

Structural and Geochemical Characterization of Gold Mineralized Quartz Veins in Belikombone Gold Prospect, Betare-oya Gold District, East Cameroon

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Abstract: Au-bearing quartz veins in the Belikombone area are confined to major NW-SE orientations in schists and N-S and ENE – WSW in other host rocks. The objectives of this work were to study field, mineralogy and geochemical characteristics of gold mineralised quartz veins. Schists (chlorite rich) and gneisses (moderately magnetic) are the dominant rock types in the area with granite occurring as intrusions within the gneisses. Gold-bearing quartz veins occur in association with metamorphosed rocks ranging in composition from quartz \pm gold to those that contain quartz \pm gold \pm sulphides \pm iron-oxides \pm sericite and mafic minerals. The mineralized veins contain Au from 0.50 to 25 ppm, while the quartz veins considered barren contain 0.20 ppm Au. The concentration of As in the quartz veins ranges from 0.30ppm to 10.70 ppm and it show positive correlation with Au in samples with high Au concentration. The Σ REE varies from 5.79 to 173.64 in the quartz veins. The investigated veins show evidence of multiphase deformation and it is associated with D₂ deformation in the area. This structural setting suggest that the emplacement of gold mineralization occurred during the late Pan African orogeny. The main alteration types are silification, sericitization and sulphidation processes.

Keywords: Gold, Quartz Veins, Geochemistry, Belikombone, Betare-oya, Cameroon

1. Introduction

Quartz vein systems in regions that have experienced deformation remain the main depository for primary Au [1]. Deformation in these regions occur in stages and the migration of hot fluids (hydrothermal) from the subsurface into the resulting structures also follow the sequence of deformation. These fluids eventually precipitate material into these structures forming lodes and veins that differ in their characteristics [1, 2]. Veins in such areas show different patterns including fault-fill veins, extensional veins and stockwork-like veins [3, 4]. Au -bearing veins are usually of polymetallic type with different textural characteristics. Some of these textures are usually associated with ore zones (mineralized veins) while others



are found in barren ones [3]. Some of the mineralized textures include overprinting and modification textures including phantom veins, ribbon and brecciated quartz, laminated quartz and comb quartz textures [3]. Goldbearing quartz veins occur in association with metamorphosed rocks ranging in composition from quartz \pm gold to those that contain quartz \pm gold \pm sulphides \pm iron-oxides \pm sericite and mafic minerals.

The Belikombone gold prospect is found within the Betare-Oya gold district of the Eastern Cameroon gold fields. General descriptive information on gold mineralization in the Eastern region has been well documented [5, 6]. In most cases of Au exploration, grade is easily assessed by crushing a quantity of quartz block samples, weighing and panning to obtain the concentrate from which the Au-grains if present,

can easily be recovered and weighed (e.g. [7-9]. As such, quartz vein systems appear to be a major tool in the study and understanding of the hydrothermal processes as well as the geochemistry of ore deposits that serves as a guide in mineral exploration [3]. Recent fieldworks have revealed an important gold bearing quartz veins in the Belikombone gold prospect and neighborhoods. Those gold occurrences are related to faults and folds, and commonly occur in shear zones.

This paper seeks to examine the field characteristics, petrography, mineralogy and geochemical composition of the quartz veins within the Belikombone area and to establish exploration prediction models. In addition to this, the possible origin of the gold bearing fluid is inferred.

The Belikombone gold district is part of the Lom Series which is a portion of the Neoproterozoic rock unit in Eastern Cameroon. The Neoproterozoic rock unit of Cameroon, north of the Congo craton is part of the Pan-African belt in Central Africa known as the Central African Fold Belt (CAFB; [10, 11], or North Equatorial Fold Belt (NEFB; [12], which was formed during the Pan-African orogeny. The Lom Series is made up of a NE-SW discontinuous narrow belt (Figure 1) comprising of a Lower and Upper Groups [13]. It represents a syn-tectonic pull-apart basin associated with the Sanaga shear zone and was deposited between 613 and 600 Ma [13, 14]. The Lower and Upper Lom Series is made up of metatuffs, volcanoclastic and sedimentary derived schists, staurolitegarnet mica schists and quartzites with local conglomerate layers [14]. The schists were metamorphosed under greenschist facies conditions [13]. The volcanoclastic rocks are felsic to intermediate in composition [12 - 14]. The lowpressure (garnet-andalusite-staurolite assemblage) regional metamorphism that was associated with transpressive tectonics in the Lom Series involved a high thermal gradient related to widespread crustal melting that produced the dominant S-type granitoids in the region [13, 14].

