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THE GEOLOGY AND GEOCHEMISTRY OF GOLD MINERALIZATION AT ATHABASCA PASS, CENTRAL ROCKY MOUNTAINS, BRITISH COLUMBIA, CANADA

by

Robert Peter Shaw

A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND REASERCH IN PARTIAL FULFILMENT OF THE REQUIREMENTS FOR THE DEGREE OF MASTER OF SCIENCE

DEPARTMENT OF GEOLOGY

EDMONTON, ALBERTA

Spring, 1990



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The undersigned certify that they have read, and recommend to the faculty of Graduate Studies and Research for acceptance, a thesis entitled THE GEOLOGY AND GEOCHEMISTRY OF GOLD MINERALIZATION AT ATHABASCA PASS, CENTRAL ROCKY MOUNTAINS, BRITISH COLUMBIA, CANADA submitted by ROBERT PETER SHAW in partial fulfilment of the requirements for the degree of MASTER OF SCIENCE.

Supervisor C

Dedicated to the memory of my Father,

Jacob Peter Shaw January 27, 1929 - July 31, 1989

to everything there is a season

ABSTRACT

Gold-bearing quartz veins were recently discovered within anchimetamorphic quartzite-rudite and pelite sequences of the Lower Cambrian McNaughton Formation at Athabasca Pass, in the Main Ranges of the central Canadian Rocky Mountains, British Columbia. Two vein types occur, namely: an early syn-tectonic, late syn- to post-metamorphic, auriferous, bedding-parallel type, generated during repeated northeast-directed compressive tectonism, and a late, post-penetrative deformational discordant type which contain only minor gold values (<500 ppb Au). Gold emplacement and discordant veining were confined to the onset of late compression leading to thrust faulting. The spatial distribution and dimensions of the veins attest to mechanically-founded lithologic anisotropy. Bedding-parallel veins are confined to pelitic rock types. Discordant veins are confined to competent quartzitic units.

Bedding-parallel vein-filling took place in two paragenetic stages, namely: a protracted, pre-gold stage (quartz ± minor white mica and pyrite) depositing over 90 % of vein material, and a late gold-bearing - post-gold stage (quartz-pyrite-gold-galena ± white mica and Fe-carbonate). Gold is associated with brecciated pelites and coeval sulfides. Discordant veins (quartz ± minor pyrite) record a single stage of vein-filling coeval with gold deposition. Minor hydrothermal alteration (pyrite ± white mica ± carbonate) of adjacent wallrocks accompanied gold emplacement.

Trace element analysis reveals distinct enrichment of Pb, As, Ba, and Fe, and marginal enrichment of Ag, Zn, and Cu in veins with respect to host rocks. Pb, As, Ba, and Fe show distinct, although non-proportional positive correlation with gold concentration in the lodes, whilst Ag, Zn, and, Cu do not. When compared with other lode-gold deposits in similar

domains, the Athabasca Pass lodes are unenriched in all trace elements except gold.

Microthermometric analyses of fluid inclusions indicates gold-stage fluids were aqueous brines containing 2 to 10 wt. % NaCl eq. and up to 12 mole % $CO_2 \pm CH_4$. Variations in fluid compositional-and total homogenization-data suggest that gradual fluid evolution accompanied gold mineralization. Gold deposition took place at between 275° and 350°C and 900 to \geq 1200 bars.

Vein pyrite δ^{34} S(CDT) values cluster between +14.2 and +16.3 ‰. Coeval galenas exhibit δ^{34} S values between +11.4 and +13.3‰. Pyrite-galena geothermometry reveals a mean temperature of mineralization of 300 ± 43°C. Comparison of the δ^{34} S values of vein pyrites, with values for pyrite porphyroblasts in the country rocks reveals that vein sulfur was derived from the McNaughton Formation.

 δ^{18} O(SMOW) values of McNaughton Formation quartzites and pelites cluster between +12.0 and +13.5‰, and +9.5 and +10.5‰, respectively. All quartz veins exhibit δ^{18} O values between +13.0 and +15.0‰. Vein inclusion fluids exhibit δ D values between -105 and -124‰(SMOW). Combined O-H-isotope data are most compatible with a source fluid involving chemically-and isotopically-evolved meteoric waters. A model involving vein deposition from fluids undergoing post-peak metamorphic cooling is proposed.

The geology and geochemistry of the Athabasca Pass gold-lodes indicates they are siliciclastic-hosted analogues of the Turbidite-Hosted class of gold deposits. Consideration of the processes involved in lode formation at Athabasca Pass suggests that gold-lodes may be more common in the Canadian Rocky Mountains than presently recognized. Some implications for their genesis are discussed.

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CHAPTER 1

General Introduction

Gold occurrences in the Rocky Mountains of Alberta and British Columbia, either lode or placer, are rare. The few documentations, (dominantly of placer occurrences) are buried within pre-1960 discussions of regional history, or broad overviews of regional geology, and provide little detail as to the nature and origin of the gold occurrences. Much of this nonchalance regarding the gold of this region likely stems from the concepts and priorities of nineteenth century prospectors who traditionally sought rich, easily-extractable placers (such as the interiors of British Columbia and Yukon Territory offered), or igneous - lode associations, a geologic improbability in the vast sequences of sedimentary rock types dominating the Canadian Rocky Mountains. Prospecting was likely further deterred by the rugged terrain, poor accessability, abundant glacial overburden, short working season, and lack of placer indicators characteristic of the Rocky Mountains. The lack of detailed geological mapping throughout much of the western ranges has only served to prolong these erroneous beliefs. In August of 1986 however, a series of auriferous quartz lodes hosted by low-grade meta-sedimentary rocks of Early Cambrian age, were discovered at Athabasca Pass, in the Main Ranges of the Canadian Rocky Mountains, British Columbia. The principle objective of this thesis is to provide detailed documentation of aspects of the geology and geochemistry of these gold occurrences.

This thesis is presented in "paper format," formally represented herein by Chapters 2, 3, and 4. Chapter 2 documents the physical

relationships of the gold occurrences utilizing traditional field-mapping and petrographic techniques and establishes the structural, mineralogical and temporal relationships of the gold-bearing lodes to their geological surroundings. Chapters 3 and 4 explore the geochemistry of the auriferous veins and their host strata, utilizing trace element, fluid inclusion (Chapter 3), and stable isotope (Chapter 4) analyses. As well, Chapter 4 considers some broader implications for the genesis of mesothermal gold-quartz veins in lithosedimentary domains. Some general implications regarding gold in the Rocky Mountains comprise the concluding statement (Chapter 5).

The Athabasca Pass gold occurrences are not, at present, economically viable mining entities. Their study however is warranted, from both an academic perspective, in terms of their applicability to the understanding of lode-gold genesis in low-grade meta-sedimentary domains, and from an economic perspective in view of exploration modelling for geologically similar, economically extractable deposits in the Canadian Rocky Mountains.

CHAPTER 2

Gold Mineralization in Lower Cambrian McNaughton Formation, Athabasca Pass, Canadian Rocky Mountains: Structural, Mineralogical, and Temporal Relationships.

Introduction

The Athabasca Pass is located on the continental divide, 60 km south-southwest of Jasper, Alberta, in the northern Park Ranges of the central Canadian Rocky Mountains (Figs. 2-1 and 2-2). The pass was pioneered in the early 1800's by David Thompson, as an alternate route through the Main Ranges to the Columbia River. A trading post was established near the Committee Punch Bowl, but the route fell into disuse by the middle of the nineteenth century. The area has no recorded history of exploration or mining. This paper describes gold-bearing quartz veins which occur in quartzose clastic rocks of the Lower Cambrian Gog Group of the Athabasca Pass.

Little information regarding the geology of the Athabasca Pass has been published. Thus preliminary documentation of the general stratigraphy, lithology, and structure in the area exhibiting gold-quartz mineralization is provided. The map presented here utilizes information gathered through personal communications with E.W. Mountjoy and coworkers, unpublished Operation Bow-Athabasca data (Price and Mountjoy, 1966; Mountjoy and Price, in preparation), and field-mapping carried out by the author during 1987 and 1988.

Previous Work

Geologic studies pertaining to the Athabasca Pass are few and of

regional scope. Recent studies include reconnaissance mapping at 1:250,000 scale by Campbell (1968) and at 1:50,000 scale by Price and Mountjoy (1970). The latter, a product of Operation Bow-Athabasca, includes a 1:50,000 scale map sheet corresponding to the NTS 83D/8 (Athabasca Pass) map sheet, which has yet to be published (Mountjoy and Price in preparation). More detailed documentation of the surrounding region include studies of structure, stratigraphy and metamorphism to the west and southwest of the Athabasca Pass by Craw (1978) and Klein and Mountjoy (1988), and to the north and northwest of the Athabasca Pass by Mountjoy and Price (1985; 1989), Mountjoy and Forest (1986), Leonard (1985), Oke (1982), and McDonough and Simony (1988).

Information on lode gold deposits within the Main Ranges and indeed, over the entire central Canadian Rocky Mountains is sparse. Sorensen (1955) notes the occurrence of auriferous quartz veins in Precambrian Miette "formation" near Tete Jaune Cache, however provides no details. All other recorded lode gold occurrences in this region are located west of the Rocky Mountain Trench. Reviews of mineral occurrences in the Main Ranges of the Rocky Mountains were given by Hedley (1954) and Little et al. (1976). Summaries of the nature and distribution of lode gold deposits in southeastern British Columbia in the vicinity of the Rocky Mountain Trench were published by Mathews (1944), Holland (1944), Sinclair et al. (1978), and Schroeter and Panteleyev (1986).

The occurrence of gold placers in the valleys of the central Rockies and contiguous foothills of Alberta has received only cursory attention in the literature. Brief mention of placer gold near the headwaters of the Fraser River was made by Sorensen (1955) and Boyle (1979), and in the Fort Steel

and Ptarmigan Creek regions by Mathews (1944). The distribution of placer gold in Alberta rivers draining the Rocky Mountains was reviewed by Giusti (1983).

Regional Geology

The Athabasca Pass lies at the boundary between the eastern Main Ranges and western Main Ranges of the central Rocky Mountains (Price and Mountjoy, 1970). The boundary at this latitude is marked by a southwestdipping thrust fault, the Chatter Creek fault (Figs. 2-2 and 2-3; Wheeler, 1963; Wheeler et al., 1972), which extends from north of the headwaters of the Fraser River (Price and Mountjoy, 1966; Mountjoy and Price, 1989), southeasterly through the study area, to the northwest of Golden. Southwest of the Athabasca Pass, the hanging wall of the Chatter Creek fault is composed of grits, pelites, psammites and carbonates of the Hadrynian Miette Group, overlain by Lower Cambrian clastic rocks of the Gog Group and, near the Rocky Mountain Trench, by un-named pelites and carbonates of Middle Cambrian age (Price and Mountjoy, 1970). This region is dominated by broad open folds comprising the Baker Glacier Syncline and Porcupine Creek Anticlinorium (Price and Mountjoy, 1970; Balkwill, 1968; Klein and Mountjoy, 1988). Northwest of the Athabasca Pass, across the head of Hugh Allan Creek, the Chatter Creek thrust sheet contains another major antiformal structure, the Fraser River Antiform (Mountjoy and Forest, 1986). Within the Chatter Creek thrust sheet, metamorphic grade increases westward from greenschist grade to kyanite-staurolite-bearing assemblages of amphibolite grade (Craw, 1978; Leonard, 1985; Klein and Mountjoy, 1988). The Purcell thrust and associated structures (Price and Mountjoy, 1970) form

the western boundary of the Chatter Creek thrust sheet.

The footwall of the Chatter Creek fault, to the north and east of the Athabasca Pass is composed of Lower Cambrian Gog Group, overlain by a series of thickly-bedded, dominantly carbonate rocks, of Middle Cambrian age. The structure of this eastern sector of the Main Ranges consists of thick, relatively flat, thrust sheets with characteristically broad, open folds (Price and Mountjoy, 1970; Mountjoy and Price, 1989). Local structural complications occur along the transition between the eastern and western Main Ranges (Wheeler, 1963; Cook, 1975). Rocks of the eastern Main Ranges have been regionally metamorphosed to pumpellyite and lower greenschist grades (Read, 1988).

Local Geology

The geology of the Athabasca Pass area is illustrated in Figures 2-3 and 2-4. To date, gold-bearing quartz veins have been found only in the lowermost formation of the Gog Group, the McNaughton.

The McNaughton Formation

The McNaughton Formation comprises a variety of mature, quartz-dominated clastic rocks of sub-greenschist metamorphic grade. The predominant rock type is a medium- to coarse-grained, moderately- to poorly-sorted, pale weathering, grey quartzite. This quartzite contains a variable amount of pelitic material, 5-10 % on average, although units grading to quartzitic pelite and pelite are not uncommon. Generally well-stratified, quartzites form sequences of 0.1 to 3 m thick beds randomly parted with pelite horizons. Most of the quartzite beds are tabular although gently undulatory and wedge-shaped beds are also present. Feldspar content is low,

generally less than 3 %. Both phyllosilicate and feldspar content are highest in the lower exposed portions of the section.

Minor conglomerate occurs as the basal portion of graded beds or as lenticular beds up to 0.3 m in thickness. Conglomerates are generally matrix supported. Subequal amounts of well rounded quartzite and vein quartz clasts, ranging up to 25 mm in greatest dimension, are contained within a poorly sorted matrix of medium to fine quartz sand and phyllosilicates, cemented by silica. Phyllosilicates average 25 % of the matrix.

Pelites comprise approximately 2-3 % of the McNaughton Formation occurring primarily as discrete, discontinuous, 5 cm to 1 m interbeds in the quartzites, or commonly, at the top of normally graded sequences. They consist almost entirely of fine-grained white micas with variable quantities of granule- to sand-sized detrital quartz grains and are generally pale green in colour. The pelites commonly contain up to 3 % very finely disseminated pyrite.

Paleoenvironmental interpretations of the McNaughton Formation include a tidally-dominated association of shallow marine shelf environments in the eastern Main Ranges (Palonen, 1976; Hein, 1984; Woberg, 1986), and fluvial braidplain to tidal complex transitions in more westerly outcrops (Young, 1979). Paleocurrent studies by Mountjoy and Aitken (1963) indicate a predominantly westward and southwestward transport of sediments, the inferred source area being the high-grade metamorphic and igneous rocks of the North American craton, with a possible minor contribution reworked from earlier-deposited sandstones (Young, 1979).

Host-Rock Petrography

Petrographic investigation of various McNaughton rock types has been undertaken by Akehurst (1964), Palonen (1976), and Young (1979).

Mineralogy

The mineralogy of the McNaughton rock types consists of three components; quartz, white mica and, potash feldspar. The modal abundance of these components in over 90 hand specimens from the Athabasca Pass is plotted in the form of a ternary diagram (Fig. 2-5). Modal composition ranges from quartzitic pelite to feldspathic quartzite. Pelitic quartzite containing 5-10 % white mica is the most common rock type. The occurrence of white mica is variable but ubiquitous. K-feldspar content is erratic with 15-20 % feldspar recorded in some horizons, but less than 3 % being more typical. Varietal minerals are rare. Those observed in thin section include epidote, rutile, zircon, augite, and muscovite. Anhedral to euhedral porphyroblasts of pyrite are common in the more pelitic McNaughton rock types, where they may comprise up to 3 % of individual horizons. Minor (<2%) recrystallized carbonate is widespread, as is the incipient replacement of K-feldspar by white mica ± carbonate (Fig. 2-6C).

Deformation - metamorphism

Microstructures observed in quartz grains are dominantly pressure solution features including grain-to-grain contact suturing, marked grain serration and embayment, and stylolite development (Figs. 2-6A, 2-6B). Undulatory extinction is variably developed, being pervasive in grain

supported quartzites but less distinct in matrix supported pelitic quartzites. The apparent lack of microscale sinks for pressure solution-derived quartz, such as quartz overgrowths and pressure shadows, is noteworthy.

White mica in the McNaughton rock types probably represents recrystallization, during low-grade regional metamorphism or deformation, of an originally allogenic clay component. Foliation defined by these micas becomes discernable as phyllosilicate contents reach 20 to 30 %. Individual phyllosilicate crystals average 0.05 mm in length, becoming coarser along solution contacts between quartz grains, within quartz grain embayments, and along stylolitic partings. The pyrite porphyroblasts and recrystallized carbonate noted earlier are also considered the products of regional metamorphism. Read (1988) infers McNaughton strata in the Athabasca Pass have been subjected to pumpellyite to lower-most greenschist facies regional metamorphism. An anchimetamorphic (ca. pumpellyite facies) regime is implied by the assemblage quartz-white mica ± pyrite ± carbonate (Frey and Kisch, 1987; Frey, 1987), and by the deformational microstructures (Kerrich, 1977; Beach, 1979) outlined above.

Structural Geology

The Chatter Creek fault and northeast-verging folds dominate the geologic structure of the Athabasca Pass. The trace of the Chatter Creek fault is not exposed in the study area. The nature and geometry of this fault is not fully understood (Mountjoy and Forest, 1986; Dechesne and Mountjoy, 1988). Recent interpretations (Dechesne and Mountjoy, 1988; Mountjoy pers. comm. 1989) indicate it might be a steeply dipping, late (out-of-sequence) thrust. This conclusion was implied by earlier mapping to the south, in the

area northeast of Golden (Cook 1975). Displacement on the fault is considered to be about 3 to 4 km, and was possibly related to late movement on the Hugh Allen (Purcell) thrust (Mountjoy, 1988).

Mountjoy (1989, personal communication) indicates that at least four discrete phases of regional deformation are recorded in the Main Ranges. Extensive, premetamorphic thrust sheets which underlie the western and eastern Main Ranges formed first (Mountjoy and Dechesne, unpublished data). These structures were later folded in regional-scale antiformal culminations such as the symmetamorphic Porcupine Creek Anticlinorium and the post-metamorphic Fraser River Antiform. Out-of-sequence thrusts, including the Chatter Creek fault, post-date these structures.

The structure of McGillivray Ridge

The eastern wall of the Athabasca Pass, known as McGillivray Ridge (NTS 83 D/8; Figs. 2-3, 2-4 and 2-7 in this study) consists of about 800 m of folded and faulted McNaughton Formation. Two sets of structures, imbricate thrust faults and mesoscopic folds, are clearly exposed on the southern-most precipice of this ridge (Figs. 2-7, 2-8). Northeast-verging imbricates of McNaughton quartzite are thrust over upper Gog Group and Middle Cambrian strata along a lower-most, northeast-verging footwall splay off the Chatter Creek fault. This detachment has been mapped to the north through the Fraser Pass by Mountjoy and Price (1989) where it duplicates McNaughton strata, and is herein termed the McGillivray Fault (Figs. 2-2, 2-3, 2-4, and 2-7). A quartz-rich mylonite up to 2 m thick is associated with this sole thrust at the base of McGillivray Ridge. The thrust dips approximately 20° southwest, striking about 145° east of north. The imbricate thrusts are of similar strike, but dip more steeply (25° to 30°) and are listric in nature,

converging asymptotically with the McGillivray Fault. The structural geometry of McGillivray Ridge is that of a leading imbricate fan (Boyer and Elliott, 1982).

Tight to isoclinal, overturned mesoscopic folds (informally referred to herein as f1) occur within the imbricate quartzite slices (Fig. 2-8). These folds are northeast-verging with axial planes that dip moderately to steeply to the southwest and fold axes which trend 140° to 150°, plunging gently to the southeast. In some instances overturned fold limbs are markedly thinned and incipient planes of detachment occur parallel to the local imbricate thrust faults (Fig. 2-8). Penetrative planar fabric is weakly developed in the more pelitic quartzites, but occurrences of intact pelite contain a southwest dipping, locally crenulated, axial planar cleavage (informally referred to herein as s1). The sparsity of crenulation development suggests s1 does not overprint a previous tectonic fabric. Crenulation is considered to be the result of the folding of a variably developed, burial-enhanced, beddingparallel schistosity. f1 folds are truncated by the imbricate faults, and are cut by late sets of conjugate shear and AC joints (Hobbs et al., 1976) which are probably related to thrust faulting. f1-style folding is not seen elsewhere in the Athabasca Pass.

Correlation of fold and thrust structures on McGillivray Ridge with the deformational phases of Mountjoy (1989, personal communication) is inhibited by a lack of detailed mapping in the region surrounding the Athabasca Pass. Imbricate faulting truncates, and therefore post-dates, folding on McGillivray Ridge. These faults are not deformed nor do they appear to reactivate earlier structures. Thus they probably formed during development of the Chatter Creek fault. Folding in the McNaughton is of uncertain age

and may be related to any of the four regional events. However, elsewhere in the Main Ranges folds associated with early, premetamorphic thrusting are confined to the vicinity of early thrusts (Mountjoy and Forest, 1986; Klein and Mountjoy, 1988; Dechesne and Mountjoy, 1988), none of which outcrop in the Athabasca Pass (Mountjoy and Price in preparation). As well, folds associated with regional syn- and post-metamorphic antiformal culminations are described as generally macroscopic, upright, and open to gentle, with associated axial planar crenulation cleavages (Mountjoy and Forest, 1986; Klein and Mountjoy, 1988), and are thus distinctly different from those on McGillivray Ridge. Folding during development of the Chatter Creek thrust and its associated footwall imbricates is suggested by the colinear nature of the McGillivray folds and thrusts, the incipient detachment of fold limbs parallel to thrust faults, the lack of similar fold styles elsewhere in the Athabasca Pass, and the apparent lack of penetrative fabric development prior to s1. The McGillivray folds may however record later tightening and overturning of earlier phase folds. Thus, although the initiation of these folds during earlier regional thrust or antiformal development cannot be precluded entirely, much of their development, and present geometry, is probably attributable to deformation leading to development of the Chatter Creek thrust.

Quartz Veins of McGillivray Ridge

Gold-quartz mineralization outcropping on the southwest slope of McGillivray Ridge is contained in a series of discrete vein structures confined to quartzites ± pelites of the McNaughton Formation. The quartzites appear unaltered on the mesoscopic scale and apparently contain

no disseminated epigenetic gold mineralization. To date over 20 veins have produced anomalous gold values. Distribution of gold within individual veins is highly erratic, grading locally from nil to over 500 g Au/tonne with visible gold observed in many cases. Channel sampling across some veins has yielded grades of between 10 and 15 g Au/tonne.

Vein type, morphology and distribution

Two categories of quartz veins are recognized on McGillivray Ridge; veins which parallel bedding planes within the McNaughton Formation, and veins which are discordant to sedimentary layering (Fig. 2-9). Only bedding-parallel veins contain high-grade gold mineralization; the discordant variety have yielded values up to *ca.* 500 ppb gold.

Bedding-parallel veins

Bedding-parallel veins vary from about 1 m in length and a few centimeters in thickness up to 50 m in length, varying between 0.7 and 1 m in thickness. They weather recessively, and locally exhibit surficial gossan due to the oxidation of pyrite. Their lateral extent is presently unknown. Inspection of bedding-parallel veins indicates that, in all cases, these structures invade bedded pelites, the remnants of which are generally seen as angular, brecciated fragments within the veins. In most cases the pelites have been entirely disrupted by veining, creating a vein-quartz supported, pelite breccia. Thus these veins essentially parallel the pelite horizons they have invaded. Vein size and distribution was fundamentally controlled by the geometry and distribution of the original pelite layers in the McNaughton Formation.

Lateral lithologic variation within pelites also appears to have exerted control on the localization and extent of bedding-parallel veining. This is suggested where individual quartz veins die out along strike, coincident with an increase in detrital quartz in the associated pelite horizon. Veining may reappear further along strike as the horizon again becomes more pelitic. Veining is not developed in horizons containing greater than approximately 30 % detrital quartz. Pelite fragments in veins generally contain less than 10 % quartz grains.

Discordant veins

Discordant veins are confined to the competent quartzitic rock types of the McNaughton Formation. Their occurrence is widespread, either as individual isolated veins, *en echelon* vein arrays, or as clustered vein stockworks which extend discontinuously for tens of meters. Individual veins are generally planar to sigmoidal in shape and range in size from centimeter-scale fracture fills to veins 3 m in length and 10 cm in thickness.

