.34	42.17	37.67	41.87	17.74	3.29
Trichoptera	0.4	0.15	0.04	0.94	0.73	10.94	2.95	0.44	2.10
Chironomidae	52•38	9.43	86.81	12.54	10.38	3.94	30.34	27.07	30.83
TOTALS	94.92	74.82	87.34	91 . 25	78.55	94.99	96•75	97•42	99•09
		·							

Another apparent example of the specificity of habitats occupied by genera is the presence of <u>Corynoneura</u> sp. at Station 1 (Rose Creek) only. Lenat (1983), showed that moderately stressed streams present an ideal environment for small grazing chironomids in that algal food and refugia are abundant and the density of competitors, especially Plecoptera, Trichoptera, Ephemeroptera and Gastropoda may be reduced. Chironomids found in stressed environments usually have a very short life cycle and are capable of quick colonization and exploitation of disturbed areas subjected to recurring acute toxicity. If pollution becomes severe, the chironomid richness will decrease so a direct correlation between pollution and chironomid populations cannot be made (Lenat, 1983).

The unique genera found at Stations 1 (Rose Creek) and 10 (Pelly River) generally represent a percentage lower than 1%. Phylum Coelenterata, Class Gastropoda and order Pelecypoda are only found at Station 1 (Rose Creek). Hydra seems to be common in ponds and very clear water (Meglitsch 1972) and Pisidium seems to be a good indicator c? a copper stressed region (Winner et al 1980). Phillips (1977) argues that the bivalve molluscs as well as the macro-algae constitute the best indicators of heavy metals pollution.

A comparison of the taxonomic group dominance from data at and downstream of the Cyprus Anvil mine area, show a refaction of the dominance held by the Chironomidae in downstream areas. In 1974 the Chironomidae were dominant as far downstream as Station 6 (Anvil Creek) In 1974 and 1976, the dominance of chironomids was still visible but only to Station 4 (Rose Creek) and in 1983 such dominance disappeared even at Station 4 (Figure 13). The dominance by the chironomidae in Rose Creek is presently only found at Station 1 (control station) and at Station 3 (seepage from ponds).

of large amount of information about the numbers and kinds of organisms. The ecological basis for the diversity indices is that in a polluted area small numbers of species are found with large numbers of individuals, while in clear waters small numbers of individuals and many different species are found (Wilhm, 1970). The diversity is affected by several

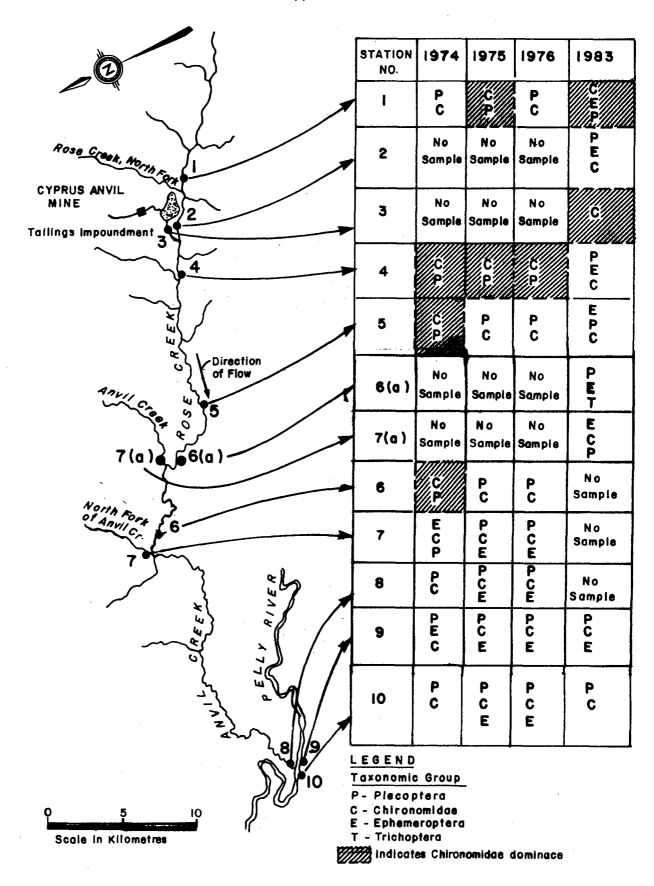


FIGURE 13 INVERTEBRATE TAXONOMIC GROUP DOMINANCE IN CYPRUS ANVIL MINE STUDY AREA

factors other than pollution and great emphasis of consistency in sampling must be shown (Hugues 1978). Depth and duration of sampling seem not to affect the diversity value but sample size, season and taxonomic level used in calculations, may change the values (Resh and Unzicher 1975).

The diversity indices calculated in this study show a peak at Station 1 (Rose Creek) with 1.07 and a second peak downstream at Station 6(a) (Rose Creek) and 7(a) (Anvil Creek) with 1.02 and 1.03 respectively. The lowest diversity was found at Station 10 (Pelly River) with 0.59 and at Station 3 (seepage) with 0.60 and both have a low evenness of 0.38 and 0.49 respectively. The diversity index at Station 10 (Pelly River) is low if compared to the density and number of species. The effect of <u>Capnia</u> sp. (60% of the population) on the diversity is obvious. At Station 3 (seepage), the low number of species as well as the low number of individuals present, is a reflection of the high mineral and lower content of dissolved oxygen (DO) of the groundwater seepage at that station.

The index values of diversity at most stations are higher than in the previous years as shown in Table 5 indicating an improvement in the benthic invertebrate community since the line operation has shut down. These results must be taken with reservation because of the variation in bottom fauna collection methodology of this study and previous others. Perkins (1983) has demonstrated that inconsistency of the results of the diversity indices is subject to variability due to differences in concentrations of copper and proposes community comparison indices as an alternative.

4.4.4 <u>Community Index.</u> The community index used in this report is calculated as the percent similarity index (Psc). The Psc is not as sensitive to rare species and weight differences in more dominant species (Brock 1977). The Psc can detect subtle shifts in community composition which can indicate long term trends of change. Table 6 shows the similarity index matrix for the 1983 monitoring program. Station 2 (Rose Creek) shows good similarity with Stations 4 and 5

TABLE 5 COMPARISON OF DIVERSITY INDICES FOR SAMPLE STATIONS SINCE 1973

STATION	1973(a)	1974(b)	1975(b)	1976(b)	1977(c)	1980(d)	1981 (e)	1982(f)	1983
-	69.0	0.64	0.64	92.0	94.0	1.2.1	0.97	1.14	1.07
2	•	í	1	ı	1	ı	ı	ı	0.93
М	0.71	f		ı	ı	ı	ı		09.0
4		0.50	0.15	0.67	0.34	0.68	0.64	0.81	0.93
īv.	0.48	0.38	0.39	0.84	0.88	0.44	0.78	•	0.87
· •	0.78	09.0	0.27	0.85	61.0	0.58	0.70	0.52	ŧ
1983 EPS 6		1		1	•	•		1	1.02
7	0.82	0.47	69*0	0.74	0.78	ı	1	ı	i
1983 EPS 7		1	•				•	1	1.03
7.0*	•	ſ	ı	ı	ı	1.01	080	0.70	ı
ω	0.57	0.45	0.23	0.52	96•0	ı	r	i .	1
σ,	0.35	0.65	0.87	0.78	0.79	•	1	ı	0.88
10	0.22	0.25	0.54	ı	96*0	ı	ı	ı	0.59
(a) Hoos,R.A. (b) Baker, S. (c) Weagle, K (d) Weagle, K (e) Wealge, K (f) Weagle, K	Hoos,R.A.W. and Holman, Baker, S.A. (1979) Weagle, K. (1981) Weagle, K. (1980) Wealge, K. (1981a)	W.N. (1973)) - Surber Sampler - Artificial sub: - Artificial sub: - Artificial sub: - Artificial sub:	Sampler lal substrate lal substrate lal substrate lal substrate	sampler sampler sampler sampler and omes sampler and omes and one of the other and omes and other	baskets baskets baskets baskets			
1983			- Artificial	- 1	sempler	baskets			

*Station just below the confluence of Anvil Creek and Rose Creek.

(Rose Creek). Stations 4, 5, 6(a) (Rose Creek) and 7(a) (Anvil Creek) show a good interstation degree of similarity (all are over 50%). Stations 9 and 10 on the Pelly River show a high degree of similarity in their benthic community (73%). Station 1 (Rose Creek), does not show a high degree of similarity with any of the downstream stations and is most similar to Station 7(a) (Anvil Creek), another station not effected by the mine site. Station 3 (seepage), which has distinctive water quality characteristics and a different chemical substrate composition, possesses a community structure with very little similarity to other stations.

4.5 Biota-Algae

The results of the analysis for extractable metals are shown in Appendix V, Table 1. Both wet and dry weight are given, however the analysis is based on the dry weight concentration to facilitate comparisons. The results involve a single sample on July 26, 1983 and a mean composed of three samples on September 15, 1983.

The single aglae sample for July possesses metal levels (Cadmium (Cd), Zinc (Zn), Calcium (Ca), Nickel Ni), Lead (Pb), Strontium (Sr), Vanadium (V) and Arsenic (As)) generally higher than the mean of the replicate samples collected in September.

The observed differences in extractable metals concentrations between July and September may be due to the age of the algae and the duration of exposure to the metals (Foster 1982, Phillips, 1977). In September the age of the algae was known to be less than 50 days because the samples were collected from the surface of the artificial substrate sampler which had been deposited 50 days previously. The age of the colonies collected in July is unknown but presumed to be older. Attempts to rinse sediment from the algal mat was made for the July samples but no such attempts were made for the September samples. The accumulation rate of heavy metals also seemed to increase inversely proportional to the temperature (Phillips, 1977). The co-existence of several heavy metals that compete for binding sites may change the

TABLE 6 THE PERCENT SIMILARITY INDEX MATRIX FOR THE 1983
MONITORING PROGRAM

STATION	2	3	4	5	6(a)	7(a)	9	10
1	36.24	10.95	43.19	40.99	39.81	46.12	40.34	20.05
2		4.24	58.32	67.15	45.25	40.60	23.21	10.84
3			3.00	12.25	11.91	2.65	2.97	3.65
4				77.75	71.02	62.23	24.53	11.11
5					55.14	50.43	22.28	11.08
6(a)						63.91	37.47	27.87
7(a)							42.64	37.59
9								73.09

uptake of an element due to the presence of a competitor metal and lead to different heavy metals levels in the algae (Phillips, 1977). For the above reasons, it is not possible to draw conclusions using the algae present as an bio-indicator of the habitat conditions due to the effect of environmental variables.

Establishing a relationship between the algae found (class Bacillariophyceae) and the environmental conditions is difficult. Foster (1982) indicates that algal diversity as measured by the number of species and their relative abundance appeared to be reduced by metal pollution, particularly at sites high in copper (Cu) and zinc (Zn). An experiment by Januszko (DeMayo and Taylor, 1981) with members of the Bacillariophyceae showed an increase of biomass from 27 g/m 3 of algae in copper treated ponds. No analysis of the periphyton community from these aspects was performed during the present study. The algae sample collected at Station 3 (seepage) was present in large quantities and maybe one species specifically adapted to exploit the unique conditions that exist at Station 3 (seepage).

4.6 Bioassay

The results of the bioassay performed on the Cyprus Anvil tailings pond seepage (Station 3) and Rose Creek control station (Station 1) are shown in Table 7.

The <u>Daphnia pulex</u> acute toxicity test performed on the control station ran for 48 hours and for 96 hours on the seepage samples. Both show an $LC_{50}>100\%$.

Experimental threshold concentration using <u>Daphnia magna</u> with various metal ions (Biesinger and Christensen, 1972) indicates that our metal effluent concentrations were always below the $48h-LC_{50}$ determined. The response of <u>Daphnia pulex</u> and <u>Daphnia magna</u> is assumed to be approximately the same under similar conditions.

The control dilution (0% treatment) had one death per test which falls well within acceptable limits to validate the test, which allows up to 20% mortality in the control (DOE, 1980).

Considering those aspects of the test, the seepage water from Station 3 on July 26, 1983 is considered of very low toxicity under the test conditions and parameters of this bioassay.

5 ENVIRONMENTAL COMPLIANCE MONITORING

5.1 Regulatory Requirements of the Cyprus Anvil Mine

The mine discharges a 'waste' within the meaning of the Northern Inland Waters Act and is required by that act to hold a water licence. In addition, certain provisions of the Fisheries Act, in particular the Metal Mining Liquid Effluent Regulations, apply to the mine's operation. Thus the mine may not discharge an effluent containing concentrations in excess of those stated in the regulations or prescribed in the licence granted by the Yukon Territory Water Board.

Water licences are obtained through application to the Yukon Territory Water Board and public hearings into each application are customary. The Board may attach conditions to any licence and among these are requirements for wastewater treatment and compliance monitoring. In March 1982, the Yukon Territory Water Board conducted a public hearing at Faro, Y.T., to consider the company's plans to utilize a new tailings facility and to amend its licence. Subsequent to granting the amendment, the Water Board required the company to collect water samples on a routine basis at the stations plotted in Figure 3 and listed in Table 3.

5.2 <u>Monitoring Data 1974-1983</u>

Appendix VI, Tables 1a, 1b and 1c, show historical water quality results for compliance monitoring sites X-3, X-14 and X-15 respectively as compiled from information in government files and consultants' reports. Station X-3 corresponds approximately to 1983 EPS Station 1 (see Table 1). Licence Station X-14 is comparable to Station 4, while licence Station X-15 is comparable to Station 4. Station X-14 when compared to Station X-3 shows an increase in the comparable parameters Ca, Mg, Mn, Na and SO₄ downstream, that is, within the influence of discharges from the mine. These discharges have been mainly

through seepages below the final or cross valley tailings dam in the long period since the mine curtailed production in June 1982, but they include surface discharges at X-5, or through large siphons. Thus the mine has had a measurable effect on receiving water quality in Rose Creek for many years and the results of the present study (Section 4) support a long-term trend.

