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Résumé de l'article

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Petrology, geochemistry, and U-Pb (zircon) age of the quartz-feldspar porphyry dyke at the Lake George antimony mine, New Brunswick: implications for origin, emplacement process, and mineralization

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ABSTRACT

A steeply west-dipping, north-northwest-trending, biotite-bearing quartz-feldspar porphyry dyke is associated with the Lake George granodiorite, a cupola of the Pokiok Batholith. These Early Devonian intrusions are spatially and temporally related to W-Mo-Au and Sb-Au mineralization in the area. The porphyry dyke yielded an age of 420.8 +5.9/-4.0 Ma (U-Pb zircon) and is inferred to be coeval with or slightly older than the Lake George granodiorite (412 +5/-4 Ma, U-Pb zircon). The variably altered dyke contains subhedral to anhedral (rounded) quartz, plagioclase, and orthoclase phenocrysts (0.5 to 30 mm; 30 to 60 vol. %), and numerous angular to subrounded xenoliths of altered intrusion carapace and quenched dyke rocks, as well as local Kingsclear Formation metasedimentary rocks. The dyke is similar to the Lake George granodiorite in phenocryst population, volcanic arc (I-type) affinity, average Zr (157 ppm), TiO₂ (0.54 %), La/Yb ratio (13), and REE (129 ppm). The average Au content is lower in the dyke (20 ppb) than in the granodiorite (32 ppb), the abundance of gold is related to minor magmatic hydrothermal chalcopyrite and pyrrhotite (< 2 %). Based on textural evidence, including autobrecciation, coupled with rheological and thermal modeling calculations for the 5 m-wide dyke, it is suggested that fluidization (vapour exsolution) of the magma was key to its rapid emplacement as a quenched glass-gas mixture (tuffisite). The S abundance and S isotopic signature of the dyke, relative to Au and Cu abundance, support reduced I-type magmatic volatiles involved in dyke emplacement and cogenetic autometasomatic alteration.

RÉSUMÉ

Un dyke de porphyre quartzo-feldspathique renfermant de la biotite, orienté vers le nord-nord-ouest et s'inclinant abruptement vers l'ouest, est associé à la granodiorite du lac George, une coupole du magmatisme du batholite de Pokiok. Ces intrusions régionales du Dévonien précoce sont spatialement et temporellement apparentées à une minéralisation de W-Mo-Au et de Sb-Au dans le secteur. Le dyke de porphyre a accusé un âge de 420,8+5,9/-4,0 Ma (U-Pb sur zircon) et on le suppose du même âge ou légèrement plus âgé que la granodiorite du lac George (412 +5/-4 Ma, U-Pb sur zircon). Le dyke altéré de façon variable abrite des phénocristaux hypautomorphes à allotriomorphes (arrondis) de quartz, de plagioclase et d'orthoclase (0,5 à 30 mm, 30 à 60 % en vol.) ainsi que de nombreux xénolites angulaires à subsphérique d'une carapace d'intrusion altérée et de roches de dykes noyées, en plus de roches métasédimentaires locales de la Formation de Kingsclear. Le dyke s'avère analogue à la granodiorite du lac George du point de vue de la population de phénocristaux, de l'affinité avec l'arc insulaire (de type intrusif), de la concentration moyenne de Zr (157 ppm), du TiO₂ (0,54 %), du ratio de La/Yb (13) et des ÉTR (129 ppm). La teneur moyenne en Au est plus faible dans le dyke (20 parties par milliard) que dans la granodiorite (32 parties par milliard); son abondance est apparentée à la présence restreinte de chalcopyrite et de pyrrhotite hydrothermales magmatiques (< 2 %). Selon les indices texturaux, notamment l'autobréchification, conjugués à des calculs de modélisation rhéologique et thermale du dyke (5 m), on peut supposer que la fluidisation (exsolution à l'état de vapeur) du magma a contribué à sa mise en place rapide sous forme d'un mélange de verre-gaz (tuffisite) noyé. L'abondance de S et la signature isotopique du S du dyke, comparativement à l'abondance d'Au et de Cu, appuient la participation réduite d'éléments volatils magmatiques de type intrusif à la mise en place du dyke et à l'altération autométasomatique cogénétique.

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INTRODUCTION

The Lake George mine is located approximately 40 km west of Fredericton, New Brunswick (Fig. 1a) and was formerly the largest producer of antimony in North America. Interest in the area has recently been revived with the discovery of gold anomalies in drill cores (Morrissy 1991; Lentz et al. 2002a). These anomalies are thought to be associated with the nearby Lake George granodiorite stock (Seal et al. 1988), which is situated 500 m northwest of the Hibbard Shaft and 350 m below surface. A strongly fractured contact aureole around the stock likely provided conduits for mineralizing fluids bearing Sb, W, Mo, Au, and base metals. Tightly folded calcareous turbiditic rocks, metamorphosed to greenschist facies (Seal et al. 1987, 1988; Caron 1996) of the Silurian Kingsclear Group form the country rock around the Lake George granodiorite, which lies approximately 350 m below the surface along the southeastern margin of the Pokiok Batholith. A contact aureole surrounding the intrusion is indicated by the presence of biotite and cordierite porphyroblasts in the pelitic rocks (Seal et al. 1987; 1988; Caron 1996).

Prior to the intrusion of the granodiorite stock, steeply dipping east-trending lamprophyre dykes were discordantly emplaced into the Silurian Kingsclear Group. The focus of the present study is a northwest-trending quartz-feldspar porphyry (QFP) dyke which also intrudes the metasedimentary rocks, although its relationship to the granodioritic stock is uncertian (Morrissy and Ruitenberg 1980; Lentz *et al.* 2002b). The QFP dyke has a north-northwest orientation (Procyshyn and Morrissy 1990), dips steeply to the west (80°), and is 3 to 5 m wide (Seal *et al.* 1988). The dyke bifurcates with a separation of less than 25 m between the branches, which merge along strike and at depth. The QFP dyke crosscuts narrow east-west lamprophyre dykes (Lentz *et al.* 2002b) but is inferred to be intruded by the Lake George quartz-hornblende-plagioclase-phyric granodiorite stock at depth (Fig. 1).