The relationships between bedding-parallel and discordant veins are illustrated in Fig. 2-9. Where the two vein systems are in contact, discordant veins usually appear as planar to sigmoidal *en echelon* vein arrays, obliquely intersecting bedding-parallel structures. They cut into, but do not offset or truncate bedding-parallel structures, nor are they truncated by them. Zones of abundant bedding-parallel veining are usually coincident with zones containing greater numbers of discordant veins.

Structural geometry and evolution of the veins

Contoured lower hemisphere stereographic probability diagrams (Fig. 2-10) project poles to both bedding-parallel and discordant vein structures, in

relation to fold axial planes and associated thrust faults on McGillivray Ridge. Orientation data were processed using the FORTRAN 77 program ORIENT (Charlesworth et al., 1989). The probability method of contour estimation has been discussed by Ramsden and Cruden (1979). Regarding these cumulative data, the following deficiencies are noted: The dominant population of northeasterly dip measurements for bedding-parallel veins is considered to be biased because the complimentary southwest-dipping limbs of mesoscopic folds on McGillivray Ridge have mostly been removed by erosion. As well, although numerous mesoscopic folds are seen in outcrop (Fig. 2-8), they are generally inaccessible to physical measurement, thus the quantity of data is limited and the overall uncertainty may be high. Despite these limitations, the geometric relationships between folding, thrusting and veining (Fig. 2-10) are considered to be representative.

Bedding-parallel veins

The distribution of poles to bedding-parallel veins (Fig. 2-10A) is essentially that of a "half" girdle reflecting the paucity of southwest-dipping fold limb data. Nevertheless, the folded geometry of the bedding-parallel veins is clearly demonstrated (Ragan, 1973) implying that bedding-parallel veining was initiated prior to, or early in, the progression of folding and thrusting on McGillivray Ridge.

Hydraulic-enhanced fracturing may be advocated to explain bedding-parallel vein formation during regional tectonic compression (Secor, 1965; Hobbs *et al.*, 1976; Fyfe *et al.*, 1978; Etheridge *et al.*, 1984). In general, under passive conditions, the greatest effective principle stress (σ_1) is compressive, being approximately normal to the earth's surface. As tectonic compression

progresses σ_1 becomes essentially horizontal and thus approximately parallel to yet-undeformed bedding. The least effective principle stress (03, also compressive) becomes approximately vertical (i.e. normal to the earth's surface), and varies as a function of the bulk density of the overlying fluidsaturated lithology, the acceleration due to gravity, the depth of overburden, and the ratio of pore fluid pressure to lithostatic pressure (Hubbert and Rubey, 1959; Secor, 1965). Should pore fluid pressures become sufficiently high, og will become tensile, and horizontal, bedding-parallel, extensional fracturing and vein filling can occur. Comparable hydraulic fracture mechanisms have often been advocated in the genesis of vein-type gold deposits, particularly of the syntectonic-metamorphic (Cox et al.,1986), and turbidite-hosted (Keppie et al., 1986) varieties (e.g. Henley et al., 1976; Kerrich and Allison, 1978; Graves and Zentilli, 1982; Goldfarb et. al., 1986; Mawer, 1986). Hydraulic fracturing within the pelite horizons of the McNaughton Formation would be favoured because these rock types form the least competent constituents in the thick, quartzite-dominated sequence. Pelite competency would be reduced by the bedding-parallel fissility and relatively higher pore-fluid pressures characteristic of pelitic rock types under low-grade metamorphic conditions (Fyfe et al., 1978; Murrell, 1985).

It is not possible to fully constrain the timing of bedding-parallel veining in the Athabasca Pass. The veins are folded, and must therefore predate or coincide with local folding (f1). As shown, much of this folding is Chatter Creek thrust-associated. However, vein initiation during earlier, regional premetamorphic thrusting, or the development of regional antiformal culminations cannot be precluded, and rather is suggested by vein textures and mineralogy (reviewed below) indicating multiple episodes

of vein reopening and deformation, and syn- to post-peak metamorphic vein infilling.

Discordant veins

The clustered nature of discordant vein poles (Fig. 2-10B) suggests that these veins are relatively undeformed planar structures (Ragan, 1973). Sigmoidal geometries are also observed. The mean strike of discordant veins is ca. east-west, with dips averaging 40°-60° north. Assuming the greatest principle stress was directed at right angles to fold and thrust structures striking N140°-150°E (i.e. compression directed ca. N50°-60°E), discordant vein orientation is not compatible with fold- or thrust-associated AC or conjugate shear fracturing (Hobbs et al., 1976; Ramsay and Huber, 1986). The consistent orientation, en echelon arrangment, and sigmoidal geometry of the veins is suggestive of lateral shear-induced (in this case sinistral) extensional fracturing, and may record a small component of shear associated with Chatter Creek deformation. Discordant veins clearly postdate f1 (Fig. 2-10). Veins are cut by unveined joints associated with late thrusting. Thrust faults contain no associated veining or alteration, suggesting veining pre-dates imbricate thrusting. Thus discordant veining is temporally constrained between late folding (f1) and early development of the Chatter Creek fault system.

Vein Petrography and Host-rock Alteration

Petrographic study of over 250 hand specimens and 80 thin- and polished-thin- sections of vein and host-rock material was undertaken in order to document vein-textural, mineralogical, and paragenetic relationships, to evaluate the degree of host-rock alteration, and to place

constraints upon the timing of vein- and gold- emplacement in relation to the local structural evolution of McGillivray Ridge. Semi-quantitative elemental analyses of specific mineral phases were obtained utilizing a Cambridge Stereoscan 250 scanning electron microscope equipped with a Kevex energy dispersive spectrometer.

Bedding-parallel veins

Figure 2-11 summarizes the mineralogy, and paragenetic and structural evolution of the bedding-parallel veins. Quartz is the dominant vein-filling phase, typically comprising over 95 % of the total vein volume. Variable, unevenly distributed amounts of sulfide, carbonate, and white mica comprise the remaining modal fraction. Brecciated fragments of host pelite may comprise a significant volume within individual veins, depending on overall vein size, and the degree of fracturing and disaggregation of the pelite horizon. The distribution of hydrothermal vein constituents generally follows that of the pelite fragments, with the quantity of sulfides, carbonate, white mica, and gold being markedly higher in zones of intense pelite brecciation.

Internal vein structures and textures

In general, the margins of bedding-parallel veins are marked by an abundance of brecciated pelite. These represent the remnants of pelite horizons which have been entirely disaggregated by quartz veining. Where pelite occurs at the top of normally graded beds, brecciation and veining occurs where pelite predominates.

Many of the macroscopic structures and textures exhibited by beddingparallel veins are illustrated in Figure 2-12. The majority of quartz was deposited as open-space fillings during one or more episodes of veinopening. Well terminated quartz crystals indicate unimpeded crystal growth
during vein filling. These crystals are usually truncated, dislocated, and
overgrown by similar quartz deposited during subsequent reopening and
filling events, creating a generally massive vein texture. Vein reopening
took place within previously deposited vein fillings, along the wallrock-vein
contact, or preferentially, along the margins of larger pelite fragments. Pelite
fragments are commonly cut by more than one generation of quartz veining.
Successive fracture fillings are spatially confined to the vein/pelite horizon
and do not cut the quartzite wall rocks. No open spaces are observed in the
veins.

Laminated textures indicative of incremental vein growth ("crack-seal filling", Ramsay, 1980; Cox and Etheridge, 1983) are locally developed (Fig. 2-13A) further suggesting multiple episodes of vein reopening.

Mineralogy

Bedding-parallel vein minerals are discussed according to their paragenesis (Fig. 2-11). Pelite fragments are considered vein constituents because the original bedded character of these pelites has been destroyed by the veining process.

Pelite fragments: Angular fragments of pelite (Figs. 2-12 and 2-13) range in size from microscopic inclusions to several centimeters in greatest dimension. These fragments are generally most abundant near the vein margins where they may constitute greater than 50 % of the vein volume. Contacts between pelite fragments and the surrounding vein-quartz matrix are sharp, with the pelite exhibiting little visible evidence of hydrothermal alteration. A weak, pervasive foliation is present in the pelite fragments,

with a distinct crenulation cleavage being noted in some cases. Neither the pelite fragments nor their foliation exhibit a preferred orientation or common alignment within the veins.

Pelite fragments in thin section occasionally reveal a thin (<ca. 0.1 mm) selvage of relatively coarse, hydrothermally recrystallized mica with individual crystals up to 0.3 mm in length. This unfoliated selvage surrounds a finer grained core of weakly foliated white mica, the product of anchimetamorphism.

Semi-quantitative energy dispersive microanalysis of phyllosilicates from 15 pelite occurrences, including both mineralized and unmineralized horizons were performed. Cations present include the essential K, Al, and Si and, consistently, between about 3 and 6 weight % Fe (±Ti). This latter component is likely responsible for the greenish tint of the micas, with Fe⁺² substituting for Al in octahedral sites (Deer *et al.*, 1962).

Many pelite fragments contain the oxidized remnants of a finely disseminated, iron-rich phase, likely pyrite, which may be seen unaltered in the center of the inclusions. Hydrous iron oxides (limonite-goethite) now occupy the fine (≤0.2 mm) anhedral to subhedral casts. Based upon its petrographic character, this pyrite is considered to be metamorphic in origin.

Quartz: This mineral accounts for greater than 90 % of the hydrothermal vein-filling constituents. On the basis of textural and optical characteristics in hand specimen and thin section, three distinct varieties of quartz, Types I, II and, III are discernible.

Type I quartz (Figs. 2-12 and 2-13) constitutes 90-95 % of the veinfilling quartz. It is commonly comprised of massive aggregates of well terminated, millimeter- to centimeter-sized crystals. Well developed, submillimeter-scale growth zonation in euhedral crystals is discernible in hand specimen. Fine-grained (≤0.05 mm) white mica enhances these growth zones in thin section (Fig. 2-13C).

Type I quartz is readily identifiable in plane-polarized light by multiple generations of healed fractures, presently demarcated by trails of fluid inclusions. In cross polarized light undulatory extinction and sub-grain development are ubiquitous. Deformation lamellae are locally well developed. Pressure solution causing grain boundary suturing is widespread, and where abundant is accompanied by incipient recrystallization along neighbouring grain boundaries. Fine-grained, recrystallized quartz exhibits distinct undulatory extinction.

Type II quartz is locally developed granular, fine-grained (1-2 mm) quartz which occupies thin (0.5-2 cm), discrete veinlets contained within earlier deposited Type I quartz (Fig. 2-12). Petrographically, Type II quartz is polygonal in form and exhibits healed fractures, undulatory extinction and subgrain development. Type I quartz grains bounding these veinlets are generally highly strained and sutured (Fig. 2-13D) thus, Type II quartz is interpreted as local, pressure solution-derived vein quartz, deposited in discrete dilational fractures created during the progressive deformation of earlier deposited Type I vein-fillings.

Type III quartz (Figs. 2-12 and 2-14D) is distinguished in hand specimen by its milky white colour and fine- to medium-grained (1-5 mm) granular texture. This variety of quartz is not confined to discrete structures, but is seen filling irregular fractures and voids of variable size and shape, most commonly originating near vein margins and extending into Type I and II vein structures. Microscopically, Type III quartz exhibits a lesser degree

of deformation than Type I. Undulatory extinction is ubiquitous but of lesser intensity. Subgrains, sutured grain boundaries and healed fracture patterns are also locally developed but to a much lesser degree.

Type III quartz is also distinctive for the greater quantities of associated white mica, pyrite and native gold relative to quartz Types I and II.

White mica: This minor vein constituent (<1%) is fine-grained (0.01-0.1 mm), occurring as disseminations and aggregates along late, partially healed fractures in all quartz types.

The origin of much of the fine disseminated and fracture dispersed white mica is considered to be in part hydrothermal, but a significant quantity (perhaps 50%) of all the white mica is considered to have been derived through brecciation and dispersal \pm recrystallization of original pelitic host-rock (Figs. 2-12 and 2-13A). It is generally not possible to petrographically distinguish these mica types.

Pyrite: Two generations of pyrite are recorded. The first, diagenetic or metamorphic pyrite in pelite has been described. The second (< 1% of total vein volume) consists of 0.05 to 5 mm subhedral to euhedral cubes which commonly exhibit finely striated crystal faces, and is hydrothermal in origin. The majority of hydrothermal pyrite occurs as disseminations, clusters and fine stringers in Type III quartz, or in the immediate vicinity of (ca. 5 mm radius), nucleated upon, or replacing pelite fragments in Types I and III quartz. Pelite-unaccompanied pyrite in Type I quartz is rare.

Post-depositional deformation of hydrothermal pyrite is uncommon. Minor fracturing is filled with quartz, and occasionally, native gold (Fig. 2-14A). Sporadic inclusions of white mica, quartz, and native gold also occur in pyrite.

Gold: Although most of the bedding-parallel veins contain gold, its

distribution within individual veins is erratic, its presence being enhanced in zones containing abundant pelite fragments, hydrothermal pyrite and Type III quartz, concentrated near vein margins.

Gold is present exclusively as native metal. Sub-equant inclusions range from 0.04 to about 2 mm in size, occurring close to, or abutting against, brecciated pelite fragments or hydrothermal pyrite crystals (Figs. 2-14B and 2-14C). Less commonly, inclusions in pyrite are seen.

Native gold also occurs as fracture fillings in Types I and III quartz (Figs. 2-14A, 2-14B, and 2-14D). In Type I quartz, gold occurs as fine dendritic fillings up to 5 mm in greatest dimension. Gold filled fractures are generally traceable to nearby (up to 1 cm distant) zones of Type III quartz mineralization. Fractures confined to Type III quartz also contain gold and usually propagate from zones rich in hydrothermal pyrite, white mica and pelite breccia. Very rarely, gold with pyrite fills fractures in Type II quartz.

Energy dispersive microanalysis of the native gold indicates it is of exceptionally high fineness. Over 100 spot analyses failed to detect the presence of any element other than gold.

Galena: Occurrences of galena are limited to sporadic, irregular, undeformed fracture fillings in Types I and III quartz, where it is commonly accompanied by native gold (Fig. 2-14D). Minor replacement by hydrous iron-oxides has taken place at the perimeter of galena fills. Although the presence of galena is a good indicator of enhanced gold concentrations, galena is actually less common than visible native gold.

Carbonate: Occurrences of small (<3 mm), anhedral to euhedral inclusions of rust-brown weathering carbonate are rare. When present, this mineral is seen abutting pelite fragments in Type I quartz, or accompanying

pyrite ± native gold in Types II and III quartz. Based upon its brownish color the carbonate is likely ankeritic in composition.

Discordant veins

Discordant veins are filled with quartz, and rarely minor pyrite (<0.5-1%). Quartz is medium-grained, massive to granular, anhedral, and exhibits fracturing, undulatory extinction, and minor pressure solution. Pyrite is fine-grained (<3 mm) and subhedral to euhedral. Discordant veins exhibit essentially one stage of vein filling. Gold values range from less than 5ppb to 500 ppb and appear to correlate with pyrite content. Visible gold has not been observed.

Hydrothermal alteration

Wall-rock alteration is notably absent adjacent to most bedding-parallel veins. Where veining was initiated at the top of graded beds minor pyritization is occasionally developed in the adjacent, unveined pelitic quartzite. Similar sulfidation is also noted where discordant vein structures cut pelitic quartzites which lack bedding-parallel vein development. Zones of sulfidation immediately adjacent to veins may carry up to 1 ppm gold.

More widespread is the incipient replacement of detrital potash feldspar by white mica ± minor iron-rich carbonate (Fig. 2-6C). The origin of these constituents is ambiguous as their occurrence is not restricted to the strata adjacent to veins. They may represent a hydrothermal assemblage associated with vein filling, but are more likely the products of regional anchimetamorphism.

Paragenesis

Two broad generations of vein filling are recorded in the bedding-parallel veins of McGillivray Ridge (Fig. 2-11). The first of these (pre-gold stage), represented by pelite fragments, Types I and II quartz and hydrothermal white mica ± minor pyrite and carbonate, records a protracted history of repeated vein opening and filling during which over 90 % of the hydrothermal vein constituents were precipitated, predominantly as open space ± laminated fillings. The second generation (gold and post-gold stages) of vein filling represents a late, volumetrically minor incursion of hydrothermal fluids during which most of the gold was deposited. Within this stage, deposition of most of the Type III quartz and pyrite ± white mica and carbonate was followed by late co-precipitation of quartz, pyrite, gold and galena. Discordant veins exhibit a single stage of vein-filling during which quartz and occasional pyrite were co-precipitated.

The sparse occurrence of s1 cleavage ± crenulation in pelite fragments indicates that pre-gold bedding-parallel vein filling was initiatied prior to cleavage development, and hence before f1. The absence of earlier phase microstructures precludes specific correlation of vein initiation with the regional deformations of Mountjoy (personal communication). However, vein textures indicative of multiple vein fillings, and the highly variable strain states observed in vein quartz suggest vein filling took place during prolonged or repeated deformation and continued during f1. The weakly deformed state of the paragenetically-late gold-bearing assemblage (quartz-pyrite-gold-galena) suggests gold deposition occurred late during or post-dating penetrative deformation (i.e. f1). The lack of veining or alteration along imbricate thrust planes suggests the Chatter Creek fault post-dates vein

filling and gold deposition.

As outlined earlier, discordant veins also post-date f1 and pre-date the Chatter Creek thrust. Thus a broad temporal relationship between discordant veins and gold emplacement is implied. The discordant veins however contain only minor gold. These observations, and the fact that the goldbearing assemblage is generally associated with pelite fragments in beddingparallel veins, suggests that fluid interaction with pelite may have triggered gold deposition. Studies by Seward (1984) indicate gold may travel as sulfur complexes in hydrothermal solutions. Disseminated, diagenetic sulfide in the McNaughton rock types may have provided sulfur for the complexing of gold by mineralizing fluids. Iron released from the pelites may have stimulated sulfidation reactions (Phillips et al., 1984; Tomkinson, 1988) which destabilized gold-sulfur complexes, resulting in the deposition of pyrite and native gold in the vicinity of pelite fragments. This hypothesis would account for the localization of gold in bedding-parallel structures, and for the lack of gold in discordant veins and quartzitic rock types. The discordant veins cut only quartzite and contain no pelite fragments. Quartzites would be intrinsically non-reactive and would not have triggered gold deposition. Thus, as well as providing dilatent zones for focussing fluid flow, pelites may have acted as chemical traps for the localization of gold mineralization on McGillivray Ridge.

Based upon its spatial association with bedding-parallel veins and auriferous nature, wallrock sulfidation likely accompanied gold-stage vein-filling. Sulfidation adjacent to some discordant veins also suggests a penecontemporaneous gold-stage-discordant vein-sulfidation relationship. The more widespread assemblage white mica \pm carbonate \pm disseminated pyrite is not uniquely associated with vein structures, and is considered the

product of regional anchimetamorphism. Notable is the mineralogical congruency between the vein (quartz-white mica ± pyrite ± carbonate), alteration (pyrite ± white mica ± carbonate), and anchimetamorphic (quartz-white mica ± pyrite ± carbonate) assemblages. This suggests subcontemporaneous vein-filling-alteration-metamorphism, and thus broad chemical and thermal equilibrium between the vein-filling fluids and the surrounding host rocks (Rose and Burt, 1979). As well, the relatively local derivation of the major vein-filling constituents is implied (Norris and Henley, 1976; Etheridge et al., 1984; Kerrich et al., 1978). With respect to regional metamorphism this mineralogical congruency, and the preservation of primary vein-filling textures and highly variable strain states in vein quartz, suggests late syn- to post-peak metamorphic vein filling. Thus vein filling may have been initiated during developent of the synmetamorphic Porcupine Creek Anticlinorium (Mountjoy and Dechesne, 1989, personal communication).

Deposit Classification

Within the Main Ranges, the gold lodes of the Athabasca Pass are unique. As well, geochemical data (trace elements, fluid inclusions, stable isotopes) are lacking. Hence, detailed comparison and quantitative process model classification would be premature. However, in terms of their setting and morphology, the Athabasca Pass lodes are considered to represent a siliciclastic-hosted analogue of the Turbidite-Hosted (Boyle, 1979 and Keppie et al., 1986; Hutchinson, 1987) class of gold deposits. Lithotectonic, metamorphic, structural, temporal, and vein textural-mineralogical analogies to turbidite-hosted deposits are clearly evident (Table 2-1).

However they are distinct in terms of host-rock composition, i.e. predominantly mature siliciclastics in the Athabasca Pass versus immature greywacke-argillite in turbidite-hosted deposits. Geochemical differences from the turbidite-hosted deposit model, such as the paucity of wallrock alteration and the mineralogical simplicity of the Athabasca Pass lodes, may be attributible to the non-reactive nature and the limited elemental spectrum of mature quartzite-dominated sequences compared with less mature turbidite-hosted rock types. Further comparison and classification must await geochemical data and/or the discovery of comparable siliciclastic-hosted lode systems.

Concluding Statement

The thick, clastic-dominated, Lower Paleozoic sedimentary sequences comprising the Main Ranges of the Canadian Rocky Mountains have previously been viewed as unfavourable environments for the search for gold-lodes (Matthews, 1944; Holland, 1944; Sinclair, et al., 1978). In light of the present study the Main Ranges may be considered as a distinct metallogenic domain containing sediment-hosted lode gold associations. Although presently unique, the Athabasca Pass lodes provide some first-order observations for lode exploration. With respect to regional deformation, gold mineralization appears to be late-stage, hence structures and veining associated with late-stage out of sequence thrust faulting (Mountjoy, 1989, personal communication) may prove significant. Locally, the important role of pelitic rock types in siliciclastic sequences such as the McNaughton Formation is emphasized. Such pelites provide ideal structural and geochemical contrasts for the localization of hydrothermal fluids and

gold in siliciclastic-dominated domains.

Table 2-1. Characteristics of the Athabasca Pass vs. typical turbidite-hosted * lode-gold deposits

| | Athabasca Pass | Turbidite-Hosted |
|----------------------------------|--|--|
| Geologic-tectonic setting: | Thick platformal clastic sequences in Rocky Mountain Thrust and Fold Belt | Thick flyschoid clastic sequences in fold- and thrust-belts ± transcurrent domains |
| Associated rock types: | Predominantly mature sedimentary rocks (pelitic quartite, quartzite, pelite, conglomerate) | Predominantly immature sedimentary rocks (grawwacke, amilitie-state crientatie) |
| Metamorpic grade: | Pumpellyite-lower greenschist | Dominantly prehnite-pumpellyite to middle greenschist |
| Host-rock age: | Lower Cambrian | Upper Proterozoic to Lower Paleozoic |
| Relative timing of veins: | Early-syn-tectonic, syn- to post peak-metamorphic | Early-syn to post-tectonic, syn- to post peak-metamorphic |
| Ore-hosting structures: | Discrete bedding-parallel ± discordant veins | Discrete bedding-parallel and discordant veins saddle reefs, fault structures |
| Alteration: | Absent to minor; silicitication, sulfidation carbonitization, sericitization | Absent to distinct; silicification, sulfidation carbonitization, sericitization, chloritization |
| Vein textures: | Open-space filling, host-rock fragments minor laminations | Open-space filling, host-rock fragments and larithations |
| Vein mineralogy: | Quartz, pyrite, white mica, galena, carbonate | Quartz, pyrite, arsenopyrite, carbonate±galena, sphalerite stibnite, scheelite, pyrrhotite, white mica, chlorite |
| Nature and distribution of gold: | Coarse-grained native gold, confined to veins, erratic grades | Coarse-grained native gold \pm sulfide associations, confined to veins \pm wallrock enrichment, erratic grades |
| Gold fineness(Au/Ag): | Very high (> 20:1) | High (average 9:1, range 3:1 to 20:1) |
| * Principle references: } | * Principle references: Keppie et al. (1986), Graves and Zentilli (1982), Hutchinson (1987). | inson (1987). |

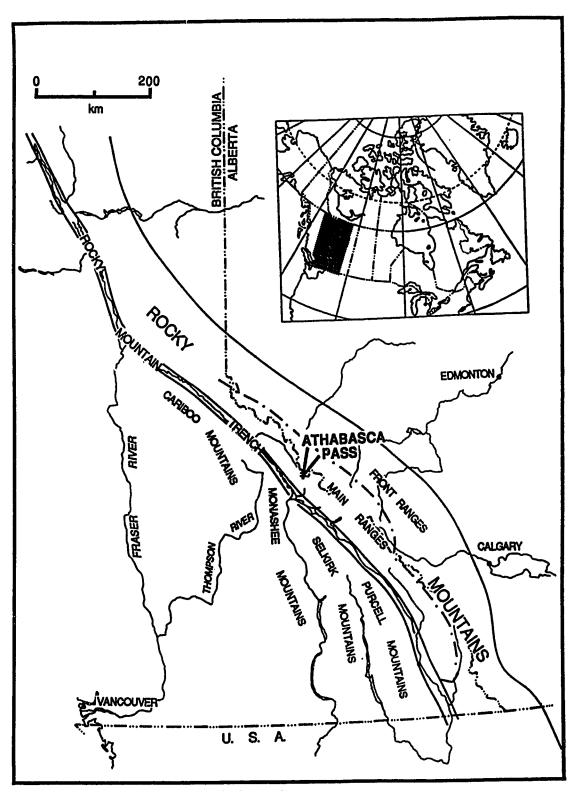


Figure 2-1. Location of the Athabasca Pass.

