Elucidating Pathfinding Elements from the Kubi Gold Mine in Ghana

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Abstract: X-ray photoelectron spectroscopy (XPS) and energy dispersive X-ray spectroscopy (EDX) are applied to investigate the properties of fine-grained concentrates on artisanal, small-scale gold mining samples from the Kubi Gold Project of the Asante Gold Corporation near Dunwka-on-Offin in the Central Region of Ghana. Both techniques show that the Au-containing residual sediments are dominated by the host elements Fe, Ag, Al, N, O, Si, Hg, and Ti that either form alloys with gold or with inherent elements in the sediments. For comparison, a bulk nugget sample mainly consisting of Au forms an electrum, i.e., a solid solution with Ag. Untreated (impure) sediments, fine-grained Au concentrate, coarse-grained Au concentrate, and processed ore (Au bulk/nugget) samples were found to contain clusters of O, C, N, and Ag, with Au concentrations significantly lower than that of the related elements. This finding can be attributed to primary geochemical dispersion, which evolved from the crystallization of magma and hydrothermal liquids as well as the migration of metasomatic elements and the rapid rate of chemical weathering of lateralization in secondary processes. The results indicate that Si and Ag are strongly concomitant with Au because of their eutectic characteristics, while N, C, and O follow alongside because of their affinity to Si. These non-noble elements thus act as pathfinders for Au ores in the exploration area. This paper further discusses relationships between gold and sediments of auriferous lodes as key to determining indicator minerals of gold in mining sites.

Keywords: gold; garnet; X-ray photoelectron spectroscopy; electron dispersive X-ray spectroscopy; chemical bonding

1. Introduction

Gold mineralization in the Kubi Gold Project of the Asante Gold Corporation near Dunkwa-on-Offin in the Central Region of Ghana is mainly controlled by the silicate type of the mineral garnet \((A_3B_2(SiO_4))_3\), where the A site represents the divalent cations (Ca, Mg, Fe, Mn)\(^{2+}\) and the B site has trivalent cations in either an octahedral or tetrahedral milieu, with \((SiO_4)^{4-}\) occupying the tetrahedral sites [1,2]. Sulfide minerals such as pyrite/marcasite (FeS\(_2\)), arsenopyrite (FeAsS), pyrrhotite (Fe\(_{1-x}\)S), and chalcopyrite (CuFeS\(_2\)) are other important pathfinder minerals for the location of the Au ore and help to understand the geology of Kubi.

Traditionally, coarse-grained concentrates are used in gold production in most large-scale mining, while the fine-grained powder sediments are considered part of mining waste (tailings). Artisanal and small-scale gold mining, without the use of mercury, also depends on the crushing or milling of Au particles into similar grain sizes, which can easily be separated by using gravity concentration methods, i.e., panning, sluicing, shaking tables, spiral concentration, vortex concentration, and centrifuges [3]. It will therefore be economically viable to extract gold and other minerals from fine-grained materials under realistic conditions.
Indicator and pathfinder theories are deployed by geologists and explorers to gain information on the location of ore deposits, using methods such as gold grain morphology (changes in Au surface shape via weathering and erosion as sediments are transported from far distances), gold grain inclusion (presence of other minerals in gold grains are used to provide information about the deposit type and mineralization associated), composition studies (to gather and examine mineral samples for information on distances and direction of transport), and geochemistry (identifying and analyzing minerals such as silver, platinum, palladium, copper, lead, iron, telluride) to test clues and patterns in order to locate ore bodies for gold [4,5].

To date, researchers have tried to understand the nature and properties of the various elements associated with gold as well as their chemical bonds and reaction mechanisms in relation to gold deposition and agglomeration [6–9]. Several studies indicate that during ore deposition in hydrothermal systems, there are surface reductions or adsorption or both in gold precipitation [10–13].

From a geological perspective, it is important to study the diffusion, flow mechanism, and chemical bonding of host (indicator) minerals to understand the dynamics and evolution of the Earth and other terrestrial planets. These features can influence the mantle convection, earth processes (erosion, weathering, landslide, volcanic eruption, earthquake, etc.), and also give information on the Earth’s thermo-chemical structure. This will enable us to understand the geodynamics of the Earth, its rheological properties (material deformation), kinetics of phase transformation, and the thermal conductivity of minerals on the planet.

Gold-associated minerals such as the sulfide group FeS₂, FeAsS, CuFeS₂, Fe₁₋ₓS, and galena (PbS) make the tiny particles of gold invisible or cause them to attach to the metallic sulfide lattices [14–18]. Pyrite is one of the most common minerals that conglomerate gold in various ore deposits; it is also part of the gold-bearing minerals (as free milling or refractory) in the Au exploration areas, alongside garnet and the sulfide group [6,7,14].