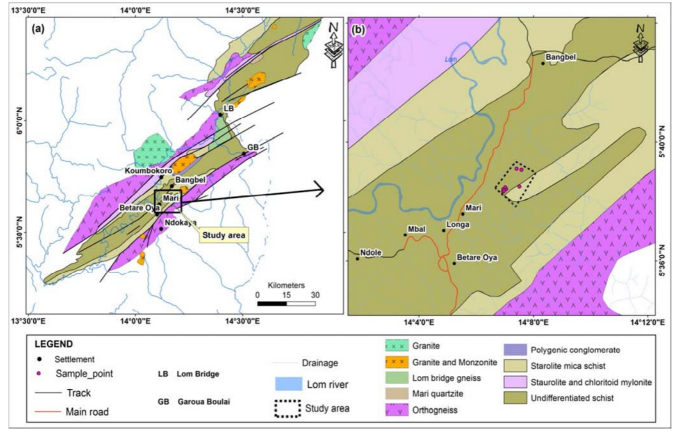


Figure 1. Regional Geologic setting of study area.

The tectono-metamorphic evolution of the Lom Series is polyphase, characterized by two stages of compressional deformation (D_1/D_2) , leading to a N50-N70 steep regional foliation without any apparent stretching lineation [12, 13]. Dextral or sinistral faults locally parallel to the main foliation characterize the later stages (D_3) of the structural evolution of the Lom Series [13-15]. Simple shear wrenching (D_3) in the Betare-Oya area is geometrically related to the NE-SW striking Betare-Oya shear zone (BOSZ) and to the Sanaga shear zone, with the mylonitic foliation spatially related to the BOSZ [15]. The host rocks are biotite schists, sericite and chlorite schists, quartzites and shales that constitute the Lom Series [6].

The Lom series is well known for its high economic potential which may be explained by the Neoproterozoic reworking and remobilization of an Archean to Palaeoproterozoic basement [13].



2. Materials and Methods

Quartz veins were collected as grab samples during structural and lithological mapping of the study area. During the lithological mapping structural measurements were taken. Thin sections were manufactured in the laboratory of the Institute of Geological and Mining Research (IGMR) at Nkolbisson, Yaounde, Cameroon, for minerals and microstructure studies. These thin sections were observed under plane and polarized light using an Olympus Cx31 electronic microscope. Petrographic images were snapped using a CMCA61300A camera attached to the microscope.

Quartz veins geochemical analyses (major, trace and rare earth elements) were obtained using Inductively Coupled Plasma Mass Spectrometry (ICP-MS) on the pulp at ALS Mineral Global Group in South Africa. Samples were weighed and dried at 100 \Box C to remove the moisture using a space warmer for 12 hours and later crushed in a jaw crusher and pulverized using ESSA® in a tungsten carbidecoated mill to obtain homogeneous samples for the analysis (www.alsglobal.com). Aqua Regia analysis was carried out on 15g of each sample pulp. The finely pulverized sample was digested in a mixture of Hydrocholoric acid (HCL) and Nitric acid (HNO₃). This acid mixture generates nascent chlorine and nitrosyl chloride which will dissolve free gold and gold compounds. A replicate was done after each analysis for quality control and quality assurance with standards and blanks inserted after every 5 samples to calibrate the equipment and ensure optimal precision of the data set obtained at a 95% confidence level.

3. Results

3.1. Petro-structural Characterization of Quartz Veins and Host Rocks

Field investigations have revealed that the prospected area is composed of metamorphic and magmatic rocks. These are the host rocks for quartz veins in the area.

3.1.1. Petrography of Host Rocks

The metamorphic rocks in the area comprise of schists and gneisses (Figure 2). Schists at Belikombone are dominated by chlorite associated with sericite (65%), quartz (25%) and oxides (10%). Chlorite occurs as thin bands with a lepidoblastic texture (Figure 3e). Gneiss which is moderately magnetic (Figure 3h) display an alternation of whitish (quartzo-feldspathic with more or less muscovite~ 5%) and dark bands of variable thickness. The dark bands are dominantly composed of magnetite, biotite, amphibole and oxides. This rock has fine to medium-grained with simple cracks which indicate possible wall rock alterations. The rock displays a granoblastic texture.