The Pokiok Batholith (Fig. 1a) has been subdivided into five units (Ruitenberg and Fyffe 1982; Lutes 1987; Whalen 1993, Whalen et al. 1996). The Hartfield tonalite is the oldest unit (U-Pb titanite, 415 ± 1 Ma), followed by the Skiff Lake granite (U-Pb zircon, 409 ± 2 Ma) and the Hawkshaw granite (U-Pb titanite, 411 ± 1 Ma) (Bevier and Whalen 1990a, b; Whalen 1993). The Lake George granodiorite stock (U-Pb zircon 412 +5/-4 Ma; McLeod et al. 2003) is chemically and petrologically similar to the Hawkshaw granite of the Pokiok Batholith, although there are some mineralogical differences (Yang et al. 2002a, b). The Allandale granite is the youngest unit (U-Pb monazite, 402 ± 1 Ma) of the Pokiok Batholith. It is possible that the Hartfield, Skiff Lake, and Hawkshaw units represent a sequential fractionation and emplacement series, although the age of the Allandale granite is too young to be related to the main intrusive episodes of the batholith.

The purpose of this study is to examine the QFP dyke in the area and to determine its relationship to the magmatic activity in the region and timing relative to the two episodes of mineralization (W-Mo-Au and Sb-Au). Previous reports suggested

that the dyke predates the granodioritic intrusion (Morrissy and Ruitenberg 1980), which is consistent with it being locally crosscut and altered by both the Hibbard Sb vein and the intrusion-related gold-bearing quartz-scheelite-molybdenite veinlets (Seal et al. 1988), although crosscutting relationships between the dyke and the stock were not observed in this study. As the granodiorite stock is linked to the exocontact W-Mo-Au mineralization (Lentz et al. 2002a), information on the QFP dyke could enhance exploration efforts for Au (W-Mo) in the area, because (1) it could indicate the presence of larger granodiorite intrusions at depth and of magmatic systems of similar petrologic association, (2) intrinsic metal abundances may reveal the presence of a favourable magmatic hydrothermal system, and (3) aspects related to dyke emplacement indirectly provide evidence of the role of volatiles in emplacement that indirectly controls the distribution of exocontact Au-W-Mo mineralization in the region.

DYKE PETROLOGY

The rock-forming minerals in the QFP dyke at Lake George were first examined in polished drill-hole cores samples (Fig. 2) to observe variations in texture and mineralogy. [Note: Sample numbers indicate diamond drill hole number and depth. For simplicity, the m (for metres) is omitted from each sample number.] The metasedimentary host rocks in contact with the dyke have preserved no macroscopic evidence of contact hornfelsing associated with dyke emplacement. Samples from the host rocks contain 30 to 60 % anhedral quartz and subhedral and anhedral feldspar phenocrysts ranging from 5 to 12 mm in length (Fig. 2a). The dyke mineralogy generally consists of porphyritic and fine-grained quartz, plagioclase, and orthoclase with accessory biotite (< 2 mm in length), chlorite, muscovite, pyrite, pyrrhotite (< 3 %), chalcopyrite, calcite, titanite, and rutile. Ilmenite, zircon, monazite, and apatite are present but recognizable only at the sub-microscopic scale through SEM-EDS analysis. The samples differ in their degree of sericitization. Based on observed abundance of sericite and chlorite, sample DDH 81-25-19.2 is the least altered and sample DDH 79-6-195.5 is most altered. Sample DDH 81-21-181.1 (Fig. 2b) has a prominent fracture running the length of the core with strong sericitization. Sample DDH 81-26-63.4 (Fig. 2c) is interesting in that it contains a fragment that has a dark homogeneous microcrystalline matrix, morphologically and compositionally unlike the sedimentary host rocks (see Scratch et al. 1984). Sample DDH 79-6-210 has dark rounded xenoliths up to 10 mm in size (10 to 15 percent) (Fig. 2d) (see also DDH 81-26-63.4). These xenoliths resemble autobrecciated altered intrusive rocks, consistent with their similar compositions (see below).

Rounded quartz occurs generally in two size populations (see Fig. 3a); phenocrysts and phenoclasts (ave 0.7 mm) with a groundmass consisting of grains <0.1 mm in size. They measure up to 6 mm, although the average is about 0.7 mm. The larger grains commonly contain melt inclusions and secondary



Fig. 1 (a) General geology of the Pokiok Batholith and surrounding area near the Lake George antimony mine, New Brunswick. (b) Map showing the subsurface Lake George granodiorite stock in relation to the QFP dyke (modified after Seal *et al.* 1988 and Lentz *et al.* 2002a).



Fig. 2 Slab photographs of QFP dyke at Lake George: (a) sample DDH 81-25-19.2 contains quartz and feldspar phenocrysts; (b) sericitic alteration in proximity to the fracture in DDH 81-21-594 as the result of feldspar alteration. Sericitization was facilitated by the fracture that behaved as a conduit for fluid transportation; (c) sample DDH 81-26-62.8 contains a fragment with a dark matrix with intense graphic texture; (d) black xenoliths in DDH 79-6-210 are unique to this sample and are intensely sericitized feldspar.



Fig. 3 Photomicrographs showing two populations of quartz from sample DDH 79-6-210 in crossed polarized light (CPL). (a) larger subhedral quartz phenoclast in contact with smaller more numerous quartz grains. (b) large quartz phenocryst (on the left) with melt and fluid inclusions from sample DDH 79-6-210 (CPL) shows where the surrounding melt was trapped by the growing quartz along the grain boundary. The low birefringence bleb (on the right) represents the remnant of plagioclase phenocrysts pseudomorphed by sericitization.

fluid inclusions, and appear to have been a nucleation point for crystal growth (see Fig. 3b).

Feldspars are present in high proportions in every sample, but typically have undergone extensive alteration to sericite. Plagioclase crystals show various degrees of sericitization, but some display relict zoning and twinning (Fig. 4; Table 1). The zoning in most of the plagioclase crystals is irregular and the edges of some grains are rounded, which is evidence for periods of both crystal growth and resorption, and (or) possible abrasion. Plagioclase has inclusions of potassium feldspar, apatite, zircon, chlorite, Fe sulphides, and rutile. Granophyric (more wormy) to graphic (more cuneiform) intergrowths between feldspar and quartz are evident in three samples (DDH 79-6-174.3, DDH 81-26-63.4, and DDH83-1-37.5; Fig. 5). Chlorite appears to be a local alteration product of reddish-brown biotite (Fig. 6). Primary biotite crystals (4.4 wt. % TiO₂ by SEM-EDS analysis) contain inclusions of zircon, apatite, pyrite, albite, interlayered muscovite, and locally trellis-like rutile needles.