Previously, X-ray photoelectron spectroscopy (XPS), scanning electron microscopy (SEM), electrochemical, and other techniques have been deployed in the study of gold deposition on pyrite and other gold-bearing minerals in aqueous solutions [7,8,16]. Pathfinder minerals for Au such as chalcopyrite and pyrite minerals in their natural states were studied using cyclic voltammetry, electrochemical impedance spectroscopy, XPS, and SEM, which also included microanalysis (SEM/EDX). In both minerals, oxygen containing Fe³⁺ and Cu²⁺ were found, while the measurements also showed alteration in both minerals at different potential values [19]. Harmer et al. [20–23] studied the rate of leaching of Cu and Fe in chalcopyrite when percolated in perchloric acid and found the leach rate to be the same after a long period of time; the final analysis was based on the surface speciation and oxidation steps in the release of Cu and Fe into solutions and their polymerization from mono-sulfide to polysulfide. Wang et al. [24] used high-resolution transmission electron microscopy and XPS to characterize the morphological change in Au nano-rods (NRs) from artisanal small-scale mining in order to understand the concentration of mercury. It was found that the low concentration of mercury that formed and was deposited onto the gold surface could be reversibly removed by an electrochemical stripping process without causing any change in terms of size or shape to the Au NRs. However, at high concentrations, Hg did not only deposit onto the gold surface, but also entered the interior of Au NRs and transformed them into irreversible gold nanospheres (Au NSs) due to the amalgamation of Au with Hg. Since the assembling of Au with Hg by small-scale artisanal miners has significant negative impacts on the health of individuals and the environment, there is a need for a more environmentally sustainable separation mechanism.

XPS has been a contributory technique [25] in the study of the products of oxidation [26] on sulfide minerals as well as in the use of synchrotron radiation [27]; it is an added advantage in explaining mineral surface chemistry (mineral geochemistry) [28–30], predominantly the primary surface conditions of materials formed by deformations and cracks [30–32]. The main advantage of XPS and resonant photoemission
using synchrotron radiation in mineralogy is their ability to define comprehensive surface-sensitive information from high-resolution spectra of samples of macroscopic sizes in different mineral phases. The distribution of oxidation products provides heterogeneity to mineral surfaces, and because most gold-bearing sulfide minerals contain other minerals and impurities with varying texture, it will be of interest to reveal the individual elements by XPS, which can then be assigned to the original host minerals. It is therefore essential to identify the existence of the different types of elements on both the bulk and impure sample surfaces using high resolution XPS, and determine the distribution of those elements across the sample surfaces [33–35].

Mikhlin et al. [36] used XPS, AFM, and scanning tunnelling microscopy/scanning tunnelling spectroscopy (STM/STs) to characterize arsenopyrite samples oxidized in Au (III) chloride solutions at ambient temperatures in order to explain the moderation of oxidation in chloride solutions. Murphy and Strongin [37] conducted surface reactivity studies on pyrite (FeS₂) and pyrrhotite (FeₓS₁₋ₓ) using different experimental techniques such as vacuum type experiments, where electron and photon spectroscopies were applied and further analyzed microscopically using infra-red spectroscopy. XPS and X-ray absorption spectroscopy techniques were also used to investigate the structure of the original pyrite and sulfide surfaces [37,38]. Acres et al. [39] applied synchrotron-based XPS, near-edge X-ray Absorption Florescence Spectroscopy, and time of flight secondary ion mass spectrometry studies on smooth and rough fracture surfaces of chalcopyrite and pyrite samples. The results showed an increase in the chalcopyrite surface roughness and an intensification in the formation of the sulfur surface in the pyrite as well as the typical chalcopyrite grain size formation [39].

Other studies on Au and associated minerals have concentrated on the valence band electronic structure using nanoparticle sizes on NiO, TiO₂, and Al₂O₃ [24,27,36,37] using ultraviolet photoelectron spectroscopy (~40 eV). Likewise, conventional Al Kα X-ray XPS (1486 eV) was used to study the electronic structure of Au nanoparticles on carbon [40–44]. The results from these studies indicate that a metal-insulator transition occurs as a function of size between Au atoms, which is attributed to quantum size effects.

In order to understand the relationships between Au and associated minerals, there is the need to study the chemical bonding and properties of elemental species in these indicator minerals. This will aid in determining the features of the elements’ concentration and diffusion during ore formation and the flow mechanism in hydrothermal systems ascribed to Au host minerals. Despite increasing knowledge that indicator minerals can host numerous elements, most geologists in the study area have mainly concentrated on host minerals for Au such as arsenopyrite, pyrite, garnet, and quartz, which occur in orogenic and sediment-hosted and disseminated gold deposits known as the Carlin type [45].

Bayari et al. [46] used X-ray florescence spectroscopy, XRD, and inductively coupled plasma mass spectrometry to identify pathfinder elements and their associated pathfinder minerals in the mineralized regolith profiles at the Bole–Nangoli gold belt in north-eastern Ghana. Moreover, Nude et al. [47] used a multivariate statistical approach to identify pathfinder elements for gold from the north-west of Ghana. In both cases, the trace elements such as Fe, Mn, Ag, As, Cu, Zn, Ni, Pb, etc., were identified and appeared to be associated with Au and suitable as pathfinder elements of Au in the study areas.