Table 8 compiles Zn, Pb and Na values for selected compliance monitoring stations on the mine's property during the June 1982 - December 1983 period. Stations X-7, X-22 and X-23 show elevated values for Pb and Zn reflecting discharges to the tailings pond systems. The main pit has been pumped continuously throughout the study period. This water at point X-22 seeps through oxidized sulphide waste rock to emerge at X-23. Downstream this flow is diluted by scavenging and circulating water discharged from the mill complex, and by natural runoff. Station X-7 data reflects the quality of the combined flows - pit pumpings, mill water and natural drainage - passing into the tailings area and the first large pond.

Flows passing over and through the old tailings areas are joined by natural groundwater flows to determine the quality of water in the large intermediate dam pond. Table 9 presents values for zinc and pH at the decant of the intermediate dam (X-4) and other stations. An average value of 1.58 mg/l for zinc at X-4 compared to 12.99 mg/l at X-7 shows that either the zinc content seen at X-7 has been reduced by some filtering process in passing through the tailings or there is an eight-fold dilution of X-7 flows from other sources flowing into the intermediate dam pond. Since the decant from this pond forms only a small portion of the flow into the cross valley pond, it is reasonable to assume that filtration under the intermediate dam is effective in reducing zinc levels to those seen at X-5, or at the equivalent stations S-8, 8 inch siphon, and S-16, 16 inch siphon. Thus the system comprising two large ponds and filtration has enabled high zinc pit pumpings to be treated to levels suitable for discharge.

Table 9 presents linear correlations between zinc concentration and pH. Significant correlations of between -0.72 to -0.86 appear

TABLE 7 Daphnia pulex BIOASSAY RESULTS OF CYPRUS ANVIL TAILING POND SEEPAGE

Acute Toxicity Tests

Water sampled July 26, 1983 at Station 1 (control) Bloassay started August 3 at 11:30 AM.

Fifty percent lethal concentration $48 \text{ h} - \text{LC}_{50} = >100\%$

	BEFORE		TIME (hours)							
CONCENTRATION	ī	EST	T=0	T=2	T=4	T=24	T=48			
	pН	DO (ppm)	11:30	13:30	15:30	11:30	11:30			
Control O# Solution	8.05	8.0	10	10	10	10	9			
20% Solution	8.05	8.0	10	9	9	9	9			
40% Solution	8.05	8.0	10	10	10	10	10			
60% Solution	8.05	8.0	10	10	10	10	10			
80% Solution	8.05	8.0	10	10	10	9	8			
100% Solution	8.05	8.0	10	10	. 10	10	10			

Water sampled July 26, 1983 at Station 3 (seepage) Bloassay started July 28 at 11:30 AM.

Fifty percent lethal concentration 96 h - LC₅₀ = >100\$

,	BEFORE		TIME (hour:)							
CONCENTRATION	· 1	EST	T=0	T=2	T=4	T=24	T=96			
		DO								
	pН	(ppm)	11:00	13:00	15:00	11:00	11:00			
Control 0% Effluent	8.14	7.6	10	10	10	9	9			
20\$ Effluent	8-14	7.6	10	10	10	9	8			
40% Effluent	8.14	7.6	10	. 10	10	9	9			
60% Effluent	8.14	7.6	10	9	9	7	7			
80# Effluent	8.14	7.6	10	10	8	7	7			
100% Effluent	8.14	7.6	10	10	9	. 8	8			

WATER QUALITY, SELECTED PARAMETERS - JUNE 1982 - DECEMBER 1983 (values extractable, mg/l)

TABLE 8

i		<u> </u>								
	STANDARD DEVIATION	4.99	2.24	26.10	2.46	47.8	35.0	22.67		
SOD 1 UM	AVG.	٧	m	37	4	54	63	43	*	
	ż S	. 70	-	2		'n	78	16		
	MAX.	25		68	=	160	%	88		
	STANDARD DEVIATION	01.0	0.039	8.52	0.11	960•0	12.13	8.16		
ZINC	AVG.	0.13	90.0	12.99	0.12	0.10	23.08	15.00	_	
	ž Z	0.04	0.02	0.28	0.03	0.01	5.12	3.02		
	MAX.	9.92	0.19	33.9	0.43	0.36	50.10	38.80		
	STANDARD DEVIATION	0.035	0.035	0.26	0.021	0.019	0.26	90•0		
LEAD	AVG.	0.05	0.04	0.23	0.04	0.04	0.16	60.0		
	M N	0.01	0.02	<0.01	0.01	0.01	<0.02	0.03		
	MAX.	0.15	0.17	4.00	0.08	0.07	1.05	0.28		
	# OF SAMPLES	18	82	19	. 81	18	14	21		
	STATION	X-2	x-x	X-7	x-10	X-14	X-22	X-23		

TABLE 9 COMPLIANCE MONITORING - FOR Zn and pH, 1982-83

rom pit below
below
below
11
ant
ant ming

only for the pond decant X-4 and groundwater monitoring stations X-21 and X-24. This result is not easily explained since the waters are not acidic and zinc-laden flows at X-7, X-22, X-23 show generally low to poor correlation. It is suggested here that the elevated zinc content of the mine or mill discharge will be reduced linearly with an increase in some constituent of the native ground and surface waters, but only within a narrow range of zinc and pH values. Station X-21 shows that there is some zinc contamination of groundwater but it is strongly attenuated compared to surface flows.

5.3 Non-Compliance Incidents 1982-1984

After the mill curtailed production in June 1982, pumpings from the pit and mill scavenging waters containing high zinc values continued to dischage into the intermediate dam pond, which at time of shut-down was markedly basic form residual mill reagents. 1982, the decant X-4 showed a zinc content of 1.55 mg/l and a pH of 6.9, and by December, the zinc had reached 4.56 mg/l. Meantime the cross valley pond water quality changed from 0.26 mg/l zinc in July to 2.06 mg/l in November. Mitigative action by the owners included sandbagging the decant at X-5 and treating the pond with massive doses of lime to elevate the pH. It was evident that during periods of mill shut-down, control of zinc levels in the ponds through filtration and seepage required the reduction of mill discharges to the two ponds and lowering of the level of the cross valley pond. This was achieved by siphoning during 1983, and no further compliance problem occurred at that point.

A second non-compliance interval occurred in the winter months of 1983-84. Routine monitoring by the company in October, 1983, at site X-2 on the north fork of Rose Creek, showed a zinc value of $1.18 \, \text{mg/l}$. A small seepage flow containing high zinc values was seen upstream of X-2. This seepage appeared to originate in water overtopping the impervious walls of Pit 2 (See Figure 3) which in turn was

caused by dumping waste rock stripped from Pit 1 into Pit 2. Control of this seepage as obtained with difficulty: elevated levels of zinc at X-2 persisted into 1984.

It is apparent that oxidation of the high sulphide ores and tailings characteristic of this mine will continue to present problems in water treatement. Mitigative measures including filtration under the dams and control of flows into the basin should succeed over the short term while mill production is curtailed. However, the long term may present a chronic problem if oxidation accelerates within the old tailings area, and if the stability of the two large tailings dams and the diversion canal deteriorates. The problem of abandonment of the site and the real potential of long term heavy metal pollution of the waters in Rose Creek remains to be addressed by the mine and the regulatory authorities.

CONCLUSION

The impact of the discharges from Cyprus Anvil mine on the aquatic environment of Rose Creek was studied in 1983. This corresponds to a period when the mill was not in operation. It was found that generally the environmental quality in Rose Creek has been improved from what had been found in previous years.

The major points found are:

- Station 2 (diversion channel) shows an increase of concentration in extractable metals (copper and zinc) which can be attributed to the North Fork of Rose Creek;
- 2. Station 3 (the seepage from the tailings pond) shows a decrease of dissolved oxygen, increases of hardness, alkalinity, filterable residue, conductivity, sulfate, manganese, sodium, copper, iron and aluminum from previous studies as well as from other results in this report;
- 3. The sediment analysis revealed an important increase of heavy metal in the middle creek station (4, 5, 6a and 7a) for the silver, arsenic, barium, copper, iron, mercury, molybdenum, lead and zinc, which might be related to the tailings spill in 1975;
- 4. The bottom fauna analysis reveals an increase in the abundance of organisms sampled over previous years which was partially caused by the sampling procedures and the improvement in water quality. The benthic invertebrate analysis shows a moderate stress at Station 3, observed by a change in species composition, and decrease in diversity. The highest similarity between the stations are between (4, 5, 6a and 7a) and (9 and 10);
- 5. The analysis of the extractable metals in the algae shows significant differences for cadmium, zinc, nickel, lead, strontium, vanadium and arsenic, between the two surveys. It is suggested that the high concentration in July is caused by bio-accumulation;
- 6. Bioassays using <u>Daphnia pulex</u> shows no acute toxic effect due to seepage from the tailings pond (Station 3);
- 7. The environmental compliance monitoring of the mine shows cases of non-compliance incidents in 1982-83 indicating high levels of zinc concentrations at the tailings pond, and in 83-84 winter months at X-2 on the north fork of Rose Creek.

REFERENCES

- Baker, S.A., Environmental Quality of Rose Creek as Affected by Cyprus

 Anvil Mining Corporation Ltd. (Survey data from 1974, 75, 76).

 Environmental Protection Service, Regional Program Report
 No. 79-25, 138 p. (1979).
- Brock, D.A., Comparison of Community Similarity Indexes. Journal-Water Pollution Control Federation, 49(12), p. 2488-2494, Dec. (1977).
- Cairns, J. and K.L. Dickson, A Simple Meethod for the Biological

 Assessment of the Effects of Waste Discharges on Aquatic Bottom

 Dwelling Organism. Journal Water Pollution Control Federation,
 p. 755-772, (1971).
- Davidge, D., Oxidation of Yukon Mine Tailings, Environment Canada, Environmental Protection Service, Regional Program Report #84-15, Whitehorse, 30p. (1984).
- Demayo A. and M.C. Taylor, <u>Guidelines for Surface Water Quality</u>, <u>Volume</u>

 1; <u>Inorganic Chemical Substances</u>, <u>Copper-</u>. <u>Environment Canada</u>,

 Inland Waters Directorate, Ottawa, 55 p., (1981).
- DOE, K.G., <u>Procedures Used By EPS Atlantic Region for Culturing and Conducting Acute Toxicity Tests with Daphnia pulex.</u> DOE, EPS, Air and Water Branch, (1980).
- Environmental Protection Agency, <u>Water Quality Criteria 1972</u>, United States, 594 p., (1973).
- Environment Canada, <u>Pollution Sampling Handbook</u>. Pacific Region Laboratory Services, Fisheries Operations and Environmental Protection Service, West Vancouver, B.C., (1976).
- Foster, P.L., Species Associations and Metal Contents of Algae from Rivers Polluted by Heavy Metals. Freshwater Biology. 12, 17-39, (1982).

REFERENCES (Continued)

- Hoos, R.A.W. and W.N. Holman, <u>A Preliminary Assessment of the Effects of Anvil Mine on the Environmental Quality of Rose Creek, Yukon.</u>
 Environmental Protection Service, Report Number EPS 5-PR-73-8, 49 p., (1973).
- Hynes, H.B.N., The Ecology of Running Waters. University of Toronto Press, 555 p., (1972).
- Hugues, B.D., The Influence of Factors Other Than Pollution on the Value of Shannon's Diversity Index for Benthic Macro-intertebrates in Streams. Water Research (12) 359-364, (1978).
- Khalaf, G. et H. Tachet, <u>Un probleme d'actualite : Revue de travaux en matiere d'utilisation des substrats artificiels pour l'echantil-lonnage des macroinvertebres des eaux courantes.</u> Bull, ecol. Tame 9 (1) 29-38, (1978).
- Khalaf, G and H. Tachet, <u>Colonization of Artifical Substrata by Macro-invertebrate in a Stream and Variations According to Stone Size</u>.

 Freshwater Biology (10), 475-482, (1982).
- Meglitsch, P.A., <u>Invertebrate Zoology</u>. Oxford University Press, 2nd Ed, New York, 834 p., (1972).
- Perkins, J.L., <u>Bioassay Evaluation of Diversity and Community Comparison Indexes</u>. Journal Water Pollution Control Federation, 55: 552-530, (1983).
- Phillips, D.J.G., <u>The Use of Biological Indicator Organisms to Monitor</u>

 <u>Trace Metal Pollution in Marine and Estuarine Environments A</u>

 Review. Environmental Pollution, 13: 281-317, (1977).

REFERENCES (Continued)

- Pielou, E.C., <u>Ecological Diversity</u>. John Wiley and Sons, New York, 151 p., (1975).
- Resh, V.H. and J.D. Unzicker, <u>Water Quality Monitoring and Aquatic Organisms</u>: The <u>Importance of Species Identification</u>. Journal Water Pollution Control Fed., 47: 9-19, (1975).
- Rosenburg, D.M. and V.H. Resh, <u>The Use of Artificial Substrates in the Study of Freshwater Benthic Macroinvertebrate in Artificial Substrates</u>. Ann Arbor Science, John Cairns, editor (1982).
- Smith, G.M., Freshwater Algae of the United States. McGraw-Hill, 2nd ed, 719 p., (1950).
- Sokal, R.S. and F.J. Rohlf, <u>Biometry</u>. W.H. Freeman and Company, 2nd ed., San Fransisco, 859 p., (1981).
- Taylor, M.C. and A. Demayo, <u>Guidelines for Surface Water Quality</u>,

 <u>Volume 1 Inorganic Chemical Substances zinc -</u>. Environment

 Canada, Inland Waters Directorate. Otawa, 52 p., (1980).
- Thurston, R.V., R.C. Russo, C.M. Felterolf, T.A. Edsall and Y.M. Barber (ed), A Review of the EPA Red Book: Quality Criteria for Water. Water Quality Section, American Fisheries Society, Bethesda M.D., 313 p., (1979).
- Weagle, K., Benthic Community Monitoring Program at Cyprus Anvil Mine, Faro, Yukon 1980. Ken Weagle Environmental Consultant Ltd. 27 p., (1980).