Sample magnetic susceptibility ranges from 1.1 to 13.5×10^{-4} (S.I. units) (see Table 2), typical of transitional reduced to oxidized magnetite-type magma (see Ishihara 1981), although most samples contain magnetic pyrrhotite rather than magnetite. Pyrrhotite and chalcopyrite are present in most samples, although probably a high temperature alteration feature, as



Fig. 4 Photomicrograph of complexly zoned plagioclase phenocrysts (CPL) in sample DDH 83-1-37.5 with variable degrees of sericitization (circled), and also showing signs of resorbtion in the irregular zoning. SEM-EDS analyses across the plagioclase are marked by black dots; the data are presented in Table 1.

	SiO ₂	Al_2O_3	CaO	Na ₂ O	K ₂ O	MgO	FeO	Total	Ab	An	Or
Sample DDH 81-26-63.4											
core	61.05	24.15	5.74	7.98	0.61	0.14	0.08	99.75	0.542	0.431	0.027
	61.23	23.92	5.34	7.97	0.80	0.12	0.10	99.48	0.554	0.410	0.037
	59.49	24.77	6.39	7.61	0.74	0.13	0.21	99.34	0.502	0.466	0.032
	60.19	24.71	6.18	7.74	0.64	0.21	0.05	99.72	0.516	0.456	0.028
	60.12	24.81	6.40	7.61	0.65	0.16	0.17	99.92	0.504	0.468	0.028
	58.32	25.48	7.32	6.93	0.47	0.12	0.21	98.85	0.452	0.528	0.020
	60.27	24.29	5.89	7.62	0.78	0.09	0.00	98.94	0.520	0.445	0.035
rim	59.77	24.97	6.57	7.46	0.58	0.15	0.20	99.70	0.494	0.481	0.025
Sample I	ODH 83	-1-37.5									
core	60.99	25.07	6.18	7.50	0.67	0.27	0.18	100.86	0.508	0.462	0.030
	61.16	25.03	6.37	7.48	0.70	0.28	0.14	101.16	0.499	0.470	0.031
	59.76	25.52	6.85	7.04	0.63	0.27	0.10	100.17	0.469	0.504	0.028
	58.99	24.79	6.55	7.25	0.67	0.28	0.00	98.53	0.486	0.485	0.030
	60.27	25.83	6.96	7.19	0.67	0.25	0.24	101.41	0.469	0.502	0.029
	58.48	26.81	8.19	6.58	0.49	0.30	0.15	101.00	0.412	0.567	0.020
	60.71	24.99	6.42	7.48	0.65	0.20	0.07	100.52	0.499	0.473	0.029
rim	61.29	24.96	6.33	7.79	0.60	0.31	0.12	101.40	0.513	0.461	0.026
Sample I	ODH 83	-1-37.5									
core	61.43	24.86	5.98	7.58	0.65	0.28	0.16	100.94	0.519	0.452	0.029
	60.41	25.56	6.80	6.96	0.66	0.24	0.00	100.63	0.467	0.504	0.029
	61.06	24.98	6.15	7.56	0.59	0.23	0.00	100.57	0.513	0.461	0.026
	59.21	25.59	7.21	6.97	0.58	0.23	0.14	99.93	0.455	0.520	0.025
rim	61.08	24.63	5.86	7.80	0.51	0.25	0.08	100.21	0.534	0.443	0.023

Table 1. SEM-EDS analyses of transects of zoned plagioclase grains from the QFP dyke at the Lake George mine



Fig. 5 Photomicrographs of graphic to granophyric intergrowth texture between quartz and plagioclase in sample DDH 81-26-63.4. (a) Granophyric texture surrounding a quartz phenocryst (CPL). (b) Graphic texture between two plagioclase phenocrysts (CPL).



Fig. 6 Photomicrograph of biotite in sample DDH 79-6-195 in plane polarized light that are partially altered to chlorite. The needle-like grains (circled) are rutile.

they coexist with alteration assemblages (as discussed later) and with biotite phenocrysts.

Euhedral titanite occurs as crystals less than 0.5 mm across, consistent with a relatively oxidized igneous system, although inconsistent with the reduced nature of the Lake George granodiorite. Apatite measures less than 0.1 mm in length and zircon is typically usually less than 100 microns, both as inclusions in biotite and plagioclase. As zircon and apatite apparently began to crystallize early, they may not be reliable thermometric indicators of dyke emplacement, but rather of maximum magma temperatures (Harrison and Watson 1984; Hanchar and Watson 2004).

Chlorite is present along fractures associated with calcite and, in some instances, contains intergrown bands of calcite. Narrow calcite veins (less than 0.2 mm wide) occupy microfractures, suggesting late paragenesis. Individual calcite grains contain pyrite inclusions. Some also contain minor pyrrhotite (<2 %) and chalcopyrite, typically tens of microns to 0.3 mm in size. Muscovite is commonly interstitial to larger plagioclase grains replacing K-feldspar phenocrysts; muscovite rosettes are common in altered parts of the dyke.

GEOCHRONOLOGY

Five single or double zircon fractions were analyzed (Table 3). On a concordia diagram (Fig. 7), they plot in a sub-concordant position (94 to 99.4 % concordant) with the exception of grains Zr3, which are only 74 % concordant with an apparent ²⁰⁷Pb/²⁰⁶Pb date of 757 Ma (see Table 3). The four remaining grains define a well-constrained upper intercept date of 420.8 +5.9/-4.0 Ma (MSWD=2.9) with a lower intercept date of 59 +85/-87 Ma. The date of 421 Ma is considered to be the age of intrusion of the QFP dyke. This age is older than the Lake George granodioritic stock (412 +5/-4 Ma; McLeod *et al.* 2003), although the ages are the same within error. The apparent date of 757 Ma from the Zr3 grains may be attributed to the presence of xenocrystic zircon incorporated from the country rock during dyke emplacement.