Up until now, there is little or no work on pathfinder elements and minerals in Ghana using XPS and EDX, and in particular, in the Kubi Gold mining area. Rock and mineral identifications leading to Au discovery in this exploration area have been based on geophysical techniques and geochemical analysis (soil/sediments sampling) in laboratories as well as geologists’ experiences in mineral identification. However, these procedures are sometimes misleading and may cause delays in projects and high financial loss to mineral exploration activities. Therefore, there is a need to introduce faster and more robust techniques for the effective identification of pathfinder elements and minerals in the mining industry.
In this present work, fine-grained sediments (mining waste) from artisanal small-scale mining sites are investigated by surface sensitive XPS and bulk-sensitive EDX to quantify their Au content and other important elements that act as indicator minerals of gold. We investigate Au samples from alluvia deposits to identify individual elements in order to gain corresponding insight into surface reaction mechanisms as well as detailed properties of precipitated Au particles observed during the hydrothermal formation of the ore deposit. These identifiable pathfinder elements are then utilized to infer the original indicator host minerals in the area.

2. Experimental Details
   2.1. Description of Study Area

Samples for the study were taken from the alluvia small-scale mining site located in the Dunkwa-on-Offin district in the central region of Ghana. Figure 1 shows the district that is located within latitudes 5°30′ and 6°02′ north of the equator and longitudes between 1° W and 2° W of the Greenwich Meridian [48]. Close to the vicinity of the Kubi Gold Barbados is the alluvia mining site, where the crushed and washed samples for the project were collected. The area is drained by a number of rivers and streams, including the Offin River as the main river source, which is 90 m above sea level, forming a boundary between the Ashanti region and the Central region with the Pra River [49]. The area consists of volcanic belts, basement rock, and sedimentary basins with a lot of igneous (volcanic) activity and two orogenic events, genuinely following the geology of Ghana and leading Au-hosted quartz veins to dip steeply in shear zones with Birimian basins of sulfur-rich minerals. The gold at the Kubi mining area is free milling (open cast mining) and occurs within a near vertical 1 km by 1 m, to 15 m thick shear confined garnet zone. The ore is estimated to be hosted by 30% garnet, 15% sulfide mineralization, and also occurs as coarse gold in late quartz veins.

In addition, the Offin river has alluvia placer Au deposits in gravels and in quartz-pebble conglomerates of the Tarkwaian deposits [48]. Within the alluvia regime are oxides or weathered rocks of the host minerals, which are mostly iron oxides and mineralized quartz [50].
Figure 1. (a) Map of Kubi village and surrounding Townships and (b) Picture of sampling area of “Kubi Gold Barbados”, Dunkwa-on-Offin. (Map and photo by G.K. Nzulu).
2.2. Behavior of Gold in Hydrothermal Systems

Figure 2a shows a schematic profile of the sampling area (Kubi soils). The saprolite (altered by hematite and limonite/goethite) that dominates the profile grades upwards into the transition and continues to the clay zone, which consist of ferruginous materials, mostly of goethite, kaolinite, and chlorite. Within the stone line horizon are fragments of quartz (pebble and cobbles) and gravels mixed with ferruginous clay that have been formed by the weathering profile. The Au in this soil is due to weathering activities and in association with the sulfide group of minerals and Fe-oxides. The smaller Au grains found a few meters from the surface occur beyond the stone line lateritic zone through chemical dissolution [51]. Thus, beyond the stone line, Au mineralization occurs with much coarser Au grains as a result of pedoturbation (soil horizon mixing) that goes downward due to the migration of fragments from the stone line, through the transition zone and to the saprolite, and vice versa.

Figure 2b shows the oxidation and reduction processes that occur when pathfinder minerals undergo chemical dissolution to deposit gold above and below the water table. Au formed on the oxides (near surface) and on external particles are usually from a mixture of chemical reactions and microbiological activity in groundwater of sedimentary origin, with a pH ranging from 6 to 9. During the flow regime within the hydrothermal system (zone of aeration), the rich sulfide minerals (pyrite group) and the oxide groups can release metastable thiosulphate ions ($S_2O_3^{2−}$) during the oxidation process to form a composite material with Au under near-neutral-alkaline pH conditions to trigger a dissolution of Au particles within the sediments/minerals [52,53]. These thiosulphate ions can decompose after a short period to aid in the oxidation of sulphate ions formed above the water table, or to aid in the reduction of dissolved hydrogen sulfide ions formed below the water table (zone of saturation) [53]. Sulphate minerals found in underground water undergo a reduction process to form bisulphide ions ($HS^{−}$) near the water table in order to assemble Au in a sedimentary environment under near-neutral pH conditions. This same chemical process can facilitate the precipitation of Ag with Au under favorable pH conditions, to deposit gold together with a small quantity of Ag [52,53]. Thus, under alkaline conditions,
the most soluble complex of the Au-Ag-S-O system is the Ag(S\textsubscript{2}O\textsubscript{3})\textsubscript{2}\textsuperscript{3−} ion, such that the separation of Ag and other elemental species from Au at high pH levels is always a possibility [54].