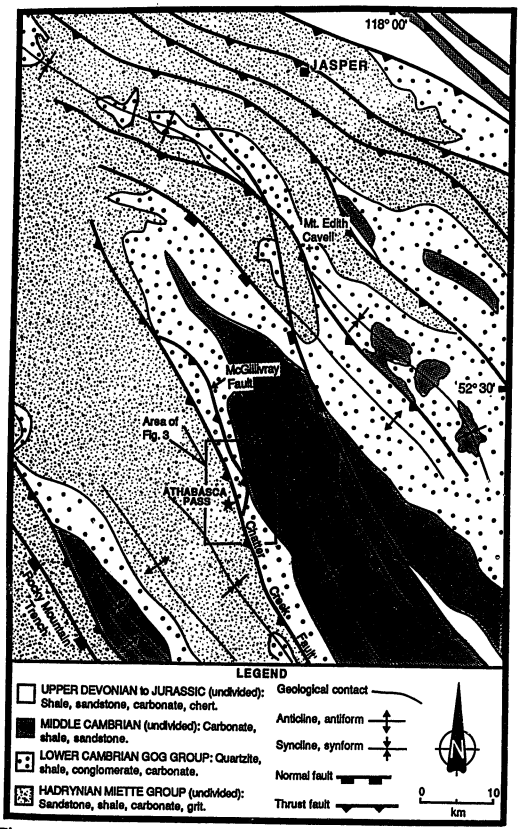


Figure 2-2. Regional geology surrounding Athabasca Pass. Modified after Wheeler et. al (1972).

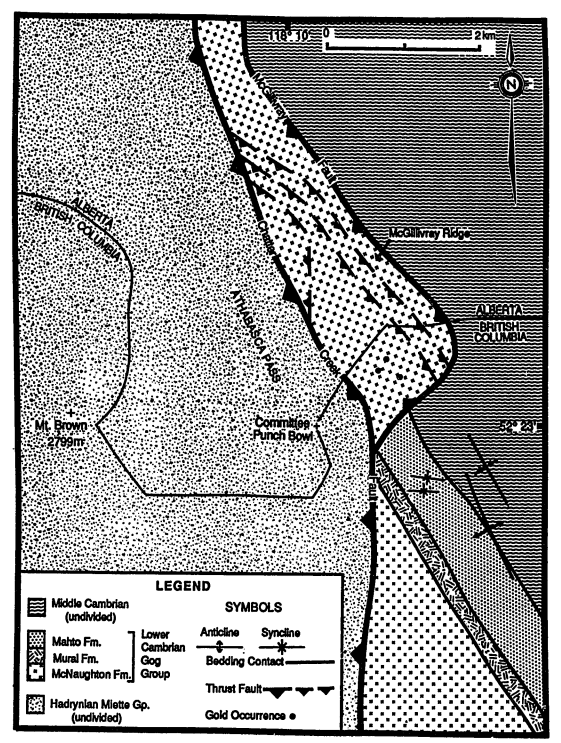


Figure 2-3. Local geology of the Athabasca Pass area (modified after Mountjoy and Price (in preparation.). Gold occurrences represent clusters of mineralized veins.

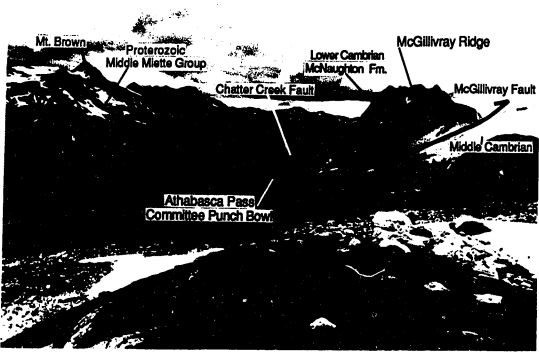


Figure 2-4. Photograph of the Athabasca Pass veiwed from the south. The Chatter Creek thrust does not outcrop in the Athabasca Pass. Its position is inferred from outcrops of Middle Miette grit to the west. Distance to McGillivray Ridge is approximately 3 km.

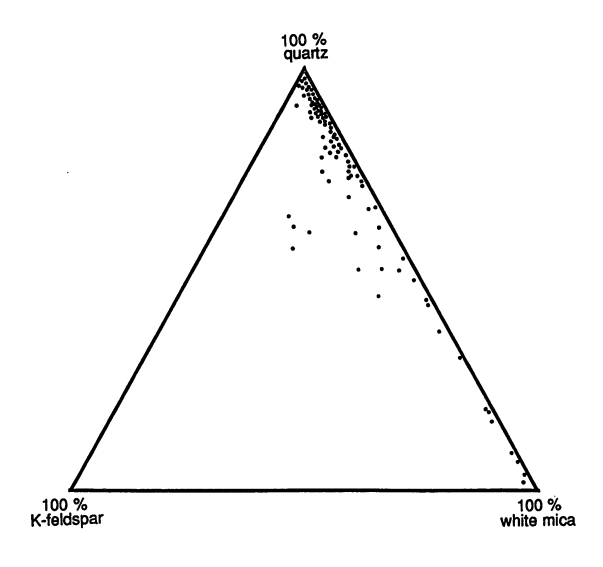


Figure 2-5. Modal composition of the McNaughton Formation rock types of McGillivray Ridge.

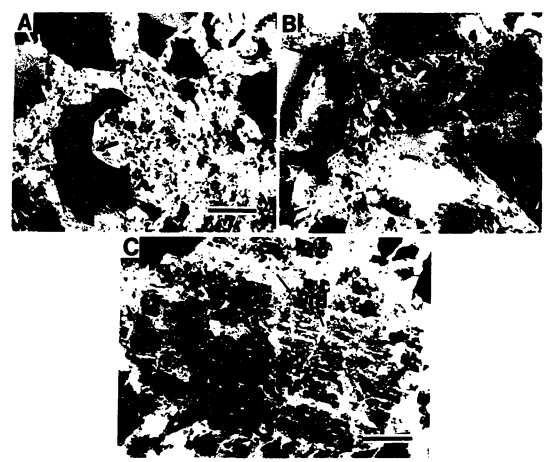


Figure 2-6. Typical deformational microstructures and alteration exhibited by the McNaughton rock types of McGillivray Ridge; transmitted light, crossed-polarized. A-B. Solution embayment of quartz grains (-) in typical pelitic quartzite. Scale bar=0.3 mm. C. Incipient replacement of a detrital potash feldspar grain by white mica (wm) and carbonate (ca); transmitted light, crossed-polars. Scale bar=0.3 mm.



Figure 2-7. View of the south end of McGillivray Ridge showing imbricate thrust geometry in the McNaughton Formation. The major thrusts are line-enhanced. Distance along McGillivray Fault is approximately 1.2 km. Location of Fig. 8 indicated by arrow.

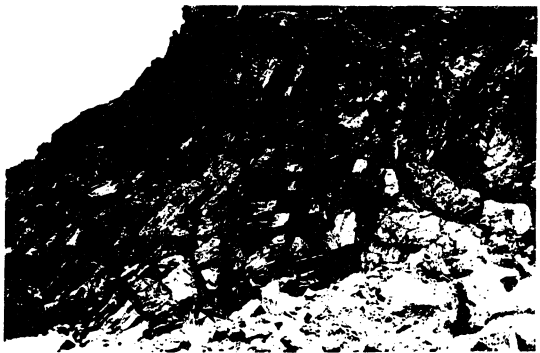


Figure 2-8. Folds in the McNaughton Formation of McGillivray Ridge at location shown in Fig. 7. View facing approximately north. Man in foreground is 1.4 m tall.

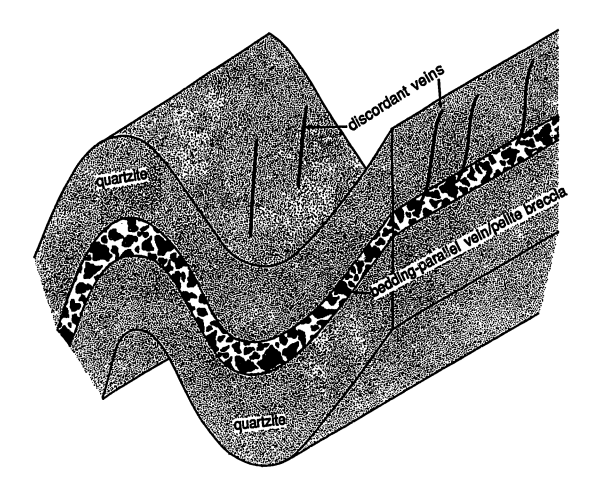
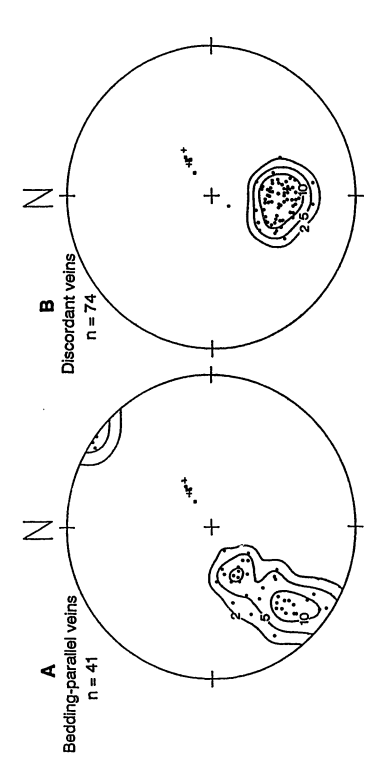


Figure 2-9. Schematic illustration of the general field relationships between bedding, folding, bedding-parallel veining, and discordant veining on McGillivray Ridge.



diagrams projecting poles to bedding-parallel (A) and discordant (B) vein structures, in relation to fold axial planes (•) and thrust faults (+) on Figure 2-10. Contoured lower hemisphere stereographic probability McGillivray Ridge. Contours at densities 2, 5, and 10.

| Pre-gold Stage | Gold Stage | Post-gold Stage |
|---|---------------|--------------------|
| quartz Type i | | |
| Type III | | |
| white mica | | |
| gold galena | | |
| carbonate | | • |
| brecciation | | |
| Time ——— Deformation ——— Folding ——— Veining | | usting——— |
| Bedding-parallel —————————————————————————————————— | | |

Figure 2-11. Mineralogy and paragenetic sequence for the auriferous bedding-parallel veins of McGillivray Ridge. Dashed lines represent possible minor quantities. Relative timing of vein formation and mineralization with respect to the deformations of Mountjoy (pers. comm.) is discussed in the text.

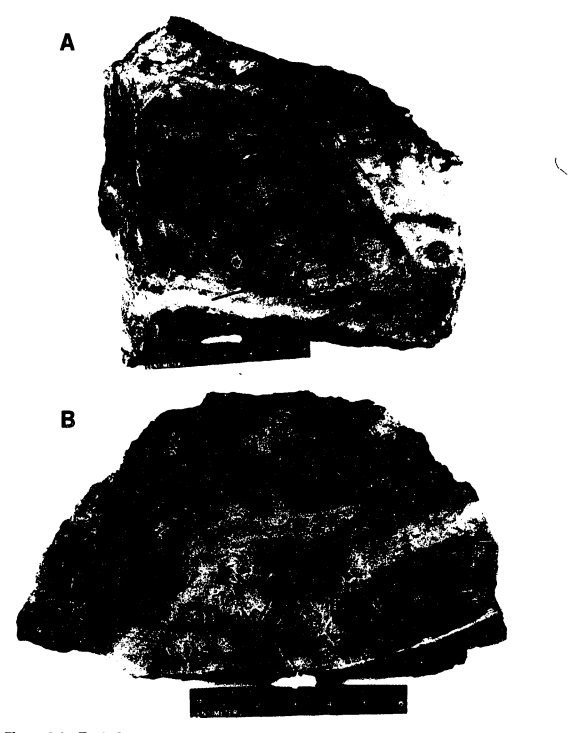


Figure 2-12. Typical textures exhibited by bedding-parallel veins from McGillivray Ridge. A. Massive Type I quartz (qz1) with pelite fragments (p) cut by milky Type III quartz (qz3) and a late discordant vein (d). B. Massive Type I quartz (qz1) containing pelite fragments (p), and a Type II quartz vein derived during progressive deformation of previously deposited Type I quartz.

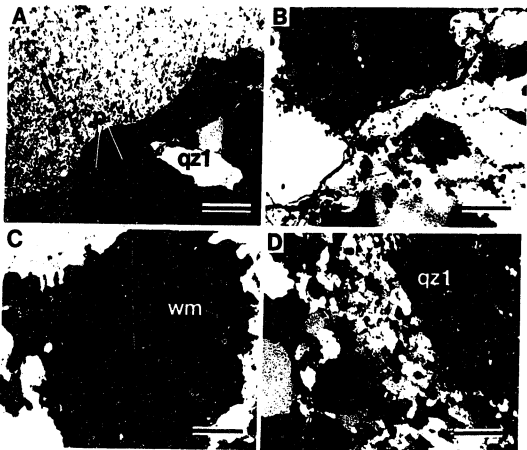


Figure 2-13. Microtextures exhibited by quartz in the bedding-parallel veins of McGillivray Ridge; transmitted light, crossed-polarized. Scale bar=0.3 mm in all photographs. A. Laminations of pelite (p) incorporated into Type 1 quartz (qz1) during incremental vein growth. B. Undulatory extinction, subgrain development, and incipient recrystallization in Type I quartz. Diagonal black fracture (lower left to upper right) was introduced during polishing. C. Co-precipitated inclusions of white mica (wm) lining a growth zone in Type I quartz. D. Relatively unstrained Type II quartz (qz2) adjacent to earlier, deformed Type I quartz (qz1).

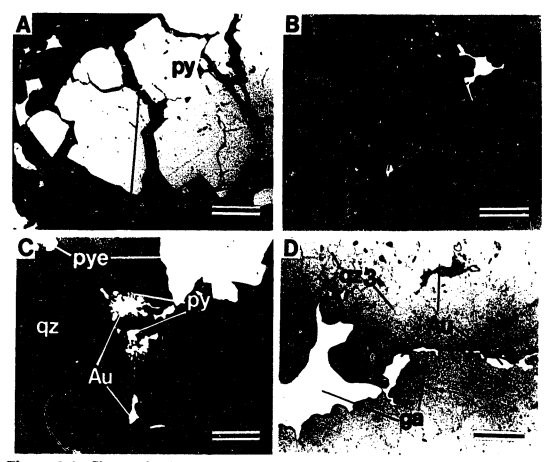


Figure 2-14. Textural relationships among the minerals from bedding-parallel veins of McGillivray Ridge; reflected plane-polarized light. Scale bar=0.5 mm in all photographs. A. Quartz (qz) and native gold (Au) filling fractures in and around pyrite (py). B. Native gold (Au) abutting pyritized pelite (p) fragments in Type III quartz. C. Subcontemporaneous native gold (Au) and pyrite (py) filling fractures in quartz (qz) with paragenetically earlier pyrite (pye). D. Native gold (Au), galena (ga) and Type III quartz (qz3) filling late fractures in Type I quartz.

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CHAPTER 3

Siliciclastic-Hosted Lode-Gold Mineralization, Athabasca Pass, Central Canadian Rocky Mountains:

A Trace Element and Fluid Inclusion Study.

Introduction

The recent discovery of gold-bearing quartz veins in Lower Cambrian siliciclastic strata, which comprise much of the Main Ranges of the central Canadian Rocky Mountains, attests to the metallogenic potential of this vast, yet sparsely prospected, lithotectonic domain. Information on lodegold deposits within this domain is lacking, the apparent first comprehensive documentation of gold-lodes in this region being that given in Chapter 1. In that paper the structural, mineralogical and paragenetic relationships of gold-quartz veins contained within the McNaughton Formation of the Athabasca Pass were established, and the hypothesis was forwarded that the Athabasca Pass lodes represented a siliciclastic-hosted analogue of the Turbidite-Hosted (Keppie *et al.*, 1986; Hutchinson, 1987) class of gold deposits.

In order to geochemically characterize the gold-lodes of the Athabasca Pass, and to place constraints upon the physico-chemical environments of vein formation, a trace-element geochemical and fluid inclusion study was undertaken.

Regional Setting

The Athabasca Pass is situated on the Continental Divide 60 km south of Jasper, Alberta, at the boundary between the eastern Main Ranges

and the western Main Ranges (Price and Mountjoy, 1970) of the central Canadian Rocky Mountain belt (Fig.3-1). The boundary in this region is marked by a southwest-dipping thrust fault, the Chatter Creek fault (Wheeler, 1963), which extends from the headwaters of the Fraser River, southeasterly through the Athabasca Pass, into the region northeast of and price, 1989; Mountjoy and Price, Golden, British Columbia in prep.). In the Atnabascon rea, this fault has transported a thick sequence of grits, graywards pelites, and carbonates of the Upper Proterozoic Miette Group over quartzites, quartz-pebble conglomerates, pelites, and carbonates of the Lower Cambrian Gog Group. Within the hanging wall of the Chatter Creek thrust sheet, regional metamorphic grade increases southwestward from greenschist grade to kyanite-staurolitebearing assemblages of amphibolite grade (Craw, 1978; Klien and Mountjoy, 1988). Rocks of the immediate footwall have been regionally metamorphosed to sub-greenschist and lower greenschist grades, decreasing to burial metamorphic facies southeastwards (Read, 1988). The occurrence of igneous rocks is rare throughout the Main Ranges.

Deformation in the Main Ranges evolved through the accretion of allochthonous terranes to the western margin of North America beginning in the Late Jurassic, which resulted in the northeastward translation and thrust-stacking of thick sequences of parautochthonous miogeoclinal sedimentary rocks derived fron the North American craton to the east, onto the flank of the craton (Price and Mountjoy, 1970; Price et al., 1985). The geologic structure of the region is dominated by complex folding patterns and by widely spaced thrust faults. Penetrative deformation is pronounced in the western Main Ranges, and less pervasive to absent in the east (Price

and Mountjoy, 1970).

Information on lode-gold deposits in the Main Ranges, and in the Canadian Rocky Mountains in general, is lacking. The mineral occurrences and metallogeny of this region have been reviewed by Hedley (1954), Little et al. (1976), and Sinclair et al. (1978). Principal deposit-types occur to the south of the Athabasca Pass area, and include stratiform lead-zinc (±silver) replacement bodies in Middle Cambrian carbonates, and generally small, quartz-carbonate vein systems containing a variety of base-metal sulfide assemblages (dominantly galena-sphalerite-pyrite-chalcopyrite). Gold production has not been reported from these occurrences. Brief mention of placer gold occurrences in the Main Ranges was made by Boyle (1979), and by Mathews (1944).

Local Geology

The general geology of the Athabasca Pass area is shown in Figure 3-1. To date, gold mineralization has been found only in the lowermost formation of the Gog Group, the McNaughton. The McNaughton Formation is composed of a series of well-stratified, mature, quartz-dominated clastic rock types (pelitic quartzite, quartzite, matrix-supported quartz-pebble conglomerate) randomly parted by discrete, discontinuous interbeds of quartz-sericite pelite. The sequence has been subjected to archimetamorphic (ca. pumpellyite facies) conditions during regional metamorphism. Penetrative planar fabric is evident in the quartz-dominated rock types as phyllosilicate contents reach approximately 30 percent. The pelites contain a variably developed, locally crenulated cleavage (s1). Evidence of igneous activity is entirely lacking.

The McNaughton Formation in the Athabasca Pass is confined to an 800 m-thick folded and faulted section known as McGillivray Ridge. Structurally, McGillivray Ridge consists of a series of northeast-verging thrust faults which truncate northeast-verging, upright to overturned, mesoscopic folds (f1) in individual imbricated slices of McNaughton quartzite. The structural geometry of McGillivray Ridge is that of a leading imbricate fan and is attributed to late-stage northeast-directed compressional tectonism.

Mineralized veins of the Athabasca Pass

Two vein-types, outcropping on McGillivray Ridge, have been identified: a folded, early syn-tectonic, syn-to post-metamorphic bedding-parallel variety recording prolonged or multiple stages of northeast-directed compression, and a post-folding discordant variety generated during minor sinistral shear associated with incipient, late-stage regional thrusting. Vein localization and geometry exhibit lithostructural control with bedding-parallel veins being confined to the less-competent pelitic rocks, and discordant veins being confined to the competent quartzitic units. High concentrations of gold are confined to bedding-parallel veins, with only a few discordant veins being slightly enriched in gold (up to ca. 500 ppb).

The mineralogy of bedding-parallel veins consists of quartz and brecciated host-pelite with minor quantities of white mica, pyrite, Fecarbonate, galena and native gold. Two broad paragenetic stages have been identified, namely: an early protracted pre-gold stage depositing quartz and white mica \pm minor Fe-carbonate and pyrite, and a late, volumetrically minor gold-bearing and post-gold stage during which quartz, pyrite, native gold and galena \pm minor white mica and Fe-carbonate were deposited. Gold

is largely associated with pelite fragments and paragenetically subcontemporaneous sulfides. Discordant veining was approximately coeval with gold emplacement, however these veins contain only quartz and occasional pyrite. It was postulated that pelites played an important chemical role in triggering gold deposition uniquely within bedding-parallel veins.

Trace Element Lithogeochemistry

In order to characterize the trace-element geochemistry of the McNaughton Formation in the Athabasca Pass, 26 samples, including (a) unveined pelitic quartzite and conglomerate, (b) unveined pelite, (c) highly anomalous (arbitrarily > ca. 15 ppm Au) gold-bearing vein, (d) slightly anomalous (arbitrarily < ca. 1 ppm Au) gold-bearing vein, (e) brecciated pelite ± minor vein material, and (f) unmineralized discordant quartz vein were analyzed by inductively coupled plasma emission spectrometry (ICPES), for 31 clements. Gold and silver contents were determined by atomic absorption spectrometry (AAS) and fire assay. All analyses were performed by a commercial laboratory (Eco-Tech Laboratories, Kamloops, B.C.). From the multi-element analyses, the element suite: arsenic-leadzinc-copper-barium-iron was selected for presentation here, as this suite contains elements commonly found in anomalous concentrations in association with lode-gold mineralization (Boyle, 1979; Nesbitt, 1988). Other elements commonly concentrated in lode gold deposits, including antimony, tungsten, boron, tellurium and thallium were found to be below their limit of detection (DL) in both vein and unmineralized host-rock material in the present study (i.e. 5 ppm, 10 ppm, 2 ppm, 10 ppm respectively), and thus are not included here. Major element analyses were not carried out.