GEOCHEMISTRY

Whole-rock major-, trace-, and rare-earth-element (REEs) analyses of the QFP dyke are presented in Table 2. The Lake George granodiorite and dyke fall just within the low SiO₂ suite of Devonian intrusions described by Whalen *et al.* (1996). ACNK [molar Al₂O₃/(CaO+K₂O+Na₂O)] averages 1.09 for the least-altered samples. Normative mineralogy calculated from the whole-rock analysis and plotted on a QAP diagram (LeMaitre *et al.* 1989) places seven of the ten samples in the granodiorite field, and two in the granite field. The normative average of these nine samples is quartz 32.3 %, corundum 2.66 %, orthoclase 18.95 %, albite 22.32 %, anorthite 12.2 %, hypersthene 7.06 %, magnetite 1.38 %, ilmenite 1.03 %, and apatite 0.28 %. The tenth sample, DDH 79-6-210, is unusual in

Table 2. Major-, minor- and trace-element analyses of the QFP dyke at the Lake George mine.

Sample	79-6-	79-6-	79-6-	81-4-	81-21-	81-21-	81-25-	81-26-	83-1-	81-14-	Average	SD	
ID	195.5	174.3	210	271.9	181.1	317.6	19.2	63.4	37.5	392.0	Average	3D	
04-PL-#	1	2	3	4	5	6	7	8	9	10			
Motion and minor elements and L Ω 1 ¹²													
SiQ.*	66 7	69.6	71.6	70.3	68.4	68.1	68.0	69.8	67.2	68.6	68.83	1 49	
510 ₂	0.58	0.56	0.58	0.42	0.55	0.62	0.56	0.40	0.58	0.57	0.54	0.07	
ALO	15 21	14.66	15.00	14.04	15.00	14.05	15.26	14.76	14.02	15.60	14.05	0.07	
AI_2O_3	15.21	14.00	15.00	14.04	15.00	14.95	15.50	14./0	14.92	15.60	14.95	0.42	
$Fe_2O_3^{-1}$	4.25	3.60	6.40	3.32	3.48	4.15	4.19	3.17	4.35	4.12	4.10	0.91	
MnO	0.05	0.06	0.02	0.05	0.06	0.06	0.08	0.06	0.07	0.06	0.06	0.02	
MgO	1.47	0.99	1.46	0.71	0.95	1.19	1.39	0.84	1.37	1.42	1.18	0.28	
CaO	3.10	2.30	0.34	2.65	3.51	3.75	2.95	2.16	2.83	2.68	2.63	0.94	
Na ₂ O	2.93	2.39	0.00	2.67	2.88	3.01	3.41	3.20	3.17	2.85	2.65	0.97	
K ₂ O	3.01	3.66	3.03	3.76	3.37	2.52	2.96	3.73	3.33	2.77	3.21	0.42	
P_2O_5	0.14	0.14	0.15	0.09	0.13	0.12	0.13	0.13	0.13	0.11	0.13	0.02	
L.O.I. ²	3.55	3.28	3.86	3.02	3.84	4.38	1.38	1.09	2.13	3.89	3.04		
Total	101.0	101.2	102.4	101.0	102.2	102.9	100.4	99.3	100.1	102.7	101.3		
ACNIZ	1 11	1 21	2.05	1.07	1.02	1.02	1.00	1 1 2	1.07	1.24	1 20		
ACINK	1.11	1.21	5.85	1.06	1.02	1.05	1.08	1.12	1.07	1.24	1.38		
Trace elem	ents ³												
Rb	116	138	123	146	146	130	121	132	123	135	131	10.2	
Cs	5	4	7	4	6	4	5	6	6	6	5	1	
Ba	466	486	277	467	386	442	432	385	413	282	404	73	
Sr	226	156	77	147	177	187	273	207	248	166	186	56	
Ga	19	17	19	17	19	18	19	18	19	20	19	1	
Та	< 0.5	1.7	< 0.5	< 0.5	1.2	1.4	< 0.5	1.1	1.2	1.0	0.9	0.3	
Nb	15	16	17	13	15	15	16	19	16	16	16	1	
Hf	4	4	4	3	3	4	4	4	4	4	4	0.3	
Zr	158	154	158	148	151	156	168	149	167	161	157	7	
Y	17	17	19	14	17	18	18	18	18	18	18	1	
Th	10.3	11.9	12.4	9.2	10.4	10.5	10.8	10.0	10.9	10.6	10.7	0.9	
U	5.8	7.1	10.1	4.5	6.5	6.1	6.7	9.5	5.7	6.1	6.8	1.7	
La	25.0	26.6	28.2	24.0	24.2	23.6	21.9	26.8	23.7	23.4	24.7	1.9	
Ce	52	51	58	49	49	48	48	56	50	49	51	3	
Nd	29	33	35	34	22	26	31	31	23	29	29	5	
Sm	3.9	3.5	4.4	3.4	3.5	3.7	3.6	4.0	3.2	3.7	3.7	0.3	
Eu	0.9	0.7	0.7	0.7	0.5	1.1	1.0	0.9	0.9	1.0	0.8	0.2	
Tb	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5			
Yb	1.9	2.2	1.9	1.3	1.9	1.9	1.9	2.0	2.0	1.9	1.9	0.2	
Lu	0.29	0.35	0.25	0.22	0.35	0.29	0.31	0.31	0.32	0.28	0.30	0.04	
Cr	30	18	28	4	23	23	22	5	23	28	20	9	
Ni	14	10	20	10	12	14	14	9	12	16	13	3	
Со	8	7	7	3	7	7	7	5	7	6	6	1	
Sc	7.3	6.6	7.7	4.7	6.5	7.3	7.5	5.5	7.3	6.9	6.7	1.0	
V	65	57	83	32	60	66	55	35	57	55.0	56.4	14.6	
Cu	130	112	136	125	148	160	58	23	108	118	112	42	
Pb	16	20	< 0.53	24	16	12	20	39	15	12	19	8	
Zn	<50	<50	60	<50	<50	<50	<50	<50	<50	<50			
W	<1	<1	<1	2	3	<1	<1	<1	<1	<1			
Mo	4	7	12	<1	11	<1	<1	<1	7	5	5	3	
Ag	<5	<5	<5	<5	<5	<5	<5	<5	<5	<5			
As	8.6	46	685	295.5	92.1	16.7	7.5	44.4	35.2	12	124	215	
Se	<3	<3	<3	<3	<3	<3	<3	<3	<3	<3			
Sb	12	30	630	28	20	33	5	10	9	23	80	193	
Br	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5	< 0.5			
Cl	55	67	77	74	66	42	79	83	43	61	65	15	
Hg	<1	<1	<1	<1	<1	<1	<1	<1	<1	<1			
Au	46	36	31	29	8	<2	3	<2	39	12	25	16	
S^4	5225	4244	4864	2459	3453	3822	1802	1062	4043	1562	3254	1447	
S ⁵	107	105	57	57	94	57	85	47	41	41	69	26	
0	107	105	57	57	77	57	00	77	- 11	11	07	20	
MS^{6}	320	1350	150	110	750	220	600	150	900	600	515	404	
δS (‰)	1.4	1.2	-0.7	0.5	0.8	0.5	3.4	1.6	1.4	0.3	1.0	1.1	

Notes: 1. Major and minor oxides are given as weight percent. 2. L.O.I. = Loss on ignition. 3. Trace element analyses all in ppm except Au which is in ppb. 4. XRF analysis of pressed pellet. 5. XRF analysis of fused disk. 6. MS = Magnetic susceptibility in SI units (x 10^5).