2.3. Sample Preparation

Sediment extracted from a depth of 10 m and containing Au samples from the artisanal small-scale mining site in Dunkwa-on-Offin was investigated in this study. The sample was collected by one of the authors (G.K.N.) with the aid of a geological hammer and placed into a sample collection bag. The composite “concentrate sample” weighing 1.90 kg, as shown in Figure 3a, was divided into two parts. One portion (1.20 kg) was refined into pure solid Au (bulk Au), while the other powder sample (0.70 kg) was subjected to panning, i.e., washing (Figure 3c) and extraction of residual sediments of Fe\textsubscript{3}O\textsubscript{4}, Fe\textsubscript{2}O\textsubscript{3}, and other minerals to obtain an untreated Au powder concentrate with a weight of 80 g (Figure 3d). This untreated powder sample was later separated into coarse-grained (0.85 g) and fine-grained (1.20 g) with the aid of a tweezer and water panning (via gravity), respectively (the residual 77.95 g containing sand and gravel was disposed). The impure powder sample (Figure 3d) was first investigated by XPS, after which a tweezer was used to pick the average-grain-sized particles as coarse-grained powder sample. The final residual sample was water-panned for tiny particles of Au, quartz, and the pyrite group as fined-grained powder samples.

The final three samples, namely solid Au (bulk), untreated Au concentrate powder, fine and coarse-grained Au concentrates in Figure 3b,e,f, respectively, in addition to a standard thin film Au sample in Figure 3g, were examined by XPS to reveal distinct features of the materials. Particle/grain sizes of the powder samples ranged from 0.05 cm to about 0.2 cm in maximum dimensions of which most were hoppered single crystals. The nature of materials indicates that most Au from this artisanal area come from weathered SiO\textsubscript{2} veins hosted by coarse-grained igneous and meta-volcanic rocks, with mineralization controlled by a garnetiferous horizon of fine-grained Au associated with sulfide minerals and occasional carbonates [50].
Figure 3. (a) Dry residual sample (tailing) from an artisanal mine site at Dunkwa-On-Offin. (b) Part of the sample refined into pure Au nugget of 1 gram and 22 carats (measured with a digital electronic Au purity Analyzer D H 300 K from VTSYIQ). (c) Panned sample of Au with other impurities, (d) untreated ore powder, (e) fine-grained concentrate Au, (f) coarse-grained concentrate powder, and (g) Au film on wafer substrate. (Photo by G.K. Nzulu).

2.4. XPS Measurements

XPS was used to analyze the elemental composition and chemistry of the powder samples (untreated, fine, and coarse-grained Au ore) and reference materials (bulk nugget and a 300 nm thick Au film with a 2.5 nm Ti buffer layer on a Si wafer substrate). The analyses were performed in an Axis Ultra DLD instrument from Kratos Analytical (UK), employing monochromatic Al Kα radiation (hν = 1486.6 eV) and operated at a base pressure lower than 1.1 × 10⁻⁹ Torr (1.5 × 10⁻⁷ Pa) during spectra acquisition. XPS depth profiles...
were acquired by sputter-etching with 0.5-keV Ar$^+$ ions incident at an angle of 70° relative to the sample surface normal. The low Ar$^+$ energy and shallow incidence angle were chosen to minimize the effect of sputtering damage in the spectra. The sample areas analyzed by XPS were $0.3 \times 0.7$ mm$^2$ and located in the center of $3 \times 3$ mm$^2$ ion-etched regions. The binding energy (BE) scale was calibrated using the ISO-certified procedure, with the spectra referenced to the Fermi edge in order to circumvent discrepancies associated with employing the C 1s peak from adventitious carbon [55]. Prior to the XPS core and valence band measurements, the structural properties were investigated [56]. After XRD analysis [56], Au, quartz (SiO$_2$), magnetite (Fe$_3$O$_4$), and hematite (Fe$_2$O$_3$) were identified in these samples.

2.5. EDX Measurements

The bulk, untreated, fine- and coarse-grained concentrate samples were analyzed using SEM/EDX operated at 20 kV. The EDX used in this study is of the Zeiss Supra 35 VP field emission SEM type whose field emission scanning electron microscopy (FESEM) is equipped with an energy dispersive X-ray spectroscopy (EDX) detector, which was used for the elemental mapping of the samples to understand the elemental distribution. Samples were prepared by mounting the macrocrystalline samples onto aluminum stubs with carbon sticky tabs.