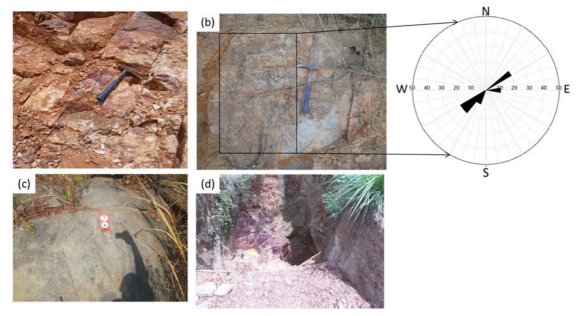


Figure 2. (a) Quartzite, (b) Schist, (c) Granite and (d) pits dug along the trend of quartz vein at Belikombone.

The magmatic rocks in the area are granites (Figure 2c and Figure 3f) which occurred as bouldering intrusions in gneiss, some of which are about 2m long and a meter high. The rocks are massive, with medium to coarse grains. The rock is made up of quartz (50%), plagioclase (8%), amphibole (7%), oxides (10%), biotite (10%) and muscovite (15%). As a whole, the rock displays a porphyroblastic-like texture showing that this rocks has been deformed.

3.1.2. Structure

The schists at Belikombone show well-developed schistosity defined by sericite. Some of the schists are crosscut by both joints and veinlets which intersect each other. Some of the granites have millimetric multi-directional quartz veins exhibiting a stockwork-like structure. These veinlets (averagely 2mm thick) also occur as individual microlines (Figure 3f), representing about 35% of the rock. They are



dominantly composed of quartz up to 95% when viewed using plane polarized light (Figure 3g). In addition to these microstructures, the rocks in the prospected area are cross cut by an important number of quartz veins. The later occur as individual or clustered veins with a general NW-SE trend and vary from few centimeter to about 7m wide. These structures have been differentiated based on their mineralogical content into mineralized and barren quartz veins (Figure 4). Most of the mineralized veins are massive with multiple fractures on their surfaces; these fractures are filled with oxides (magnetite) (Figure 4a) found in thin pellicles. Mineralogically these quartz veins have approximately 90% quartz grain, 5% sulphide and 5% oxides and show numerous textural features such as porphyroblastic and granoblastic. The relationship between mineralised veins and the host schists suggest an epigenetic style of mineralisation. These veins vary considerably in thickness and often exhibit significant vertical and lateral continuity (Figure 4a). They vary from few centimeter to 5m wide with strike values ranging from N006E to N165E (mean strike of N78E). The dip of the mineralized veins varies from subhorizontal (9°) to vertical (89°) to the NW and SE (Figure 5).

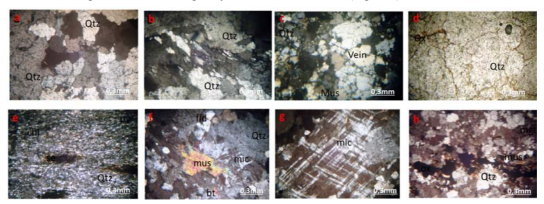


Figure 3. (a-d) Photomicrograph of representative rock samples from the study area. (a – d) Quartzite, (e) Schist, (f) Granit, (g) Microcline and (h) Gneiss.

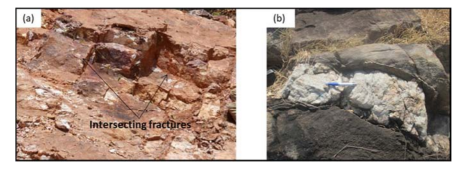
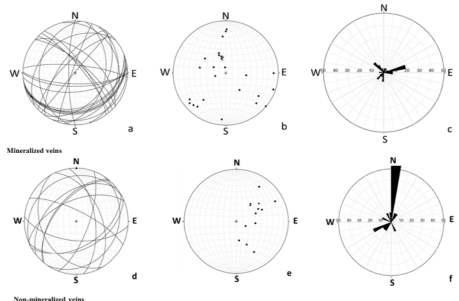


Figure 4. (a) Mineralized vein dissected by intersecting fractures (b) Non-Mineralized vein.



Non-inmeralized venis

Figure 5. β , π , and Rose diagrams of Mineralized and non-mineralized veins.