Results

Figure 3-2 portrays data from ICPES and AAS analyses (also see Appendicies 1 and 2). For comparative purposes, histograms of average concentrations of the same element-suite for similar unmineralized rock types (quartzite, argillite/slate, conglomerate, greywacke) drawn utilizing data compiled from the literature are also included. Comparing the traceelement concentrations of the unmineralized McNaughton rock types of Athabasca Pass (Figs. 3-2A, 3-2B) with estimates of average element contents in comparable rock types (Figs. 3-2G, 3-2H) it is evident that although consistently somewhat lower than average in trace element contents, the Athabasca Pass rock types are not significantly different from the average comparable rock-types from other localities. This apparent depletion is perhaps attributable to the fact that the McNaughton Formation in the Athabasca Pass is a very mature clastic sequence composed dominantly (ca. 97 %) of quartz-rich (averaging >90 % quartz) rock types which only rarely contain significant quantities of varietal minerals or of feldspar. The remaining few percent of strata comprising the McNaughton Formation at this locality are pelites composed of fine-grained greenish-white mica with variable proportions of detrital quartz. Thus, in view of this mature, almost monomineralic (hence limited elemental) spectrum relatively low abundances of trace elements are understandable. Also, the McNaughton rock types have been subjected to anchimetamorphic conditions, during which a variable degree of recrystallization of allogenic quartz and clay

constituents has taken place. Although unmineralized samples were collected over 200 m from the nearest gold-bearing veins and the samples appeared visibly unaltered, some trace element mobilization/depletion during recrystallization cannot be precluded. Finally, some discrepancy may have been introduced by the inclusion of data from the literature for samples which vary in mineralogical maturity and hence exhibit different trace-element chemistry from the McNaughton rock types. The effect of such variation is evidenced by the higher trace element contents exhibited by less mature sedimentary rocks such as greywacke, as shown in Figure 3-21.

Athabasca Pass rock types

Focusing upon the Athabasca Pass data, the following observations are noteworthy. In general, vein and vein-associated rock types (Figs. 3-2C, 3-2D, 3-2E, 3-2F) show a variable, yet distinct, enrichment of most trace elements in the suite with respect to the unmineralized host rocks (Figs. 3-2A, 3-2B). Gold shows the greatest enrichment in all vein samples. Silver, zinc and copper show negligible to minor (near level-of-detection, therefore possibly only apparent) enrichments, while lead, arsenic and barium show enrichment between approximately two and ten times their concentrations in the host rocks. The behaviour of iron is somewhat more complex with the mineralized rock types showing enrichment over the quartzose host rocks (Fig. 3-2A) and depletion with respect to the unmineralized pelitic rocks (Fig. 3-2B). The vein samples were selected to contain as few pelite fragments as possible, thus the iron in these samples is contained dominantly in the form of pyrite (and secondary Fe-oxides) whilst that in

the pelitic rocks would be structurally contained within the phyllosilicate minerals of the rock.

In order to identify and portray sympathetic variations in traceelement concentrations with respect to gold, data from the vein samples used to generate the averages shown in histograms 3-2C (highly anomalous bedding-parallel veins) and 3-2D (moderately anomalous bedding-parallel veins) were plotted in the form of a concentration versus element line graph (Fig. 3-3). Eight samples representing four orders of magnitude of gold concentration (i.e. 0.4 ppm to 573 ppm Au) were selected for comparison. From these data it is evident that silver, zinc and copper are of essentially constant concentration in the vein structures, regardless of gold content. Lead, arsenic and barium show relatively good (albeit non-proportional) positive correlation with gold content. Positive lead correlation is in accord with the observation that galena is paragenetically sub-contemporaneous with native gold, its presence often being indicative of higher gold grades. Arsenic is likely contained in isomorphous solution within pyrite which is also broadly coeval with gold (instrumental neutron activation analyses of vein pyrite separates contained up to 1800 ppm arsenic (Appendix 2)), No discrete arsenide phases have been identified in the veins. Barium is likely present substituting for potassium in the white micas (± pelite fragments) contained within the vein structures. Hydrothermal white mica deposition is again demonstrably coeval with gold emplacement.

Points A and B, shown on Figure 3-3, denote the silver content (8.8 ppm) of sample 524 and the zinc content (147 ppm) of sample 599 respectively. These points represent isolated, anomalously high (1 to 2 orders of magnitude) concentrations of silver and zinc with no apparent correlations within the general data trend and are interpreted to suggest the

presence of minor mineral phases (Ag and Zn sulfides?) which have not been detected during hand specimen and thin section petrography.

Of interest is the overall low concentration of silver, and the apparent lack of gold-silver correlation within the mineralized veins of the Athabasca Pass. Numerous works documenting gold fineness and Au/Ag ratios across essentially the entire spectrum of hypogene gold-bearing deposit types (see Boyle, 1979 for a literature review of gold fineness and Au/Ag ratios) have shown that silver is almost ubiquitously alloyed with native gold, and that silver-bearing sulfides and/or sulfosalts are common in the ore assemblages of many gold deposits. With respect to the general turbidite-hosted class of gold deposits, both gold fineness and Au/Ag ratios are, in general, relatively high (averaging approximately 850-900 fine and 4:1 to 9:1 respectively). Energy dispersive analyses of native gold grains from the Athabasca Pass lodes failed to reveal the presence of silver (or of any other trace metal) in quantities detectable by this method (DL call 1-2 %). In order to further investigate gold-silver relationships in the Athabasca Pass lodes, 30 samples of gold-bearing vein material were analysed for gold and silver by AAS. The results of this survey, portrayed in simple scatter-plot form in Figure 3-4 emphasize the sparse concentration of silver and the lack of gold-silver correlation. These data support the homogeneity and high fineness of the native gold implied by the energy dispersive analyses. Also, the few isolated concentrations of silver indicated by AAS further suggest the occasional presence of a minor silver-bearing phase in the Athabasca Pass lodes.

Discussion of trace element data

An integrated discussion of these data, and data derived from the fluid inclusion study will be undertaken following presentation of the latter. At this point some conclusions regarding the trace element distributions will be emphasized.

Of importance is the observation that, although the mineralized rock types of the McNaughton Formation are enriched in trace-elements with respect to their host rocks, they are, when compared with Turbidite-Hosted, and lode-gold systems in general, conspicuously non-enriched with respect to all trace elements, excepting gold (e.g. Boyle, 1979; Foster et al., 1986; Steed and Morris, 1986). This characteristic is observed directly in the relatively simple mineralogy exhibited by the mineralized Athabasca Pass structures, and may be attributable to 1) the paucity of trace elements in the source regions and channel-ways of the mineralizing fluids leading to only sparse concentrations within depositional structures, or 2) the inability of the fluids responsible for vein-filling to mobilize, transport, or deposit trace elements and hence create anomalous concentrations.

With respect to gold, the arithmetic average of 22 ppb Au for the unmineralized quartzitic rock types of the Athabasca Pass (Fig. 3-2A) falls central within the range given by Boyle (1979) and Crocket (1974) for average quartz sandstone/conglomerate (31 ppb) (Fig. 3-2G) and greywacke (3 to 13 ppb, Fig. 3-2H). A more extensive survey of unmineralized rock types in the Athabasca Pass involving 60 analyses of McNaughton quartzite indicates gold values range between limit of detection (5ppb) and 115 ppb, with 8 samples containing over 30 ppb Au (Appendix 3). Thus, although the McNaughton rock types contain, in general, "average" quantities of gold, it

is evident that they may locally contain anomalously high gold concentrations. The nature of this gold is open to speculation (Boyle, 1979). Samples containing sulfide in any form were avoided during this survey, precluding potential epigenetic gold concentrations in this form. If one considers the McNaughton Formation to represent a thick, mature, quartzitic sequence deposited under prolonged fluvial braidplain to shallow marine conditions (Young, 1979; Palonen, 1976; Woberg, 1986), the gold contained within these rocks would potentially be of a detrital nature. Greater concentrations within the host rocks may represent low-grade, syngenetic placer accumulations, derived during the erosion of Precambrian North American basement rocks to the east (Burwash, 1951; Mountjoy and Aiken, 1963). In comparison with studies investigating local rock types as the potential source for lode-contained gold in sedimentary domains (e.g. Glasson and Keays, 1976; Graves and Zentilli, 1982; Henley et al., 1976; Crocket et al., 1982), the unmineralized McNaughton rock types of the Athabasca Pass apparently contain ample gold which could have been concentrated by mobilization into lode-type structures. The mobilization of detrital gold, likely of enhanced fineness due to the preferential leaching of silver and other metals in the placer regime (Desborough, 1970; Boyle, 1979), could account for the very high fineness of gold now contained within the Athabasca Pass lodes.

Fluid Inclusion Study

To investigate the fluid compositional and P-T regime, and to gain insight into the processes involved in gold-quartz mineralization in the Athabasca Pass, a detailed study of fluid inclusions in mineralized vein

quartz was undertaken. A brief outline of the vein-quartz paragenetic relationships is summarized here in order to clarify the paragenetic relationships between gold-stage mineralization and the fluid inclusions examined in this study (see chapter 1 for a complete discussion of vein paragenesis).

On the basis of textural, optical and paragenetic characteristics, observed in hand specimen and in thin sections, three varieties of quartz, Types I, II and III have been defined within the gold-bearing veins of the Athabasca Pass. Type I quartz, comprising 90 to 95 percent of most veins, represents a prolonged, paragenetically early, pre-gold stage of multiple vein infillings, and has been subject to penetrative deformation during fl folding. Deformational characteristics including undulatory extinction, deformation lamellae, and sub-grain development have been overprinted by episodes of late, variably healed, brittle fracturing. The gold-stage assemblage native gold ± pyrite ± galena ± minor white mica and Fecarbonate is commonly seen where late fractures intersect fragments of pelitic host rock contained within the veins. Type II quartz occurs as discrete dilational fracture-fillings in Type I quartz. It is a minor vein filling phase (≤ 2 %) considered to have been derived through pressure solution of local, Type I quartz during progressive deformation. Type III quartz is seen filling diffusely bound fractures and voids of irregular size and shape in Types I and II quartz. It is paragenetically late and is petrographically distinguishable by its relatively unstrained appearance (minor undulatory extinction and brittle fracturing). Emplacement of the gold-stage assemblage is subcontemporaneous with this latter quartz type.

Methodology

Analyses were made on fluid inclusions in 12 samples containing Types I and III quartz from high-grade gold-bearing veins outcropping on McGillivray Ridge. Thirty doubly-polished quartz-plates were prepared from these samples. Microthermometric measurements were performed using a U.S. Geological Survey gas-flow heating-freezing stage which was calibrated using liquid nitrogen, the triple points of pure CO₂ and H₂O, and the Merck standard 9800 (melting point 200°C). Analytical reproducibilities were ± 0.2°C for freezing and ± 4°C for heating to temperatures of approximately 300°C.

Classification of fluid inclusions

Due to the pervasively deformed nature of much of the quartz vein material, and to the multiple generations of fracturing, vein filling, and fracture healing (precluding establishment of fluid inclusion paragenesis), fluid inclusions were classified primarily on their mode of occurrence in relation to the gold ± galena ± pyrite assemblage, and subsequently on their appearance upon cooling to 0-10°C. Thus, two general types of fluid inclusions, Type A and Type B have been defined. Type A inclusions were studied in most inclusion plates but bore no physically traceable link to gold mineralization. A full range of physical and microthermometric data were recorded for Type A inclusions, but these data were utilized only as a general survey of the various fluid inclusion types sensu lato observed in the Athabasca Pass gold-quartz lodes. Little interpretive weight is placed on Type A data, due to their uncertain, likely varied origin. Type A inclusions are considered secondary or pseudosecondary (Roedder, 1984) with respect to

gold stage mineralization.

Fortunately, abundant coarse-grained visible gold is observed in the Athabasca Pass lodes. Thus, 15 inclusion plates representing 9 different samples of vein material containing the assemblage native gold ± galena ± pyrite, hosted by Types I and III quartz were prepared. Clearly visible groups of inclusions were located in relatively undeformed quartz surrounding and filling embayments in native gold and galena. Type B inclusions are defined by this occurrence (Fig. 3-5). Most Type B inclusions analysed were less than 100 µm distant from native gold. Those fluid inclusions exhibiting any evidence of necking, leakage or (rarely) erratic liquid to vapor-phase ratios were not used. Type B inclusions may be classified as secondary and/or pseudosecondary with respect to Type I quartz, but relative to Type III quartz and the gold ± galena ± pyrite assemblage, they were considered to be primary and hence representative of the fluids responsible for gold mineralization in the Athabasca Pass.

Description of Fluid Inclusions

Type A: This fluid inclusion type occurs in both Types I and III quartz, distributed as isolated inclusions, random inclusion clusters, and along late and partially healed fractures. Type A inclusions generally contain two phases at room temperature, H2O-rich liquid and vapor, although 3 phases (H2O-rich liquid, carbonaceous fluid and vapor) are occasionally observed. Disregarding inclusions which have obviously undergone post-entrapment modification (necking, leakage etc.), Type A inclusions are generally subequant to ellipsoidal in shape (although irregular and negative crystal shapes were also recorded), and range in

greatest dimension from 4 to 45 μ m, averaging 10.5 μ m. Phase ratios for Type A inclusions vary between approximately 5 and 30, with most containing between 10 and 20 volume percent vapor. Vapor-rich Type A inclusions are rare. Those observed exhibit no phase changes between -140° and 300°C and are likely "empty", the result of natural decrepitation or leakage (Roedder, 1984). Daughter phases were not observed in any Type A fluid inclusions.

Type B: This fluid inclusion type is also found in both Types I and III quartz, but based upon petrographically established relationships is considered primary with respect to the gold assemblage. Two varieties of Type B fluid inclusions were defined based upon the number of phases observed upon cooling to 0-10°C.

Type B1. Two phase H₂O-rich liquid + vapor ± (not visible) CO₂-bearing fluid inclusions. This inclusion type ranges in size from 4 to 24 μm, averaging approximately 10 μm in greatest dimension. Generally, subangular to sub-rounded and sub-equant to prismatic in shape, Type B1 inclusions occur as isolated individual fluid inclusions or randomly arranged inclusion clusters. Freezing experiments indicated Type B1 inclusions contain H₂O-rich brine. Clathrate formation (melting at less than 10°C) indicated the presence of small quantities of CO₂ in about one half of all Type B1 inclusions, but a discrete liquid CO₂ phase was not observed. Such inclusions may contain up to 3 mole percent CO₂ at pressures between ca. 10.4 and 45 bars (Hedenquist and Henley, 1985).

Type B2. Three phase H₂O-rich liquid + carbonaceous liquid + vapor-bearing fluid inclusions. Based upon habit and mode of occurrence, at room temperature Type B2 fluid inclusions are indistinguishable from

Type B1. Slow cooling, usually below about 15°C nucleated a third, CO₂-rich vapor phase, defining Type B2 inclusions. Phase changes recorded during freezing experiments (including heating up to the homogenization temperature of CO₂) indicated the presence of variable (generally small) but ubiquitous quantities of additional volatiles and dissolved salts. CO₂ in all Type B2 inclusions homogenized exclusively to the liquid state.

As noted, based upon appearance at room temperature, Type B1 and B2 fluid inclusions are indistinguishable. Both types may occur in a single inclusion cluster, but generally one type is dominant over the other (ratios generally 2 10:1). Phase ratios of liquid to vapor in Type B1 inclusions and H₂O-rich liquid to CO₂-rich liquid in Type B2 inclusions are similar varying only slightly between different samples and being uniform with individual samples and inclusion clusters. Visual estimates of phase volumes obtained using a graduated ocular lens, and diagrams illustrating degree-of-fill and volume percentages (Roedder, 1984; Shepherd et al., 1985) for various fluid inclusion shapes, range from 9 to 20 and average 12 to 18 volume percent vapor (Type B1) or liquid CO2 (Type B2), Such visual estimates are subject to potentially large errors due to the uncertainties and assumptions made in estimating the depth dimension of fluid inclusions (Bodner, 1983; Roedder, 1984; Shepherd et al., 1985). Neither daughter crystals nor vapor-rich inclusions were observed during Type B inclusion study.

Microthermometric analysis of fluid inclusions

In order to minimize the effects of phase metastability during freezing-heating runs, all phase changes were observed during the heating cycle. Data recorded through phase changes included; 1) the temperature of

final CO₂ melting (Tm_{CO2}, observed in Type B2 inclusions), 2) the temperature of final ice melting (Tm_{ice}, observed in Types A, B1 and B2 inclusions), 3) the temperature of final clathrate melting (Tm_{Clath}, observed in Types A, B1 and B2 inclusions), 4) the temperature of CO₂ homogenization (Th_{CO2}, observed in three phase Type A inclusions, and Type B2 inclusions), 5) the temperature of decrepitation (T_{decrep}, observed dominantly in Type B2 and occasionally Type B1 and Type A inclusions) and, 6) the temperature of total phase homogenization (Th_{total}, observed where possible in all inclusion types).

Estimates of aqueous fluid salinities (expressed as weight % NaCl equivalent) were derived from Tm_{ice} and Tm_{clath} . data utilizing phase relations and techniques described by Potter *et al.* (1978; H₂O-NaCl equivalent system), and by Bozzo *et al.* (1975) and Collins (1979; H₂O-CO₂-NaCl system). Additional constraints were placed upon bulk fluid composition and phase densities utilizing Tm_{CO2} , Th_{CO2} and Th_{total} data in conjunction with phase relations in the H₂O-NaCl system (summarized by Crawford, 1981 and Roedder, 1984), and in the H₂O + CO₂ \pm NaCl \pm CH₄ systems (summarized by Hollister and Burruss, 1976; Eurruss, 1981; Pichavant *et al.*, 1982; Roedder, 1984; and Seitz *et al.*, 1987). Microthermometrically-derived analytical results for all Type A fluid inclusions studied are listed in Appendix 4. Data for all Type B inclusions are graphically portrayed in Figures 6 and 7.

Type A. Due to the unconstrained origin of Type A inclusions, data from these types will not be reviewed in detail here other than to note the overall similarity between inclusion Types A and B (see Appendix 4 for comparison with Type B inclusions). The more varied phase ratios,

salinities and temperatures of total homogenization observed in Type A inclusions likely reflect a potentially more varied origin or some post-trapping modification, but the general congruence between many Type A and Type B1 inclusion data suggests many Type A inclusions may share an origin similar to that of the Type B inclusions.

Type B1. This inclusion type is essentially defined by the system H2O + NaCl ± CO2. NaCl-equivalent salinities calculated from aqueous fluid freezing-point depression (Tmice) data for those inclusions showing no evidence of the presence of CO₂ (i.e. clathrate formation) range from 7 to 10, averaging 8.5 wt. percent NaCl equivalent (Fig. 3-6A). Hedenquist and Henley (1985) note that the presence of small quantities of dissolved CO₂ (< ca. 1 mole %) in aqueous solutions may go undetected, enhancing freezing point depression, hence the above salinity estimates are considered as maxima. Where clathrate formation was detected, coexisting aqueous solution, liquid CO2 and vapor phase were not present upon clathrate melting, thus accurate salinity estimates based upon clathrate melting are precluded. Such estimates are erroneously high (Collins, 1979). Nevertheless, salinities were calculated from Type B1 Tmclath. data, providing approximate maxima, and hence constraining the salinity of clathrate-forming Type B1 inclusions to values somewhat less than those calculated (i.e. ca. 4 to 10, averaging 5 wt. % NaCl equivalent). This implies that Type B1 inclusions containing enough CO2 to form clathrate (between ca. 1 and 3 mole %; Hedenquist and Henley, 1985) are distinctly less saline than those which do not form clathrate (Fig.3-6A).

Th_{total} data for Type B1 inclusions vary from approximately 190° to 290°C with most data clustering in the 210 to 225°C range (Fig. 3-6B). Based

upon an average salinity of 8.5 wt. percent NaCl equivalent and negligible CO₂ content, this translates to fluid densities of approximately 0.9 to 0.92 g cm⁻³ (Haas, 1976). Significant differences in Th_{total} between clathrate-forming and non-clathrate forming Type B1 inclusions were not observed.

Type B2. This fluid inclusion type is compositionally approximated by the system H₂O + carbonaceous fluid + NaCl. Consistent depression of the CO₂ freezing point (Tm_{CO2}) below the pure CO₂ invariant point (-56.6°C) indicates the presence of small quantities of additional volatile components in Type B2 inclusions. Tm_{CO2} measurements range from approximately -60° to -57°C clustering about -58°C (Fig. 3-7A), and are interpreted to indicate Tm_{CO2} depression due to the presence of minor quantities of CH₄ (Hollister and Burruss, 1976; Burruss, 1981; Roedder, 1984). Other gases including higher hydrocarbons, N₂, SO₂, and/or H₂S may also be present (Roedder, 1984). Due to the preferential partitioning of CH₄ into the vapor phase, Tm_{CO2} does not uniquely define the bulk carbonaceous phase CH₄ content (Burruss, 1981); such data must be used in conjunction with Th_{CO2}.

ThCO2 data for Type B2 inclusions are summarized in Figure 3-7B. Carbonaceous phase liquid-vapor homogenization temperatures range from approximately -7° to +25°C, clustering in the +23°C region. Carbonaceous liquid-vapor homogenization occurs exclusively to the liquid phase. Applying the graphical methods of Heyen et. al. (1982) for estimation of CH₄ content from combined TmCO2 and ThCO2 data, the carbonaceous phase CH₄ content of Type B2 inclusions ranges from 2 to 16 mole percent, averaging ca. 8 mole percent CH₄. Combined with typical values for TmCO2 and ThCO2 of -58°C and +18°C respectively, this translates to a

typical bulk carbonaceous fluid phase density of approximately 0.7 g cm⁻³. It is noted, however, that Tm_{CO2} vs. Th_{CO2} measurements in Type B2 fluid inclusions do not vary in a linear fashion, and samples exhibiting similar Tm_{CO2} do not necessarily homogenize at the same Th_{CO2}. Such inconsistencies might be attributed to variations in fluid density and/or variations in fluid composition (Seitz et. al., 1987).

Aqueous phase salinity estimates for Type B2 inclusions (Fig. 3-7C) were derived using the relationship between final clathrate melting (Tmclath.) and salinity given by Bozzo et. al. (1975). As discussed by Hollister and Burruss (1976), and Collins (1979) the presence of additional volatile compounds such as CH₄ shifts Tm_{clath}, to higher temperatures, due to the formation of mixed CO₂-CH₄ hydrates, hence counteracting the effect of Tmclath, depression by NaCl. However, the effect of CH4 on clathrate decomposition temperatures is considered to be minimal in this study. Clathrate decomposition in Type B2 inclusions is tightly clustered between 7.5° and 8°C, rarely extending above 8.5°C, and shows no sympathetic increase with increasing CH4. As well, recent study of CO2-CH4 hydrates by Seitz et. al., (1987) implies negligible partitioning of CH4 into clathrate at low CH4 abundances. Furthermore, Tmice data for Type B2 inclusions imply residual fluid (hence maximum) salinities of between 6 and 10 wt. percent NaCl equivalent, thus salinities averaging between 3.5 and 6 wt. percent NaCl equivalent derived from Type B2 Tmclath. data appear reasonable.

Figure 3-7D portrays T_{decrep}, and Th_{total} (exclusively to liquid) data derived from Type B2 fluid inclusions. Th_{total} data were obtained for less than ca. one third of Type B2 homogenizations attempted, with temperatures acquired ranging from 210° to 315°C, and most data falling

between 225° and 245°C. More commonly, decrepitations were witnessed for Type B2 inclusions, generally in the temperature range 150° to 220°C. It was observed that prior to decrepitation, most of these inclusions exhibited only minor bubble contraction hence the average homogenization temperature observed (ca. 230°C very represent a relatively low value with respect to the entire Type B2 inclusion population.

The lack of published empirical data pertaining to the compositional and volumetric poperties of the H₂O-NaCl-CO₂-CH₄ system at high temperatures precludes quantitative assessment of bulk fluid density in Type B2 fluid inclusions via the non-destructive analytical techniques used in this study. The predictive capabilities of theoretical equations of state developed in recent years are restricted in general to three component systems (e.g. H₂O-CO₂-CH₄; Jacobs and Kerrick, 1981; H₂O-CO₂-NaCl; Bowers and Helgeson, 1983a), and extrapolation to the fourth component is relatively unconstrained (Holloway, 1981). However, it is well established that addition of even small quantities of MaCl and/or CH₄ to the complimentary three component system will significantly affect the P-V-T properties of the bulk fluid by overall expansion of the two phase (liquid-vapor) field to higher temperatures and pressures (e.g. Bowers and Helgeson, 1983b; Hollister and Burruss, 1976).