3. Results and Discussion

3.1. Core-Level XPS

Figure 4 shows the XPS survey spectra of Au in the form of thin film, bulk, and powder (untreated, fine-grained, and coarse-grained) concentrate samples, which are vertically offset for clarity. The observed peaks of the thin film are identified as Au 4f$^{7/2,5/2}$ at 84 and 87 eV, Au 4d$^{5/2,3/2}$ at 335 and 353 eV, Au 4p$^{3/2,1/2}$ at 546 and 642 eV, and Au 4s at 762 eV. The impure powder sample was first investigated by XPS, after which a tweezer was used to pick the average-grain-sized particles as coarse-grained powder sample. The final residual sample was water-panned for tiny particles of Au, quartz, and the pyrite group as fined-grained powder samples.
Figure 4. XPS survey spectra of film, bulk, and powder (untreated, fine, and coarse-grained concentrate) Au. The broken line connects the band edges as well as the apparent spin-orbit-splitting maxima. VB denotes valence band.

In the Au bulk sample, we can also identify all the Au peaks as well as peaks arising from Ag $3d_{5/2,3/2}$ at 368 and 374 eV, Ag $3p_{3/2,1/2}$ at 573 and 604 eV, and Ag $3s$ at 719 eV. In addition, there are C $1s$ and O $1s$ peaks at 285 and 532 eV, respectively. In the untreated concentrate powder sample, in addition to the Au peaks, we find a Ti $3p$ peak at 37.50 eV, Ti $2p_{3/2,1/2}$ peaks at 458 and 461 eV, respectively, and an N $1s$ peak at 397 eV, as listed in Table 1 and consistent with reference data [57].

Table 1. Elements identified in the analysis of XPS core levels for Au bulk and powder (untreated, fine-grained, and coarse-grained) Au concentrate samples.

| Element | Spectral Line | Formula | Energy (eV) | Parent Mineral |
|---------|---------------|---------|-------------|----------------|
| Au      | 4s            | Au      | 762.0       | Gold           |
| Au      | $4p_{3/2,1/2}$| Au      | 546.0/642.0 | Gold           |
| Au      | 4d            | Au      | 335.0       | Gold           |
| Au      | $5p_{1/2}$    | Au      | 57.20       | Gold           |
| Au      | $4f$          | Au      | 84.0/87.0   | Gold           |
| Ti      | 3p            | TiO$_2$ | 37.50       | Hematite, garnet, and other silicate minerals |
| Ti      | 2s            | TiO$_2$ | 561.00      | Hematite, garnet, and other silicate minerals |
| Ti      | $2p_{3/2,1/2}$| TiO$_2$ | 458.0/464.19| Hematite, garnet, and other silicate minerals |
| Si      | 2p            | SiO$_2$ | 102         | Quartz         |
| Si      | 2s            | SiO$_2$ | 153.0       | Quartz         |
| S       | 2p            | S       | 163         | Au-S/FeS$_2$   |
Table 1. Cont.

| Element | Spectral Line | Formula | Energy (eV) | Parent Mineral |
|---------|---------------|---------|-------------|----------------|
| S       | 2s            | S       | 231         | Au-S/FeS₂      |
| Ag      | 3p₁/₂,3/₂     | Ag      | 604.0–573.0 | Silver         |
| Ag      | 3d₃/₂,5/₂     | Ag      | 368.0–374.0 | Silver         |
| Ag      | 3s            | Ag      | 719.0       | Silver         |
| N       | 1s            | N       | 397.0       | Nitrogen       |
| O       | 1s            | O       | 532.0       | Oxides         |
| C       | 1s            | C       | 285.0       | Carbon/graphite|
| Fe      | 2p₃/₂         | Fe/Cu   | 706.90      | Iron/FeS₂      |
| Fe      | 2p₁/₂         | Fe/Cu   | 720.0       | Iron/FeS₂      |
| Hg      | 2p            | Hg      | 104         | Hg             |
| Hg      | 4p₁/₂         | Hg      | 577         | Hg             |

In the present experiment, sulfur is very diluted in the samples due to the preparation method that makes the detection of sulfur by XPS very challenging (low sensitivity). Only a very weak S 2p peak around 165 eV was observed in untreated powder, while in the fine-grained sample it was hardly discernable. In the other samples, it was below the signal-to-noise level. The S 2s cross-section at 231 eV is even lower than the S 2p signal.

The Fe in the powder (untreated, fine, and coarse-grained concentrates) and Au bulk samples has 2p₃/₂,1/₂ peaks located at 706.90 and 720 eV, respectively, consistent with the literature [58,59] and in agreement with reference data [57]. The Fe 2p₁/₂ peaks are closely located at the same BE of 719 eV as the Ag 3s peak [57].

The measured XPS data of Au 4f XPS spectra of the Au film, Au bulk, and untreated concentrate powder of Au. For the Au film and bulk samples, the 4f₅/₂ peaks are located at 87.63 eV and the 4f₇/₂ peaks are located at 83.95 eV, with a 3.6 eV spin-orbit splitting. These binding energies are similar to the literature values of 87.60 and 84.0 eV [57] for pure Au.