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that it plots as in the quartz-rich granitoid field and has much lower CaO and Na_2O contents (0.41 and 0.22 %, respectively), suggesting that the sample has been strongly altered; it contains altered xenoliths of similar composition (see also sample 81-26-63.4).

The Nb versus Y and the Rb versus Y+Nb discrimination diagrams (Figs. 9a, b) (Pearce *et al.* 1984) show that the dyke has a volcanic-arc, I-type granite affinity, similar to the Lake George granodiorite stock (Yang *et al.* 2002a). Normalized to primitive mantle and chondrite compositions (Fig. 10a, b), the dyke shows negative Ba (ave. 404 ppm), Nb (ave. 16 ppm), TiO₂ (ave. 0.54 wt. %), and Sr (ave. 186 ppm) anomalies.

Multi-element (spider) and chondrite-normalized REE patterns for the QFP dyke and proximal plutonic units (Fig. 10a, b) are similar with notable negative Ba, Nb, P, and weak Ti anomalies. The spider diagrams also show that sample DDH-81-21-181.1 (fractured) and sample DDH 81-26-63.4 (dark glassy matrix) are similar in composition to the other samples despite textural differences. The QFP dyke most closely resembles the Lake George granodiorite and the Hawkshaw granite (Fig. 11a, b). The La/Yb ratios for these units are also similar (13.0, 14.4, 15.2, respectively), whereas the ratios for the Hartfield tonalite (16.5) and the Skiff Lake granite (14.6)are relatively higher. The lower LREE and HREE abundances in the QFP dyke, granodiorite stock, and Hawkshaw granite, relative to more primitive phases of the Pokiok Batholith, suggest low temperature fractionation process (e.g., Miller and Mittlefehldt 1982).

The Rb, Ba, and Sr ternary diagram (Fig. 12) shows the differentiation trend in the Pokiok Batholith. Based on whole-rock compositions, Sr content should decrease, whereas Ba increases within a differentiating intermediate to felsic system (el Bouseily and el Sokkary 1975). The Rb-Ba-Sr diagram shows that the Hartfield tonalite is the most primitive (oldest) and the Allandale granite is the most differentiated unit of the batholith, although the latter unit likely represents a separate magmatic event, based on its much younger age. However, this diagram displays some scatter for the Lake George granodiorite and QFP dyke samples, attributed to alteration.

Although zircon saturation thermometry was not used due to the possibility of inheritance, it is notable that the Hartfield tonalite and the Skiff Lake and Hawkshaw granite units exhibit decreasing Zr and TiO_2 (Fig. 13), consistent with fractional crystallization. This variation is consistent with the saturation behaviour of zircon in crystallizing granitic magmas (Hanchar and Watson 2004), and the decreasing REE contents noted above.

The average S content determining using pressed pellet XRF analysis is 3250 ± 1450 ppm; compared to the results from the fused bead method, the data indicate that S is dominantly tied up in sulphide minerals. The average δ^{34} S for the bulk S contents of the QFP dykes is $1.0 \pm 1.1 \%$ (Table 2); this narrow range near typical mantle values in a reduced I-type intrusion (Ohmoto and Goldhaber 1997) is consistent with a high-T magmatic origin for the sulphide minerals, with negligible fractionation evident (see detailed discussion in Yang 2005).

Au (<2 to 46, average 20 ppb), W (<1 to 3 ppm), and Mo (<1 to 11, average 7 ppm) contents in the QFP dyke are substantially lower than averages for the Lake George granodiorite (32 ppb, 9, and 7 ppm, respectively; Yang et al. 2002a). The average Cu abundance (112 \pm 42 ppm) in the QFP dyke is notably higher than in the Lake George granodiorite (average 41 \pm 38 ppm). S abundance co-varies with both Au and Cu (Fig. 14), supporting a deuteric origin for the S and metal enrichment; these data are consistent with the observations of Stirling (1979) of S enrichment in the most quenched contact zones of a similar dyke. Average Sb (80 ± 193 ppm) and As $(124 \pm 215 \text{ ppm})$ contents are significantly higher in the dyke than in the Lake George granodiorite, indicative of alteration, but they are unrelated to S, Cu, and Au abundance based on their low correlation coefficients (<0.3). However, all of these values are above crustal averages (Wedepohl 1995) and point to enrichment associated with magmatic volatiles related to emplacement of the QFP dyke.

GEOTHERMOMETRY

In terms of magma geothermometry, apatite saturation temperature (AST) calculated using P_2O_5 abundance and compositional corrections was estimated for the Lake George granodiorite to be $902 \pm 14^{\circ}$ C. This temperature is interpreted to represent a melt temperature with apatite as a liquidus phase (Yang *et al.* 2002b). Similarly, apatite also occurs as a liquidus

Table 3. U-Pb data for the QFP dyke (DDH 81-21) at the Lake George antimony mine, New Brunswick.