Most of these veins are brecciated with fractured surfaces which are filled with iron oxide. The main gangue mineral in the quartz vein consists of exclusively xenomorphic quartz crystals (more than 98% of the veins) with an equigranular texture. The proportion of the vein -forming minerals varies from veins that contain only quartz to those containing quartz \pm sulphides \pm iron-oxides \pm sericite. In the sulphide bearing veins, the sulphides occur as disseminations with medium crystal sizes. The quartz crystals that show undulatory extinction are aligned suggesting a post deformation.

The barren veins (Figure 4b) are characterized by oblique joints. Barren veins vary from few centimeter to 8m wide with strike values ranging from N20E to N175E (mean strike of N97E). The dip of these veins ranges from 21° to 74° to the NW or the SE and E or W directions. Most of these veins are massive with very minor fractures on their surfaces. Structural data of these veins shows that the veins in the prospected area are variably oriented; however the N-S and ENE - WSW trends appear to be the dominant ones (Figure 6a-c)

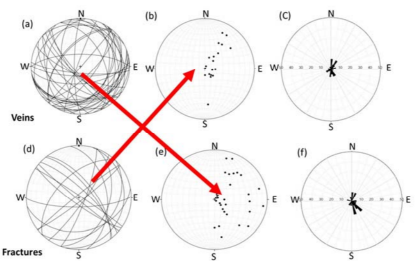


Figure 6. β , π , and Rose diagrams of veins and fractures in the Belikombone gold district. $a=\beta$ diagram of veins, $b=\pi$ diagram of veins, c=Rose diagram of veins, $d=\beta$ diagram of fractures, $e=\pi$ diagram of fractures and f=Rose diagram of fractures.

In this area most fracture surfaces have an average dip of 56° predominantly to the west with minor displacement which crosscut the quartz veins and the fractures are parallel (NE-SW). Stereoplots (β -diagrams and π -diagrams) show that the fracture surfaces are variably oriented (Figures 6d and 6e). As a whole, two principal sets of fractures have been recorded over the prospect: the N045-089E and N030-041W defining two broad directions for this deformation including NE-SW and NW-SE respectively (Figure 6f).

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3.2. Geochemistry of Quartz Veins

The whole rock geochemical composition of quartz veins is shown in Table 1. In mineralized veins, the concentration of Si₂O ranges from 83.54 to 99.43 wt%, Ti₂O from 0.003 to 0.264 wt%, Al₂O₃ from 0.1 to 8.96 wt%, Fe₂O₃ from 0.11 to 6.26 wt%, Mn₂O from 0.002 to 0.557wt%, P₂O₅ from 0.006 to 0.19 wt%, K₂O from 0.01 to 0.075 wt%, Na₂O from 0.01 to 0.56 wt%, CaO from 0.022 to 0.286 wt%, MgO from 0.02 to 1.74 wt%, ClO from 0.006 to 0.045 wt%.

Table 1. Whole-rock ICP-MS analyses of quartz vein samples.

Sample types Sample names	Mineralized samples				Barren samples					
	QV01	QV02	QV05	QV08	QV03	QV04	QV06	QV07	QZT	SCH
Major elements (%)									
Si ₂ O	83.54	89.81	92.22	99.24	97.49	99.22	99.43	94.56	99.23	74.25
Ti ₂ O	0.26	0.05	0.01	0.00	0.03	0.01	0.00	0.14	0.00	0.10
Al ₂ O ₃	8.96	1.49	2.23	0.27	0.94	0.34	0.10	0.90	0.29	13.68
Fe ₂ O ₃	3.56	6.26	2.59	0.11	0.66	0.13	0.21	3.12	0.14	1.09
Mn ₂ O	0.03	0.56	0.10	0.00	0.04	0.00	0.00	0.08	0.00	0.01
MgO	1.74	0.02	1.23	0.02	0.03	0.02	0.02	0.36	0.02	0.08
CaO	0.29	0.02	0.23	0.03	0.04	0.03	0.06	0.13	0.03	0.36
Na ₂ O	0.56	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	3.41
K ₂ O	0.02	0.06	0.03	0.08	0.03	0.10	0.01	0.02	0.09	5.81
P_2O_5	0.01	0.19	0.17	0.01	0.01	0.01	0.01	0.10	0.01	0.03
ClO	0.05	0.01	0.01	0.01	0.02	0.01	0.02	0.01	0.02	0.01
LOI	0.95	1.35	1.08	0.19	0.69	0.16	0.09	0.49	0.15	0.99