For comparison, in the Au powder sample, the 4f₅/₂ is located at 87.77 eV and the 4f₇/₂ peak at 84.09 eV. The Au 4f peaks are asymmetric, while the shape of the untreated Au powder concentrate, and the coarse-grained Au powder concentrate spectrum broadens with tails toward higher BE. The chemical shift of 0.14 eV towards higher BE for the powder samples suggests that there is a charge transfer from Au towards the other elements. The high-energy shift can be attributed to the screening effect in comparison to the pure bulk Au and suggests total charge-transfer from Au towards certain elements.

From the spectra in Figure 5, the Au 4f region has well separated spin-orbit components of 3.6 eV, which is close to the Au metal (Δ = 3.7 eV) and is in agreement with the literature [62]. The Au 4f₇/₂ peak at 84.0 eV serves as a useful BE reference for the Au metal [57] and is expected to shift (BE shift) to Au nanoparticle sizes for smaller cluster samples.
Figure 5. Au 4f core-level XPS spectra of the Au film, Au bulk, fine-grained Au concentrate, and coarse-grained Au concentrate. The dashed line connects the 4f peaks as well as the apparent spin-orbit-splitting maxima. The spectra containing lower Au concentrations were scaled-up by factors of 3 and 30 for comparison.

The spectra from Figure 6 shows the Ag 3d_{5/2,3/2} core level XPS of the bulk and untreated Au powder, fine-grained Au, and coarse-grained Au concentrate samples. The peaks for the Ag 3d_{5/2,3/2} levels are located at 374.0 eV and 368.0 eV, respectively, with respect to the Fermi level (E_F) and in reasonable agreement with the literature data [57].

Figure 6. Ag 3d_{5/2,3/2} core level XPS spectra of the bulk Au and powder (unprocessed and processed) Au samples. The dashed line connects the peaks as well as the apparent spin-orbit-splitting maxima.
The Ag 3d_{5/2,3/2} spectra in Figure 6 show well-separated spin-orbit components ($\Delta_{\text{metal}} = 6.0$ eV); the peaks display asymmetric peak shapes for the Ag metal and also show loss features in the higher BE side of each spin-orbit component of the Ag metal [63]. The bulk Au and fine-grained Au powder concentrate, show higher intense peaks due to the high number of Ag atoms present in the samples, interacting with unfilled electron levels of the valence band above $E_F$. The lower intense peaks and the loss of features for the coarse-grained Au concentrate sample suggest a negligible number of Ag atoms present in the material. The smaller BE shifts observed in the spectra suggest the presence of oxides in the materials causing the Ag 3d_{5/2,3/2} peaks to broaden with respect to metal peaks as a result of a difference in electronegativity (lattice potential, work function changes, and extra-atomic relaxation energy), which then causes a charge transfer from the metal atoms to the ligands [64–66].

The reason that the Au and Ag alloy forms an “electrum” in the bulk and powder samples is due to their partly filled d bands (d-hole count effect) and having the same fcc structure makes it possible to exhibit relativistic effects where both bands display a correct spin-orbit-splitting signature. The structural changes that occur during the formation of the electrum affect the d-hole count in both Ag and Au and increase with decreasing Au concentration, which provides a strong correlation with the ostensible Au 5d spin-orbit splitting, lower d-band energy, and the Au 4f_{7/2} BE shift [67].