REFERENCES (Continued)

- Meagle, K., An Examination of the Impact of the Discharges from Cyprus

 Anvil mine on the Aquatic Environmental Quality of Rose Creek,

 Yukon Territory (1977 and 1978). Environment Canada, Environmental

 Protection Service, Regional Program Report 81-25, Whitehorse
 59 p., (1981).
- Weagle, K., Benthic Community Monitoring Program at Cyprus Anvil Mine Faro, Yukon 1982. Ken Weagle Environmental Consultant Ltd. 13 p., (1982).
- Wetzel, R.G., <u>Limnology</u>. Saunders College publishing, Philadelphia, 743 p., (1975).
- wilhm, Jerry. L., Range of Diversity Index in Benthic Macro-invertebrate Populations. Journal of Water Pollution Control Federation, 42(5): R221-R224, (1970).
- winner, R.W., M.W. Boesel, M.P. Farrell, <u>Insect Community Structure as</u>
 an Index of Heavy Metal Pollution in Lotic Ecosystems. Can J.
 Fish. Aquat. Science, 37: 647-655, (1980).
- Yake, W.E. and R.K. James, <u>Setting Effluent Ammonia Limits to Meet In-stream Toxicity Criteria</u>. Journal Water Pollution Control Federation 55 (3) 303-309, (1983).

ACKNOWLEDGEMENTS

The author wishes to thank D. Davidge, I. Soroka, J. Cornell, and T. Tanner for their assistance in the collection and lab analysis of the field data. A special thanks to R. McCandless for the writing of section 5.0 on the environmental compliance monitoring and to George Mackenzie-Grieve for the editing of the report.

APPENDICES

-

APPENDIX I

COLLECTION, PRESERVATION, ANALYSIS OR
IDENTIFICATION METHODS AND WATER
QUALITY CRITERIA

WATER SAMPLE COLLECTION, PRESERVATION AND ANALYSIS METHODS TABLE 1 APPEND1X 1

PARAMETER	DETECTION LIMIT	COLLECTION AND PRESERVATION PROCEDURE ¹	ANALYTICAL PROCEDURE	METHOD SECTION ²
Temperature	0.1.0	in situ temperature reading.	Standard Centigrade Thermometer	
<u>8</u> <u>u</u>		in situ flow measurements using a Price-type current meter.	Cross-section of the stream was measured and the velocity of flow was calculated using the standard Price-type Current Meter Method ³ .	
Dissolved Oxygen	1.00 mg/l	in situ measurement. The instru- ment was calibrated in the field under water saturated air condition.	YS! Dissolved Oxygen Meter	
£		Small allquots of sample were taken and read soon after collection. No preservative. Instrument was callbrated using 7.0 buffering solution.	Potentiometric	080
Conduct Iv I ty	0.2 umhos/cm	In situ measurement. Laboratory measurement, specific conductivity at 25°C. No preservative. The measurement was taken from the same as NH3 below.	YSI Conductivity Meter Model 33 Radiometer Conductivity Meter (CDMC).	44

METHOD SECTION² 058 040 130 900 104 9 980 070 Piatinum-Cobait Visual Compar-Filtration, drying and weigh-Filtration, drying and weigh-Phenol Hypochlorite-Colori-Acid-persulphate, Autoclave Diazotization-Colorimetric-ANALYTICAL PROCEDURE ing of residue on filter Potentiometric Titration Nephelometric Turbidity WATER SAMPLE COLLECTION, PRESERVATION AND ANALYSIS METHUDS (continued) metric-Automated ing of filtrate Digestion Automated ison containers. Each container was preservatives. Stored at 4°C. Single samples collected in 2 COLLECTION AND PRESERVATION rinsed 3 times with sample litre linear polyethylene before it was filled. No PROCEDURE 1 Same sample as NH3. DETECTION 0.005 mg/l 0.005 mg/1 0.005 mg/l LIMIT 5 (colour 0.1 (FTU) 10.0 mg/l 5.0 mg/l as CaCO3 1.0 mg/1 units) Total Alkalinity Total Phosphate Non-Filterable Residue (NFR) PARAMETER Residue (FR) APPENDIX I Filterable Turbidity Ammon i a NH3-N T PO4-P Nitrate NO₂-N Colour

TABLE 1

SECTION² METHOD 122 300 072 024 Thiocyanate-Combined Reagent-Inductively Coupled Argon Cadmium Copper Reduction ANALYTICAL PROCEDURE Barium Chloranilate -UV Colorimetric Automated Spectrophotometric WATER SAMPLE COLLECTION, PRESERVATION AND ANALYSIS METHODS (continued) Colorimetric Plasma COLLECTION AND PRESERVATION Pre-Each bottle was rinsed 3 times with samples linear served to a pH <1.5 using sample before filling. Single or triplicate collected in 200 mi polyethylene bottles. PROCEDURE1 mi concentrated HNO3. Same sample as NH3. Same sample as NH3. Same sample as NH3. DETECTION LIMIT 1.0 mg/l 0.01 mg/1 0.5 mg/l - /gm 0.050 0.004 0.005 0.005 0.005 0.001 0.001 0.001 0.05 0.05 TABLE 1 **PARAMETER** Extractable APPENDIX 1 Sulphate SO₄ Chloride Cl Ni trate Metals NO3-N

METHOD SECTION² 350 330 Flame Atomic Absorption Spectro-ANALYTICAL PROCEDURE Hydride Generation - ICAP WATER SAMPLE COLLECTION, PRESERVATION AND ANALYSIS METHODS (continued) photometry COLLECTION AND PRESERVATION Same as sample metals. Same sample as metals. Same sample as metals. Same sample as metals. PROCEDURE1 DETECTION LIMIT 0.00050 0.0005 0.005 0.002 0.002 0.001 - John 0.05 0.01 0.02 0.01 0.2 TABLE 1 Metals (continued) PARAMETER Extractable APPENDIX 1 B 5 8 As

Ę
2
÷
Selie : tuco
5
3
MFTHODS
5
YS
A.A.A.
9
A
O
ΥAΤ
ER
KE?
٠
Ö
ECT
7.10
Ö
£
Š
ER
WATER SIMPLE COLLECTION.
ABLE 1 WATER SIMPLE COLLECTION, PRESERVATION AND AMAINSIS M
-
TABLE
TABLE
-
Σ
AP, LIND
AP,

METHOD SECTION ²	330	340		
ANALYTICAL PROCEDURE	Flame Atomatic Absorption Spectrophotometry	Flame Atomic Emission Spectro- photometry	The sum of the ICAP results for Mg \times 4.116 and Ca \times 2.497 reported as mg/1 CaCO3.	
COLLECTION AND PRESERVATION PROCEDURE ¹	Same sample as metals.	Same sample as metals.	Same sample as metals.	a (1976). ironment (1979).
DETECTION	0.0005 mg/l	0.01 mg/1	0.030 mg/l as CaCO ₃	As described in Environment Canada (1976). As described in Department of Environment (1979).
PARAMETER	Ag	¥	Total Hardness	1 As described i 2 As described i

SEDIMENT COLLECTION, PREPARATION AND ANALYSIS METHODS TABLE 2 APPEND (X 1

PARAMETER	COLLECT I ON/PREPARAT I ON	ANALYSIS	METHOD CODE ¹
All Parameters	Creek and River Stations: Sediment samples were collected using an aluminum shovel to scoop sample into pre-labelled Whirl-Pak bags. Three samples were taken at each station. Samples were kept cool and were frozen (-19°C) as soon as possible.		
Mercury Hg (Total)	Sample was freeze-dried for 48 hours to remove water. Sample was sieved through a size 100 mesh (.15 mm) stainless steel sieve. The portion passing through was analyzed for mercury. Sample was completely oxidized by digestion with $\rm H_2SO_4$ and $\rm H_2O_2$.	Mercury Monitor	370
Metals (Leachable) Al, B, Ba, Ba, Ca, Cd, Co, Cr, Cu, Fe, Mg, Mn, Mo, Na, Ni, P, Pb, Si, Sn, Sr, Ti, V, Zn	Same as Mercury except portion passing through was analyzed for metals. Sample was leached with HC and HNO3. The sample was heated for 3 hours.	Inductively Coupled Argon Plasma (ICAP)	320

SEDIMENT COLLECTION, PREPARATION AND ANALYSIS METHODS (continued) TABLE 2 APPENDIX I

PARAMETER	PREPARATION	ANALYSIS	METHOD CODE ¹
Metals (Leachable) (continued)			
As	Same as other metals.	Hydride Generation ICAP	350
P P	Same as other metals.	Flame Atomic Absorption Spectrophotometry	330
¥	Same as other metals.	Flame Atomic Emission Spectrophotometry	340
Particle Size	Sample was freeze-dried.	Standard Sieving Operation	078
1 Department of Environm Service, Fisheries	partment of Environment, Department of Fisheries and Ocea Service, Fisheries and Marine Service (1979).	Department of Environment, Department of Fisheries and Oceans, <u>Laboratory Manual</u> , Environmental Protection Service, Fisheries and Marine Service (1979).	

BOTTOM FAUNA COLLECTION, PRESERVATION AND IDENTIFICATION METHODS TABLE 3 APPENDIX I

IDENTIFICATION AND ENUMERATION	Bottom fauna was sent to Dr. C. Low, Consulting invertebrate Biologist, Nanalmo, B. C. for identification to genus, species if possible, and enumeration.	
LABORATORY PROCEDURES	Bottom fauna was removed from other material and stored in a labelled vial containing 70% methanol.	
FIELD COLLECTION, SAMPLING PROCEDURES AND PRESERVATION	Barbecue Chicken Basket: creek and river samples were faken using Barbecue Basket (17 x 25 cm) filled with local rocks. The rock size range was from 1 cm to 3 cm. No attempt was made to wash the rocks to remove the periphyton and biological film. The baskets stayed in the stream for 50 days. Three samples were taken at each station.	

IDENTIFICATION	The identification was made from a subsample of the July 26 survey and kept in a solution of Lugols iodine. The identification to the family was made by EPS Whitehorse staff.
LABORATORY PROCEDURES	For the July samples, all the chironomids were separated and rinsed with tap water into a 450 um mesh selve before the sample was frozen. The samples were shipped to EPS Laboratory in West Vancouver. The September samples were frozen without sorting or rinsing and shipped to Vancouver for extractable metals analysis.
FIELD COLLECTION, SAMPLING PROCEDURES AND PRESERVATION	The filamentous algae were collected by tearing the plant from the substrate near the attachment point. The algae was gently washed to remove the excess of an orange precipitate (Iron sulfate). The sample was put in a Whirl Pak TM bag and kept cool. Inclusion of chironomids was avoided to the extent practical.

WATER QUALITY CRITERIA FOR DRINKING WATER AND AQUATIC LIFE TABLE 5 APPENDIX 1

SUBSTANCE	RECOMMENDED LEVEL(S) FOR DRINKING WATER	REFERENCE (S)	RECOMMENDED LEVEL(S) FOR AQUATIC LIFE	REFERENCE(S)
Physical				
Colour Pt. Counts	15	· _		
Udour and taste	0	-		
Turbidity J.T.U.	•	-		
Chemical				
Alkalinity mg/L	Not considered a public			
(Total)	health problem	4	>20	m
Aluminum (Al) mg/L	Not considered a public			
	health problem	7	0.1	S
Ammonia (NH3-N) mg/L	0.5 (Total)	4	0.02 (un-lonized)	n
Antimony (SD) mg/L				
Arsenic (As) mg/L	0.05	-	0.05	2 (total)
Barium (Ba) mg/L	1.0	_	5.0	_
Boron (B) mg/L	1.0	-		
Cadmium (Cd) mg/L	0.005	-	0.0002	2 (total)
Calcium (Ca) mg/L	75–200	7		
Chloride (CI)mg/L	250	-		
Chromium (Cr) mg/L	0.05	-	0.04	2 (total)
Cobalt (Co) mg/L				
Conductivity @ 25°C	Depends on dissolved			
(umhos/cm)	salts	7	150-500	9
Copper (Cu) mg/L	1.0	-	0.002	2 (total)
Cyanide (CN) mg/L	0.2	-	0.005	n
Dissolved oxygen				•
(% saturation)	Near 100\$	4	25.0 mg/L	n

WATER QUALITY CRITERIA FOR DRINKING WATER AND AQUATIC LIFE (continued) TABLE 5 APPENDIX 1

SUBSTANCE	RECOMMENDED LEVEL(S) FOR DRINKING WATER	REFERENCE(S)	RECOMMENDED LEVEL(S) FOR AQUATIC LIFE	REF ERENCE (S)
Fluoride (F) mg/L	1.5	-	1.5	7
Hardness (Total)				
as mg/L CaCO3	80-100	-		
Iron (Fe) mg/L	0.3	_	1.0	2
Lead (Pb) mg/L	0.05		0.005 (soft H20*)	2 (total)
			$0.01 \text{ (hard H}_2^{0*})$	2 (total)
Magneslum (Mg) mg/L		4		
Manganese (Mn) mg/L	0.05	-	1.0	7
Mercury (Hg) mg/L	0.002		0.0001-0.0002	2 (total)
Molybdenum (Mo) mg/L				
Nickel (NI) mg/L	0.25	2	0.025 (soft H20*)	2 (total)
			$0.25 \text{ (hard H}_20*)$	2 (total)
Nitrate (NO3-N) mg/L	10	-		
Nitrite (NO2-N) mg/L	0.001			
pH units	6.5 - 8.5	-	6.5 - 9.0	۲
Phosphorus (P) mg/L				
(Total)			0.020 to prevent algae	in.
Potassium (K) mg/L				
Residue: Filterable mg/L			70 - 400 with a maximum	
(Total dissolved solids)	1000	4	of 2000	ø
Residue: Non-Filterable				
(mg/L)				;
Selenium (Se) mg/L	0.01	-	0.01	2 (total)
Silica (SI) mg/L				
Silver (Ag) mg/L	0.05	-	0.0001	2 (total)
Sodium (Na) mg/L	20			
Strontium (Sr) mg/L	10	-		
Sulphate (SO ₄) mg/L	500			