Grain	Desc.	Wt. (mg)	U (ppm)	<u>Th</u> U	Pb* (ppm)	Comm. Pb (pg)	²⁰⁶ Pb ²⁰⁴ Pb	²⁰⁶ Pb ²³⁸ U	$(2 \sigma \%)$	$\frac{\frac{207}{Pb}}{^{235}U}$	$(2 \sigma \%)$	²⁰⁷ Pb ²⁰⁶ Pb	$\overset{\pm}{(2\sigma\%)}$	²⁰⁶ Pb/ ²³⁸ U	Ages ²⁰⁷ Pb/ ²³⁵ U	²⁰⁷ Pb/ ²⁰⁶ Pb	Conc. %
Zr 1	2Pr	0.015	284	0.23	18.3	10	869.0	0.0662	0.35	0.5042	0.36	0.05525	0.16	413	415	422	97.8
Zr 2	1Pr	0.009	117	0.35	7.8	8	539.5	0.0664	0.31	0.5047	0.45	0.05511	0.36	415	415	417	99.4
Zr 3	2Rd	0.018	127	0.29	11.4	6	2303.4	0.0924	0.27	0.8209	0.60	0.06446	0.54	569	609	757	74.1
Zr4	1Pr	0.01	455	0.30	29.5	7	2549.1	0.0652	0.28	0.4951	0.28	0.05507	0.20	407	408	415	98.0
Zr 5	2Pr	0.03	471	0.37	30.2	9	2925.4	0.0629	0.30	0.4784	0.29	0.05513	0.12	393	397	417	94.1

Pr = prismatic; Rd = Rounded; 1 = 1 zircon; 2 = 2 zircons; Pb* = Radiogenic Pb



Fig. 7 U-Pb concordia diagrams for five zircon fractions from the QFP dyke at Lake George (DDH 81-21)(see Table 3 and Appendix). Data point error ellipses are reported at 2σ .



Fig. 8 QAP diagram (LeMaitre et al. 1989) based on CIPW norms for the QFP dyke at Lake George, New Brunswick.



Fig. 9 Trace-element geotectonic discrimination diagrams (after Pearce *et al.* 1984; Christiansen and Keith 1996). a) Nb versus Y diagram and b) Rb versus Nb+Y including averages from units of the Pokiok Batholith compared to the QFP dyke and Lake George granodiorite (see Yang *et al.* 2002a). Data from the Pokiok Batholith are from Whalen (1993).

phase in the QFP dyke (this study) and in units of the Pokiok Batholith (i.e., Hartfield, Skiff Lake, Hawkshaw, and Allandale granitoids) (Lutes 1987; Whalen 1993; Yang *et al.* 2002a, b). Thus, we calculated liquidus temperatures for these granitoids using AST (Harrison and Watson 1984) information which may help constrain their genetic relations. Average AST values for the QFP dyke, Hartfield, Skiff Lake, Hawkshaw, and Allandale granitoids are 902°, 909°, 955°, 958°, and 982°C, respectively. The similarity in AST between the dyke and the units of the batholith suggests that they may be genetically related, consistent with compositional similarities. AST decreases from



Fig. 10 (a) Primitive mantle-normalized, trace-element spider diagrams for the QFP dyke. The primitive mantle values are from McDonough and Sun (1995). (b) Chondrite-normalized rare-earth-element spider diagram for the QFP dyke at Lake George. The chondrite-normalizing values are from McDonough and Sun (1995).

Fig. 11 (a) Primitive mantle-normalized, trace-element spider diagram for phases of the Pokiok Batholith, QFP dyke, and Lake George granodiorite. The primitive mantle values are from McDonough and Sun (1995). (b) Chondrite-normalized rare-earth-element patterns for the Pokiok Batholith, QFP dyke, and Lake George granodiorite. The chondritenormalizing values are from McDonough and Sun (1995).

the Hawkshaw granite (958°C) to the QFP dyke and stock (902°C), consistent with fractional crystallization (cf. Yang *et al.* 2002a).

DISCUSSION

Textural constraints on crystallization

No change is perceptible in groundmass texture or grain size in the dyke at depth relative to near-surface samples, al-





Fig. 12 Rb-Ba-Sr ternary diagram of el Bouseily and el Sokkary (1975) for the QFP dyke at Lake George plotted with the five units of the Pokiok Batholith and Lake George granodiorite (data from Whalen 1993; Yang *et al.* 2002a).



Fig. 13 Zr versus TiO_2 plot of the QFP dyke at Lake George, as well as average and range (SD) for related intrusive units in the area (data from Whalen 1993; Yang *et al.* 2002b). The vector is the interpreted fractionation array, with Zr/TiO₂ ratios illustrated.



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Fig. 14 S versus Au and Cu abundances in the QFP dyke at Lake George. The r values are calculated Pearson Product correlation coefficients.

though the dyke margins exhibit more auto-brecciation. The groundmass is recrystallized and hydrothermally altered to a variable degree, probably reflecting the autometasomatic effects of deuteric fluids, as well as the slightly later contact metamorphic and metasomatic effects of intrusion of the Lake George granodiorite and antimony vein mineralization in the area. Phenocrysts of subhedral to rounded feldspar and quartz are common in both the dyke and Lake George granodiorite (Yang et al. 2002b). Rounded phenocrysts suggest that quartz and plagioclase were partially remelted (depressurization) and (or) were abraded mechanically during rapid emplacement. The subrounded quartz phenocrysts form composite crystals that are fused together locally. No evidence of gas cavities is preserved. The altered granitic fragments (0.5 to > 10 cm in diameter) near the QFP dyke margins are subangular to rounded and possibly recrystallized; these fragments probably represent the fractionated roof zone to this intrusion and (or) autobrecciated products of the margin of the dyke itself, as they have the same bulk composition as the dyke. Although angular to rounded wall rock fragments exist, they were avoided during sampling.

The oscillatory normal zoning in plagioclase (Fig. 4) (An 43 to 53) is interpreted as the result of diffusion-controlled growth locally interrupted by magma replenishment and possible resorption in the subvolcanic magma chamber before dyke emplacement. Zoning of plagioclase was obscured during periods of uneven resorption of the grains, which resulted in a rounded zoning pattern. SEM-EDS analysis across zoned plagioclase grains (Fig 4) in sample DDH 81-62-63.4 shows variations in Ab and An within each subzone, reflecting variation in crystallization and/or melt diffusion rates of Na and Ca (Table 1). The centres of the plagioclase phenocrysts contain an unzoned homogeneous core, indicating that during this period crystallization was slower to allow diffusion across the grain. The textures present in the plagioclase from the Lake George granodiorite stock are similar to those in the QFP dyke (Seal et al. 1988; Yang et al. 2002a).