Sample types	Mineralized samples				Barren samples					
Sample names	QV01	QV02	QV05	QV08	QV03	QV04	QV06	QV07	QZT	SCH
Major elements (%))									
Total	99.97	99.83	99.91	99.97	99.99	100.03	99.96	99.92	99.99	99.82
As	9.60	9.70	4.20	2.50	0.40	10.70	1.50	1.20	0.30	0.60
Au	25.00	4.30	0.50	3.70	0.20	0.20	0.20	0.20	0.20	0.20
Ag	0.01	0.22	0.64	0.04	0.06	0.51	2.03	0.25	0.02	0.03
Ba	10.00	10.00	10.00	10.00	10.00	10.00	10.00	10.00	10.00	10.00
Со	10.70	27.30	4.10	1.10	1.50	0.30	2.50	1.20	1.00	1.30
Hf	0.05	0.13	0.70	0.32	0.09	0.16	0.66	0.66	0.27	0.19
Мо	0.50	1.00	0.30	0.20	0.20	0.20	0.30	0.20	0.20	0.20
Rb	0.10	0.20	0.20	0.80	0.30	0.20	1.00	0.40	2.00	0.10
Sb	0.39	0.30	0.17	0.12	0.26	0.18	0.16	0.16	0.25	0.13
Sc	0.10	0.10	0.20	0.10	0.10	0.10	0.10	0.10	0.10	0.10
Та	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01
Th	0.20	0.20	0.20	0.20	0.20	0.20	0.20	0.20	0.70	0.20
U	0.05	0.05	0.05	0.07	0.05	0.05	0.05	0.08	0.06	0.05
W	0.06	0.05	0.05	0.05	0.05	0.05	0.15	0.05	0.05	0.05
Zn	43.00	9.00	6.00	5.00	7.00	4.00	3.00	8.00	4.00	12.00
Zr	0.50	0.50	0.50	0.50	0.60	0.50	0.90	0.50	0.80	0.50
Cu	1.60	5.30	3.50	22.80	9.60	20.50	12.90	8.60	33.80	27.40
Pb	14.00	2.80	1.10	0.40	1.80	0.20	0.60	2.00	2.40	1.20
Cr	727.00	42.00	51.00	53.00	605.00	34.00	725.00	29.00	429.00	547.00
Ni	21.10	44.60	8.00	4.60	5.80	2.30	7.30	5.10	3.80	5.20
Sr	0.90	1.00	1.00	1.00	2.00	0.30	1.00	3.30	1.70	0.80
V	2.00	1.00	1.00	1.00	2.00	1.00	2.00	1.00	1.00	1.00
Nb	0.09	0.05	0.05	0.05	0.11	0.05	0.07	0.05	0.06	0.08
Mn	0.09	1.44	3.98	0.31	0.47	2.16	7.12	5.45	0.12	0.42
La	0.50	2.20	1.30	84.60	1.10	7.80	3.30	18.30	51.90	49.40
Ce	0.06	0.06	0.43	1.51	0.51	0.48	0.10	0.78	4.70	0.84
Pr	11.40	11.80	0.92	0.32	19.25	4.24	0.46	1.91	0.56	0.06
Nd	0.20	2.20	1.50	71.00	1.50	7.40	3.50	15.90	44.10	43.00
Sm	0.20	0.20	0.20	0.20	0.20	0.20	0.20	0.20	0.20	0.20
Eu	0.50	1.00	0.30	0.20	0.20	0.20	0.30	0.20	0.20	0.20
Gd	5.87	5.86	0.56	0.61	10.85	3.14	0.09	1.44	0.57	0.08
Tb	0.04	0.09	0.05	0.02	0.01	0.01	0.03	0.01	0.01	0.00
Dy	0.08	0.48	0.18	8.33	0.56	1.06	0.36	3.77	4.63	4.75
Y	0.05	0.05	0.15	0.15	0.08	0.05	0.06	0.44	0.36	0.07
Но	0.01	0.03	0.13	1.53	0.08	0.03	0.00	0.73	0.98	0.95
Er	0.04	0.32	0.04	4.48	0.31	0.76	0.34	1.90	2.91	2.71
Tm	0.04	0.03	0.01	0.64	0.04	0.11	0.04	0.27	0.48	0.41
Yb	0.01	0.03	0.01	0.04	0.04	0.01	0.04	0.27	0.48	0.41
Lu	0.03	0.04	0.03	0.04	0.01	0.01	0.03	0.01	0.01	0.01
ΣREE	19.08	0.02 24.46	5.79	173.64	34.73	25.71	8.95	45.88	111.62	102.70
		24.46					8.95 7.95		102.23	
ΣLREE SHREE	18.73 0.35	25.52 1.14	5.21	158.44	33.61	23.46 2.25		38.73	9.39	93.78 8.92
ΣHREE SIDEE/SHDEE	0.35 53.51	1.14 20.46	0.58	15.20 10.42	1.12 30.01	2.25 10.43	1.00 7.95	7.15	9.39 10.89	8.92 10.51
ΣLREE/ΣHREE K/Rb	0.24		8.98	0.09			7.95 0.01	5.42		10.51 58.12
		0.32	0.13		0.11	0.52		0.06	0.04	
K/Ba Ba/Bh	0.00	0.01	0.00	0.01	0.00	0.01	0.00	0.00	0.01	0.58
Ba/Rb	100.00	50.00	50.00	12.50	33.33	50.00	10.00	25.00	5.00	100.00
Rb/Sr	0.11	0.20	0.20	0.80	0.15	0.67	1.00	0.12	1.18	0.13
Co/Ni	0.51	0.61	0.51	0.24	0.26	0.13	0.34	0.24	0.26	0.25
Hf/Sm	0.25	0.65	3.50	1.60	0.45	0.80	3.30	3.30	1.35	0.95
Nb/La	0.18	0.02	0.04	0.00	0.10	0.01	0.02	0.00	0.00	0.00
Y/Ho	5.00	0.45	3.75	0.10	0.80	0.21	0.55	0.60	0.37	0.07
Th/U	4.00	4.00	4.00	2.86	4.00	4.00	4.00	2.50	11.67	4.00
(La/Yb)n	11.32	37.36	29.44	1436.77	74.73	529.87	44.84	1243.16		
(La/Sm)n	1.56	6.87	4.06	264.15	3.43	24.35	10.30	57.14		
(Gd/Yb)n	158.30	118.53	15.10	12.34	877.81	254.04	1.46	116.50		