In order to semi-quantitatively estimate bulk fluid densities and compositions of Type B2 inclusions used herein, the bulk density of the carbonaceous phase was adjusted to a "CO₂-equivalent" density using the method of Swanenberg (1979). This data, in the form of an adjusted Th_{CO2} (the addition of ca. 1°C per mole % CH₄ (Arai et al., 1971)), was then inserted along with Tm_{clath}, and Th_{total} data for individual inclusions

into an iterative computer program (Lynch, 1988) which calculates XCO₂, bulk molar volume, and pressure (on the two phase boundary) of H2O-CO2-NaCl fluid inclusions from salinity, ThCO2, and Thtotal data based upon the method of Parry (1986). This method applies calculations and phase relationships presented by Bodner (1983) and Bowers and Helgeson (1983a,b) to three phase H2O-CO2-NaCl fluid inclusions, and requires only accurate freezing and heating measurements. The method is advantageous in that the accuracy of the XCO2 estimate obtained is not related to a volumetric estimate of the CO2 phase as is required for other methods (e.g. Burruss, 1981; Bodner, 1983), but only to the accuracy of the temperature data generated through heating and freezing experiments. Thus uncertain, potentially large inaccuracies related to the visual estimation of phase volumes in fluid inclusions (Bodner, 1983; Roedder, 1984; Shepherd et al., 1985) are avoided. Applying the above outlined method to appropriate Type B2 fluid inclusions (i.e. those for which Tmclath., ThcO2, and Thtotal data were obtained, see Table 3-1), compositional estimates range from 4 to 11 averaging 7 to 8 mole percent CO2 "equivalent" with bulk fluid densities averaging approximately 0.9 to 0.95 g cm⁻³. XCO₂ within individual fields of inclusions is noted to be relatively constant. Application of a graphical procedure given by Schwartz (1989) yielded comparable results.

Summary of Fluid Inclusion Data

Two general types of fluid inclusions, two phase Type B1, and three phase Type B2 have been identified as coeval with gold mineralization in the quartz lodes of the Athabasca Pass. Three broad fluid compositions are preserved in these inclusions. Type B1 H₂O-NaCl±CO₂-bearing inclusions

may be subdivided into two varieties, namely: 1) those which form gas hydrates upon cooling and thus contain between approximately 1 and 3 mole percent CO₂ (Hedenquist and Henley, 1985), and 2) those which do not form gas hydrates and thus contain less than approximately 1 mole percent CO₂. Based upon Tm_{iCe} data, Type B1 inclusions exhibit salinities between 5 and 10 wt. percent NaCl equivalent with clathrate-forming inclusions being somewnat less saline. Th_{total} data for Type B1 inclusions are relatively well constrained in the 215° to 225°C range.

Type B2 H₂O-NaCl-CO₂ ±CH₄-bearing inclusions exhibit aqueous phase salinities between approximately 1.5 and 6, averaging 3.8 wt. percent NaCl equivalent. Expressed on a "CO₂-equivalent" basis (Swanenberg, 1979), Type B2 inclusions contain between 4 and 12 averaging approximately 8 mole percent total carbonaceous phase (dominantly CO₂ containing 2 to 16 mole % CH₄). Th_{total} data for Type B2 inclusions is limited due to frequent decrepitation prior to homogenization. Th_{total} measurements obtained cluster between 225° and 245°C and are considered to represent a minimum Th_{total} value. Inclusion decrepitation generally occurred between 150° and 220°C.

Type B1 and B2 fluid inclusions are petrographically indistinguishable at room temperature, being of similar size and shape, exhibiting comparable apparent phase ratios, and occupying an equivalent paragenetic position with respect to the native gold \pm galena \pm pyrite assemblage.

Interpretation

The constant phase ratios and relatively well-constrained

compositional and Thtotal data exhibited by Type B1 and Type B2 fluid inclusions suggests entrapment of homogeneous one phase fluids accompanied gold emplacement in the Athabasca Pass lodes. Although Types B1 and B2 inclusions appear paragenetically sub-contemporaneous, it is evident from their relative spatial distribution, contained phase ratios, and the general P-V-T-X relationships in the H2O-NaCl-CO2 (±CH4) system that the observed fluid compositions were likely not generated through fluid immiscibility processes (effervescence, unmixing, etc; see Ramboz et al. 1982). One feasible alternative is the mixing of two compositionally distinct fluids, in this case a moderately saline, relatively cooler aqueous brine (Type B1), mixed with a warmer H2O-CO2±CH4-bearing fluid of low salimity (Type B2). This interpretation, however, is precluded by a general lack of fluid mixing trends in the overall data set. For instance fluid inclusions of intermediate bulk composition are relatively rare, and correlation between Thtotal and salinity, and/or mole percent CO2 and salinity is lacking. Furthermore, as noted earlier, inclusion fields containing compositionally mixed fluids are not observed. Clathrate-forming Type B1 inclusions studied here may in part represent these mixtures, but the data are too sparse to be conclusive.

A second alternative process involves the evolution of the gold-mineralizing fluid from a relatively warmer, CO₂-rich aqueous fluid (Type B2) to a cooler CO₂-poor aqueous brine (Type B1). This interpretation also suffers from the general lack of progressively developed, intermediate fluid compositions. However, it better accounts for the absence of compositionally mixed fields of fluid inclusions; individual inclusion clusters tend to be isocompositional because they have trapped homogeneous fluids at unique instances during the progressive

mineralizing event. This interpretation implies that, within the gold depositional stage, Type B1 inclusions were trapped somewhat later than Type B2, an inference suggested by the lower Th_{total} temperatures recorded for Type B1 inclusions. Clathrate-feeming Type B1 inclusions may again represent intermediate fluid compositions.

P-T conditions of fluid entrapment

The fluids trapped as inclusions in the auriferous quartz lodes of the Athabasca Pass exhibit no evidence of immiscibility, hence only minimum pressure and temperature conditions during mineralization are provided by Thtotal and fluid compositional data. Thtotal for the majority of Type B1 and Type B2 inclusions fall between 210° and 245°C with Type B2 data averaging approximately 15°C higher than Type B1 (ca. 230°C versus ca. 215°C respectively). Pressure estimates at Thiogal (i.e. on the 2 phase solvus) for H2O-NaCl-"CO2 equivalent"-bearing Type B2 fluid inclusions, obtained utilizing the technique and phase relationships described by Parry (1986) and Bowers and Helgeson (1983 a,b) respectively are listed in Table 3-1. Because of frequent decrepitation prior to Thtotal, the quantity of data is limited. Regardless the estimates obtained for individual samples (four are represented) are consistent, with minimum (i.e. Thtotal) pressure estimates ranging from approximately 1000 to 1200 bars. Such estimates are in accord with the general tendency for Type B2 inclusions to decrepitate prior to homogenization implying internal inclusion pressures in excess of approximately 1000 bars (Leroy, 1979; Roedder and Bodner, 1980). As well, moderately high ambient pressures would be required to prevent the unmixing of Type B2 H2O-NaCl-CO2 ± CH4 fluids (Takenouchi and

Kennedy, 1965; Bowers and Helgeson, 1983a,b). In the absence of an independent geobarometer or geothermometer, further constraint of the pressure and temperature range of trapping cannot be derived from Type B2 inclusion data alone.

Type B1 fluid inclusions, as discussed earlier, were trapped subcontemporaneously with or (if progressive fluid evolution is considered) slightly later than Type B2 inclusions. If the apparent lack of fluid unmixing, and the generally consistent minimum fluid pressure estimates derived for Type B2 inclusions may be considered to represent relatively otolic during inclusion entrapment, then the pressure of trapping estimates for Type B2 inclusions should also apply to Type B1 inclusions. Hence, because phase separation takes place at higher pressures in carbonaceous phase-bearing brines than in brines containing no carbonaceous phase (e.g. Takenouchi and Kennnedy, 1965; Haas, 1976), a semi-quantitative, pressure correction may be derived for Type B1 inclusions. Consider an average Type B1 fluid composition (aqueous liquid + ca. 8.5 wt. % NaCl eq.) which homogenizes at 215°C. A pressure of trapping in the vicinity of 900 to 1200 bars is reasonable if the fluids are assumed to have been trapped coeval with or slightly later than Type B2 (decrepitation was also observed prior to Thtotal for a number of Type B1 inclusions, supporting trapping pressures in this range). Thus, interpolating pressure correction data for the H2O-NaCl system given by Potter (1977), addition of approximately 80° to 110°C for the pressure range and average fluid composition outlined above yields pressure-corrected Thtotal-temperatures for Type B1 inclusions of 295° to 315°C. A similar temperature increment must also apply to Type B2 inclusions assuming

penecontemporaneous Type B¹ and B2 inclusion trapping. Thus pressure-corrected Th_{total} temperatures for Type B2 inclusions fall in the 310° to 330°C range.

It must be emphasized that the corrected P-T conditions of inclusion trapping proposed above are conservative, minimum approximations as they are based upon a minimum estimate of trapping pressure, i.e. on the two phase boundary for Type B2 inclusions. In view of the lack of fluid immiscibility, and the frequency with which inclusion decrepitation was observed, fluid pressures at trapping were likely significantly above this solvus. As well, the expansion of the two phase field due to the presence of minor CH4 (Hollister and Burruss, 1976) has not been taken into account. With respect to the proposed pressure corrections, these estimates are based upon average fluid composition and Thtotal measurements for which there are in fact ranges. Compensating for fluid compositional and Thtotal variation involves corrections of ± 20°C for the temperatures of entrapment proposed above.

It is concluded that the P-T conditions prevalent during gold mineralization in the Athabasca Pass were in the range of 295° to 330° \pm 20°C at pressures in the order of 900 to \geq 1200 bars. The absence of independent, quantitative geobarometric-thermometric techniques applicable to the Athabasca Pass gold-quartz lodes has been noted above. However, it is noteworthy that P-T constraints proposed above are in agreement with the broader constraints imposed by the regional anchimetamorphic-pumpellyite facius (ca. 200° to 350°C and 1 to 4 kb (Turner, 1981; Kisch, 1987)) conditions under which the Athabasca Pass lodes are considered to have been emplaced.

Discussion

Based upon a wide range of geological parameters, including lithotectonic setting, metamorphic regime, and vein temporal, structural, textural and mineralogical relationships, the gold-quartz lodes of the Athabasca Pass were previously classified as a siliciclastic-hosted analogue of the Turbidite Hosted class of gold deposits (Keppie et al., 1986; Hutchinson, 1987). Based upon integrated geological-geochemical parameters turbidite-hosted deposits have been termed mesothermal in character.

A comparison between fluid compositional and P-T constraints derived for the Athabasca Pass lodes and data summarized for a number of turbidite-hosted gold deposits is presented in Table 3-2. From this table, it is evident that the fluid compositional data for the Athabasca Pass lodes is typical of turbidite-hosted deposits, i.e. essentially low salinity brines containing generally low, but variable, quantities of CO2, with minor quantities of other volatiles, dominantly CH₄ (± N₂). As shown, P-T constraints proposed for gold-stage mineralization in the Athabasca Pass in terms of both minimum Thtotal and pressure corrected scenarios are also in the general range for those derived from turbidite-hosted deposits, and mesothermal gold-lodes in general. With respect to fluid processes and gold depositional mechanisms, it is notable that many authors studying turbidite-hosted gold deposits consider, as is documented herein, fluid immiscibility to be absent from or unimportant with respect to the gold mineralizing event (Paterson, 1986; Steed and Morris, 1986; Goldfarb et al., 1986; Kontak et al., 1988).

In view of a fluid and P-T regime in the Athabasca Pass typical of mesothermal turbidite-hosted gold deposits, a comparable suite of associated trace elements accompanying gold mineralization might be expected.

However as documented earlier, in terms of both the sparse variety and overall abundance of trace elements in the Athabasca Pass lodes, this is not the case. A fundamental difference between the Athabasca Pass and turbidite-hosted gold deposits in general involves the nature of the host rocks i.e., mature, quartz-dominated ± pelitic clastic rocks in the Athabasca Pass versus a wide variety of generally immature volcano-sedimentary clastic rock types in turbidite-hosted deposits. The general paucity of trace elements in quartz-dominated clastic rocks accounts for lower concentrations of trace-elements, as well suggests the relatively local derivation and control of lode trace element concentrations by the quartzitic rock types hosting the mineralized Athabasca Pass structures. Such an inference would also suggest that the finids respond as a for mineralization in the Athabasca Pass were either dominantly interstratigraphically derived, or that their circulation was relatively restricted to the quartz-dominated rocks of the McNaughton Formation or the Gog Group (a regionally correlatable stratigraphic unit in excess of 1 km thick throughout stratigraphic of the Main Ranges of the Rocky Mountains). Interestingly, Miette Group strata which underly the McNaughton Formation and outcrop immediately to the west of the Athabasca Pass (Fig. 3-1) contain quartz veins which host a variety of sulfide minerals including pyrite, chalcopyrite, galena and arsenopyrite. Host rocks include thick turbidite-lain sequences of grit, greywacke, semi-pelite and slate; immature rock types from which a variety of trace elements would be more readily available.

With respect to the interstratigraphic derivation of fluids from the host rocks of the Athabasca Pass abundant pressure solution and recrystallization of allogenic clays to white mica in the host quartzites, pebble-conglomerates and pelites of the McNaughton Formation have been documented. Both of these processes have been attributed to regional anchimetamorphism in the Main Ranges of the Rocky Mountains, and notably provide a potential fluid source (and mode of gold-trace element mobilization) from which mineralized lodes may be derived. Similar processes have frequently been advocated in the genesis of turbidite-hosted, and mesothermal-metamorphic vein-type gold deposits. (e.g. Boyle, 1979; Cox et al.,1986; Graves and Zentilli, 1982; Kerrich and Allison, 1978; Paterson, 1986; Tomkinson, 1988). As discussed previously, the general congruency between the regional metamorphic, host-rock exteration, and vein mineralogical assemblages would further imply the relatively lociderivation of the vein-filling constituents in the Athabasca Pass. Steble isotopic analysis of host-rock and vein-constituents may provide a more complete evaluation of potential fluid sources for the Athabasca Pass lodes.

Concluding Statement

The gold metallogenic potential of low-grade meta-sedimentary fold belts such as Victoria, Australia, and Nova Scotia, Canada has been widely recognized. These and other districts, which have founded the classical turbidite-hosted lode gold deposit-type, have in recent years, become of renewed interest in exploration programs (e.g. Kontak and Smith, 1988; Watchorn et al., 1988; Wilson el al., 1988). The siliciclastic-hosted lode-gold system documented herein may be viewed as an analogue of the turbidite-hosted class of gold deposits, thus the potential of siliciclastic domains should not be overlooked in the broad context of turbidite-hosted, and mesothermal lode-gold exploration programs. The Main Ranges of the

Rocky Mountains may be viewed as a viable region for the exploration for such deposits.

Table 3-1. Mole % ${\rm GO_2}$ and pressure of trapping estimates for Type B2 fluid inclusions.

| Sample no. | Tm _{CO2} , ℃ | Th _{CO2} , ℃ | CH ₄ carb. Mole % | Salinity, ² wt.%'Ol eq. | Th _{total} , ℃ | CO2 ³ Mole % | Pressure, bars |
|------------|--------------------------|--------------------------|---------------------------------|---------------------------------------|----------------------------|----------------------------|-------------------|
| PE-A3 | -57.3 | 22.5 | 5 | 5.6 | 253 | 10 | 1084 |
| . = , | -57.8 | i9.5 | 7 | 4.1 | 221 | 6 | 1107 |
| | -58.0 | 19.4 | 8 | 3. <u>Q</u> | 226 | 6 | 1108 |
| | -58.2 | 14.1 | 9 | 3.2 | 220 | 8 | 1101 |
| | -58.1 | 17.9 | 8 | 3.4 | 229 | 8 | 1106 |
| | -57.7 | 22.9 | 5 | 3.9 | 231 | 6 | 1097 |
| | -57.0 | 21.6 | 4 | 4.3 | 243 | 11 | 1117 |
| | -58.3 | 22.4 | 9 | 3.8 | 232 | 5 | 1097 |
| PB-A11 | -57.5 | 23.6 | 5 | 3.9 | 229 | 6 | 1103 |
| | -57.8 | 23.4 | 5 | 4.5 | 225 | 5 | 1142 |
| | 4 | 23.9 | 9 | 3.9 | 232 | 10 | 1094 |
| | 4 | 23.0 | 6 | 3.8 | 219 | 5 | 1098 |
| | ,7.2 | 22.4 | 5 | 3.8 | 231 | 8 | 1089 |
| | -57.5 | 22.3 | 5 | 3.9 | 234 | 8 | 1090 |
| PB-A16 | -57.9 | 16.8 | 8 | 4.3 | 244 | 11 | 1094 |
| | -58.1 | 15.8 | 8 | 3.0 | 236 | 11 | 1102 |
| | -58.1 | 17.5 | 8 | 2.8 | 244 | 11 | 1096 |
| | -57.7 | 20.6 | 6 | 3.6 | 242 | 10 | 1095 |
| | -57.9 | 18.5 | 8 | 4.1 | 239 | 9 | 1102 |
| PB-A24 | -57.3 | 24.5 | 8 | 5.1 | 211 | 4 | 1107 |

Hadmated from the graphs of Heyen et al., 1982.

²Calculated from relations given by Bozzo et al., 1976.

^{3,4}Calculated using the method of Parry, 1986.

Table 3-2. Fluid inclusion characteristics of the Athabasca Pass vs. typical turbiditehosted * and mesothermal ≹lode-gold deposits.

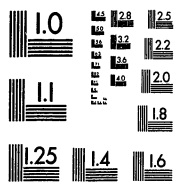
| | Athabasca Pass | Turbidite-Hosted | Mesothermal |
|--|--|----------------------------------|-----------------------------------|
| Th total, °C: | 190-315 | range~200-4େଣ | |
| Est. mole $\%$ CO $_2$: | 0-11 average~8 | range 0-20 generally <15 | X |
| Dominant additional volatiles: | CH ₄ ± N ₂ | CH ₄ ± N ₂ | ±CH ₄ , N ₂ |
| Est. salinity, wt.% NaCl eq.: | ≥10 | 2-0 | 0-4 |
| Inferred temperature and pressure of gold deposition: | 275°-350°C, ≥900 bars | 290°-360°C, 500-3000 bars | 250°-350°C, >700 bars |
| Principle references: *Goldfarb et al. (1986), *Kontak et al. (1988), *Paterson (1986), *Steed and Morris (1986), † Nesbitt (1988) | arb et al. (1986),*Kontak et Nesbitt (1988) | al. (1988),*Paterson (198 | 36), |

*Steed and Morris (1986), † Nesbitt (1988).

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MICROCOPY RESOLUTION TEST CHART NATIONAL BUREAU OF STANDARDS STANDARD REFERENCE MATERIAL 1010a (ANS) and ISO TEST CHART No. 2)

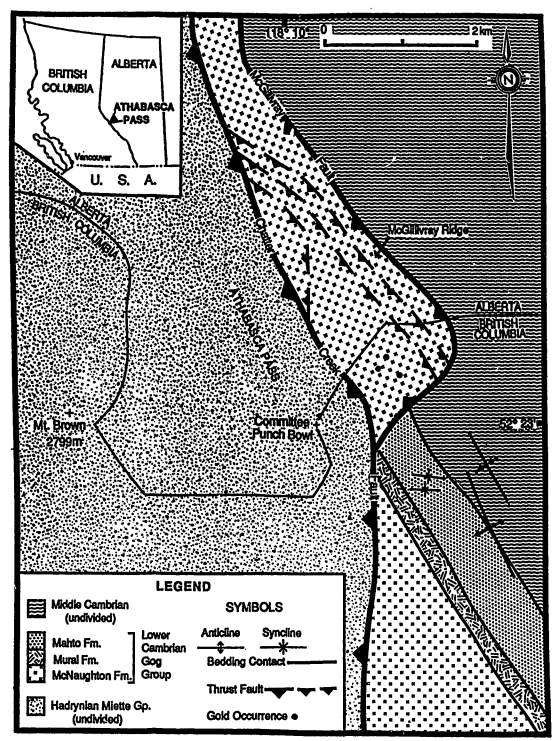
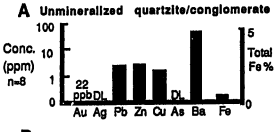
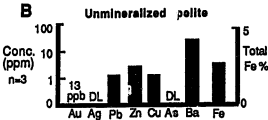
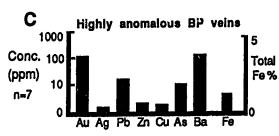


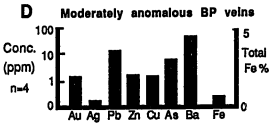
Figure 3-1. Location and general geology of the Athabasca Pass area. Data compiled from Mountjoy and Price (in prep.) and thesis data. Gold occurrences represent clusters of mineralized veins.

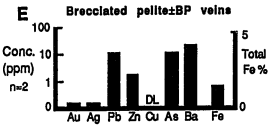
Athabasca Pass Rock Types

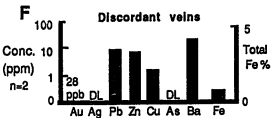




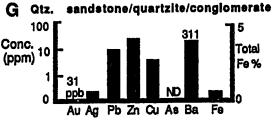


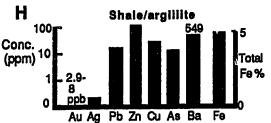






Literature-Compiled Data





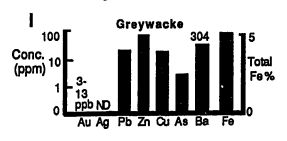
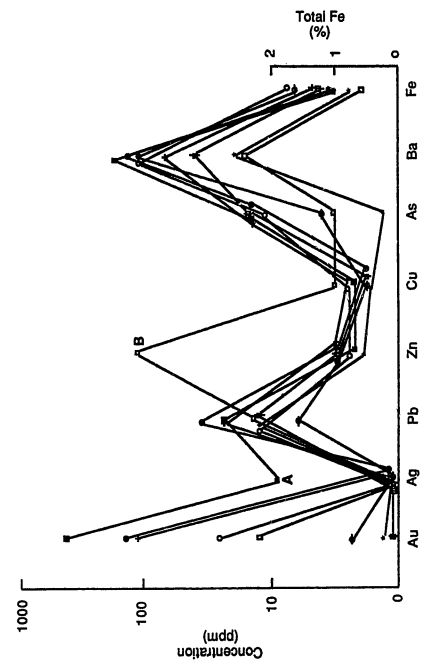


Figure 3-2. Selected trace element data for the Athabasca Pass and "average" comparable lithologies. Data for the Athabasca Pass lithologies represents the arithmetic mean for the number of samples (n) indicated. Detection limits in present study: (ICPES) Pb-2ppm, Zn-1ppm, Cu-1ppm, As-5ppm, Ba-1ppm (AAS) Au-5ppb, Ag-0.1 ppm. Abbreviations: BP=Bedding-Parallel, ND=No Data, DL=Detection Limit. Principle references for compiled data; Wedepohl (1974) and Boyle (1979).



vein material (Figs. 3-2C and 3-2D) from the Athabasca Pass. Symbols denote individual samples. Figure 3-3. Trace element variations exhibited by 8 samples of Au-bearing bedding-parallel Points A and B are referred to in the text.

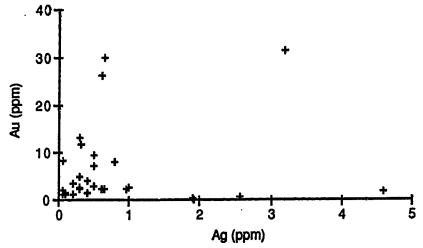


Figure 3-4. Scatter plot illustrating the poor correlation between gold and silver for 30 samples of auriferous bedding-parallel vein material from the Athabasca Pass.