It was concluded earlier that the source magma for the granodioritic intrusion and QFP dyke crystallized amphibole

and biotite during its ascent (Yang et al. 2002b). The absence of amphibole in the dyke suggests that the amphibole phenocrysts were directly replaced by biotite, i.e., biotite became a liquidus phase at the expense of hornblende in the presence of evolving melt at lower temperature (see Naney 1983), or fractionation removed amphibole from the dyke-forming magma. If amphibole partitioned higher amounts of HREE, their absence in the dyke would be indicated by a noticeable decrease in HREE (Barker 1979). The expected relative decrease in HREE is not evident between the granodioritic intrusion and the dyke (Fig. 12), suggesting that the lack of amphibole was not due to fractionation. The geochemical similarity between the QFP dyke and the Lake George granodiorite stock suggests that the parental magma of the dyke was hornblende bearing. As can be seen on the chondrite-normalized REE diagram (Fig. 12b), a lower abundance of total REE and LREE is present in younger units of the Pokiok Batholith. The decrease in LREE with fractionation supports apatite (and titanite \pm monazite) as liquidus phases, possibly enhanced by an association with ferromagnesian phases.

Reddish-brown (reduced) biotite was observed as inclusions within quartz phenocrysts, although it was not observed in plagioclase; chlorite (the main product of biotite alteration) was also observed as inclusions in quartz. These observations support the interpretation that biotite was present when these phenocrysts crystallized, although they look sutured and anhedral possibly due to destabilization during emplacement/ depressurization.

The euhedral titanite crystals in the dyke are of igneous origin, as are the rare ilmenite crystals. The inclusions in the titanite are likely the result of hydrothermal alteration, although some may have been entrapped during episodes of crystallization and regrowth. The zoning and resorption surfaces in titanite suggest an evolving magmatic system that may have resulted from changes in melt composition due to fractionation or could indicate that Ti enrichment occurred on more than one occasion. Regardless of possible Ti enrichment, the presence of ilmenite, titanite, rutile, and Ti-rich biotite appear to reflect high Ti contents in the dyke-forming magma. The assemblage titanite-magnetite-quartz-pyrrhotite may be a late auto-oxidation and sulphidation product of vapour exsolution (Candela 1991), as it is not consistent with high Fe (-Ti) biotite (Wones 1981).

Magmatic pyrrhotite phenocrysts may have crystallized under reducing conditions in the magma, although they would have segregated quickly and lowered the base-metals and gold abundance. Presently, the S abundance in the QFP dyke is almost 10 times greater than the S solubility in granitoid magmas at 1000°C (Ohmoto and Goldhaber 1997). Therefore pyrrhotite formation at late stages of crystallization related to alteration with H₂S-bearing magmatic fluids better explains the S abundances and isotopic signatures observed. Gold distribution and abundance, relative to that of S and other metals, in igneous rocks provides important information about the potential for associated gold mineralization (cf. Müller and Groves 2000 and references therein).

Dyke emplacement constraints

The presence of quartz and two feldspar phenocrysts in the stock and dyke implies high $P(H_2O)(>3 \text{ kb}; \text{Martin and Bonin})$ 1976) in the magma before final emplacement. Considering the lack of marginal quenching and flow foliation, as well as the rheology of the phenocryst-rich siliceous magma (discussed below), it is probable that exsolved gases from the magma (fluidization) enhanced dyke emplacement (Clarke 1992); fluidization is consistent with the angular to subrounded nature of xenoliths and the rounding and fragmentation of phenocrysts that are now best described as phenoclasts. The fine-grained groundmass and minor granophyric (wormy) to more prevalent graphic (cuneiform) texture resulted from quench crystallization (low degrees of supercooling) of the dyke, possibly enhanced by vapour exsolution and depressurization (see Fenn 1986; Lentz and Fowler 1992). The highest level dyke samples exhibit microcrystalline texture consisting of intergrown fibrolites rimming quartz and feldspar phenocrysts, reflecting greater degrees and rates of supercooling or devitrification (Stirling 1981; MacLellan and Trembath 1991), akin to aplite.

Calculated cooling rates using a conductive thermal cooling model (HEAT, Wohletz and Heiken 1991) for this vertical 5 m-wide QFP dyke (initial 1000°C liquidus temperature, see below) in a low geothermal gradient (20°C/km) reveals that the dyke margins quench to less that 500°C in less that 1 day at 1 to 3 km depth, and that complete solidification (< 760°C at 1 kb) occurs in less than 10 days at 1 km, and 30 days at less than 3 km (equivalent to 1 kb). These figures are consistent with the cooling rates and quench textures in high-level felsic dykes (Stirling 1979). These extremely fast cooling rates in upper crustal thermal gradients should limit the extent of dyke emplacement from its source magma chamber considering melt and effective melt viscosities (cf. Weinberg 1996; Baker 1998).

The calculated viscosity (Shaw 1972) for this magma composition (no crystals) is 4.6 PaS at the calculated liquidus

temperature (980°C) and 1 kb, which decreases to 7.5 PaS at 700°C (1 kb, 100 MPa). However, the effective viscosity using the Einstein-Roscoe equation (McBirney and Murase 1984) increases dramatically with crystal size and abundance. The effective magma viscosity for 50 vol. % crystals ranges from 24.0 PaS (1 mm diameter) to 200 PaS (1 cm diameter). However, dyke emplacement pressures are probably considerably less. At 200 bars (20 MPa) vesiculation begins based on conservative water contents of 1.7 wt. %, with critical melt/vapour ratio at 50 bars (>70 vol. % gas) with a bulk density of 0.7 g/cm³; these represent minimum estimates and gas vesiculation probably began much earlier considering the high degree of crystallization (> 30 vol. %) associated with decreasing temperature, so 200 bars (20 MPa) is a reasonable minimum estimate for H_2O exsolution. The bulk density of the crystallized magma is calculated as 2.5 g/cm³. The melt-vapour density is approximately (0.7 g/cm^3) at the critical fragmentation pressure (< 20 MPa) (magmatostatic). It therefore seems reasonable to expect that a vesiculating hydrous phase was involved in the emplacement of this QFP dyke. Passive to forcible dyke emplacement occurs via fracture propagation (Shaw 1980), possibly enhanced by fluidization (low density, low rheology melt-vapour mixture); the aspect ratio of this dyke (5 m wide by > 1 km in length, and > 500 m depth) is also consistent with this emplacement mechanism.

It is possible that the QFP dyke was emplaced along a pre-existing fault or fracture, based on its compositional uniformity, planar attributes, and evidence of auto-brecciation within the dyke (Richards 2001). Initial fracture propagation can occur along a fault (Weinberg 1996), although no offsets are evident relative to the earlier east-trending lamprophyre dykes at the map scale. The lack of evidence for quench crystallization on the margins of the dyke suggests that the degree of undercooling was extreme, possibly even as a glass-gas mixture that subsequently devitrified to produce the observed microcrystalline textures and alteration. The shallow level of dyke emplacement into host rocks at less that 300°C (max. 50°C/km for 6 km or 2 kb) requires a rapid emplacement mechanism (> 100 m/day) given the thermal modelling constraints.