From the XPS data, the Fe 2p in the sample corresponds to pure Fe metal, with the 2p_{3/2} peak at 710.9 eV and the spin-orbit splitting at 13.6 eV. These values are close to those obtained from the coarse-grained Au concentrate sample, with $2p_{3/2} = 706.9$ eV and spin-orbit components ($\Delta_{\text{metal}} = 13.1$ eV). The Fe 2p peaks in Figure 7 show an asymmetric shape for metal, but the Fe 2p_{3/2} spectrum is not well-resolved to exhibit multiplet splitting. A high BE shift due to screening in comparison to pure metal shows that there is charge transfer from the metal atoms to the ligands. If the Fe in the samples originated from the oxide zone (FeO and Fe$_2$O$_3$, and Fe$_3$O$_4$), the peaks would have significantly shifted towards higher BE, but the actual shift towards lower BE (~710.9 eV) suggests that the peaks originated from pure Fe metal or pyrite (FeS$_2$), as opined by Biesinger et al. [68]. Iron compounds may possess high or low spin, with Fe (III) compounds exhibiting high-spin, which leads to a complex multiplet-split [68]. Fe (II) compounds, however, have both high and low spin, with spectra from the low spin lacking multiplet splitting and are attributable to the marcasite (FeS$_2$) or pyrite group of compounds [68]. According to Uhlig et al. (2001), for air-exposed FeS$_2$, the Fe$^{2+}$ is in a low-spin configuration and the broad structure towards higher BE (710.5 eV) of the main Fe 2p_{3/2} peak is due to the oxidation of the Fe$^{3+}$ states of the surface [69]. For the smaller marcasite peak, the higher BE (708.4 eV) is due to the electronic-deficiency of Fe$^{2+}$ sites, which are formed by the breaking of Fe–S bonds [69]. This higher BE (708.4 eV) and the lower BE for marcasite FeS$_2$ is in close agreement with the peak at (706.9 eV) from the XPS spectra in Figure 7, which is consistent with reference data [70] for FeS$_2$. It can be deduced that the deficient main Fe 2p_{3/2} peak in the samples from the Kubi Gold project contains pure metal Fe and FeS$_2$, since FeS$_2$ is one of the main indicator minerals in the mining area [56]. As telluric iron is extremely rare, we assume that the pure metallic Fe observed by our XPS measurements originates from iron oxides that are readily reduced in the argon monomer sputter cleaning process.
The assigned peak values of Ti in Table 1 are in good agreement with reference data [62] and consistent with the literature [71–74]. The abundant titanium in the powder sample is believed to be contained in fine-grained aggregates of hematite [75–77], which is a gangue mineral in this mining area [56]. The silicate garnet minerals that control Au mineralization in Kubi contain some percentage of TiO$_2$ [59]; on the other hand, other occasionally encountered silicate minerals such as biotite, hornblende, and the abundant silica (quartz), as well as some oxide groups in this area containing some TiO$_2$ are responsible for the significant Ti in the untreated Au sample. The assigned peak values of Si in Table 1 are in close agreement with reference data [62] and consistent with the literature [78,79]; it is also believed to be contained in the SiO$_2$, which is another indicator mineral in the mining area. Finally, the assigned peak positions of Hg, a trace element in the untreated powder sample that occurs at binding energies of 104 eV and 577 eV, overlap with Si 2p and Ag 3p peaks. These values are in agreement with reference data [62] and consistent with the literature [80,81].

As shown in Table 2, quantitative XPS analysis shows that the Ag content in the Au bulk nugget is about 6 at.%, while the C, O, and Fe contents are 3 at.%, 2 at.%, and 1 at.%, respectively. In the fine-grained Au powder concentrate sample, the Ag content is at.% half of that of the bulk sample, while the levels of C and O are significantly higher.

| System             | Au4f (at.%) | Ag3d (at.%) | Cls (at.%) | O1s (at.%) | Fe2p (at.%) | Mn (at.%) |
|--------------------|-------------|-------------|------------|------------|-------------|-----------|
| Au bulk            | 88          | 6           | 3          | 2          | 1           | -         |
| Fine-grained Au    | 32          | 4           | 7          | 32         | 24          | 3         |
| Coarse-grained Au  | 6           | 1           | 27         | 34         | 26          | 6         |
3.2. Chemical Bonding

Figure 8 shows valence band XPS spectra of an Au film, Au bulk, unprocessed ore powder, and processed (coarse and fine grained) ore powder of Au in comparison to calculated Au spectra. The agreement with the calculated peak positions in the valence band is improved when experimental lattice parameters ($a = 4.078$ Å) are used, while the theoretical (relaxed) lattice parameter of 4.156 Å is larger and results in peak positions that have extremely low BE.

The valence band spectra in Figure 8 show the coordination complex Au (transition metal) with an octahedral geometry where the $d$ orbitals split into a $t_{2g}$ set (lower energy) and an $e_g$ set (higher energy) [82,83]. From the spectra, the $e_g$ set moves towards higher BE due to the high involvement of the $d$ orbitals in the $e_g$ set in a metal–ligand sigma ($\sigma$) interaction. These orbitals in the $e_g$ are directed towards the axis where they are opposed by strong repulsive forces from ligands; hence, they acquire higher energy than the $t_{2g}$, thus favoring covalent bonding in $e_g$ towards higher energy [83]. The shift in BE along the $e_g$ set for the powder sample (coarse-grained) is an indication of the covalent bonding of Au with the oxides (ligands) in the powder samples and these extend beyond the $e_g$ to higher BE. Au has a face-centered cubic (fcc) crystal structure, and its bonding is a mixture of covalent, ionic, and metallic bonds. Due to symmetry considerations, the covalent contribution consists of Au $e_g$-ligand $2p$ ($pd-\sigma$), Au $t_{2g}$-ligand $2p$ ($pd-\pi$), and Au-Au $t_{2g}$ ($dd-\sigma$) bond regions that can be observed in the valence band.

The lower energy $t_{2g}$ orbitals do not favor sigma interactions with ligands. The $t_{2g}$ symmetry favors the participation of $\pi$ interactions with ligands due to their weaker interactions in metal complexes [83]. This means that the $t_{2g}$ orbitals are completely metal based and contribute to metallic bonding. From the valence band spectra, the pronounced
peaks at \( t_{2g} \) for Au film, bulk, and fine-grained powder indicate metallic bonding with other metals, while the coarse-grained powder show weaker peaks for metallic bonding due to the high level of oxidizing materials.