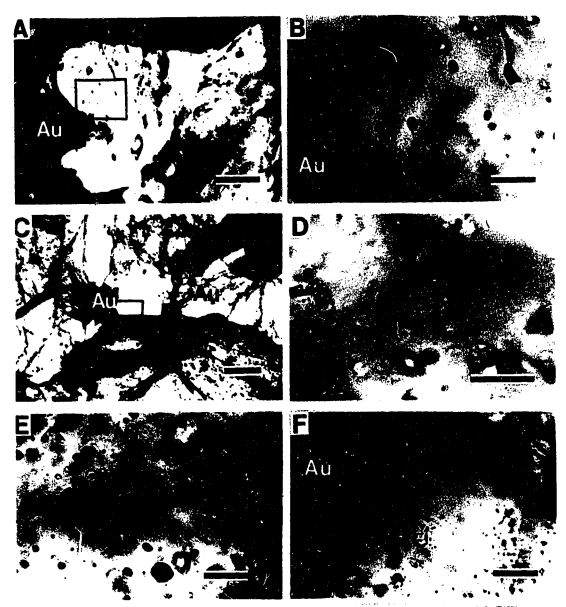
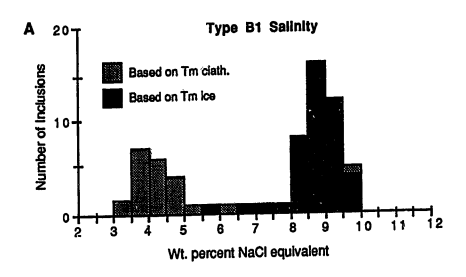


Figure 3-5. Type B fluid inclusions in quartz from some Au-bearing veins on McGillivray Ridge. All photographs taken at room temperature (ca. 22° C). A. Quartz-filled embayment in native gold (Au). Rectangle approximates area in photograph B. Scale bar = 80μm. B. Type B1 fluid inclusions adjacent to native gold (Au). Scale bar = 20μm. C. Gold-filled (Au) fractures in quartz. Rectangle approximates area in photograph D. Scale bar = 80μm. D. Type B1 fluid inclusions adjacent to native gold. Scale bar = 25μm. E. Numerous Type B2 fluid inclusions adjacent to native gold (Au). Scale bar = 25μm. F. Solitary Type B2 fluid inclusion adjacent to native gold (Au). Scale bar = 80μm.



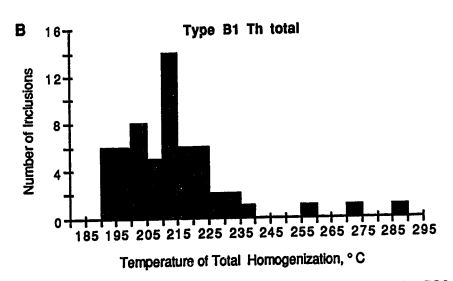


Figure 3-6 Microthermometric data for Type B1 H2O-NaCl±CO2 -bearing fluid inclusions contained within auriferous vein quartz from the Athabasca Pass. Salinity estimates based upon clathrate melting are approximate maxima.

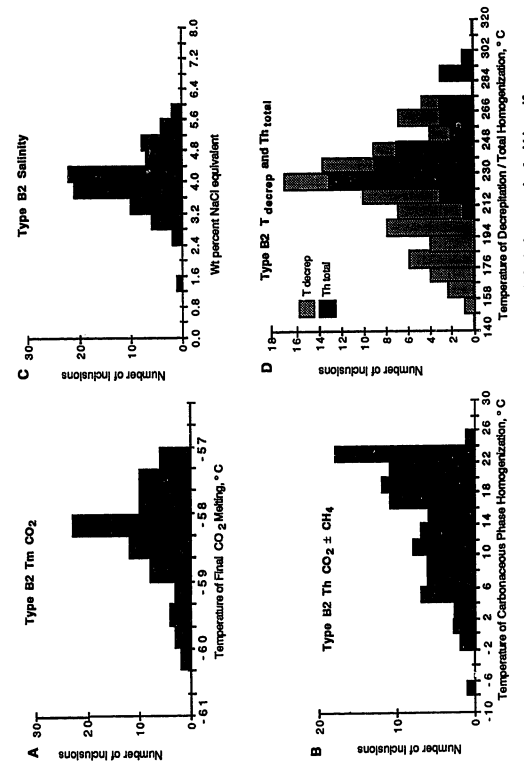


Figure 3-7. Microthermometric data for Type B2 H2O-NaCl-CO2±CH4-bearing fluid inclusions contained within auriferous vein quartz from the Athabasca Pass.

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CHAPTER 4

Origins of Metamorphogenic Lode Gold Deposits: Implications of Stable Isotope Data From the Central Rocky Mountains, Canada.

Introduction

The origins of hydrothermal fluids which generated mesothermal gold-quartz mineralization in low-grade meta-sedimentary ± plutonic domains ("Turbidite-Hosted gold deposits", Keppie et al., 1986; Hutchinson, 1987, Nesbitt, in press) have, in recent years, become a subject of renewed controversy (eg. Nesbitt et al., 1986, 1987; Goldfarb, et al., 1987; Kerrich, 1987). The root of much of this controversy stems from the definition of the isotopic (O-H) signatures of naturally occurring waters (Craig, 1961; Taylor, 1974, 1979), and from the increased application of stable isotope (O-H-C-S) systematics to the study of this mineral deposit type. It is becoming increasingly evident through integrated geological-geochemical studies (e.g. Mitchell et al., 1981; Nesbitt et al., 1986; Weir and Kerrick, 1987; Shelton et al., 1988) that isotopically evolved waters of meteoric origin may play a significant role in what, based upon field-petrographic relationships, and often cursory geochemical (fluid inclusion, trace element, and/or isotopic) studies, has traditionally been considered a "metamorphogenic" style of gold mineralization (Boyle, 1979; Nesbitt, in press).

The stable isotope relationships of gold-bearing quartz veins recently discovered at Athabasca Pass, in the Canadian Rocky Mountains, provide a new insight into the genesis of gold-quartz veins in low-grade metasedimentary domains.

The Gold-Bearing Quartz Veins of the Athabasca Pass Regional setting

The Athabasca Pass is located 65 km south-southwest of Jasper, Alberta, in the Main Ranges of the central Canadian Rocky Mountain thrust and fold belt (Fig. 4-1). This lithotectonic domain, physiographically bound by the Rocky Mountain-Tintina Trench system to the West, and by the Foothills and Interior Plains region to the east, consists almost entirely of clastic and carbonate strata. Metamorphic grade decreases from amphibolite grade in the west to low-grade burial metamorphic facies east of the deformation front. Igneous rocks are rare in the Main Ranges. Mid-Cretaceous to Early Tertiary Cordilleran deformation in the Main Ranges caused the northeastward translation and thrust-stacking of parautochthonous miogeoclinal strata derived from the North American craton to the east, onto the western flank of the craton (Price and Mountjoy, 1970; Price et al., 1985). The geologic structure of the region is dominated by complex folds and by widely-spaced thrust faults. Penetrative deformation is pronounced in the western Main Ranges, and less pervasive to absent in the east (Price and Mountjoy, 1970).

Information on lode-gold deposits in the Main Ranges, and in the Canadian Rocky Mountains in general, is sparse. The mineral occurrences and metallogeny of this region have been reviewed by Hedley (1954), Little et al. (1976), and by Sinclair et al. (1978). Principal deposit-types occuring to the south of the Athabasca Pass include stratiform lead-zinc (±silver) replacement bodies in Middle Cambrian carbonates, and small, quartz-carbonate vein systems containing a variety of base-metal sulfide

assemblages (dominantly galena-sphalerite-pyrite-chalcopyrite). Gold production has not been reported from these occurrences. Brief mention of placer gold occurrences in the Main Ranges was made by Mathews (1944), Sorenson (1955), and by Boyle (1979).

Local geology

Aspects of the geology and geochemistry of the Athabasca Pass gold-quartz lodes have been discussed in detail in Chapters 2 and 3. These mineral occurrences consist of numerous 0.5 cm- to 1 m-thick auriferous, bedding-parallel quartz veins contained within an 800 m-thick, folded and thrust-faulted section of Lower Cambrian quartzite with minor pelite, and quartz-pebble conglomerate known as the McNaughton Formation (Fig. 4-1) This is the lowermost formation of the Gog Group, a regionally extensive stratigraphic unit in excess of 1 km thick throughout much of the Main Ranges. The area has been metamorphosed to sub-or lowermost greenschist grade. Igneous rocks are not known in the area.

Native gold, accompanied by minor pyrite and galena, was deposited late during prolonged, syn-tectonic, syn- to post-metamorphic bedding-parallel vein formation. Thin, incompetent, iron-rich pelites provided structural-chemical traps for the localization of hydrothermal fluids and for the precipitation of native gold. Discordant veins which formed penecontemporaneously with gold emplacement cut only competent quartzites and lack gold and sulfides. Hydrothermal alteration adjacent to some veins includes pyrite ± ankerite ± white mica. However, this alteration assemblage is not uniquely confined to the proximity of veins and is seen widely distributed within the regional metamorphic suite.

Trace element analysis of auriferous veins and unmineralized host rocks revealed distinct enrichment of Pb, As, Ba, and Fe, and marginal enrichment of Ag, Zn, and Cu in veins, with respect to host rocks. When compared with other lode-gold deposits in similar domains, the Athabasca Pass lodes are conspicuously unenriched in all trace elements except gold. This situation has been attributed to the mineralogical maturity of the McNaughton Formation quartzites.

Microthermometric analyses of fluid inclusions in vein quartz indicated gold-stage fluids were aqueous brines containing 2 to 10 wt. % NaCl eq. and up to 12 mole % $CO_2 \pm CH_4$. Variations in fluid compositional- and total homogenization-data suggest that gradual fluid evolution (decreasing CO_2 and temperature, increasing salinity) accompanied gold mineralization. Gold deposition took place at between 275° and 350°C and 900 to \geq 1200 bars.

Combined field, petrographic, trace element and fluid inclusion evidence indicates the Athabasca Pass lodes to be typical of mesothermal gold-quartz lodes in meta-sedimentary domains. This evidence implies the relatively local derivation of vein-filling constituents, and it was suggested in Chapter 3 that the lodes may have been derived through devolitalization and deformation processes associated with low-grade regional metamorphism. To evaluate the possible role of metamorphic fluids in the genesis of the Athabasca Pass lodes, stable isotope analyses of vein and country-rock minerals were undertaken.

Stable Isotope (S-O-H) Study

Experimental techniques

Sulfur was extracted from hand-picked vein and country-rock sulfides

using the technique of Ueda and Krouse (1986). Oxygen from vein and host-rock silicate minerals was extracted using the BrF5 technique of Clayton and Mayeda (1963). Fluid inclusion waters from bedding-parallel and discordant vein quartz were extracted by thermal decrepitation under vacuum at 1100°C. Production of hydrogen gas from the released waters was accomplished by reaction with zinc metal at 450°C (Coleman et al., 1982).

Mass spectrometric results are reported in standard δ notation relative to the Cañon Diablo Troilite (CDT) standard for sulfur, and the Standard Mean Ocean Water (SMOW) standard for oxygen and hydrogen. Analytical reproducibilities were \pm 0.2 per mil for sulfur, \pm 0.2 per mil for oxygen, and \pm 5.0 per mil for hydrogen.

Sulfur isotope study

Twenty-nine samples of sulfide-bearing unmineralized country-rock and gold- mineralized quartz vein were selected. Thirty-seven sulfide separates including seven pyrite separates from the Hadrynian Miette Group, seven pyrite separates from the Lower Cambrian McNaughton Formation, 22 separates from mineralized veins within the McNaughton Formation (including 16 pyrites and 6 galenas), and a single pyrite separate from Middle Cambrian strata were obtained. Petrographic studies indicate that pyrite occurring within Miette Group, McNaughton Formation and Middle Cambrian strata is metamorphic in origin, whereas sulfides occurring within mineralized veins in the McNaughton Formation are hydrothermal in origin, and were coeval with gold mineralization.

Results

δ 34 S values for unmineralized country-rock pyrites (Fig 4-2) range from +1.1 to +8.9 per mil and +12.2 to +16.9 per mil for the Miette Group, and the McNaughton Formation separates respectively. The Middle Cambrian separate produced a value of +36.7 per mil. McNaughton Formation vein pyrites produced values ranging from +13.7 to +19.9 per mil (with 14 of 16 samples falling between +14.2 and +16.3‰). Vein galenas returned values ranging from +11.4 to +13.3 per mil.

Sulfur isotope geothermometry

Pyrite and galena have not been observed in physical contact in the McNaughton Formation quartz veins. Thus it is difficult to assign specific pairs for geothermometry. However, based upon their paragenetic associations, these sulfides were interpreted as coeval. The observed $\delta^{34}S$ enrichment of pyrite over galena (Fig. 4-2) indicates that isotopic equilibrium with respect to sulfur was approached (Bachinski, 1969; Kajiwara and Krouse, 1971). Thus it was deemed that useful paleotemperatures might be derived from pyrite-galena separates. Two pyritegalena "pairs" have Δ^{34} S (pyrite - galena) values of 2.6 per mil and 3.0 per mil, thus yielding apparent equilibrium isotope paleo-temperatures of 353 \pm 53°C and 309 \pm 44°C (Ohmoto and Rye, 1979). Alternatively, as the distribution of $\delta^{34}S$ for vein sulfides exhibits relatively narrow ranges, suggestive of contemporaneous precipitation, geothermometric calculations utilizing the arithmetic means for the δ^{34} S values plotted in Figure 4-2 may be useful. Thus, the mean $\Delta^{34}S$ (pyrite - galena) becomes 3.1 per mil and a paleo-temperature of 300 \pm 43 °C is derived. Notably all paleo-temperatures

calculated here are in agreement with pressure corrected paleo-temperatures of 275 - 350°C inferred by gold-stage fluid inclusions in quartz.

$\delta^{34}S(\Sigma S)$ of the hydrothermal fluids

In order to estimate $\delta^{34}S(\Sigma S)$ of the hydrothermal fluid, and in turn to speculate upon the source of sulfur for the auriferous veins of the McNaughton Formation, the dominant sulfur species in the fluid must be known. Sulfur speciation is controlled by fO2, pH and by temperature. An assumed paleo-temperature of 325°C is reasonable, based upon fluid inclusion- and pyrite-galena- geothermometry. As oxide minerals are absent from the veins, and methane is present in fluid inclusions, fO2 was clearly low, and did not vary outside the stability field of pyrite. The pH was likely buffered to near neutral (ca. 5.8 at 325°C) as evidenced by the partial replacement of K-feldspar by white mica in the wall rocks, and by the presence of hydrothermal white mica in the auriferous veins. Under such conditions, the dominant sulfur species in solution would have been H2S, and $\delta^{34}S_{H_2S} \approx \delta^{34}S_{(\Sigma S)}$ can be assumed (Ohmoto and Rye, 1979; Ohmoto, 1986). Using isotopic equilibrium constants between sulfide minerals and H₂S given by Ohmoto and Rye (1979), the δ^{34} S(Σ S) of the fluid from which vein sulfides precipitated would have been in the range +13 per mil to +16 per mil.

Sulfur source

Abundant pyrite porphyroblasts occur within the country rocks of the Athabasca Pass. These country-rock pyrites fall within distinct δ^{34} S ranges, characteristic of the stratigraphic horizon in which the pyrites occur (Fig. 4-2). McNaughton Formation vein sulfides exhibit approximately the same

range of δ^{34} S values as their country-rocks (i.e. +11.4 to +19.9 per mil and +11.8 to +16.9 per mil respectively). As stated earlier, the δ^{34} S(Σ S) of the fluid from which vein sulfides were deposited was +13 to +16 per mil. H₂S in this mineralizing fluid may have been derived through the hydration of country-rock pyrite. Ohmoto (1986) indicates that H₂S generated by such reactions will have a δ^{34} S signature essentially identical to the reactant pyrite. Thus the congruency between δ^{34} S values for McNaughton Formation vein sulfides, mineralizing fluids, and host-rocks implies that sulfur was locally (i.e. interformationally) derived, and that the surrounding Miette Group or Middle Cambrian strata did not contribute sulfur to the gold-bearing veins.

Oxygen isotope study

Thirty samples, representing local McNaughton Formation quartzites and pelites, bedding-parallel quartz veins (including paragenetically early (Type I), and late (Type III) quartz) and unmineralized discordant veins were analysed. Quartzites were sampled proximal to, and distal from both vein types. Pelites were sampled from cataclasts within mineralized veins and as intact, unveined strata.

Results

 δ^{18} O of McNaughton Formation quartzites (7 samples) range from +12.0 to +13.5 (averaging +12.9‰, Table 4-1) Values are uniform, regardless of sample proximity to quartz veins. Likewise, pelites (6 samples) exhibit a narrow range in δ^{18} O (+9.5 to +10.5, averaging +10.0 ‰) with the δ^{18} O of veined and unveined pelite being statistically identical. All 17 pre-gold and

gold-stage vein quartz samples returned $\delta^{18}O$ values between +13.0 and +15.0 per mil. Early, Type I quartz, late Type III quartz, and discordant vein quartz averaged +14.4 \pm 0.9, +14.0 \pm 1.0, and +14.5 \pm 0.5 per mil respectively.

Temperature of metamorphism

Oxygen isotope studies of regional metamorphic domains (see Taylor, 1979; Valley, 1986) reveal a tendency toward total isotopic homogenization and equilibration, due to the lengthy time periods involved in regional metamorphism. McNaughton Formation quartzites and pelites exhibit narrow $\delta^{18}O$ ranges, independant of their proximity to, or the degree of, quartz veining. Hence it may be assumed that these rocks have approached isotopic homogenization, and have retained their metamorphic $\delta^{18}O$ signatures throughout the mineralizing event. An approximate paleotemperature of metamorphism may therefore be derived utilizing the quartz - muscovite isotopic geothermometer and $\Delta^{18}O$ (quartz-white mica) from McNaughton Formation quartzites and pelites (these rock types are essentially monomineralic). Mean $\delta^{18}\text{O}$ values of ± 12.9 (quartzites) and +10.0 (pelites) per mil yield a Δ^{18} O value of +2.9 per mil. Equations given by Field and Fifarek (1985) provide an approximate paleo-temperature of 370°C. Retrograde isotopic resetting during cooling is unlikely at moderate temperatures, hence 370°C likely represents near peak-metamorphic conditions. This temperature is in good agreement with the archimetamorphic (sub- to lowermost greenschist facies) regime implied by the metamorphic mineral assemblages, and by deformational microstructures in the McNaughton Formation.

Interpretation and $\delta^{18}O$ of the hydrothermal fluids

Segregation vein minerals formed during regional metamorphism generally exhibit δ^{18} O values similar to their host-rocks (Taylor, 1979; Kerrich, 1987). The overlap between the δ^{18} O of quartzites (range +12.0 to +13.5 ‰) and bedding-parallel quartz veins (range +13.5 to +15.0 ‰) at Athabasca Pass suggests that the veins were, in part, deposited from fluids which were isotopically- and thermally-equilibrated with the McNaughton Formation. However, the trend toward higher δ^{18} O values for the quartz veins suggests that the deposition of the bulk of the vein material, including gold and sulfides, took place from fluids which were not isotopically equilibrated with their wallrocks, or alternatively, that wall-rock equilibrated fluids cooled somewhat prior to/during vein deposition, hence depositing vein quartz enriched in 18 O. Direct lateral secretion of the veinforming components during peak metamorphism is precluded by the trend toward higher δ^{18} O values in the veins (Kerrich, 1987).

Deposition from non-equilibrated fluids can be ruled out by observations implying broad chemical-thermal equilibrium between veins and host-rocks (e.g. the paucity of wallrock alteration and the mineralogical congruency between the metamorphic suite \pm alteration and vein assemblages), and by trace-element and fluid inclusion data which suggest the relatively local derivation of the vein-filling constituents. The involvment of additional, externally-derived fluids and fluid mixing is precluded by the narrow range of δ^{18} O values, and by the lack of mixing trends in both the O- and S-isotope data sets.

Alternatively, the cooling of isotopically equilibrated fluids could account for the apparent vein - wall-rock disequilibrium. For example,

utilizing mean-, and maximum- and minimum- δ^{18} O values for McNaughton Formation quartzites and pelites, and an approximate paleotemperature of metamorphism of 370°C, aqueous fluids in equilibrium with wall-rocks would have had a δ^{18} O value of +8.1 (±0.9 ‰ for quartzites, ±0.5 ‰ for pelites; Matsuhisa et. al., 1979 (quartz-water); Friedman and O'Neil, 1976 (muscovite-water)). As this fluid cooled toward/during gold stage deposition (ca. 350-300°C, based upon fluid inclusion and sulfur isotope geothermometry), quartz exhibiting δ^{18} O values in the range 13 to 15 per mil would be deposited. This interpretation is supported by fluid inclusion data which suggests the gradual cooling of fluids during vein formation, and by structural, vein-textural, and paragenetic evidence which supports progressive late-syn to post-peak metamorphic vein filling. Thus vein deposition from relatively local aqueous fluids which were isotopically equilibrated with their wall-rocks prior to vein formation best accounts for the McNaughton Formation vein - host-rock O-isotope relationships at Athabasca Pass. The ultimate source of these fluids may be indicated by their H-isotope composition.

Hydrogen-isotope study

Fluid inclusion extracts from 6 auriferous bedding-parallel vein quartz separates (4 early, Type I, and 2 late Type III) from the McNaughton Formation were analysed. Such fluids represent samples of the fluids from which the vein components were precipitated (Roedder, 1984). Detailed petrographic investigations indicate abundant fluid inclusions in quartz are associated with gold-stage mineralization at Athabasca Pass.

Results

All fluid inclusion fluids exhibited δD values between -105 and -124 per mil (averaging ca. -115 ‰, Table 4-1). No systematic differences between Type I and Type III quartz, or discordant vein quartz fluids were noted. Variations outside analytical uncertainty limits may be accounted for by minor fluctuations in the δD value of the mineralizing fluid during the progressive mineralizing event, or by minor contamination by fluids, trapped during late deformation, which were unrelated to vein formation.

Interpretation and δD of the hydrothermal fluids

The H- (versus O-) isotope compositions of the Athabasca Pass vein fluids are plotted in Fig. 4-3. The isotopic (O-H) composition of other naturally occurring waters (magmatic, metamorphic, and meteoric) are also shown. It appears that the δD values of the Athabasca Pass fluids lie significantly below the magmatic- and metamorphic-water fields (ca. 30-60 % more negative). Hence large, negative δD shifts must be explained if either of these fluid-types played a significant role in the formation of the Athabasca Pass gold-quartz lodes.

The involvement of magmatic fluids in vein formation is precluded by the regional paucity of igneous rocks. In order for fluids of metamorphic derivation to be involved, negative δD shifts of ≥ 45 per mil must be invoked. Such large shifts may, under certain circumstances, be attributible to either fluid evolution-, or to fluid mixing- processes. The evolution of fluids as a result of fluid - wall-rock interactions which cause negative δD (fluid) shifts, is unlikely, as most mineral-water fractionation factors for hydrogen are negative (i.e. the fluid phase becomes enriched (Field and

Fifarek, 1985)). Other processes such as boiling or fluid unmixing also tend to enrich, not deplete, the fluid phase in D. The mixing of metamorphic fluids with isotopically lighter fluids at Athabasca Pass is precluded by the relatively narrow ranges, and by the lack of mixing trends exhibited by the associated O-, S-, and H-isotope data sets.

Thus the involvement of significant quantities of magmatic or metamorphic fluids in the formation of the auriferous quartz veins at Athabasca Pass cannot be postulated. Rather, the fluid source for these veins must have involved evolved waters of meteoric origin.

Present-day meteoric water from the Athabasca Pass has a δD value of approximately -135 per mil (Craig, 1961; Nesbitt and Muehlenbachs 1989). Assuming the isotopic composition of meteoric water has not shifted significantly during the last 150 Ma (Taylor, 1979), vein fluids averaging -115 per mil at Athabasca Pass record a positive δD (fluid) shift of up to 20 per mil (possibly somewhat less given the uncertainty in the H-composition of ancient meteoric waters (i.e. ± 10 ‰, Taylor, 1979)). Positive H-isotope shifts of this magnitude may be reconciled through either of two processes, namely: 1) the mixing of minor amounts of isotopically-heavy fluids (in this case metamorphic) with isotopically-light meteoric waters. or 2) by fluid - wall-rock isotope exchange reactions at moderate- to low-water to rock (w/r) ratios. The first possibility is precluded by well constrained S-, O-, and H-isotope data, and by the lack of fluid mixing trends. The involvment of minor amounts of metamorphic water however, cannot be ruled out completely, especially during the early stages of vein formation. Alternatively, the second case of fluid - wall-rock isotope exchange reactions, as discussed below, could account for both the H-, and O-isotope

data sets from the Athabasca Pass, without the involvment of metamorphic fluids.