WATER QUALITY CRITERIA FOR DRINKING WATER AND AQUATIC LIFE (continued) TABLE 5 APPENDIX 1

	R	RECOMMENDED LEVEL(S) FOR DRINKING WATER	REF ERENCE (S)	RECOMMENDED LEVEL(S) FOR AQUATIC LIFE	REFERENCE (S)
11n (Tin (sn) mg/L	Not present in natural waters	7		
Titar	Titanium (Ti) mg/L				**************************************
Total (T	Total Inorganic Carbon (TIC)				
Total (TC	Total Organic Carbon (TOC)	5.0	٦		
Vanad	Vanadium (V) Zinc (Zn) mg/L	5.0	-	0.030	I
* *	Soft water has a total hardness 95 mg/L as ${\sf CaCO_3}$ (Reference 6).	s less than 95 mg/L as CaCO3.		Hard water has a total hardness of more than	to co
REFER	REFERENCES:				
<u>.</u>	Health & Welfare Canada, Guldelines for Canadian Drinking Water Quality 1978,	delines for Canadian Drink	ing Water Quality 1	978, Supply and Services, Canada (1979)	Canada (1979).
2.	Inland Waters Directorate, <u>Guidelines for Surface Water Quality</u> , Vol. 1, Inorganic Chemical Substances. Environment Canada, Ottawa (1979, 1980).	uidelines for Surface Wate, Ottawa (1979, 1980).	er Quality, Vol. 1,	Inorganic Chemical Substand	• \$60
ň	Thurston, R.V., K.C. Russo, C.M. Fetteroff Jr., T.A. Edsail, and Y.M. Barber Jr. (Eds.), <u>A Review of the EPA Red Book: Quality Criteria for Water</u> . Water Quality Section, American Fisheries Society, Bethesda, MD, 313p. (1979).	4.C. Russo, C.M. Fetteroff Jr., T.A. E Quality Criteria for Water. Water Qu (1979).	dsall, and Y.M. Bar nallty Section, Amer	, T.A. Edsail, and Y.M. Barber Jr. (Eds.), <u>A Review of the EPA R</u> Water Quality Section, American Fisheries Society, Bethesda, MD,	f the EPA Red thesda, MD,
					-

WATER WALITY CRITERIA FOR DRINKING WATER AND AQUATIC LIFE (continued) TABLE 5 APPENDIX I

REFERENCE(S)	he Yukon and e Chairmen,	ntation Procedures	Operations and	cond Edition by			
RECOMMENDED LEVEL(S) FOR AQUATIC LIFE	Anonymous, <u>Guidelines for Establishing Water Quality Objectives for the Territorial Waters of the Yukon and Northwest Territories</u> . Report of the Working Group on Water Quality Objectives to the Chairmen, Water Boards, Yukon and Northwest Territories, July (1977).	Ontario Ministry of the Environment, Water Management - Goals, Policies, Objectives and Implementation Procedures of the Ministry of the Environment. (1978).	Pacific Region Laboratory Services, Fisheries Operations and Jancouver, B.C. (1976).	a. Publication No. 3-A Second Edition by	. •		
REFERENCE (S)	ality Objectives for the Working Group on Water	gement - Goals, Policie (1978).	_	Control Board, Water Quality Criteria.			
RECOMMENDED LEVEL(S) FOR DRINKING WATER	Guidelines for Establishing Water Quality Objectives for the Northwest Territories. Report of the Working Group on Water Water Boards, Yukon and Northwest Territories, July (1977).	nistry of the Environment, Water Mana of the Ministry of the Environment.	ution Sampling Handbook Protection Service, Wes				
SUBSTANCE	Anonymous, <u>Guidelines for Establi</u> Northwest Territories. Water Boards, Yukon and	Ontario Ministry of the Double of the Double of the Ministry	Environment Canada, <u>Pollution Sampling Handbook</u> . Environmental Protection Service, West	California State Water Resources McKee and Wolf. (1963).			
	.	ۍ.	٥				

APPENDIX II

WATER QUALITY DATA

WATER QUALITY - PHYSICAL AND CHEMICAL PARAMETERS APPENDIX II TABLE 1

1 × 1	FLOW m3/s
97 9.0 10.2	
t i	t i
5.0 11.3	
25 7.0 4.7	
6.0 4.8	
0 10.0 9.7	
1 4 2.	4.5
0 10.0 9.9	
4	4.5
1	

x - Arithmetic mean S.D. - Standard deviation

APPENDIX II TABLE 1 WATER QUALITY - PHYSICAL AND CHEMICAL PARAMETERS (continued)

83/9/15 \$\begin{array}{c c c c c c c c c c c c c c c c c c c	STATION	DATE	S. S.	FLOW m ³ /s	TEMP (°C)	0.0 (mg/1)	\$ D.O. SATURATION (\$)	UN SITU	₹ ₹	IN SITU CONDUCTIVITY (umhos/cm)	LAB CONDUCT IV I TY (umhos/cm)	COLOUR (colour units)	TURBIOITY (FTU)
83/9/15 X - 1.5 11.8 65.5 - 7.9 155 272 83/7/26 X 4.33 8.5 10.6 91.8 8.2 8.2 165 251 83/9/15 X - 2.5 12.0 99.3 - 8.0 140 253 83/9/15 X - 2.5 12.0 89.3 - 8.0 140 253 83/9/15 X - 2.5 12.0 97.6 8.1 8.1 150 228 83/9/15 X - 3 12.5 93.0 - 8.0 135 239 83/9/15 X - 12 8.6 79.9 8.0 8.1 10.0 - 5.5 83/9/16 X - 12 8.6 62.0 - 6.3 2.3 1 83/9/15 X - 14 11.0 10.7 8.0 8.1 1	\$	83/1/26	S XI	4.50	5 .	10.3	4.19	7.9	7.9	170	241	5.0	0.0 0.0
83/9/15 X - 12.5 10.6 91.8 8.2 8.2 165 251 83/9/15 X - - - - - - 0.0 - 0.0 83/9/15 X - - 2.5 12.0 89.3 - 8.0 140 253 83/9/15 X - - - - 0.0 - 0.0 83/9/15 X - - - - 0.0 - 0.0 83/9/15 X - - - - - 0.0 - 0.0 83/9/15 X - - - - 0.0 - 2.39 1 83/9/15 X - - - - - 0.0 - 2.59 1 83/9/15 X - - - - - - - - 2.39		83/9/15	s.D.		1.5	11.8	85.5	1 1	0.0	155 I	272	0.0	0.0
83/9/15 \overline{X} - 2.5 12.0 89.3 - 8.0 140 253 83/1/26 \overline{X} - - - - - 0.0 - 0.0 83/3/1/26 \overline{X} - - - - - 0.0 - 0.0 83/3/1/26 \overline{X} - - - - 0.0 - 2.39 1 83/3/1/26 \overline{X} - 12.5 93.0 - 0.0 - 2.1 2.39 1 83/3/1/26 \overline{X} - 12.5 93.0 62.0 - 0.0 - 2.1 2.1 2.1 2.1 2.1 2.1 2.2 2.1 2.1 2.1 2.2 2.1 2.2 2.1 2.2 2.1 2.2 2.1 2.2 2.1 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2	78	83/7/26	× × S.D.	4.33	8.5	10.6	91.8	8.2	8.2 0.0	165	251	5.0	0.0
83/9/15		83/9/15	S.D.	1 1	2.5	12.0	89.3	, ,	0.0	140	253	7.5	0.10
83/9/15 X - 3 12.5 93.0 - 6.0 - 2.9 83/7/26 X - 12 8.6 79.9 8.0 8.1 200 245 83/9/15 X - - - - 0.0 - 3.5 83/9/15 X - - - - - - 3.5 83/9/15 X - 14 11.0 107 8.0 8.1 168 230 83/9/15 X - - - - - - 0.0 - 0.0 83/9/15 X - - - - - 0.0 - 0.0 - 0.0 83/9/15 X - - - - - - 0.0 - 0.0 - 0.0 - 0.0 - 0.0 - 0.0 - 0.0 - 0.0 <td>60</td> <td>83/1/26</td> <td>× × S.D.</td> <td>9.75</td> <td>10</td> <td>11.0</td> <td>97.6</td> <td>8.1</td> <td>0.0</td> <td>150</td> <td>228 0•0</td> <td>5.0</td> <td>0.80</td>	60	83/1/26	× × S.D.	9.75	10	11.0	97.6	8.1	0.0	150	2 28 0•0	5.0	0.80
83/9/15		83/9/15	S.D.	1 (m ı	12.5	93.0	1 1	0.0	135	239	0.0	0.50
83/7/26 \overrightarrow{x} - 14 11.0 107 8.0 8.1 168 230 83/9/15 \overrightarrow{x} - 4 12.2 90.7 - 8.0 140 245 S.D 0.00 - 0.00	٥	83/7/26	8.b.	111	12	9 0	79.9	0. 1	8.1	500	245	0.0	0.0
$83/7/26$ \overline{x} - 14 11.0 107 8.0 8.1 168 230 0.0 - 0.0 - 0.0 0.0 0.0 - 0.0 0.0 83/9/15 \overline{x} - 4 12.2 90.7 - 8.0 140 245 0.0 - 0.07 - 0.0			S.D.				0.70		0.0	-	6.36	0.0	1.95
x - 4 12.2 90.7 - 8.0 140 245 S.D 0.07 - 0.0	0	83/7/26	S.D.		4 1	11.0	107	0 8 1	0.0	168	230	0.0	1.80
		83/9/15	S*D•	1 1	4 1	12.2	7.06	1 1	8.0 0.07	140	245 0.0	0.0	1.55 0.78

x - Arithmetic mean S.D. - Standard deviation

WATER UNALITY - PHYSICAL AND CHEMICAL PARAMETERS (continued) TABLE 1 APPLNUIX 11

					
C1 (mg/1)	<0.50 0.06 0.60	0.00	1.50	<0.50 0.0 <0.50 0.0	<0.50 0.0 <0.50 <0.50
SO4 (mg/1)	8.4 0.5 10.4	10.0	372 10•7 468 10•6	49.9 0.8 59.3 0.35	47.8 0.9 55.3 0.35
NH3 ⁻ N (mg/1)	0.0070	0.0050	0.740 0.2 0.449	0.0240 0.002 0.0360 0.006	0.0180 0.0006 0.0310 0.002
NO ₃ -N (1 /gm)	0.020	0.00	0.050 0.006 0.023 0.0035	0.030 0.0 0.023 0.003	0.030 0.0 0.031 0.007
NO ₂ -N (mg/1)	0.00000	0.0000	0.0050 0.0 0.0130 0.0007	<0.0050 0.0 0.0100 0.0	<0.0050 0.0 0.0100 0.0
T.P04_P (mg/1)	0.0060	<0.0050 <0.0050 <0.0050 <0.0050 <0.0050 <0.0050	<0.00500.00.00900.004	<0.0050 0.0 <0.0050 0.0	<0.0050 0.0 <0.0050 0.0
T. Hard. (mg/∣ Caco ₃)	50.2 0.4 72.4	72.7 0.6 85.3 0.28	344 3.1 354 2.8	103 1.0 122 0.7	106 1.5 123 1.4
T. Alk. (mg/l CaCO ₃)	56.0	64.0 0.0 74.5	136 0.0	77.0 0.0 88.0	79.0 0.0 89.5 0.7
FR. (mg/l)	83.0 1.0	106 2.9 114 4.95	670 33.0 714 2.12	9.5 9.5 170 23.3	177 10.8 158 2.0
N.F.R. (mg/1)	6.0 0.6 5.0	6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	0.0 0.0 0.0 0.0 0.0	<5.0 0.0 <5.0 <5.0	<6.7 2.9 186 257
S S.D.	ı× ç ı× ;	I× S I× S	I× S. I× S. D. S.	S.D.	1× °C 1× °C 8
DATE	83/7/26	83/7/26	83/7/26	83/7/26	83/7/26
STATION	-	2	۲	4	īv

x = arithmetic mean
S.U.= standard deviation

APPENDIX II TABLE 1 MATER QUALITY - PHYSICAL AND CHEMICAL PARAMETERS (continued)

STATION NUMBER	DATE	l× % O•D•	N.F.R. (mg/1)	ER. (mg/1)	T. Alk. (mg/l Ca∞ ₃)	T. Hard. (mg/l $c = \infty_3$)	T.PO4_P (mg/1)	NO ₂ -N (mg/1)	NO ₅ N (mg/l)	NH ₅ TN (1/gm)	504 (mg/1)	(1/gm)
9	83/1/26	S XI	<7.0 3.0	162	0.0	104	<0.0050	0.0	0.040	0.0080	42.0	0.0
	83/9/15	S.D.	430	202 55.8	88.0	119	<0.0050 0.0	0.0110	0.045	0.0190	50.0	0.0
7.9	83/1/26	× °°°	<55.0 0.0	165 2.0	113	125	0.00000	<0.0050	0.040	<0.0050	18.0	<0.50 0.0
	83/9/15	S.D.	110	176	112	132	0.0080	0.0100	0.065	0.0070	20.3	0.0
ω	83/1/26	S. XI	<3.7 2.1	152 5.2	86.0	104	0.0060	<0.0050	0.020	0.0080	27.6	<0.50 0.06
	83/9/15	S.D.	54.0 39.6	168	93.0	116	0.0000	0.0110	0.038	0.0080	31.7	<0.50
Ø	83/1/26	i× °	<7.7>2.1	168	87.0	120	0.0160	<0.0050	<0.010 0.0	0.0070	36.2	<0.50 0.0
	83/9/15	S.D.	30.0	172	99.5	131	0.0090	0.010	0.010	0.0090	37.8	<0.50
0	83/1/26	l× S.	5.7	157	1.5	10. 6.0	0.0100	<0.0050	0.020	0.0110	30.4	<0.50 0.0
	83/9/15	S.D.	<6.0 0.7	163	93.0	118	0.0070	0.010	0.022	0.0090	32.9 0.07	0.0

 \bar{x} = arithmentic mean S.D.= standard deviation

APPENDIX II TABLE 2 WATER QUALITY-EXTRACTABLE METALS (mg/1)