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In the barren quartz veins, most of the major elements have concentrations less than 0.01 wt%, except for Al_2O_3 Fe₂O₃ and CaO which have higher concentrations. However, these veins have high silica concentrations. Therefore, the quartz veins are highly siliceous. Al_2O_3 content varies from 0.1 to 8.96 wt%, Fe₂O₃ ranges from 0.13 to 6.26 wt% while MgO varies from 0.02 to 1.74 wt%. The concentrations of other major oxides are very low.

The mineralized veins (QV01, QV02, QV05, and QV08) contain Au from 0.50 to 25 ppm, while the quartz veins considered barren contain 0.20 ppm Au. It is worth noting that samples QV01 (25 ppm of Au) and QV02 (4.30 ppm Au) were recovered from the quartz veins that outcropped on the surface and are about 10 meters apart. The gold concentration in the barren veins exceed the background (5ppb) and the Clarke's concentration (4ppb) in the study area. The low concentration of Au in the barren quartz veins shows that gold mineralization is not evenly distributed throughout the area.

The concentrations of other trace elements in the mineralized veins such as Zn, P, Ni, V, Sr, Pb, U, Rb, As and Cr varies from 3.0 to 43ppm, 10 – 40 ppm, 4.6 – 44.6ppm, 1-2ppm, 0.92ppm, 0.4-14ppm, 0.05-0.07ppm, 0.1-1ppm, 0.4-9.7ppm and 29 - 727ppm respectively. The concentrations of B, Ba, Ta and Tl are the quite equivalent in both mineralized and barren quartz veins. The concentration of As in the quartz veins ranges from 0.30ppm to 10.70 ppm and it is worth noting that As show positive correlation with Au in samples with high Au concentration. The concentrations of (Co+Ni+Cu+Zn) - Fe-Mn in quartz veins were plotted on a ternary diagram differentiate between submarine hydrothermal and hydrogenous deposits [16]. The data plotted indicate that the quartz veins bearing gold are of hydrothermal origin (Figure 7). However only the concentration of As (4ppm) exceeded the backgrounds limit for non-mineralized veins. The concentration of Pb was below the limit (20.7 ppm).

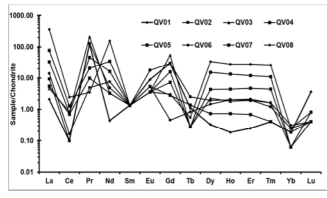


Figure 7. Chondrite - normalized REE patterns of the Quartz vein samples.