Evolution of the hydrothermal fluids

The generation of O-enriched meteoric waters through isotope exchange reactions with wall-rock minerals during crustal fluid circulation, and the involvment of these fluids in the formation of lode gold deposits has been demonstrated by many authors (see Taylor (1979), Ohmoto (1986), Kerrich (1987), and Nesbitt (in press) for examples). Recently Nesbitt (1988), and Nesbitt and Muehlenbachs (1989) have demonstrated, through the consideration of crustal permeabilities, that meteoric waters may penetrate to great depths in the crust (possibly to the brittle-ductile transition zone), and in this process become greatly enriched in ¹⁸O. Concomitant positive δ D-shifts are generally not as marked (depending on w/r ratio), due to the low hydrogen content of most rocks. Integrated process models involving the deep circulation and isotopic-chemical evolution of meteoric waters, and their involvment in lode gold-forming systems have been given by Nesbitt (1988; and, in press).

As stated, meteoric water at Athabasca Pass has a δD value of ca. -135 per mil. The equilibrium $\delta^{18}O$ value in this area is thus about -18 per mil (Craig, 1961; Dansgaard, 1964). Utilizing the $\delta^{18}O$ (fluid) value of ca. +8 per mil calculated earlier, vein quartz at Athbasca Pass records a $\delta^{18}O$ (fluid) shift of ca. +26 per mil. $\delta^{18}O$ (fluid) shifts of this magnitude can be produced through O-isotope exchange reactions between fluid and wall-rocks under conditions of T>300°C, and with w/r ratios on the order of 0.1 (see Field and Fifarek (1985)). Additionally, under such conditions,

accompanying δD (fluid) shifts would be +15 to +25 per mil; approximately equal to those observed in the Athabasca Pass vein fluids (i.e. ≤20 ‰). A curve depicting the evolution of the fluids which deposited the gold-quartz veins at Athabasca Pass (using T≈ 350°C and a w/r ratio of ca. 0.1), is given in Fig. 4-3. Nesbitt and Muehlenbachs (1989) have documented meteoric water O- and H-isotope shifts of this magnitude, under similar conditions, for a large number of quartz vein ± gold occurrences in the Canadian Cordillera.

Thus it is evident that a fluid source for the gold-quartz veins at Athabasca Pass is most simply, and most completely explained by invoking the involvement of evolved meteoric waters. Contribution from a metamorphic fluid is not required. These findings may be more broadly applicable to the generation of mesothermal gold-quartz deposits in low-grade meta-sedimentary domains elsewhere in the world.

Implications for metamorphogenic lode gold deposits

A metamorphic origin for the fluids which generated mesothermal, turbidite hosted (ca. greywacke or slate-belt hosted) gold-quartz veins in low-grade meta-sedimentary domains has been proposed by numerous authors in recent years (e.g. Norris et.al., 1976; Graves and Zentilli, 1982; Paterson, 1982, 1986; Cox et.al., 1986; Mawer, 1986; Sandiford and Keays, 1986; Tomkinson, 1988: Kontak and Smith, 1989; Seccombe and Hicks, 1989; McKeag et. al., 1989). In general these fluids are considered to have been generated through prograde metamorphic devolitalization reactions and deformational processes. Ore-and gangue constituents are mobilized by ascending fluids through fluid - wall-rock interactions, and the fluids are focussed into structural conduits (e.g. fault zones, dialatent bedding-planes,

fold hinges) where mineral deposition takes place. Field relationships suggesting a lack of temporally-related igneous intrusions, gold distribution related to regional metamorphic isograds, the limited development of wall-rock alteration, and fluid overpressuring (hydrofracturing) during vein formation are well documented. Fluid inclusion studies reveal vein-filling at paleo-temperatures of *ca.* 250 to 350°C, from moderately saline fluids (≤8 wt % NaCl eq.). The presence of high density CO₂ ± CH₄ suggests ambient pressures of ≥1000 bars. Augmented by only cursory S- or O-isotope studies and/or trace-element distribution data, these factors are generally considered to be indicative of metamorphogenic processes in lode gold formation. H-isotope analyses however, are conspicuously absent from all of the studies cited above.

With respect to the auriferous quartz lodes at Athabasca Pass, field, petrographic, and fluid inclusion relationships essentially identical to those documented in the studies cited above have been documented. Additionally, mineralogical and trace-element, and S-, and O-isotope evidence supports the relatively local derivation of the vein-filling constituents, and suggest broad chemical- thermal- and isotopic-equilibrium between the country rocks and the mineralizing fluids at Athabasca Pass. Such lines of evidence are all suggestive of a metamorphogenic origin for the Athabasca Pass lodes, and based upon them alone, such an interpretation would seem inescapable. However, as stated, a metamorphogenic origin is clearly incompatible with the H-isotope data presented herein. The formation of such lodes can only be reconciled by a fluid which involved meteoric water. As outlined by Nesbitt (1988; and, in press) models for mesothermal lode gold mineralization involving evolved

meteoric waters versus metamorphic fluids are very similar. The meteoric water model however is consistent with all of the data generally cited in support of the metamorphic model, as well as being in agreement with the δD data.

Concluding statment

It is now apparent that an accurate evaluation of the potential fluid sources contributing to the formation of mesothermal, turbidite hosted lode gold deposits should not be performed without first aquiring H-isotope data. In the absence of H-isotope data, the involvment of an evolved meteoric water source-fluid cannot be ruled out. Indeed, considering the paucity of H-isotope data pertaining to turbidite hosted gold lodes, such involvment may be much more common than presently recognized. The application of fully integrated O-H-isotope studies, and the critical role of H-isotope data in the evaluation of potential source fluids for turbidite hosted lode gold deposits, is again re-emphasized.

Table 4-1. Oxygen and hydrogen isotope data for mineralized and unmineralized quartz veins and McNaughton Formation host rocks at Athabasca Pass, B.C., Canada.

| | | | | |
|--------------------------|--------------------------------|--|--|---------------------------------|
| Description & Sample No. | δ ¹⁸ O (‰, SMOW) | Average £ ¹⁸ O (‰, SMOW) | δ ¹⁸ Ο _{Η2} Ο (‰, SMOW) | δD _{H2} O (‰, SMOW) |
| Quartzites | | | | |
| PBG-A1 | 12.8 | | | |
| PBG-A4 | 13.5 | | | |
| PBG-A5 | 13.5 | | 8.1 ± 0.9 [#] | |
| PB-632 | 12.9 | 12.9 | 8.1 ± 0.9" | |
| PB-633 | 12.0 | | | |
| PBM-A1 | 13.3 | | | |
| PBM-A2 | 12.4 | | | |
| Pelites | _ | | | |
| PB-A11-P1 | 10.4 | | | |
| PB-A11-P2 | 9.7 | | | |
| PB-A16-P1 | 10.5 | 40.0 | 8.1 ± 0.5 [#] | |
| PB-A17-P1 | 10.0 | 10.0 | 6.1 ± 0.5 | |
| PB-A18-P1 | 10.2 | | | |
| KEA-P1 | 9.5 | | | |
| Bedding-parallel quartz | | | | |
| vein (Type I qtz.) | = | | | -124 |
| PB-A4-1 | 13.5 | | | -124 |
| PB-A8-1 | 14.3 | | | -105 |
| PB-A11-1 PB-A16-1 | 14.6 13.3 | | | -118 |
| PB-A17-1 | 15.0 | | | -114 |
| PB-A18-1 | 14.9 | | | • • • • |
| KEA-1 | 14.8 | | | |
| Gam-1 | 14.8 | | | |
| | | | | |
| Bedding-parallel quartz | <u> </u> | 14.2 | 8.2 ± 1.4 | |
| vein (Type III qtz.) | اميد | | | -109 |
| PB-A4-3 | 14.4 | | | -109 |
| PB-A8-3 | 14.3 13.0 | | | |
| PB-A11-3 | 1 | | | |
| PB-A15-3 | 13.6 15.0 | | | |
| PB-A16-3 PB-A18-3 | 14.0 | | | |
| | 14.0 | | | |
| Discordant veins | 14.7 | | | |
| PBG-A5-V1 | 14.7 | | | |
| PB-V632 | 14.7 | | | |
| PB-V633 | 14.6 | | | |

[#] Calculated using a temperature of 370°C derived from quartz - muscovite geothermometry.

^{*} Calculated using a temperature of 325°C derived from fluid inclusion and sulfur isotope geothermometry

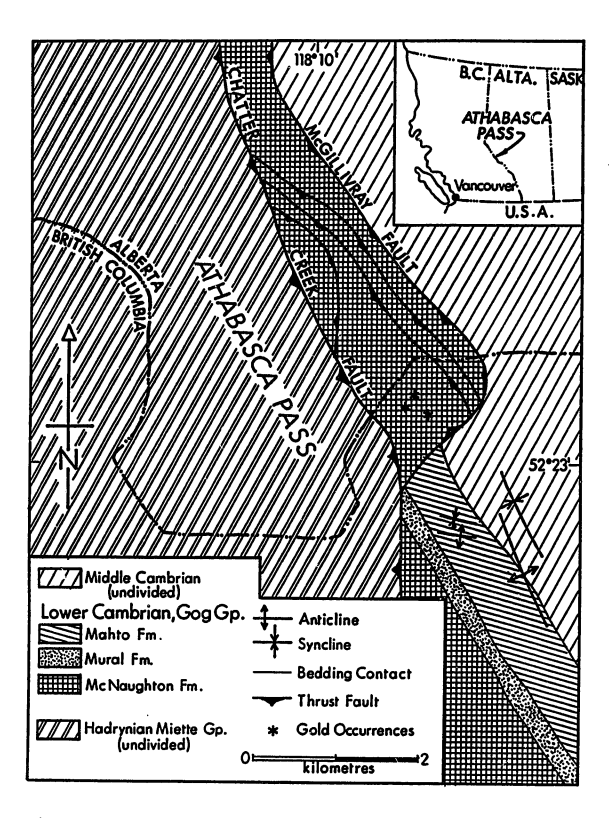


Figure 4-1. Location and general geology of the Athabasca Pass (Modified after Mountjoy and Price, in prep.). Gold occurrences represent clusters of mineralized veins.

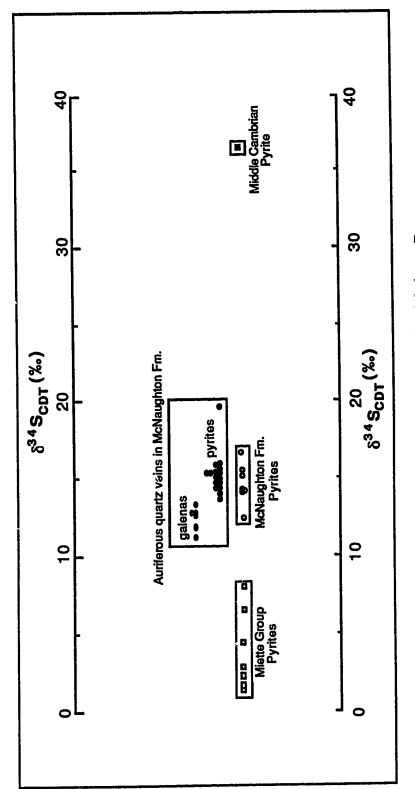


Figure 4-2. δ^{34} S values for sulfides from the various rock types in the Athabasca Pass.

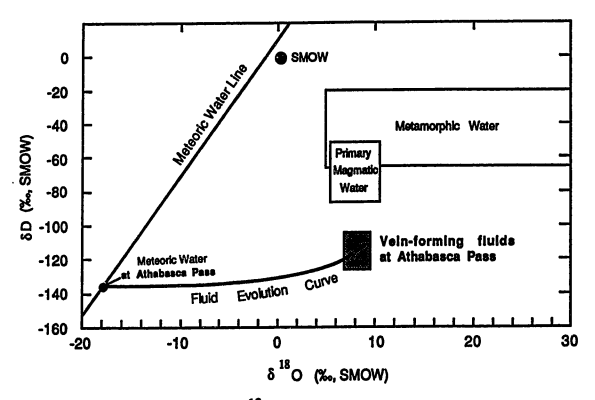


Figure 4-3. Plot of δD vs. $\delta^{18}O$ of the hydrothermal fluids which generated auriferous quartz veins in the McNaughton Formation at Athabasca Pass. Fluid evolution curve depicts the O- and H-isotope shift of meteoric waters due to water - rock interactions at T=350°C and water to rock ratios of ca. 0.1.

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CHAPTER 5

Concluding Statement

In the preceeding chapters, it has been demonstrated that the auriferous veins at Athabasca Pass are typical cases of mesothermal, gold-quartz mineralization in low-grade meta-sedimentary domains. It is evident that a combination of regional- and local-scale controls have contributed to the genesis of these gold-bearing veins. All of these controls will not be reiterated here. However, because of its ubiquitious nature, the role of meteoric water, its involvment in, and implications for, lode gold formation in the Main Ranges of the Rocky Mountains will here be considered further.

As discussed, (Chapters 3 and 4), the marked chemical-isotopic evolution of meteoric waters (i.e. addition of dissolved salts, CO₂, metal-complexes etc.) must take place prior to/during mesothermal gold mineralization. Such evolution is the result of fluid - rock interactions, at moderately deep crustal levels, and moderate water to rock ratios. The structural, mineralogical and temporal relationships at Athabasca Pass place some constraints on these processes in the Main Ranges.

It has been established (Chapter 2) that the formation of the Athabasca Pass lodes post-dated both penetrative deformation and peak metamorphism in the Main Ranges. Hence, two major structural-metamorphic events pre-dated mineralization, namely 1) the formation of the large, pre-metamorphic thrust faults which underlie the Main Ranges, and 2) the pre- to syn-metamorphic rise of major antiformal culminations such as the Porcupine Creek Anticlinorium. These earlier events are

considered important as they would have provided a regional scale plumbing system for the pre-metamorphic influx, and subsequent synto post-metamorphic chemical (thermal-isotopic) evolution of meteoric waters prior to gold mineralization. During these events, meteoric waters could have penetrated to intermediate crustal levels. Localized equilibration of these fluids with their wall-rocks, and the scavenging of vein components could have taken place as metamorphism progressed. The structural initiation of fluid conduits could also have been induced at this time.

Post-dating metamorphism, structures associated with the development of late, out-of -sequence thrust faults such as the Chatter Creek thrust (i.e. dilatant bedding planes, en echelon tension fractures, incipient thrust faults) would have provided localized reservoirs for the previously evolved fluids. Other local controls, such as the availability of gold and sulfur, the presence of effective chemical-structural trapping mechanisms, and the existence of temperature -pressure - fO_2 - pH etc. conditions conducive to gold precipitation would determine whether or not those veins were auriferous.

Thus, in the broadest sense, the involvment of meteoric waters in lode gold genesis at Athabasca Pass suggests that gold lodes in the Main Ranges may be more widespread than presently recognized. The stratigraphic - structural - lithochemical - and mineralogical controls on gold mineralization at Athabasca Pass, which have been recognized during the course of this study, provide some discriminating guide parameters which may aid in the future exploration for other gold-quartz lodes in the Main Ranges and elsewhere in the Canadian Rocky Mountains.

APPENDICES

Appendix 1. Results of ICPES multielement analyses of various rock types from the Athabasca Pass

| Unmineralized 10 02 15 4 3 41 4 | Sample # | Classification | Au | Ag | As | Pb | Zn | Cu | Ва | Fe(%) |
|--|----------|----------------------------|-------------|--------------|----------------------|------------|------|-----|----------|-------|
| quartzite/conglomerate '80 <0.2 <5 <2 2 3 7 7 15 0.2 <5 <2 2 3 7 7 15 0.2 <5 5 2 2 3 7 7 15 0.2 <5 5 2 2 3 7 7 15 0.2 <5 5 2 4 2 2 3 7 1 | 720 | lominariland | 01. | 0.2 | 15 | 4 | ო | 7 | ક્ષ | 029 |
| Unmineralized *5 < 0.2 < 5 < 2 3 7 7 7 5 | 180 | | 8 | <0.2 | ~ | % | ત | က | ଷ | 0.34 |
| ## 175 | 084 | qualizitate wilgibilielate | | 50.5 | ۸. | 4 | ო | N | ₹ | 0.31 |
| Unmineralized '85 <0.2 <5 < 9 < 4 < 2 < 0.2 <5 < 9 < 4 < 2 < 0.2 <5 < 2 < 4 < 2 < 0.2 <5 < 0.2 <5 < 3 < 1 < 0.2 <5 < 0.2 <5 < 0.2 <5 < 0.2 <5 < 0.3 <1 < 0.2 <1 < 0.2 <5 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 < 0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 <0.3 <1 < | 086 | | . | 0.2 | , S | Q | Q | က | <u>용</u> | 0.48 |
| Unmineralized **5 < 0.2 < 5 | 105 | | 15 | 0.2 | ß | ω | 4 | ત | ₹ | 2.46 |
| Unmineralized *65 < 2 3 1 *10 0.2 < 5 2 3 1 Unmineralized *85 0.2 < 5 8 5 < 1 Pellia pellia *20 < 0.2 < 5 6 6 2 2 1 Highly anomalous 26.2 0.4 20 58 4 2 8 bedding-parallel 124.8 0.4 20 42 4 3 veins 41.9 0.2 15 10 3 3 Moderately anomalous 570 0.2 < 5 21 147 5 bedding-parallel *480 0.2 < 5 21 147 5 Brecciated pelitet *17.0 < 0.2 2 0 18 4 2 Brecciated pelitet *480 0.2 2 0 18 4 2 Brecciated pelitet *45 0.2 20 8 4 1 3 Discordant vein *7 < 0.2 2 0 6 6 4 1 2 Brecciated pelitet *45 0.2 20 8 6 4 1 3 Discordant vein *45 0.2 5 16 20 3 | 603 | | ស្ព | <0.5 | ~ | 01 | 4 | ત | 8 | 0.39 |
| Unmineralized '85 0.2 <5 2 4 3 Unmineralized '20 0.2 <5 2 2 1 Pelite '20 <0.2 <5 2 2 1 Highly anomatous 26.2 0.4 20 58 4 2 bedding-parallel 573.7 8.8 20 42 4 3 Veins 41.9 0.2 15 10 3 3 Moderately anomatous '570 0.2 <5 24 1 3 bedding-parallel '480 0.2 <5 21 147 5 Brecciated pelitett '170 <0.2 20 6 4 2 Brecciated pelitett '170 <0.2 20 6 4 2 Discordant vein '170 <0.2 20 6 4 2 Discordant vein '170 <0.2 20 6 6 4 1 3 A 20 22 20 18 4 2 Brecciated pelitett '170 <0.2 20 6 6 4 2 Discordant vein '170 <0.2 20 6 6 4 2 Brecciated pelitett '170 <0.2 20 18 4 2 Discordant vein '170 <0.2 5 16 20 3 | 504 | | . | <0.2 | v Q | 81 | ო | - | 8 | 0.35 |
| Unmineralized "85 0.2 <5 8 5 <1 9 1 | 629 | | 유 | 05 | , S | 8 | 4 | က | ω | 0.54 |
| Highly anomalous 26.2 0.4 20 58 4 2 10 3 41.9 condant veins 27.3.7 8.8 20 42 4 3 3 3 4 2 15.8 0.2 15 10 3 3 3 15.8 0.2 15 10 3 3 3 15.8 0.2 15 10 2 15 10 2 15 10 2 15 10 10 10 10 10 10 10 10 10 10 10 10 10 | 9 | | 104 | 0 | , rc | α | r. | 7 | Ю | 278 |
| Highly anomalous 26.2 | 0 40 | Onmineralized | 3 § | , i v | , rê | 25 | ۰ ۵ | - | R | 0.35 |
| Highly anomalous 26.2 0.4 20 58 4 2 4 3 4 9 5 4 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 | 601 | eilled | Q. S | <0.2 2.02 | .6 | , N | 5 | ო | 47 | 3.69 |
| bedding-parallel 124.8 0.4 20 42 4 3 veins 573.7 8.8 20 42 3 3 41.9 0.2 15 10 3 3 71.5 0.4 20 28 4 2 71.5 0.4 20 28 4 2 71.5 0.2 <5 6 6 4 2 Noderately anomalous 570 0.2 <5 10 2 veins 540 0.2 5 24 1 3 Brecciated pelitett 17.0 <0.2 20 6 4 2 bedding-parallel vein 480 0.2 20 18 4 2 Brecciated pelitett 17.0 <0.2 20 6 4 1 Brecciated pelitett 17.0 <0.2 20 6 4 1 Brecciated pelitett 29 0.2 15 28 4 2 Discordant vein 45 <0.2 5 16 20 3 | 1000 | | 26.2 | 0.4 | 20 | 20 | 4 | N | 245 | 50.1 |
| Moderately anomalous *573.7 8.8 20 42 3 3 Woderately anomalous *570 0.2 *5 6 4 2 Pedding-parallel *490 0.2 <5 | 7 2 | <u>0</u> | a 701 | 7 | 0 | 4 | 4 | ო | 8 | 127 |
| Moderately anomalous *570 0.2 5 6 4 2 2 71.5 0.4 20 28 4 2 2 71.5 0.4 20 28 4 2 2 71.5 0.2 <5 21 14.7 5 bedding-parallel *480 0.2 <5 10 2 8 4 2 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 | 520 | ng-paramen | 679.7 | ά | 0 | 3 | . 64 | ო | 250 | 1.0 |
| Moderately anomalous *570 0.2 <5 0.4 20 28 4 2 71.5 0.4 20 28 4 2 71.5 0.4 20 28 4 2 71.5 0.2 <5 21 14.7 5 5 bedding-parallel *480 0.2 <5 10 2 8 4 1 3 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 | 926 | | 2.5.4 | 3 6 |) Y | į | , e | ო | 180 | 1.79 |
| Moderately anomalous *570 0.2 <5 21 147 5 5 5 6 6 2 8 4 2 2 6 6 1 147 5 5 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 | 8// | | - 4 - 4 | 9 6 | , r | | 4 | N | 160 | 1.86 |
| Moderately anomalous *570 0.2 <5 21 147 5 Moderately anomalous *570 0.2 <5 10 2 3 bedding-parallel *480 0.2 5 24 1 3 veins *440 0.2 20 18 4 2 Brecciated pelitet *172 <0.2 20 6 4 1 bedding-parallel vein *45 <0.2 5 16 20 3 | 0 0 | | 5 7 5 4 | 3 6 | 0 0 | , g | 4 | N | K | 0.78 |
| Moderately anomalous *570 0.2 <5 10 2 3 bedding-parallel *480 0.2 5 24 1 3 veins *440 0.2 20 18 4 2 Brecciated pelitet *172 <0.2 20 6 4 1 bedding-parallel vein *45 <0.2 5 16 20 3 | 0 0 | | | ; c |) | 7 | 147 | ις. | ম | 0.58 |
| Moderately anomalous *570 0.2 <5 10 2 3 bedding-parallel *480 0.2 5 24 1 3 veins *29 0.2 15 28 4 2 veins *440 0.2 20 18 4 2 Brecciated pelitet *172 <0.2 20 6 4 1 bedding-parallel vein *45 <0.2 5 16 20 3 Discordant vein *1 <0.2 5 16 20 3 | n n | | D. | 40.K | ? | i | • | 1 | i | |
| bedding-parallel *480 0.2 5 24 1 3 4 2 veins *440 0.2 20 18 4 2 8 4 2 8 4 2 8 4 2 8 4 2 8 4 2 8 4 2 8 4 2 8 4 2 8 4 2 8 4 2 8 8 8 8 | 597 | Moderately anomalous | .570 | 0.2 | , C | 9 | 8 | ო | 8 | 0.75 |
| Velins 2.9 0.2 15 28 4 2 Velins *440 0.2 20 18 4 2 *A40 0.2 20 18 4 2 Brecciated pelite± *170 <0.2 | 767 | modeling porollol | 480 | 0 | ល | 2 | - | ო | 4 | 0.74 |
| Brecciated pelite± *170 <0.2 20 18 4 2 Brecciated pelite± *170 <0.2 20 6 4 1 bedding-parallel vein *45 <0.2 5 19 125 1 Discordant vein *1 <0.2 5 16 20 3 | - CO 0 | Decomptantal | 6 | 0 | 15 | 88 | 4 | 01 | 185 | 1.60 |
| Brecciated pelite± *170 <0.2 20 6 4 bedding-parallel vein *45 <0.2 5 19 125 Discordant vein *1 <0.2 5 16 20 | 907 | CHIEA | 4 | 0.5 | 20 | 8 | 4 | N | 8 | 1.17 |
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| bedding-parallel vein *45 <0.2 5 19 125 Discordant vein *1 <0.2 5 16 20 | 806 | Brecciated pelitet | 2 | 7.0V | 2 | > ! | • | . , | 3 1 | |
| Discordant vein *1 <0.2 5 16 20 | 009 | bedding-parallel vein | . 45 | <0.5 | ស | <u>6</u> | 125 | - | Ð | 97.1 |
| Discordam ven | 803 | | | <0.2 | ß | 5 | 8 | ო | સ | 0.79 |
| 7 t 65 Z0 8. | 632 | Discordant vein | . iz | 05 | Ċ, | 4 | 81 | က | 32 | 0.42 |

All data recorded in parts per million (ppm) except where indicated.
• indicates parts per billion (ppb).
Fe = % total iron.