STATION	DATE	s × s	Ag Ing/I	AI # /gm	As mg/l	B mg/1	Ba mg/1	Be 176m	Ca mg/1	Cd mg/1	CO 1/6m	Cr mg/+	Cu mg/4
27, <u>-</u> 9	83/7/26	l× .0. ►	<0.00050	<0.050	<0.00050	0.0180	0.026	<0.0010 0.0	15.1 0.15	0.00060	<0.0050 0.0	<0.0050	0.0020
	83/3/12	× 100 ×	0.00	0.01	0.0	0.0	0.0007	0.0	0.07	0.0	0.0	0.0	0.001
7	83/9/15	v °C srx	0.0	0.0006	0.0001	0.0120	0.0	0.0	0.17	0.0	0.0	0.0	0.0006
n	83/7/26	.d.s × x × s.D.	0.0	0.0	0.00	0.0	0.075	0.00	0.14 106 0.58	0.0 <0.00050	0.0 <0.0050 0.0	0.0	0.005 0.0010
	83/9/15	S.D.	<0.00050	0.060	0.00200	0.0050	0.103	<0.0010 0.0	111	<0.00500	<0.0060	<0.0050	0.0270
7/4	83/7/26	ا× څ	<0.00050	0.0	0.00070	0.0290	0.045	0.00	30.6 0.31	<0.00050	<0.0050	<0.0050	0.0020
	83/9/15	S × i	<0.00050	<0.050	<0.00050	0.0120	0.052	<0.0010	37.0	<0.00050	<0.000.0 0.0007	<0.0050	0.0150
,) ,	83/7/26	S. D.	<0.00050	<0.050	<0.00050	0.0140	0.046	<0.0010 0.0	31.6	<0.00050 0.0	<0.0050	<0.0050	0.0020
	83/9/15	l× °C	<0.00050	<0.050	<0.00050	0.0120	0.076	<0.0010 0.0	37.2	<0.00050	<0.0050	<0.0050	0.0120

x - Arithmetic mean S.D. - Standard deviation

APPENDIX II TABLE 2 WATER QUALITY-EXTRACTABLE METALS (mg/1) (confinued

	× °O°	Ag mg/1	A1 mg/1	. As	8 1,00	B S	Be mg/1	Ca mg/1	DO E	8 2	o j	3 2
1	× ≪ S•D•	<0.00050 0.0	<.060 · 0.02	<0.00050	0.0040	0.063	<0.0010 0.0	31.0	<0.00050 0.0	<0.0050	0.00000	0.0020
- 1	S.D.	0.00050	0.0	<0.00050 0.0	0.0190	0.068	0.0	35.9	<0.00050	<0.0050	<0.0050	0.0110
	8.0•	<0.00050	<.050 0.0	<0.00050 0.0	<0.0030	0.060	0.0010	34.9 0.4	<0.00050	0.0	<0.0050	0.0020
	S.D.	0.0	0.120	<0.00050	0.0190	0.070	0.0010	57.4	<0.00050 0.0	0.00050	<0.0050	0.0100
83/1/26	l× °0°	<0.00050	0.090	0.00060	0.0150	0.058	0.0010	30.4 0.32	0.0	0.0	<0.0050	0.0030
Ì	S.D.	<0.00050	0.090	<0.00050	0.0190	0.067	0.00	34.4	<0.00050	<0.0050	<0.0050	0.0110
83/7/26	l× °°	<0.00050	0.080	0.00070	<0.0040	0.077	0.0	32.3 0.35	<0.00050	<0.0050	<0.0050	0.0030
1	S.D.	0.00050	0.070	<0.0005.	0.0180	0.108	0.0010	36.05	<0.00050	0.0	0.0050	0.0120
83/1/26	S.D.	<0.00050	0.080	0.00070	<0.0100 0.02	0.062	0.0010	30.6 0.25	<0.00050	<0.0050 0.0	<0.0050	0.0000
	S.D.	<0.00050	0.070	<0.00050 0.0	0.0120	0.060	0.0010	34.3	<0°000000 0°0	<0.0050	<0.0050	0.0110

x - Arithmetic mean S.D. - Standard deviation

APPENDIX II TABLE 2 WATER QUALITY-EXTRACTABLE METALS (mg/l) (continued)

STATION	DATE	ı× S.	Fe mg/l	Х П	Mg mg/l	Mn 1 / g m	oM I ∕gm	eN 1 /gm	iN 1 \Qm	Pb mg/l	dS 1 ∕g ≡	Se + /6 =	is 1. Qm
-	83/7/26 83/9/15	ıx öğ lx ö	0.170 0.0021 0.160 0.010	0.730 0.015 0.698 0.0	2.8 0.06 3.9	0.014 0.002 0.019 0.003	<0.0050 0.0 <0.0050 0.0	0.0 0.06 1.6	0.030	<0.0010 0.0 0.0020 0.0007	<0.050 0.0 <0.050 0.0	<0.050 0.0 <0.050 0.0	2.9 0.0 4.0 0.07
7	83/7/26	ı× ö ı× ö	0.160 0.005 0.127 0.0007	0.610 0.014 0.626 0.0	4.6 0.06 5.2 0.0	0.024 0.0006 0.038 0.0007	<0.0050 0.0 <0.0050 0.0	1.3 0.06 1.8 0.0	<0.020 0.006 0.030 0.030	<0.0020 0.0006 0.0040 0.001	<0.050 0.0 <0.050 0.0	<0.050 0.0 <0.050 <0.050	3.9 0.06 4.8 0.0
w :	83/7/26	l× .∪	0.057 0.005 0.456 0.06	4.01 0.09 4.27 0.0	17.7 0.17 17.0 0.07	2.60 0.06 2.90 0.02	<0.0050 0.0 <0.0050 0.0	74.7 1.2 76.6 0.7	<0.020 0.0 <0.020 0.0	<0.0010 0.0 <0.0010 0.0	<0.050 0.0 <0.050 0.0	<0.050 0.0 <0.050 0.0	4.6 0.06 5.0
4	83/7/26	Ix s. o.s.	0.150 0.0021 0.149 0.0007	0.990 0.016 1.08 0.0	6.2 0.1 7.0 0.07	0.260 0.006 0.444 0.002	<0.0050 0.0 <0.0050 0.0	8.2 0.06 11.6 0.07	<0.020 0.006 <0.020 0.0	<0.0010 0.0 0.0020 0.0007	<0.050 0.0 <0.050 0.0	<0.050 0.0 <0.050 0.0	4.1 0.06 4.9 0.07
īv	83/7/26		0.130 0.006 0.136 0.001	1.03 0.02 1.08 0.0	6.3 0.1 7.0 0.07	0.220 0.004 0.384 0.002	<pre><0.0050 0.0 <0.0050 0.0 0.0 0.0</pre>	8.0 0.15 10.9 0.14	0.030	<pre><0.0010 0.0006 0.0020 0.0007</pre>	<0.050 0.0 <0.050 0.0	<0.050 0.0 <0.050 <0.050 <0.050	4.2 0.0 0.0

x - Arithmetic mean S.D. • Standard deviation

TABLE 2 WATER QUALITY-EXTRACTABLE METALS (mg/1) (continued) APPENDIX 11

	Fe Fe F	× \$€	Mg 1 1 Jg€	Mg/I	A DE	2 N	1.0m	4 P	\$\$ 1/6 _m	88 E	S1.
0.120		1.03	6.3	0.135	<0.0050	1.1	<0.020	<0.0010 0.0	<0.050 0.0	<0.050 0.0	4.2
0.137	0 0	1.08	6.8	0.270	40.0050	9.4	<0.020 0.0	0.0020	<0.050 0.0	<0.050	4.9
0.106	000	0.930	9.0	0.0006	0.0050	1.0 0.0	0.040	<0.0010 0.0	0.050	<0.090 0.0	0.0
0.277	000	0,997	9.0	0.039	0.0050	0.0	<0.020	0.0020	0.050	<0.050 0.0	4.7
0.182	- 0	1.12	6.6 0.06	0.023	0.0050	3.9	0.030	0.00010	<0°020 0°0	050,050	4.0
0.190	- 0	1.14	6.9	0.072	<0.0050	4.9	<0.020	0.0010	0.050	<0.050	0.0
0.205	000	0.770	9.4	0.0013	<0.0050 0.0	2.0	0.050	<0.0010 0.0	0.050	<0.050 0.0	2.9
0.160	0.0	0.773	9.8	0.012	<0.0050	2.6	<0.020	0.0010	<0.050	0.050	3.4
0.138	1.01	0 0	7.3	0.020	0.0050	3.2	0.050	0.00010	<0.050 0.0	<0.050	8°0°0
0.156	-0	1.03	7.7	0.050	<0.0050	4.1	<0.020	0.0010	<0.050	<0.050 0.0	4.4

x -Arithmetic mean S.D. - Standard deviation

APPENDIX II TABLE 2 WATER QUALITY-EXTRACTABLE METALS (mg/l) (continued

Zn I/gm	0.0300 0.0006 0.133 0.015	0.0460 0.004 0.293 0.01	<pre><0.0020 0.0 0.140 0.140 0.104</pre>	0.0580 0.002 0.105 0.012	0.0460 0.005 0.117 0.06
> 26 m	0.0	0.0 0.0 0.0 0.0 0.0	0.00	<0.010 0.0 <0.010 0.0	<0.010 0.0 <0.010 0.0
11 1/g#	0.0050	0.0040	<0.0020 0.0 0.0030	0.0040 0.006 0.0020 0.0020	<0.0020 0.0 <0.0020 0.7
Sr +/gm	0.078	0.094 0.002 0.106	0.352 0.004 0.335 0.003	0.122 0.001 0.133 0.001	0.12 0.003 0.133 0.002
S 1/gm	0.0	0.0	0.0	<0.010 0.0 <0.010	<0.010 0.0 <0.010 0.0
l× %	I× S. I× S.	I× °° I× °°	I× ° I× °	l× s l× s	Ix so ix so
DATE	83/7/26	83/7/26	83/7/26	83/7/26	83/7/26
STATION	-	2	m	4	rv.

x - Arithmetic mean S.D. Standard deviation

WATER QUALITY-EXTRACTABLE METALS (mg/l) (continued) TABLE 2 APPENDIX 11

			·		
			·		
		·			
				·	
•					
ì					
Zn I/gm	0.050 0.010 0.072 0.016	0.034 0.020 0.031 0.004	0.036 0.004 0.049	0.043 0.020 0.040 0.008	0.040 0.020 0.038 0.014
у 1/gm	60.01 0.0 60.01	0.0	<0.01 <0.0 <0.01 <0.0	0.0 0.0 0.0	0.0 0.0 0.0 0.0
11. 1/gm	<0.0020 0.0006 <0.0020 0.0	0.0060 0.0060 0.0060	0.0070 0.0006 0.0050	0.0050	0.0060 0.0060 0.0020 0.0
,		•			
Sr mg/ł	0.120 0.001 0.127 0.0007	0.106 0.002 0.104 0.0007	0.106 0.002 0.109	0.141 0.002 0.142 0.001	0.115 0.001 0.117 0.002
_ =	= = = =	= = = -	= = = =	= = = =	5
Sn I/gm	0.0	0.0	6.01 0.0 0.0	0.00	0.01 0.0 0.0
S, D.	S.D.	lx o lx o	S. D. S.	lx so lx so	۶. ا ای ای ا
i v					
DATE	83/7/26	83/7/26	83/7/26	83/7/26	83/7/26
STATION	& &	7a 8			
STATION	ν .		, ω	6	0

x Arithmetic mean S.D. Standard deviation

APPENDIX III

SEDIMENT DATA

APPENDIX III TABLE 1 SEDIMENT PARTICLE SIZE - MEAN AND STANDARD DEVIATION

JULY 25, 1983 AND SEPTEMBER 15, 1983

			Р	ERCENT CO	MPOSITION		
STATION		<.063 mm	.06315 mm	•15-•25 mm	•25-•5 mm	•5-1 • 0 mm	>1 • 0 m m
NUMBER	x S.D.	SILT &	V. FINE SAND	FINE SAND	MED I UM SAND	COARSE SAND	GRAVEL
1	× S.D.	0•6 0•8	3.6 2.1	6.1 4.1	. 8.7 3.6	9.4 3.9	71 - 4
	3.0.	0.0	2 • 1	4 • 1	J•0	3.9	14.4
2	×	1.0	3.5	2.1	5.7	16.7	70.7
	S.D.	1.3	1.4	1.3	3.7	4.0	11.0
3	×	2.3	6.3	3.4	6.2	7.2	74.2
	S.D.	1.9	0.7	0.1	0.7	0.5	0.8
	_						
4	× S.D.	3.4 3.6	8.8 8.6	7.4 5.7	7.9 0.8	4.8 0.6	67.9
	3.0.	J•0	0.0	5.7		0.6	18.2
5	×	0.6	1.8	2.2	8.6	16.8	70.3
	S.D.	0.1	0.9	0.8	0.4	0.5	2.7
6 a	x	1.3	3.3	2.9	9.2	27.2	56.3
	S.D.	0.4	0.07	0.5	0.4	0.4	3.5
_	_						
7 a	∝ S•D•	3.1 1.6	26.0 2.6	38.1	29 -1	3.3	0.7
	3.0.	1.0	2.0	1.3	4.3	1.2	0.1
8	x	0.1	0.3	1.5	13.8	24.8	59.8
	S.D.	0.0	0.07	0.2	0.6	4.2	3.9
9	×	1.5	2.0	3.6	7 • 8	4.7	90 6
			0.4		. 0.8		80.6 1.9
10		23.4		9.9	3.6		1.6
}	S.D.	10.3	10.7	2.3	1.1	0.8	0.07

x - arithmetic mean

S.D. - standard deviation

25366.7 4839.8 66400.0 Fe mg/kg 3927.3 26600.0 282.8 3262.4 25633.3 25450.0 29300.0 848.5 6657.3 53050.0 6178.5 66050.0 35166.7 8697.4 58066.7 919.2 Cu mg∕kg 4.0 114 45.4 47.6 3.5 130 13.5 4.0 26.1 54.2 3.2 58.9 0.7 05 119 49.9 ار ار kg 45.5 42.7 7.64 1.13 13.0 59.7 52.9 1.9 34.6 5.6 36.7 68.5 4.6 43.3 1.7 ag∕kg S 14.2 12.9 4.5 17.3 17.0 14.0 4.0 33.6 5.2 34.9 34.8 20.3 Gd kg 0.70 0.85 0.06 1.20 3.20 1.57 0.75 0.17 0.85 1.50 © Mg∕kg 42.4 9010 6390 6770 480 9730 8770 1400 22600 6860 7450 284 9440 1360 7030 128 7560 688 **39**,kg 0.50 0.55 0.50 0.65 0.40 0.55 0.55 0.12 0.37 Ba mg/kg 11.0 482 85.8 64.3 46.0 28.3 233 3600 80 506 1940 3040 651 2970 368 2860 474 1490 148 As mg/kg 0.0 109 62.0 21.3 89.0 2.8 14.7 3.2 7.6 22.5 33.3 20.5 14.6 12.7 18.4 118 AI mg/kg 0009 15800 212 1258 16400 17000 16800 0009 14300 15200 14900 14100 404 2334 1131 Ag mg/kg 0.29 0.38 1.16 1.42 ! 1 0.51 ! ! 1 1 0.24 -| | S.D. S.S. S.D. S.D. S.D. S. S. S ×I S. 0. S.U. S.D. S.b. JХ ١× 83/1/26 83/9/15 33/9/15 83/9/15 83/1/26 33/9/15 83/9/15 83/7/26 83/7/26 83/7/26 DATE STATION NUMBER 4) 5