The ΣREE varies from 5.79 to 173.64 in the analyzed quartz veins (Table 1) with notable LREE (range from 5.21 to 158.44) and HREE (range from 0.35 to 15.20) fractionation, while the quartzite and schist samples content is 111.62 and 102.70 respectively. The (La/Yb)_N, (La/Sm)_N



and $(Gd/Yb)_N$ values ranges from 11.32 - 1436.77, 1.56 - 264.15 and 1.46 - 254.04 respectively. Chondrite normalized plots of the REE in the quartz veins is shown in Figure 8. The normalized pattern shows negative Ce, Nd, Tb and Yb anomalies with positive La, Pr, Ce and Gd anomalies.

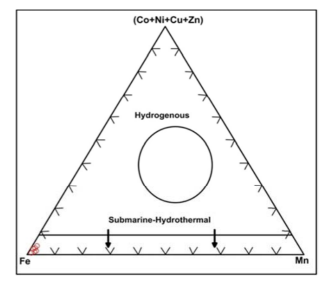


Figure 8. (Co+Ni+Cu+Zn)-Fe-Mn ternary plot of quartz veins used to differentiate submarine and hydrothermal and hydrogenous deposits.

4. Discussion

In many gold fields, gold is hosted by quartz veins [8, 17] and such is the case with the Belikombone gold prospect. The mineralized quartz veins observed showed cross-cutting relationships implying different stages of vein emplacement in the area. The fact that the mineralized quartz veins crosscut the schist in the area also indicate epigenetic style of mineralization. The poles to quartz veins plotted on the stereographic projection and rose diagrams of the quartz veins show that the quartz veins in the schists are generally trending NW-SE (Figure 6), and are concordant to the foliations in the schist, which also have a NW-SE trend. Studies within the Betare Oya gold district [15] to which Belikombone belong, generally trend in the NW-SE direction. The N-S and ENE - WSW trends being the dominating ones for the veins and thus gold mineralization within the Belikombone prospect is structurally controlled. The concordant nature of the quartz vein in this area to the foliation in the schist country rock indicates that the emplacement of the quartz was syngenetic. The emplacement of the hydrothermal quartz veins in the Belikombone prospect can globally be related to the NE-SW trending structures Lom basin structure (Figure 1), which provided pathways for hydrothermal fluid flow [18]. Hydrothermal fluids are controlled by a wide variety of pre-existing structures particularly foliations, shear zones and fractures [19, 20] and such structural relationships are common in orogenic zones, particularly in shear related deformation [21, 22]. Subsequent plutonism or metamorphism may cause the reactivation of these structures [23].

The mineralogy of the studied quartz veins is dominantly composed of quartz \pm sulphides, some sericite and iron oxides in the form of hematite or magnetite (mostly as fracture infill). According to Wegner et al. [24] massive quartz veins are formed during multiple episodes of vein opening, fluid interaction and vein sealing events, while brecciated quartz veins [25] are the result of alteration of earlier formed veins and represent a later event. The differences in terms of texture and mineralogy of the massive and brecciated quartz in the area suggest that they were formed during different hydrothermal events. The fact that most mineralized quartz veins are brecciated shows that hydro-fracturing (fluids assisted fracturing) is likely to have been the process that led to the formation of the breccias and was triggered by variations in fluid pressure [26] and according to Craw et al. [27], breccias are indicators of brittle deformation and have been considered typical to be of shallowly-formed veins. This could also be seen in the Belikombone quartz veins and implies that the quartz veins were formed at shallow depths and probably associated with brittle deformation. Figure 9 summarized the mineral paragenesis of quartz vein containing gold obtained from the study area. The mineral sequence of the study area comprises initial host rock formation which later experienced fracturing. This is subsequently followed by the introduction of hydrothermal mineralizing fluid composing of quartz, gold, sulphide and magnetite. Some primary minerals like biotite and amphibole in the quartz veins were later altered to chlorite and sericite. The development of quartz, sericite, and pyrite alteration minerals in the quartz veins confirms that silicification, sericitization, and sulphidation processes are associated with gold mineralization.

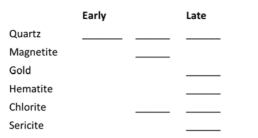


Figure 9. Summarized mineral paragenesis of quartz vein containing gold obtained from this study.