Appendix 2. Instrumental neutron activation elemental analysis of vein and country-rock pyrite separates from the Athabasca Pass area: Sample descriptions and results.

| Sample number | Description |
|---------------|--|
| PB-A1 | Pyrite from auriferous quartz vein on McGillivray Ridge. |
| PB-A4 | Pyrite from auriferous quartz vein on McGillivray Ridge. |
| PB-A11 | Pyrite from auriferous quartz vein on McGillivray Ridge. |
| PB-A16 | Pyrite from auriferous quartz vein on McGillivray Ridge. |
| PB-A16 | Pyrite contained within pelite fragment from auriferous quarty vein on McGillivray Ridge. |
| PB-A5 | Metamorphic pyrite contained within McNaughton Formation quartzite froก McGillivray Ridge. |
| PB-A6 | Metamorphic pyrite contained within McNaughton Formation quartzite from McGilllvray Ridge. |
| MB17-A26 | Metamorphic pyrite contained within Miette Group grit from Mount Brown. |
| MB9-A24 | Metamorphic pyrite contained within Miette Group pelite from Mount Brown. |

Results

| 0 | | Element | |
|------------------|-------------|-------------|-------------|
| Sample number | Au (ppm) | Ag (ppm) | As (ppm) |
| PB-A1 | 93.30 | <8 | 990 |
| PB-A4 | 16.30 | <12 | 1100 |
| PB-A11 | 551.00 | <13 | 740 |
| PB-A16 | 1.95 | <14 | 1800 |
| PB-A16 | 3.82 | <11 | 2200 |
| PB-A5 | 2.02 | <9 | 80 |
| PB-A6 | 0.15 | <10 | 490 |
| MB17-A26 | <0.02 | <8 | 550 |
| MB9-A24 | 0.08 | <7 | 870 |

Analyses performed by Dr. E.L. Hoffman, Activation Laboratories, Branford, Ontario.

Appendix 3. Gold content of unmineralized, quartz-dominated McNaughton Fm. rock types of McGillivray Ridge, Athabasca Pass, B.C.

| Sample # | Quartz ¹ (%) | White Mica ¹ (%) | K-Feldspar ¹ (%) | Au ² (ppb) |
|----------|----------------------------|--------------------------------|--------------------------------|--------------------------|
| 026 | 85-90 | 10-15 | 3-5 | 10 |
| 027 | 95-99 | ර ් | <1 | 15 |
| 028 | 80-85 | 15-20 | <3 | <5 |
| 029 | 90-95 | 5-10 | ્ લ | 5 |
| 030 | 65-70 | 30-35 | 3-5 | 5 |
| 031 | 65-70 | 30-35 | 3-5 | 10 |
| 032 | 90-95 | 5-10 | <3 | <5 |
| 033 | 45-50 | 45-50 | હ્યું | <5 |
| 034 | 90-95 | 5-10 | 3-5 | <5 |
| 035 | 95-99 | <5 | <1 | 20 |
| 036 | 65-70 | 30-35 | ⊲3 | <5 |
| 037 | 50-60 | 40-50 | 43 | <5 |
| 038 | 85-90 | 10-15 | ઢ | <5 |
| 039 | 75-80 | 20-25 | 5-10 | <5 |
| 040 | 90-95 | 5-10 | ⊲3 | <5 |
| 041 | 75-80 | 20-25 | 3-5 | 15 |
| 042 | 75-80 | 20-25 | 5-10 | 5 |
| 043 | 85-90 | 10-15 | _ | 10 |
| 044 | 75-80 | 20-25 | 5-10 | 35 |
| 045 | 90-95 | 5-10 | હ | 5 |
| 046 | 90-95 | 5-10 | ્ ક | <5 |
| 047 | 90-95 | 5-10 | 3-5 | 5 |
| 048 | 85-90 | 10-15 | 3-5 | 5 |
| 049 | 75-80 | 20-25 | 5-7 | 10 |
| 050 | 80-85 | 15-20 | 3-5 | 10 |
| 051 | 85-90 | 10-15 | 3-5 | 5 |
| 052 | 90-95 | 5-10 | ⊲3 | <5 |
| 053 | 85-90 | 10-15 | 5-7 | <5 |
| 054 | 85-90 | 10-15 | <3 | 15 |
| 055 | 90-95 | 5-10 | <3 - 10 | 10 |
| 076 | 70-75 | 25-30 | 5-10 | <5 |
| 077 | 70-75 | 25-30 | প্র | <5 |
| 078 | 90-95 | 5-10 | 3-5 | <5 |
| 079 | 85-90 | 10-15 | <1 | 75 |
| | | (Cont) | | |

Appendix 3. (con't)

| Appendix 5. (55 | | | | | | |
|-----------------|----------------------------|--------------------------------|--------------------------------|--------------------------|--|--|
| Sample # | Quartz ¹ (%) | White Mica ¹ (%) | K-Feldspar ¹ (%) | Au ² (ppb) | | |
| 080 | 75-80 | 20-25 | 5-10 | <5 | | |
| 081 | 80-85 | 15-20 | <3 | 80 | | |
| 082 | 70-75 | 25-30 | ⊲ | 10 | | |
| 083 | 95-99 | ර ් | <1 | 10 | | |
| 084 | 90-95 | 5-10 | 3-5 | <5 | | |
| 085 | 80-85 | 15-20 | 3-5 | 15 | | |
| 086 | 80-85 | 15-20 | 3-5 | 40 | | |
| 087 | 80-85 | 15-20 | <3 | 10 | | |
| 088 | 80-85 | 15-20 | 3-5 | 5 | | |
| 089 | 90-95 | 5-10 | ⊲ડ | <5 | | |
| 090 | 60-65 | 35-40 | 5-7 | 25 | | |
| 091 | 85-90 | 10-15 | 3-5 | 5 | | |
| 092 | 85-90 | 10-15 | ઢ | 5 | | |
| 093 | 90-95 | 5-10 | <3 | 5 | | |
| 094 | 75-80 | 20-25 | ઢ | 10 | | |
| 096 | 70-75 | 25-30 | 10-15 | 20 | | |
| 097 | 50-60 | 40-50 | ઢ | 25 | | |
| 098 | 85-90 | 10-15 | 3-5 | 30 | | |
| 099 | 70-75 | 25-30 | ⋖3 | 25 | | |
| 100 | 80 -85 | 15-20 | 3-5 | 115 | | |
| 101 | 75-80 | 20-25 | 10-15 | 10 | | |
| 102 | 75-80 | 20-25 | 5-7 | 10 | | |
| 103 | 80-90 | 10-20 | 3-5 | 10 | | |
| 104 | 90-95 | 5-10 | <3 | 20 | | |
| 105 | 85-90 | 10-15 | 3-5 | 15 | | |
| 106 | 75-80 | 20-25 | 3-5 | 30 | | |

¹ Estimated modal percentage of mineral observed in hand specimen.

² Determined by atomic absorbtion spectrometry, detection limit 5 ppb Au.

Appendix 4. Microtherometric data for Type A fluid inclusions in quartz from auriferous velns on McGillivray Ridge, Athabasca Pass, B.C.

| Sample # | Quartz type | Tm CO2 (°C) | Tm Ica (°C) | Wt. % NaCleq. (Tm lce) | Tm Clath. (°C) | Wt. % NaCl eq. (Tm Clath.) | Th CO2 (°C) | T decrep. (°C) | Th total (°C) |
|-------------|----------------|----------------|----------------|---------------------------|-------------------|-------------------------------|-------------|-------------------|---------------|
| PB-A2 | 1 | | | • | 7.9 | 4.1 | • | • | 360 |
| | • | • | - | • | 7.9 | 4.1 | • | • | 350 |
| | | • | -5.8 | 8.9 | • | • | • | • | 251 |
| | | | -4.3 | 6.9 | • | - | • | • | 205 |
| | | • | -4.3 | 6.9 | • | • | - | • | 280 |
| | | • | -4.1 | 6.6 | 8.6 | 2.8 | • | - | 187 |
| | | • | -3.8 | 6.1 | 8.8 | 2.4 | • | • | 187 |
| | | - | • | • | 8.6 | 2.8 | • | • | 197 |
| | | • | -4.5 | 7.2 | 8.6 | 2.8 | • | - | 207 |
| | | • | -4.8 | 7.6 | 8.6 | 2.8 | • | • | 215 |
| | | • | -3.8 | 6.1 | • | • | • | • | 233 |
| | | • | -4.3 | 6.9 | • | • | • | • | 253 |
| | | • | - | • | • | 3.0 | • | • | 266 |
| | | • | -3.3 | 5.4 | 8.5 | 3.0 4.5 | • | 001 | 194 |
| | | • | • | • - | 7.7 | 4.5 | • | 261 | - 320 |
| PB-A3 | 1 | • | -5.6 | 8.7 | • | - | • | • | 300 |
| | | • | -3.8 | 6.1 | • | - | - | • | 289 |
| | | • | -3.6 | 5.8 | • | - | • | • | 304 |
| | | • | -5.5 | 8.5 | • | - | - | • | 235 |
| PB-A4 | ı | • | -4.2 | 6.7 | • | - | • | _ | 206 |
| | | • | -5.1 | 8.0 | • | • | • | _ | 227 |
| | | • | -5.1 | 8.0 | • | - | : | • | 225 |
| | | • | -5.1 | 8.0 | • | - | • | • | 222 |
| | | • | • | • | 8 | 3.9 | <u>.</u> | | |
| | | • | -4.2 | 6.7 | | • | • | - | 225 |
| | | • | -3.8 | 6.1 | • | • | | • | 244 |
| | | • | -5.1 | 8.0 | - | | _ | - | 231 |
| | | • | -4.5 | 7.2 | _ | • | • | | 251 |
| | | • | -4.1 | 6.6 | _ | _ | - | 315 | 282 |
| | | • | • | - | _ | - | • | 315 | |
| | | • | - | | _ | • | - | • | 307 |
| | | - | -5.0 | 7.9 10.5 | _ | • | • | • | 315 |
| | | • | -7.0 | | - | • | _ | | 283 |
| | | • | - -5.5 | 8.5 | • | • | | • | 287 |
| | 111 | - | | 6.5 | • | • | - | • | 251 |
| | 1111 | • | • | - | • | • | • | • | 252 |
| | | - | -3.6 | 5.8 | 7.5 | 4.9 | • | • | 308 |
| | | - | -5.4 | 8.4 | • | • | • | • | 281 |
| | | _ | | 5.8 | 6.9 | 5.9 | • | • | 293 |
| | | _ | -3.6 | • | • | • | • | 215 | • |
| | | - | | • | • | • | • | 236 | • |
| | | _ | • | • | 7.6 | 4.7 | • | • | 285 |
| | | _ | -5.6 | 8.7 | 8.2 | 3.6 | 20.8 | - | • |
| | | - | -5.2 | | 8.3 | 3.4 | - | • | • |
| | | _ | -5.2 | | • | • | • | • | • |
| | | | -4.2 | | 8.1 | 3.8 | • | • | - |
| | | • | -4.7 | 7.4 | • | • | • | • | 253 |
| | | -58 | • | - | 8.3 | 3.4 | • | 177 | • |
| PB-A1 | 1 1 | | -5.2 | | 8.3 | 3.4 | • | 220 | • |
| | • | | -9.5 | - | 9.0 | 2.0 | 22.1 | | • |
| | | - | | | (Con't) | | | | |

Appendix 4.(con't)

| Sample # | Quartz type | Tm CO2 (°C) | Tm los (°C) | Wt. % NaCl eq. (Tm loe) | Tm Clath. | Wt. % NaCl eq. (Tm Clath.) | Th CO2 (°C) | T decrep. (°C) | Th total |
|---|----------------|----------------|----------------|----------------------------|------------|-------------------------------|-------------|-------------------|------------|
| PB-A11 | | | -5.0 | 7.9 | • | • | • | - | 241 |
| ייייייייייייייייייייייייייייייייייייייי | • | • | -4.3 | 6.9 | - | • | - | • | 257 |
| | | • | -4.8 | 7.6 | • | - | • | - | 264 |
| | | • | -4.8 | 7.6 | - | • | • | • | 305 |
| | | • | • | - | 8.8 | 2.4 | - | • | 163 |
| | | • | - | - | 8.1 | 3.8 | • | • | 292 |
| | | • | -4.8 | 7.6 | • | • | • | - | 259 |
| | | • | -6.0 | 9.2 | • | • | • | • | 276 |
| | | • | • | - | 3.6 | 11.2 | • | d | • |
| | | • | • | • | 8.1 | 3.8 | • | • | 266 |
| | | • | -5.0 | 7.9 | • | - | • | 248 | - |
| PB-A17 | 1 | - | -4.2 | 6.7 | • | • | - | 306 | • |
| | | • | -4.8 | 7.6 | - | • - | • | 282 | • |
| | | • | • | • | 7.6 | 4.7 | • | 176 | - |
| | | • | -3.8 | 6.1 | - | • | • | 196 | • |
| | | • | -4.0 | 6.4 | • | • | • | 228 | - 045 |
| | | • | -5.3 | 8.3 | - | • | • | • | 245 |
| | | - | | - | - | • | • | • | 250 |
| | | • | -5.5 | 8.5 | • | • | • | • | 252 |
| | | • | -5.7 | 8.8 | • | • | • | 045 | • |
| | | • | -5.6 | 8.7 | • | • | - | 245 | • |
| | | - | -5.6 | 8.7 | - | - | • | 234 | • 050 |
| | | - | -6.8 | 10.2 | • | - | • | • | 250 265 |
| | | • | -6.4 | 9.7 | - | - 5.9 | • | • | 265 265 |
| | | • | - | - | 6.9 | | • | • | 270 |
| | | • | -5.6 | 8.7 | 8.6 | 2.8 | • | - | 267 |
| | | - | -5.6 | 8.7 | - 0 1 | - 3.8 | • | 190 | - |
| | | • | -5.7 5.9 | 8.8 | 8.1 8.1 | 3.8 | - 15.7 | 190 | _ |
| | | • | -5.8 | 8.9 | 0. I - | 3.0 - | 13.7 | 190 | 172 |
| | | • | -4.8 | 7.6 | - | _ | _ | _ | 161 |
| PB-30 | l | • | • | - | 8.1 | 3.8 | _ | _ | 160 |
| | | • | - -5.3 | 8.3 | 8.4 | 3.2 | • | 190 | - |
| | | • | -5.3 -6.2 | 9.5 | 7.9 | 4.1 | • | 270 | - |
| PB-A31 | H. | - | -0.2 | 5.J - | 7.9 | 4.1 | • | - | 258 |
| 1 5-701 | - | _ | -5.9 | 9.1 | 7.8 | 4.3 | | • | 267 |
| | | | -4.1 | 6.6 | 8.7 | 2.6 | • | • | 280 |
| | | • | -2.1 | 3.5 | 8.7 | 2.6 | - | - | 223 |
| | | | -4.1 | 6.6 | 8.2 | 3.6 | - | • | 220 |
| | | • | -4.2 | 6.7 | 8.1 | 3.8 | - | - | - |
| | | • | -4.2 | 6.7 | 8.1 | 3.8 | • | • | - |
| | | - | -3.8 | 6.1 | - | • | - | - | 245 |
| | | • | • | • | - | - | - | - | 205 |
| | | • | -3.4 | 5.5 | • | - | - | • | 222 |
| | | • | -3.2 | 5.2 | - | - | • | - | 225 |
| | | • | -4.1 | 6.6 | - | - | - | - | 229 |
| | | • | -3.7 | 6.0 | - | • | - | • | 237 |
| | | | -4.1 | 6.6 | | | | | 221 |

Appendix 5. Sulfur-Isotope data for country rock and vein sulfide minerals from the Athabasca Pass area, B.C.

| | 34.0 | | | | | | | | |
|-----------|---------------|------------------|-------------------|--|--|--|--|--|--|
| Sample | Mineral | * Host lithology | δ ³⁴ S | | | | | | |
| number | analysed | | ‰, CDT | | | | | | |
| | COUNT | TRY ROCKS | | | | | | | |
| PBG-A4 | pyrite | McN quartzite | 15.3 | | | | | | |
| PBG-A5 | pyrite | McN quartzite | 12.4 | | | | | | |
| PBG-A5r | pyrite | McN quartzite | 11.4 | | | | | | |
| PBG-A5r | pyrite | McN quartzite | 12.2 | | | | | | |
| PBG-A5r | pyrite | McN quartzite | 12.4 | | | | | | |
| PBG-A5r | pyrite | McN quartzite | 11.8 | | | | | | |
| PBG-A5r | pyrite | McN quartzite | 11.8 | | | | | | |
| PBG-A6 | pyrite | McN quartzite | 14.1 | | | | | | |
| PB-A15 | pyrite | McN pelite | 14.2 | | | | | | |
| PBM-A20 | pyrite | McN pelite | 14.0 | | | | | | |
| PBP-A3 | pyrite | McN pelite | 16.9 | | | | | | |
| PBP-A7 | pyrite | McN pelite | 15.3 | | | | | | |
| PBE-A14 | pyrite | MC dolomite | 36.7 | | | | | | |
| MBG-A12a | pyrite | MG pelite | 3.0 | | | | | | |
| MBG-A12b | pyrite | MG pelite | 2.8 | | | | | | |
| MBG-A9 | pyrite | MG pelite | 1.1 | | | | | | |
| MBG-A16 | pyrite | MG pelite | 1.5 | | | | | | |
| MBG-A17 | pyrite | MG grit | 8.9 | | | | | | |
| MBG-A18 | pyrite | MG grit | 4.3 | | | | | | |
| MBG-A13 | pyrite | MG grit | 7.1 | | | | | | |
| VEIN SULF | IDES from the | MCNAUGHTON FORM | MATION | | | | | | |
| PBM-A1 | pyrite | Type III qtz. | 19.9 | | | | | | |
| PBM-A2a | galena | Type I qtz. | 12.8 | | | | | | |
| PBM-A2b | pyrite | Type I qtz. | 15.8 | | | | | | |
| PBM-A3 | galena | Type I qtz. | 11.4 | | | | | | |
| PBM-A4 | pyrite | Type III qtz. | 15.2 | | | | | | |
| PBM-A7 | pyrite | Type III qtz. | 16.6 | | | | | | |
| PBM-A8 | pyrite | Type III qtz. | 15.7 | | | | | | |
| PBM-A9 | galena | Type I qtz. | 12.8 | | | | | | |
| PBM-A10a | pyrite | Type III qtz. | 15.1 | | | | | | |
| PBM-A10b | pyrite | Type I qtz. | 16.3 | | | | | | |
| PBM-A11a | galena | Type I qtz. | 12.8 | | | | | | |
| PBM-A11b | pyrite | Type I qtz. | 14.4 | | | | | | |
| PBM-A11c | pyrite | Type III qtz. | 15.4 | | | | | | |
| PBM-A12a | pyrite | Type I qtz. | 14.7 | | | | | | |
| PBM-A12ar | pyrite | Type I qtz. | 15.3 | | | | | | |
| | (Co | on't) | | | | | | | |

Appendix 5 (con't).

| Sample number | Mineral analysed | *Host lithology | δ ³⁴ S ‰, CDT |
|---|--|---|--|
| PBM-A16a PBM-A16ar PBM-A16b PBM-A16b PBM-A16c PBM-A17a PBM-A17b PBM-A17b | pyrite pyrite pyrite galena galena galena pyrite pyrite pyrite | Type III qtz. Type III qtz. Type III qtz. Type I qtz. | 16.3 15.1 15.5 13.3 14.2 11.6 14.8 15.0 15.3 15.1 |
| PBM-A18a PBM-A18b PBM-A18br PBM-A19 PBM-A22 | pyrite pyrite pyrite pyrite pyrite | Type III qtz. | 15.0 15.3 16.3 13.7 |

*Lithologic Abbreviations:

McN = McNaughton Formation

MC = Middle Cambrian strata

MG = Hadrynian Miette Group strata

Type I qtz. = Paragenetically early bedding-parallel vein quartz
Type II qtz. = Paragenetically late bedding-parallel vein quartz

APPENDIX 6

Formulae Used for Fluid Inclusion and Stable Isotope Calculations

Chapter 2

Fluid inclusion study

Calculation of salinity for H2O - NaCl-bearing fluid inclusions (Potter et al., 1978):

$$W_s = 0.00 + 1.76958 \theta - 4.2384 \times 10^{-2} \theta^2 + 5.2778 \times 10^{-4} \theta^3 (\pm 0.028)$$

Where;

W_s = weight percent NaCl in solution.

 θ = the freezing point depression of ice, in °C.

Calculation of salinity for H₂O - NaCl - CO₂-bearing fluid inclusions (Bozzo et. al., 1973):

$$W_s = 15.52023 - 1.02342 \text{ (Tm}_{clath}) - 0.05286 \text{ (Tm}_{clath})^2$$

Where;

W_s = weight percent NaCl in solution.

Tm_{clath} = Final melting temperature of clathrate in °C.

Chapter 3

Sulfur isotope study

Sulfur isotope (pyrite - galena) geothermometry (Ohmoto and Rye, 1979):

$$T = ((1.01 \pm 0.04) \times 10^3) + \Delta^{1/2} (\pm 25^\circ)$$

Where;

T = temperature in °Kelvin.

 $\Delta = \delta^{34} S$ (pyrite) - $\delta^{34} S$ (galena).

Calculation of fluid (H2S) - pyrite sulfur isotope fractionation (Ohmoto and Rye, 1979):

$$\delta^{34} \text{S (pyrite)} - \delta^{34} \text{S (H}_2 \text{S)} = ((0.40 \pm 0.08) + \text{T}^2) \times 10^6$$

Where;

T = temperature in °Kelvin.

APPENDIX 6 (con't)

Oxygen isotope study

Oxygen isotope (quartz - white mica) geothermometry (Field and Fifarek, 1985):

T (°K) =
$$(0.98 \times 10^3) + (\Delta - 0.58)^{1/2}$$

Where;

$$\Delta = \delta^{18} O$$
 (quartz) - $\delta^{18} O$ (white mica).

Calculation of fluid - mineral oxygen isotope fractionation:

Quartz - H2O:

$$\delta^{18}$$
O (quartz) - δ^{18} O (H₂O) = ((3.34 x 10⁶) + T²)) - 3.31 (Matsuhisa *et. al.*, 1979).

Muscovite - H₂O:

$$\delta^{18}$$
O (muscovite) - δ^{18} O (H₂O) = ((2.38 x 10⁶) + T²)) - 3.89 (Friedman and O'Neil, 1976).

Where;

T = Temperature (°K)