CONCLUSIONS

Although seemingly simple, the QFP dyke has numerous features that help to constrain the emplacement of the magma and understand the role of such subvolcanic bodies in forming the mineralization in the Lake George area.

1) The QFP dyke yielded an age of 420.8 +5.9/-4.0 Ma by U-Pb zircon geochronology. Therefore, it is likely genetically related to units of similar age in the Pokiok Batholith, and possibly the Lake George granodiorite stock.

2) The QFP dyke has petrochemical features of a reduced I-type granodiorite, like the Lake George granodiorite stock. Therefore, it is probably a fractionation product of a deeper level unit of the Pokiok Batholith.

3) The presence of two feldspar phenocryst phases supports

the existence of a deep level (> 3 kb, 300 MPa) magma chamber feeding this dyke. Oscillatory zoning in rounded plagioclase phenocrysts/phenoclasts reveals a dynamic system, which caused melting/resorption. This system was likely caused by several episodes of magma mixing or the introduction of fresh magma pulses from the source.

4) Amphibole, which is present in most of the other Pokiok granitoid units, reacted with the fractionated melt to form biotite prior to dyke emplacement. This change explains the similar REE patterns and abundances throughout the Pokiok Batholith, even in the absence of amphibole.

5) Autobrecciation along the margins of the dyke, subrounded fragments, subhedral to rounded and fragmented plagioclase and quartz phenocrysts (phenoclasts), and the autometasomatic effects of deuteric fluids support fluidization (vapour-magma emulsion) as a key emplacement mechanism for the dyke, i.e., analogous to tuffisite. Thermal modelling coupled with integration of the estimated magma properties affecting rheology of this particular system is consistent with these observations.

6) The presence of pyrrhotite (>10 times sulphur saturation levels), which accounts for the range in magnetic susceptibilities, with chalcopyrite (with minor gold) in the QFP dyke was probably produced by high-T subsolidus reactions of deuteric fluids with the quenched walls of the dyke as it was emplaced; this alteration is manifested as alteration in the microcrystal-line groundmass. The sulphur isotopic signature ($\delta^{34}S = 1.0 \pm 1.0 \%$) and S abundance (3250 ± 1450 ppm) indicate that magmatic volatiles were directly involved during dyke emplacement (autometasomatic) and by inference that Cu and Au were associated with these volatile components.

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APPENDIX: ANALYTICAL METHODS

Ten drill-hole core samples from 15 to 25 cm in length were collected from the diamond-drill holes along the length of the QFP dyke on the Lake George mine property. These samples were taken from vertical depths between 37.5 m (123 ft.) to 392 m (1286 ft.) depending on the drillhole intersection. Core exhibiting significant weathering or alteration was avoided. The samples were crushed, then pulverized using a soft steel mill at the University of New Brunswick. Portions of each sample were sent to Actlabs for instrumental neutron activation analysis (INAA), which provided As, Au, Ag, Ba, Br, Ce, Co, Cs, Cr, Eu, Hf, Hg, La, Lu, Mo, Nd, Ni, Rb, Sb, Sc, Se, Sm, Ta, Tb,

Th, U, W, Yb, and Zn. The remainder of the pulped samples were analysed for Al₂O₃, CaO, Fe₂O₃ (total), K₂O, MgO, MnO, Na₂O, P₂O₅, SiO₂, and TiO₂ by fused disk and As, Ba, Cl, Cr, Cu, Ga, Nb, Ni, Pb, Rb, Sr, S, Th, U, V, Y, Zn, and Zr by pressed pellet X-ray fluoresence analysis at Memorial University (see Longerich 1995) (see Table 2).

The certified standard SY-4 was included as our internal standard for analytical quality control. The percent difference was calculated from the results obtained from both laboratories; where an element was analysed by both methods the result that was closer to the standard was used, choices which are indicated in Table 1. The percent error in the trace-element analyses is generally high where the standard had low abundance of the analysed element. If the quantity of the trace elements in the standard was higher than 10 ppm, then the percent error is generally less than ten percent.

Sulphur isotope analyses were performed on bulk rock samples at the University of Ottawa using the method detailed in Yang (2005).

Ten polished thin sections were analysed optically to document petrology, with follow up using the JEOL 6400 scanning electron microscope (SEM) and backscattered electron (BSE) imaging at the Electron Microscopy Unit at the University of New Brunswick. Individual grain analysis was conducted using an energy dispersive spectrometer (EDS).

For the U-Pb ID-TIMS dating technique, a heavy mineral separate was prepared from a 10 kg rock sample. The sample was pulverized using a heavy-duty hydraulic rock splitter, jaw crusher and swing mill. Mineral separation involved the use of a Wilfley Table, heavy liquids (bromoform and methylene iodide) and a Frantz Isodynamic Separator. Analyses were performed at Memorial University of Newfoundland, Canada. Normal transmitted and reflected light microscopy, as well as SEM back-scattered or cathodoluminescence (CL) imagery were used to determine the zircon internal structures prior to analysis. Handpicked zircons were abraded (Krogh 1982), then washed in dilute nitric acid and ultra-pure acetone. Single grains or small populations of zircons (max. 2 grains) were then placed into 0.35 ml Teflon vials together with HF and few drops of HNO3 and a mixed 205Pb-235U spike. These Teflon vials were then placed in a Parr Container for several days at 210 °C (Parrish 1987). The samples were measured on a Finnigan MAT262 mass spectrometer equipped with an ion-counting secondary electron multiplier. A detailed account of the entire analytical technique was been given by Dubé et al. (1996).

Total Pb blanks over the period of the analyses range from 5 to 1 pg and a value of 5 pg was assigned as the laboratory blank (206 Pb/ 204 Pb = 18.97, 207 Pb/ 204 Pb = 15.73, and 208 Pb/ 204 Pb = 39.19). The calculation of common Pb was carried out by subtracting blanks and then assuming that the remaining common Pb has a Paleozoic composition determined from the model of Stacey and Kramers (1975). Data were reduced using PbDat (Ludwig 1993). Analytical uncertainties are listed at 2σ and age determinations were processed using Isoplot/Ex (Ludwig 2000).