Gold concentrations in the Belikombone quartz veins vary considerably (0.2 - 25.0ppm). The concentration of gold in sample QV01 is quite high to qualify as a potential ore grade. Economic occurrence of gold generally consists of very small amounts of dispersed gold or gold-silver alloys. Even in the well-known ore of the Witwatersand deposit in South Africa, the average concentration of gold is only about 16000ppb [18]. Where there are considerable variations in gold concentrations, there is the possibility of an existence of high-grade ore shoots within a largely low-grade gold system [6]. Most quartz vein samples that have high gold content equally have high arsenic concentration and the appreciable As signature further confirms that the quartz veins were formed in an orogenic setting [28]. The association of Au with As therefore implies that As could be utilized as an



effective geochemical pathfinder element for gold exploration in the study area. The concentration of Sb (0.12 to 0.39 ppm) is below the minimum concentration of 0.59ppm in unmineralized materials (1 ppm) in all the samples analyzed and it is known to be associated with gold deposit. Au, Ag, As, Sb, Pb, Zn, W, and Mo in quartz veins and host rocks display great positive correlations which are obvious between Au and Ag, Pb, Zn, and Sb with Au, suggesting that the enrichment of Au, Ag, As, Sb, Pb, and Zn may have synchronously taken place. In addition, hightemperature elements such as W and Mo show positive correlations with Au. Anomalies of W and Mo are often correlated with the activity of magmatic hydrothermal fluids [29]. The trace element composition of quartz veins possibly reflects the element features of ore-forming fluids [29]. It can be concluded that ore-forming fluids of the Belikombone deposit were enriched in sulphophile elements such as Au, Ag, As, Sb, Pb and Zn and simultaneously concentrated hightemperature elements such as W and Mo. At the same time, enrichment in W and Mo reflects multi-stage mineralization or overprinting of magmatic hydrothermal fluids.

Recent studies suggest that hydrothermal fluids enriched in Cl can effectively concentrate light REE but deplete HFSE, whose values of Hf/Sm, Nb/La and Th/La are usually less than 1; and hydrothermal fluids enriched in F can synchronously concentrate light REE and HFSE, whose values of Hf/Sm, Nb/La and Th/La are usually more than 1 [30]. According to the analytical results (Table 1), quartz veins and host rocks of the Belikombone gold prospect are obviously enriched in light REE and universally depleted in HFSE, with the values of Hf/Sm, Nb/La and Th/La being less than 1. This suggests that the hydrothermal fluids responsible for the emplacement of quartz veins in the studied area were enriched in Cl.

Field and petrographic evidences from the Belikombone area indicate the occurrence of gold bearing quartz veins hosted in schists, granites and gneisses. Similar observations elsewhere e.g. Bini Yauri [31] and Maru [18] concluded that magmatic fluid or recirculated groundwater may be part of the ore constituents at some stage of vein evolution and accompanied alteration. The ore fluid may largely have been derived from fracturing and metamorphic dewatering [31] of the host rocks. There are very few auriferous Archean greenstone belts that contain gold provinces without nearby intrusions of roughly the same age. Whether or not any of these igneous bodies are the source of fluids and metals that become a part of the gold-forming system is often a highly debated issues; the other obvious scenario is that both melts and fluid may be product of the same deep-crustal or even mantle generated, thermal event [20, 32].

5. Conclusion

Au-bearing quartz veins in the Belikombone area are confined to major NW-SE orientations in schists and N-S and ENE – WSW in other host rocks. These veins are characterized by partially and completely filled fractures and foliation planes. This therefore constitute the main mineralization style within the area. Both the mineralized and non-mineralized (barren) quartz veins varieties occur in this area. The studied quartz veins are dominantly composed of quartz \pm sulphides, some sericite and iron oxides in the form of hematite or magnetite and were formed during different hydrothermal events. Hydro-facturing is likely to have been the process that led to the formation of brecciaed veins.

The investigated veins show evidence of multiphase deformation as indicated by the mineralogical, textural and cross-cutting relationships among veins. The emplacement of the quartz veins is associated with D_2 deformation as indicated by similarity in the generalized strike direction of quartz veins and host rocks. It postdates regional metamorphism and fracturing. This structural setting suggest that the emplacement of gold mineralization occurred during the late Pan African orogeny. The quartz veins have simple mineralogy dominated by quartz and sulphides. The main alteration types are silification, sericitization and sulphidation processes.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

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