SLUIMENT - LEACHABLE METALS (mg/kg)

TABLE 2

APPENDIX 111

3° 40 0

x - arithmetic mean S.D. - standard deviation

APPENDIX III TABLE 2 SEDIMENT - LEACHABLE METALS (mg/kg) (continued)

83/7/26	STATION	DATE	S × i	Ag mg/kg	A1 mg/kg	As mg/kg	Ba mg/kg	Be mg/kg	Ca mg/kg	Cd mg/kg	So. mg/kg	Cr mg/kg	Cu mg/kg	Fe mg/kg
83/9/15 X	63	83/7/26	ı× °°	11	14500	71.7	5990	0.60	6730 620	1.73	22.9	39.7	224 98.8	39500
$83/7/26 \qquad \overline{x} \qquad \qquad 14100 \qquad 62.7 \qquad 5620 \qquad 0.40 \qquad 6180 \qquad 1.50 \qquad 2.00$ $83/9/15 \qquad \overline{x} \qquad 1.32 \qquad 12600 \qquad 79.0 \qquad 1410 \qquad 0.40 \qquad 6230 \qquad 0.60 \qquad 2.00$ $83/9/15 \qquad \overline{x} \qquad 1.32 \qquad 12600 \qquad 79.0 \qquad 4140 \qquad 0.40 \qquad 6230 \qquad 0.60 \qquad 2.00$ $83/9/15 \qquad \overline{x} \qquad \qquad 12800 \qquad 17.0 \qquad 1240 \qquad 0.20 \qquad 16500 \qquad 1.10 \qquad 1.10 \qquad 1.10$ $83/9/15 \qquad \overline{x} \qquad 0.83 \qquad 15000 \qquad 12.3 \qquad 1550 \qquad 0.35 \qquad 15000 \qquad 1.90 \qquad 1.90$ $83/9/15 \qquad \overline{x} \qquad \qquad 10200 \qquad 12.3 \qquad 1550 \qquad 0.25 \qquad 15000 \qquad 1.90 \qquad 1.90$ $83/9/15 \qquad \overline{x} \qquad \qquad 116 \qquad 0.6 \qquad 826 \qquad 0.06 \qquad 656 \qquad 0.20$ $83/9/15 \qquad \overline{x} \qquad \qquad 13400 \qquad 11.5 \qquad 2950 \qquad 0.04 \qquad 22200 \qquad 1.00$ $83/9/15 \qquad \overline{x} \qquad \qquad 13400 \qquad 11.0 \qquad 864 \qquad 0.30 \qquad 16300 \qquad 1.30 \qquad 1.30$ $83/9/15 \qquad \overline{x} \qquad 0.27 \qquad 15200 \qquad 1.70 \qquad 0.90 \qquad 1.30$ $83/9/15 \qquad \overline{x} \qquad 0.27 \qquad 15200 \qquad 1.3.5 \qquad 768 \qquad 0.50 \qquad 554 \qquad 0.14$		83/9/15	ı× °C.	1.12	15400	30.5	3660	0.65	5800 721	0.55	18.4	41.4	120	35400 424
83/9/15 X 1.32 12600 79.0 4140 0.40 6230 0.60 2.50 0.09 778 15.6 1560 0.00 42.4 0.00 42.4 0.00 42.4 0.00 42.4 0.00 42.4 0.00 45.00 1.10 1.10 1.20	78	83/1/26	S.D.	11	14100	62.7	3620 1470	0.40	6180	0.06	21.7	38.6 1.5	119	50100
$83/7/26 \times 12800 17.0 1240 0.20 16500 110 1$ $83/9/15 \times 1300 12.3 1550 0.35 15000 190 1$ $83/9/15 \times 10200 12.3 1550 0.35 15000 190 1$ $83/7/26 \times 10200 12.3 3830 0.27 25500 100 1$ $83/9/15 \times 118 12600 11.5 2950 0.04 22200 100 1$ $83/9/15 \times 13400 11.0 864 0.30 16300 1.30 1$ $83/9/15 \times 1010 1.7 79.7 0.00 874 0.00 1$ $83/9/15 \times 0.27 15200 13.5 768 0.50 15600 0.90 1$ $83/9/15 \times 0.27 15200 13.5 768 0.50 15600 0.90 1$ $83/9/15 \times 0.27 15200 13.5 768 0.50 15600 0.90 1$ $83/9/15 \times 0.27 15200 13.5 768 0.50 15600 0.90 1$ $83/9/15 \times 0.27 15200 13.5 768 0.50 15600 0.90 1$)	83/9/15	S.D.	1.32	1 <u>2</u> 600 778	79.0	4140	0.40	6230	00.0	25.5	34.2	116	54700
83/9/15 \overline{x} 0.83 15000 12.3 1550 0.35 15000 1.90 1 83/7/26 \overline{x} 10200 12.3 3830 0.27 25500 1.00 83/7/26 \overline{x} 1.18 12600 11.5 2950 0.04 22200 1.00 83/9/15 \overline{x} 1.18 12600 11.5 2950 0.04 22200 1.00 83/9/15 \overline{x} 1.18 12600 11.5 2950 0.00 37.7 0.14 83/9/15 \overline{x} 1.17 130 11.0 864 0.30 15300 1.30 1350 1.30 15500 0.00 354 0.14	60	83/7/26	۱× ۵۶	1 1	12800	17.0	1240 222	0.20	16500 4590	1.10	12.9	52.5 2.8	28.8	23000
83/7/26	and y	83/9/15	ı× ç	0.83	15000 636	12.3	1550 424	0.35	15000 141	1.90	13.8 1.9	85.9 4.6	30.7	27800 707
83/9/15	o	83/1/26	ı× ç	11	10200	12.3	3830 826	0.27	25500 656	1.00	6.4	44.0 5.5	23.4	21700
83/7/26		83/9/15	i× °C	1.18	12600	11.5	2950 156	0.00	22200	1.00	8.0	44.1	26.2	22900
x 0.27 15200 13.5 768 0.50 15600 0.90 S.D. 0.02 636 0.7 33.2 0.00 354 0.14	6	83/1/26	S.D.		13400	11.0	864 79.7	0.30	16300 874	1.30	10.9	38.2 2.6	32.3	21300
		83/9/15	۱× ۵۰	0.27	15200 636	13.5	768 33 . 2	0.50	15600 354	0.90	12.5	1.0	33.8 1.5	22600

x - arithmetic mean S.D. - standard deviation

5.3

3.5

<2.0

<2.0

0.0

0.0

<2.0 0.0

Sn mg/kg

4.3

<2.0

2.7

4840 37.9 mg/kg 4260 2790 127 4220 163 2*3*90 700 4210 164 2190 863 2280 4590 238 2690 608 873 S 775 50•2 681 53.3 Pb mg/kg 338 87.5 682 31.8 37.0 97.6 36.0 34.5 124 22-1 368 198 554 1040 997 50**.**8 983 30**.**6 ₽ mg/kg 15.3 40.4 53.0 36.0 68.2 60 1010 1010 000 1040 1150 184 985 932 Ni mg/kg 47.0 26.5 6.0 40.0 2.8 33.0 31.5 69.0 12.0 57.5 37.7 41.0 4.4 27.7 425 77.8 360 10•0 mg/kg 28.7 425 35.4 565 35.4 11.5 28.3 11.5 14.1 413 520 647 206 470 407 483 Mo mg∕kg 7.20 10.8 6.60 6.07 7.40 1.80 2.90 4.80 3.77 0.07 SEDIMENT - LEACHABLE METALS (mg/kg) (continued) Mn mg/kg 21.2 544 148 2030 1340 904 248 509 872 855 844 161 866 270 1240 428 1070 186 Mg mg∕kg 8670 1370 7140 7120 6180 5860 834 6420 982 5410 233 7120 8670 1960 70007 367 601 1800 ж mg/kg | | | | 2410 240 2205 2300 113 1925 375 | | | | | | Hg mg/kg 0.25 0.21 0.45 1.24 1.15 0.20 0.05 0.70 0.29 0.86 0.11 0.97 S. S. s. S. S. C. S. D. % ×1 S ×I s. U.s S.U. S xI S.U. S.D. l× t× TABLE 2 83/9/15 83/1/26 33/9/15 33/7/26 83/9/15 83/1/26 83/9/15 83/1/26 83/7/26 83/9/15 DATE APPENDIX 111 STATION NUMBER S ~

x - arithmetic mean S.D. - standard deviation

APPENDIX III TABLE 2 SEDIMENT - LEACHABLE METALS (mg/kg) (continued)

STATION	I DATE	l× °	Hg mg/kg	х ву/к <u>в</u>	Mg mg/kg	Mn mg/kg	Mo mg/kg	Na mg/kg	Ni mg/kg	Вд/кд	Pb mg/kg	S.I mg/kg	Sn mg/kg
68	83/1/26	ı× ç	1.26		5930	2780 945	8.17	547 230	32.0 8.2	1020	585 107	8270 6010	6.0
	83/9/15	۱× ۹۰	0.80	2260	5730	666	7.40	390	33.0	1100	899 157	2950	<2.0
78	83/1/26	l× ° 1	1.02		6150	1040	1.3	397	27.3	924 65.9	510 75.8	4520	0.0
	83/9/15	S.D.	0.88	1750	5060	762 67.9	10.8	14.1	22.0	1100	504	2860 56.6	0.0
ω	83/1/26	ı× °°	0.13	11	8460 1320	497 59 . 2	3.00	380	37.7	1340 83.9	46.0	4620 315	5.0
	83/9/15	S.D.	0.12	2020	9100	992	00.9	470	49.0	1330	54.0	3120 148	11.0
0	83/7/26	ı× ö ı× ö	0.02	2760	14500 755 12800	9.2	3.80 0.46 4.90	220 20.0 20.0	36.0	2050 98.7 1670	3.5	4980 296 3380	3.0
5	83/1/26	l× °C °S	0.18		9490	575	3.10	340	40.0	1150	42.3	5120	2.3
	83/9/15	l× °C.	0.15	2650 184	9250	745 36.8	4.30	340	40.5	1100	44.5	3540 42.4	0.0

x - arithmetic mean S.D. - standard deviation

APPENDIX !!! TABLE 2 SEDIMENT - LEACHABLE METALS (mg/kg) (continued) 1000 992 215 49•3 300 29.0 282 967 19.8 Zn mg/kg 327 11.3 46.0 3.5 868 298 304 √ mg⁄kg 43.5 48.5 46.0 2.8 1.5 42.5 45.3 1.4 45.3 6.8 44.7 1.2 34.7 Tí mg/kg 746 98•3 666 43.4 735 53.5 692 543 20•6 21.7 579 0.0 36.1 619 745 277 121 692 mg/kg 68.4 58.0 4.6 56.9 108 83.4 58.9 3.9 57.6 1.6 62.7 56.7 5. 64.8 አ S.D. \$. S. D. i× ° s. S. s.b. ∢ s. 0. s s s. v. S.D. S.D. S.D. ١× ł× 83/9/15 83/1/26 83/1/26 83/1/26 83/9/15 83/1/26 83/9/15 83/9/15 83/7/26 83/9/15 DATE STATION NUMBER Ŋ 7

x - arithmetic mean S.D. - standard deviation

APPENDIX III TABLE 2 SEDIMENT - LEACHABLE METALS (mg/kg) (continued)

STATION	DATE	l× ở	Sr mg/kg	TI mg/kg	v mg/kg	Zn mg/kg
68	83/1/26	l× S.	71.5	758	45.7	1020 20.8
	83/9/15	S.D.	55.7	748	45.0	750 144.2
78	83/7/26	l× 3	60.5 9.1	717	44.3	977 1.27
	83/9/15	s.D.	57.1 7.2	746	42.5	778 38.2
.	83/7/26	ا× ق	68.5 9.3	859 22.5	56.3	133 12.9
	83/9/15	8.D.	72.0	1180 49.5	66.5	156 3.5
6	83/7/26	l× co	100	457	64.7 3.5	142 5.1
	83/9/15	×× S.D.	87.6 0.07	470 38.9	1.4	169 2.8
01	83/7/26	۱× د.	69.1	519 51•3	58.7	180 11.6
	83/9/15	۶. S. C.	1.6	602 26 . 2	63.5	202 10.0

x - arithmetic mean S.D. - standard deviation

APPENDIX IV

BOTTOM FAUNA DATA

APPENDIX IV TABLE 1 TAXONOMIC LIST OF THE BOTTOM FAUNA AT CYPRUS ANVIL, SEPTEMBER, 1983

1. 2. 3. 4. 5.	Order: Family: Phylum: Phylum: Class: Order: Family: Family:	Hydroida Hydridae Hydra sp. Nematoda Annelida Oligochaeta Haplotaxida
2. 3. 4. 5.	Phylum: Phylum: Class: Order: Family:	Hydra sp. Nematoda Annelida Oligochaeta
2. 3. 4. 5.	Phylum: Class: Order: Family:	Nematoda Annelida Oligochaeta
3. 4. 5.	Phylum: Class: Order: Family:	Annelida Oligochaeta
4.	Class: Order: Family:	Oligochaeta
4.	Order: Family:	
4.	Family:	Hanlotaxida
4.		p
5.	Esmilus	Enchytraeidae
5.	ганті і у :	Naididae
		Paranals sp.
6.		Nais sp.
	Family:	Tublflcldae
	Phylum:	Arthropoda
	Class:	Crustacea
	Order:	Cladocera
	Family:	Daphnidae
7.		Daphnia rosea
	Order:	Copepoda
	Family:	Cyclopidae
8.		Cyclops sp.
	Family:	Diaptomidae
9.		Diaptomus sp.
10.	Order:	Ostracoda
	Class:	Arachnoidea
11.	Subclass:	Arachnida
12.	Order:	Acari
	Class:	Insecta
	Order:	Collembola
13.	Family:	Isotomidae
	Order:	Plecoptera
	Family:	Capniidae
14.		Capnia sp.
	Family:	Chloroperiidae

APPENDIX IV TABLE 1 TAXONOMIC LIST OF THE BOTTOM FAUNA AT CYPRUS ANVIL, SEPTEMBER, 1983 (continued)

15.		Kathroperia sp.
16.		Alloperia sp.
	Family:	Nemouridae
17.		(Amphinemura sp. or Amphinemoura sp.)
18.		Prostola sp.
19.		Zapapa sp.
	Family:	Periodidae
20.		Cultus sp.
21.		isoperia sp.
22.		Kogotus sp.
23,	•	Megarcys sp.
	Family:	Pteronarcydae
24 👡		Pteronarcella sp.
	Family:	Taeniopterygidae
25.		Brachyptera sp.
	Order:	Ephemeroptera
	Family:	Baetidae
26 💊		Baetis sp.
	Family:	Ephemerellidae
27 💊		Ephemerella sp. unld.
28.		Ephemerella sp. juve.
29 👡		Ephemerella dodds1
30.		Ephemerella Inermis
31.		Ephemerella Infrequens?
32		Ephemerella proserpina
	Family:	Heptageniidae
53.		Cinygmula sp.
	Family:	Siphionuridae
34.	-	Ameletus sp.
	Order:	Trichoptera
	Family:	Brachycentridae
35.		Brachycentrus sp.
	Family:	Glossosomatidae
36 📞	•	Agapetus sp.
-	Family:	Hydrop sych I dae
37 📞	•	Arctopsyche sp.
		•

APPENDIX IV TABLE 1 TAXONOMIC LIST OF THE BOTTOM FAUNA AT CYPRUS ANVIL, SEPTEMBER, 1983 (continued)

	Family:	Hydroptilidae
38.		Hydroptila sp.
	Family:	Lepidostomatidae
39.		Lepidostoma sp.
	Family:	Limnephilidae
40.		Hesperophylax
41.		Onocosmoecus sp.
42.		Unidentify Limnephilidae
	Family:	Rhyacophilidae
43.		Rhyacophila sp.
	Order:	Diptera
44.	Family:	Ceratopogonidae
45.		Palpomya sp.
46.	Family:	Chironomidae (pupae)
47.		Brillia sp.
48.		Cardiociadius sp.
49.		Corynoneura sp.
50.		Cricatopus sp.
51.		Diamesa sp.
52.		Euklefferiella sp.
53.		Heterotrissociadius sp.
54.		Microspectra sp.
55.		Monodiamesa sp.
56.		Paralauterborniella sp.
57.		Phalnopsectra sp.
58.		Polypedilum sp.
59.		Procladius sp.
60.		Tribelos sp.
61.	Family:	Empididae pupae
62.		Chelifera sp.
63.		Clinocera sp.
	Family:	Muscidae
64.		Limnophora sp.
65.		Lispe sp.
	Family:	Psychodidae
66.		Pericoma sp.
	Family:	Simuliidae
67.		Cnephia sp.
68.		Ectemnia sp.
69.		Prosimulium sp.
		•

APPENDIX IV TABLE 1 TAXONOMIC LIST OF THE BOTTOM FAUNA AT CYPRUS ANVIL, SEPTEMBER, 1983 (continued)

	Family:	Tîpulidae	
70.		Antocha sp.	
71.		Tipula sp.	
	Order:	Coleoptera	
72.		Unid. larva	
	Phylum:	Mollusca	
	Class:	Gastropoda	
	Order:	Basommatophora	
	Family:	Lymnaeidae	
73.		Lymnacea sp.	
	Order:	Ctenobranchiata	
	Family:	Valvatidae	
74.		Valvata sp.	
	Order:	Pelecypoda	
	Family:	Sphaerlidae	
75.		Pisidium sp.	
٠			
		•	
		e e	
		•	
٠		•	

CYPRUS ANVIL MACRO-INVERTEBRATE DATA - SEPTEMBER 1983 TABLE 2 APPENDIX IV

				S	STATION				
Taxonomic	-	2	2	4	5	6 9	7.0	6	10
Group	* \$	*	*	*	**	*	**	**	**
1. Hydra sp.	23 (0.82)	,	,	•	,		,	,	•
* 2. Nematoda	7	•	2	4	9	_	9	2	n
* 3. Enchytraeldae	,_	ı		7	7	2	'n	01	0
4. Paranais sp.	2 (0.07)	,	,	,	1	ı	•	,	ı
5. Nais sp.	16 (0.57)	ı	,	1	1	1	1	1 (0.05)	ı
* 6. Tubificidae	5	ı	,		ı	,	•	_	ı
* 7. Daphnia rosea	9	,	,	-	-	•	ı	1	ı
* 8. Cyclops sp.	13		2	_	2		ı	ı	ı
* 9. Diaptomus sp.	_	_	,	-	-	ı	1	ı	,
*10. Ostracoda	7	,	,	•	-	•	1	,	,
*11. Acari	7	_	,	4	10	ı	16	20	&
*12. Arachnida	-	,	,	,	,		,	ı	
*13. Isotomidae	2	,	,	ı	ı		,	ı	1
14. Capnia sp.	157 (5.63)	45 (1.41)	2 (0.09)	70 (1.36)	50 (0.93)	438(18.80)	135 (6.65)	882(42.71)	4172(59.74)
15. Kathroperia sp.	,	,	,	18 (0.35)	18 (0.33)	14 (0.60)	1 (0.05)	15 (0.73)	1
16. Alloperia sp.	31 (11.11)	,	,	37 (0.72)	70 (1.30)	14 (0.60)	13 (0.64)	1 (0.05)	2 (0.03)
17. Amphinemura sp.	78 (2.79)	105 (3.29)	,	208 (4.04)	38 (0.71)	42 (1.80)	14 (0.69)	1 (0.05)	10 (0.14)
18. Prostola sp.	6 (0.21)	11 (0.34)	,	38 (0.74)	40 (0.74)	4 (0.17)	2 (0.10)	3 (0.15)	4 (0.06)
19. Zapada sp.	119 (4.26)	296 (9.26)	3 (0.13)	1495(29.05)	791(14.71)	305(13.09)	201 (9.90)	10 (0.48)	136 (1.95)
20. Cultus sp.	35 (1.25)	,	,	1	2 (0.04)	3 (0.13)	1 (0.05)	20 (0.97)	5 (0.07)
	7 (0.25)	102 (3.19)		145 (2.82)	195 (3.63)	55 (2.36)	5 (0.25)	3 (0.15)	32 (0.46)
22. Kogotus sp.	6 (0.21)	,	,	1 (0.02)	4 (0.07)	37 (1.59)	47 (2,32)		1 (0.01)
	,	2 (0.06)	,	21 (0.41)	24 (0.45)	29 (1.24)	1 (0.05)	,	•
24. Pteronarcella sp.	,	,	,	2 (0.04)	10 (0.19)	19 (0.82)	•	7 (0.34)	5 (0.07)
25. Brachyptera sp.	16 (0.57)	717(22.43)	2 (0.09)	97 (1.88)	63 (1.17)	29 (1.24)	18 (0.89)	135 (6.54)	24 (0.34)
26. Baetis sp.	572(20.49)	743(23.25)	4 (0.18)	1461 (28.39)	1899(35.32)	430(18.45)	340(16.75)	129 (6.25)	90 (1.29)
27. Ephemerella sp. unid.	3 (0.11)	,	,	3 (0.06)	1 (0.02)	,		,	1
28. Ephemerella sp. juven.	19 (0.68)	4 (0.13)		1	1 (0.02)	3 (0.13)	13 (0.64)	1 (0.05)	19 (0.27)
29. Ephemerella doddsl	,	1 (0.03)	,	1 (0.02)	3 (0.06)	5 (0.21)	7 (0.34)	ı	•
30. Ephemerella inermis	32 (1.15)	11 (0.34)		23 (0.45)	12 (0.22)	28 (1.20)	163 (8.03)	6 (0.29)	23 (0.33)
Tacobacha for a feet of the fe			,						

* Taxonomic level not included in the Shannon-Weiner diversity index (H') for bottom fauna,

and the calculation of the percent similarity index (PSc). *Numbers of organisms sampled at each station. *Percent abundance of organisms sampled at each station.

CYPRUS ANVIL MACRO-INVERTEBRATE DATA - SEPTEMBER 1983 (Continued)

TABLE 2

APPENDIX IV

				S	STATION				
Taxonomic	-	2	2	4	2	99	7.0	6	01
Group	4 \$		* *	# %	*	*	* *	* *	×
someonial elleramond . If	(1170) \$	1 (0.03)	•		•	•	•	34 (1 66)	10, 07, 01
32. Enhanced la procernina		;	,	1	1 (0.02)	3 (0.13)	8 (0.30)	10000	
Cinygaula sp.	76 (2.72)	43 (1.35)	ı	380 (7.38)	403 (7.49)	4	314(15.47)	60 (2.91)	21 (0.30)
	11 (0.39)	4 (0.13)	,	2 (0.04)	1 (0.02)		5 (0.25)	136 (6.59)	
35. Brachycentrus sp.	1 (0.04)	1 (0.03)	ı	8 (0.16)	23 (0.43)	5 (0.21)	11 (0.54)	4 (0.19)	
36. Agapetus sp.	1 (0.04)		ı	37 (0.72)		1 (0.04)	•	1	2 (0.03)
37. Arctopsyche sp.	•	2 (0.06)	1 (0.04)	ı	13 (0.24)	222 (9.53)	49 (2.41)	3 (0.15)	7 (0.10)
38. Hydroptila sp.	•	•	ı	1	•	ı	1	2 (0.10)	1
39. Lepidostoma sp.	•	•	1	•	•	ı	ı	•	1 (0.01)
40. Hesperophylax sp.	2 (0.07)	1	•	1	•	1 (0.04)	1	1	2 (0.03)
41. Onocosmoecus sp.	1	ı	•	•	•	1	1		(60.0) 9
*42. Limnephilidae unid.	•	•	1		7	'n	-	_	13
43. Rhyacophila sp.	7 (0.25)	2 (0.06)	ı	5 (0.06)	3 (0.06)	26 (1.12)	ı	1	1 (0.01)
*44. Ceretopogonides	-	•	1	•	-	ı	1	1	
45. Palpomya sp.	1		•	•		ı	1 (0.04)	1 (0.05)	1 (0.01)
eednd o	65	4	n		_	'n	38	50	368
47. Brillia sp.	2 (0.07)	43 (1.35)	090(47.79)	4 (0.08)	•	ı	1	2 (1.00)	25 (0.36)
48. Cardiociadius sp.	35 (1.25)	102 (3.19)	15 (0.66)	99 (1.92)	112 (2.08)	34 (1.46)	63 (3.10)	8 (0.39)	27 (0.39)
49. Corynoneura sp.	57 (1.33)	1	ı	•	•	ı	ı	•	,
50. Cricotopus sp.	76 (2.72)	(16.0) 62	10 (0.44)	7 (0.14)	2 (0.04)	11 (0.47)	417(20.54)	324(15.69)	145(24.99)
51. Diamesa sp.	1 (0.04)	1	1 (0.04)	•		ı	1	1	1 (0.01)
52. Euklofferiella sp.	ı	53 (1.66)	ı	180 (3.50)	77 (1.43)	5 (0.21)	18 (0.89)	4 (0.19)	7 (0.10)
53. Heterotrissociadius sp.	(52.73)	(16-1)	19 (0.83)	334 (6.49)	338 (6.29)	42 (1.80)	118 (5.81)	164 (7.94)	223 (3.19)
54. Microspectra sp.	179(27.91)	7 (0.22)	165 (8.02)	20 (0.39)	26 (0.48)	ı	1	14 (0.68)	72 (1.03)
55. Monodiamesa sp.	1	1 (0.03)	7 (0.51)	1	•	ŀ	1	1	ı
56. Paralauterborniella sp.	•	1	ı	•	•	ı	ı	1	1 (0.01)
57. Phaenopsectra sp.	2 (0.07)	5 (0.16)	655(28.72)	1	•	ı	1	1	1 (0.01)
58. Polypedilum sp.	1	1	1	•		ı	ı	10 (0.48)	12 (0.17)
	569(13.22)	ſ	1	1 (0.02)	3 (0.06)	1	ı	33 (1.60)	57 (0.53)
60. Tribelos sp.	1 (0.04)	ı	1	1	ı	ı	1	,	3 (0.04)

* Taxonomic level not included in the Shannon-Weiner diversity index (H') for bottom fauna,

and the calculation of the percent similarity index (PSc). # Numbers of organisms sampled at each station. % Percent abundance of organisms sampled at each station.

83 (Continued)
EPTEMBER 1983
S
DATA -
INVERTEBRATE
MACRO-
ANVIL
CYPRUS
TABLE 2

					STATION				
Taxonomic	-	2	3	4	5	69	78	6	10
Group	*	*	*	**	**	*	**	**	*

epdnd epninger	1	1	,	7	/7		•		,
		1		1 (0.02)	7 (0.13)	•	1	•	,
	2 (0.07)	1	,	,		,	ı	,	11 (0.16)
64. Limnophora sp.	ı	1 (0.03)	254 (11.14)	ı	•	,	,	•	•
65. Lispe sp.	1 (0.04)	1	3 (0.13)	,	,	,	,	•	,
66. Pericoma sp.	10 (0.36)	ı	1 (0.04)	3 (0.06)	8 (0.15)	4 (0.17)	16 (0.79)	,	31 (0.44)
67. Cnephia sp.	5 (0.18)	128 (4.01)	,	32 (0.62)	25 (0.46)	7 (0.30)	8 (0.39)	6 (0.29)	
68. Ectemnia sp.	1	.1 (0.03)	•	6 (0.12)	3 (0.06)	,			,
69. Prosimulium sp.	30 (1.07)	675(21.12)	•	391 (7.60)	1091 (20.29)	94 (4.03)	38 (1.87)	47 (2.28)	18 (0.26)
70. Antocha sp.	7 (0,25)	ı	31 (1,36)	18	19 (0.35)		3 (0.15)	. 1	1 (0.01)
71. Tipula sp.	1 (0.04)	ı		,	1 (0.02)			,	
*72. Coleoptera larva	,	•	,	,		-	•	•	,
73. Lymnacea sp.	2 (0.07)	,	•	,	,		,	,	,
74. Valvata sp.	42 (1.50)	,	,	•	•	,		_	_
							1.	1	
ods mildian sp.	(0.04)			•		,	,	1.	,
7 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -									_
N lotal of individuals	7907 (100)	\$203 (100)	2203 (100) 2289 (100)	5187 (100)	5187 (100) 5436 (100)	2336 (100)	2094 (100)	2135 (100)	7407 (100)
N _m Number of Individuals per m ³ -density	170,000	187,310	133,860	303,333	317,895	136.608	122.456	124.854	433,158
	`								
Ng Total of Individuals									
bottom fauna	1672	3196	2281	5146	5377	2320	2030	2065	6984
G Total of genera identi-									
, pei	14	92	11	2	32	72	56	&	37
H' Diversity index	1.07	0.93	09.0	0.93	0.87	1.02	1.03	0.88	0.59
J' Evenness	99•0	0.66	0.49	0.62	0.58	0.71	0.73	09*0	0.38
* Taxonomic level not included in the		Shannon-Well	ner diversit	y Index (H'	Shannon-Weiner diversity index (H') for bottom fauna,	fauna,			

and the calculation of the percent similarity index (PSc). $\rlap{/}{t}$ Numbers of organisms sampled at each station.

% Percent abundance of organisms sampled at each station.

ATTACHED ALGAE DATA

APPENDIX V TABLE 1 ATTACHED ALGAE - EXTRACTABLE METALS - STATION 3

12.00

	DATE	i× °°	Al mg/kg	As mg/kg	Ba mg/kg	Be mg/kg	Ca mg/kg	Cd mg/kg	Co mg/kg	Cr mg/kg	Cu mg/kg	Fe mg/kg	Mg mg/kg
+64	83/1/26		373	28.0	46.7	<0.050	3670	2.20	•	1.60	5.4	9030	294
Welght	83/9/15	ı×	269	12.7	33.6	<0.050	1500	0.26	23.4	1.47	4.97	8690	288
		S.D.	67.6	2.6	9•9	•000	204	0.03	5.8	0.2	0.15	872	25.0
٥	83/1/26		2040	150	255	<0.30	20100	11.8	ı	00.6	29•0	49300	1610
Weight	83/9/15	۱×	1450	02	180	<0.23	8090	1.50	126	8.00	27.0	46900	1570
		S•D•	323	10.0	22.0	90.0	451	0.2	24.2	0.1	2.65	1180	218

	DATE		Mn mg/kg	Mo mg/kg	Na mg/kg	Ni mg/kg	Pb mg/kg	Sb mg/kg	Si mg/kg	Sn mg/kg	Sr mg/kg	Ti mg/kg	v mg/kg	Zn mg/kg
±9 <u>₹</u>	83/1/26		264	<0.20	140	0•9	5.0	<2.0	13.0	2.8	12.9	7.9	3.7	56.4
₩6!gh+	83/9/15	I×	559	<0.23	134	2.7	5.9	<2.3	12.3	2.9	6.11	6.2	1.7	25.5
		S.D.	47.5	90•	21.1	9•0	0.4	9.0	12	0.8	0.7	1.6	0.2	2.9
à	83/7/26		1440	<0.10	750	33.0	29.0	<10.0	70.0	15.0	70.2	43.4	20•0	308
₩eight	83/9/15	ı× ở	3020 116	0.0	733 162	14.6	16.3	<10.0 0.0	70.0	15.7 5.7	33.0 1.6	33.3 1.6	0.0	137

x - arithmetic mean S.D.- standard deviation

ENVIRONMENTAL COMPLIANCE MONITORING DATA

TABLE 1a CAWC - HISTORICAL WATER QUALITY DATA (Values in mg/L unless otherwise stated, metals extractable) - Station x-3 = EPS 1

NH3	1	1	ı	0.350	0.0050	0.330	0.0070	9600*0	
S	-	1	ı	0.010	060*0	<0.020	ı	ı	
TOTAL	102	73	72	r	9/	ſ	50	72	
TOTAL ALKAL IN ITY	95	. 65	19	•	75	106	56	62	ker 1979).
TURB I DI TY (FTU)	36.0	73.0	5.0	r	ñ.0	2.0	0.2	0.1	tation 1 (Ba
COLOR	40.0	40.0	20•0		10.0	5.0	10.0	7.5	79-25 for St
COND. (umhos/am)	193	160	129	105	174	130	07	8	gram Report
7. 7.	691	536	46.0	F	19.0	0	<5.0	<5.0	egion Pro
FLOW (m ³ /s)	1	1	•	•	1	1	166.0	ì	aton EPS Re
TEMP C.	1	ı	1	Ξ	7	-	σ	7	present
Ħ.	7.7	7.9	7.5	7.4	8.0	7.1	7.3	7.7	-om data
DATE	1974 Average	1975 Average	1976 Average	Augus† 16 & 19 1980	Aug. 26/80	Sep † /81	July 26, 1983	Sept. 15, 1983	stermined fr
SOURCE	*EPS/DIAND	*EPS/DIAND	*EPS/DIAND	weagle STA R-1 (CAMC)	EPS STA 3	CAMC (Klohn Leonoff)	EPS	EPS	* Averages determined from data presentaton EPS Region Program Report 79–25 for Station 1 (Baker 1979).

DIANU - Department of Indian Affairs and Northern Development CAMC - Cyprus Anvil Mining Corporation

EPS - Environmental Protection Service

CAMC - HISTORICAL WATER QUALITY DATA
(Values in mg/L unless otherwise stated, metals extractable) (Continued)
- Station x-3 TABLE 18

											-
SOURCE	DATE	As	8	હ	æ	Mg	£	D Z	£	80 8	Zn
*EPS/DIAND	1974 Average	ı	ı	0.010	3.10	1	0•180	1	0.020	13.2	0.28
*EPS/DIAND	1975 Average	ı	r	0.020	. 80	ı	0.270	ı	060*0	23.8	0.12
*EPS/DIAND	1976 Average	ı	1	<0.010	0.48	1	0.080	1	0.020	8.7	90.0
Weagle STA R-1 (CAMC)	Augus† 16 & 19 1980	•	18.0	0.030	0.30	4.00	0.120	3.0	0.51	1	61.0
EPS STA 3	Aug. 26/80	<0.015	22.5	<0.010	0.16	4.80	0•050	2.2	<0.00	10.8	0.04
CAMC (Kiohn Leonoff)	Sep1/81	0.200	26.0	0.010	0.16	7.51	060*0	3.0	0.0200	ı	0.50
EPS	July 26, 1983	<0*0020	15.1	000ء	0.17	2.80	0.014	1.0	<0.0010	8.4	0.03
FPS	Sept. 15, 1983	<0.0050	22.2	0.017	0.16	3.90	0.019	1.6	0.0020	10.4	0.133
* Averages de	* Averages determined from data presentaton EPS	m data prese	entaton EPS	Region Progr	am Report	79-25 for S	Region Program Report 79-25 for Station 1 (Baker 1979).	(er 1979).			-
EPS - Enviro DIAND - Depar CAMC - Cyprus	EPS - Environmental Protection Service DIAND - Department of Indian Affairs ar CAMC - Cyprus Anvil Mining Corporation	oction Servic Lian Affairs Ig Corporatic	ce and Northe on	EPS - Environmental Protection Service DIAND - Department of Indian Affairs and Northern Development CAMC - Cyprus Anvil Mining Corporation	*						

(Values in mg/L unless otherwise stated, metals extractable) - Station x-14, EPS CAMC - HISTORICAL WATER QUALITY DATA TABLE 1b

NH3	1	1	ŧ	0.240	0.104	0.024	0.036	
8		r	0.188	0.010	0.050	r	ı	n canal
TOTAL HARDNESS	88	95	121	,	119	103	122	day diversio
TOTAL ALKAL I N I TY	7.3	65	16	1	100	7.7	88	tegion Program Report 79-25 for Station ! (Baker 1979). This station was located d/s of decant and u/s of present day diversion canal
TURB I D I TY	78	64	7	•	2	0.10	0.10	Region Program Report 79~25 for Station 1 (Baker 1979). This station was located d/s of decant and u/s of press
∞L0R	75	22	92	1	23	ī.	ľν	79-25 for the designation 19-25 for the designation of the designation
COND. (umhos/cm)	275	279	407	221	400	180	170	gram Report 7 ion was locat
л Х	252	267	11.0	1	<5.0	<5.0	< 5. 0	egion Pro This stat
FLOW (m ³ /s)	1	. 1	,	ı	ı	3.11	ı	Щ.
TEMP C°	,	•	•	10	9	01	4 .5	presenta Station
₽.	7.7	7.7	7.7	7.9	7.8	7.7	7.9	om date ted from Sreek.
DATE	1974 Average	1975 Average	1976 Average	Augus† 16, 1980	Aug• 26/80	July 26, 1983	Sept. 15, 1983	Averages determined from da 3-day average calculated fro confluence with Rose Creek.
SOURCE	*EPS/DIAND	*EPS/DIAND	*EPS/DIANU	Weagle STA R-3 (CAMC)	**EPS STA 1	EPS STA 4	EPS STA 4	* Averages determined from data presentaton EPS **3-day average calculated from Station 3 data. confluence with Rose Creek.

EPS - Environmental Protection Service DIAND - Department of Indian Affairs and Northern Development CAMC - Cyprus Anvil Mining Corporation

(Values in mg/L unless otherwise stated, metals extractable) (Continued) - Station x-14 CAMC - HISTORICAL WATER QUALITY DATA TABLE 15

SOURCE	DATE	As	స్తో	3	ē.	Ž,	Æ	8	£	804	Zn
*EPS/DIAND	1974 Average	0.18	ı	0.091	1.6	•	0.50	4	0.08	70.3	0.08
•EPS/DIAND	1975 Average	1	30.0	0.021	3.3	6. 8	61.0	ı	0.14	105	0.08
*EPS/DIAND	1976 Average	0.0010	35.7	0.038	0.44	5.7	ı	1	0.07	<u>#</u>	0.18
Weaste STA R-1 (CAMC)	Augus† 16, 1980	1	32.0	0.01	0.34	7.0	0.80	20	0.08	ı	0.20
**EPS STA 1	Aug. 26/80	<0.015	35.0	<0.010	0.71	7.6	1.4	33	080*0>	0•16	0.07
EPS STA 4	July 26, 1983	0.00070	30.6	0.002	0.15	6.2	0.26	ω	<0.0010	49.9	90•0
EPS STA4	Sept. 15, 1983	<0.00050	37.0	0،015	0	7.0	0.44	12	0.0020	59.3	0•10
* Averages d **3-day aver confluenc	* Averages determined from data presentaton EPS **3-day average calculated from Station 3 data. confluence with Rose Creek.	om data presk ed from Stat Creek.	entaton EPS Ion 3 data:		Program Report 79-25 for staton was located d/s of	79-25 for ted d/s of	Station 1 (Baker 1 decant and u/s of	Region Program Report 79-25 for Station 1 (Baker 1979). This staton was located d/s of decant and u/s of present	1979). present day diversion canal	n canal	
EPS - Enviro DIAND - Depa CAMC - Cypru	EPS - Environmental Protection Service DIAND - Department of Indian Affairs and Norther CAMC - Cyprus Anvil Mining Corporation	ection Servid dian Affairs ng Corporatio	ce and Northe on	srn Developmen†	ţ.						

CAMC - HISTORICAL WATER QUALITY DATA (Values in mg/L unless otherwise stated, metals extractable) - Station x-15, EPS TABLE 1c APPENDIX VI

SOURCE	DATE	£	Tight o	FLOW (m ³ /s)	A R	COND. (umhos/cm)	COLOR	TURBIDITY	TOTAL ALKAL INITY	TOTAL HARDNESS	8	NH ₅
*EPS	Sept. 16, 1974	7.9	-		6.5	172	S	28	16	11	ı	1
*EPS	Aug/Sept Average 1975	7.5	5/3		32.0	215	v	vo	.	86	í	•
*EPS	July/Aug Average 1976	7.7	8/10		5.0	01	8	4	76	69	<0.030	•
EPS STA 5	July 26, 1983	1.1	5	3.02	6.1	180	٠,	0.1	67	106	1	0.018
EPS STA 6a	Sep ↑ 15, 1983	7.9	4.5	ı	•	170	7.5	0.1	8	123	ı	0.031
* Averages	* Averages determined from data presentaton EPS Region Program Report 79–25 for Station 1 (Baker 1979).	om data	present	aton EPS Re	egion Pr	ogram Report	79-25 for 9	Station 1 (B	aker 1979).			
	Ers - Environmental Profection Service	ect ion	99LA I C0									

CAMC - HISTORICAL WATER QUALITY DATA (Values in mg/L unless otherwise stated, metals extractable) (Continued) - Station x-15 TABLE 1c

SOURCE	DATE	As.	0 U	3	F	M O	£	G Z	g	* 0s	Zn
€FPS	Sept. 16, 1974	•	1	<0.010	0.51	1	0.44	1	0.04	9	90•0
	Aug/Sept Average 1975	ŧ	28.0	<0.020	1.8	9.	•	ı	<0.020	•	0.03
\$45 *	July/Aug Average 1976	<0.020	18.0	0.010	0.33	0.9	ı	•	0.04	t	90.0
EPS STA 5	July 26, 1983	<0.00050	31.6	0.0020	0.13	6.3	0.22	60	<0.0010	48.8 8.8	0.05
EPS STA 68	Sept. 15, 1983	<0*00020	37.2	0.012	0.14	7.0	0.38	Ξ	0.0020	55.3	0.12
		:									
* Averages	* Averages determined from data presentaton EPS Region Program Report 79~25 for Station 1 (Baker 1979).	om data presk	entaton EP	'S Region Prog	gram Report	79-25 for	Station 1 (B	aker 1979).			<u></u>
EPS - Envi	EPS - Environmental Protection Service	ection Servio	8								