The spectra results in Figure 8 for the Au film, bulk, fine, and coarse-grained indicate that the BEq of the Au 5d\(_{3/2}\) component of the valence band remains almost unaffected, while the Au 5d\(_{5/2}\) shifted away from the \( E_F \) for a decreased Au sample size, and the top of the d-band is observed to move away from the \( E_F \). Thus, the Au 5d\(_{5/2}\) of fine and coarse-grained samples due to the small grain size shifted away from the \( E_F \) as well as the top of the d-band [44,84,85].

The spectra show that for smaller Au powder samples (fine and coarse-grained concentrates), there is an increase in the BE side of the asymmetry component, while the Auger yield tends to decrease [55]. The spectra of the powder samples show disturbance peaks towards the Auger yield that are attributed to the state of oxidation the samples are subjected to. This shift towards higher BE of the Au 4f\(_{7/2}\) core-level for smaller cluster size (powder samples) is related to valence band narrowing, ascribes a general shift, and is interpreted to be dependent on the initial electronic state structure and the final relaxation state processes as a result of photoemission [84,86,87].

For the Au (film, bulk, fine, and coarse-grained) samples, the valence band width and the binding energies of the two Au 5d and 4f\(_{7/2}\) peaks depend solely on the mean coordination number [44,83,84]. The 4f photoemission from Au split into two distinct peaks, namely Au 4f\(_{7/2}\) and Au 4f\(_{5/2}\) with different BE as a result of spin-orbit coupling effects corresponding to final states, with angular momentum of \( j_+ = + 1/2 = 7/2 \) and \( j_- = + 1/2 = 5/2 \), respectively [88,89]. It is seen from the Au 4f spectra that the Au 4f\(_{7/2}\) -to-4f\(_{5/2}\) peak ratios are not the same but vary between the four different samples. The useful information one can obtain from the study of the BE shift of Au 4f\(_{7/2}\), full-width at half-maximum (FWHM) of the Au 4f\(_{7/2}\) peak and valence band width, is the spectra response to average sample size in terms of peak broadening and shift towards higher BE, as seen in the Au bulk and powder samples [90–92].

### 3.3. EDX Results

Examination under the EDX of the Au powder samples (fine and coarse-grained) show that more than 50 at.% of the Au is present in the grains and these are associated with the pathfinder minerals garnet, hematite, and quartz. The EDX results in Figure 9 and Table 3 also reveal other elements such as Al and Na whose peaks are not easily identified by the surface-sensitive XPS measurements since light elements (N, O, C, and surface oxides including Fe\(_2\)O\(_3\), Fe\(_3\)O\(_4\), and SiO\(_2\)) are removed by the differential sputtering of Ar\(^+\) ions, while bulk-sensitive EDX does not require any surface cleaning.

| Table 3. Elements identified in the EDX analysis of bulk Au and Au powder (fine-grained and coarse-grained) concentrate samples. |
|---|---|---|---|
| **Sample** | Bulk Au (at%) | Powder Fine-Grained (at.%) | Powder Coarse-Grained (at.%) |
| **Element** | | | |
| C | 11.7 | 9.50 | 23.3 |
| O | 7.44 | 14.4 | 22.1 |
| Si | 1.17 | 0.58 | 2.55 |
| Fe | 1.82 | 4.39 | 3.40 |
| Ag | 6.39 | 6.40 | 4.58 |
| Au | 71.2 | 64.2 | 42.6 |
| Al | 0.38 | 0.57 | - |
| Hg | - | - | 1.55 |
Figure 9. EDX spectra of bulk Au, fine-grained, and coarse-grained Au powder concentrate samples.

The Fe in these alluvia Au samples is believed to be contributions from the presence of pyrite, pyrrhotite, chalcopyrite, and the “gangue mineral” magnetite [56]. According to Moslemia et al. [93] and Corkhill et al. [94], the oxidation of minerals such as Fe$_3$O$_4$ and CuFeS$_2$ (pyrite group of minerals) under conditions such as pressure, temperature, pH, particles size, concentration, and electrochemical mechanisms results in the release of ferrous ions and other constituents [93,94]. Hence, it can be inferred that the Fe in these samples is from pure elemental Fe and FeS$_2$. In our analysis, we have prepared the samples (fine and coarse-grained) from the untreated powder so that almost all sulfur-containing grains were discarded. This is due to the fact that the sulfide minerals are washed away during panning and washing since they are much lighter than the Fe and Au metals that settle at the bottom of the pan.

In contrast to previous observations by XRD [56] where the crystal structures were identified, we find that there are significant changes in the chemical bonds in the investigated materials. The XPS analysis confirms that there is covalent bonding between the Au atoms and ligands in particular for the coarse-grained, while the Au bulk and fine-grained powder indicate metallic bonding with other metals, as shown in the valence band spectra in Figure 8. The ratio between metallic and covalent bonding depends on the level of oxidizing materials present in the vicinity of the Au atoms. These oxidizing materials from the near surface (oxide zones) consist of the gangue minerals Fe$_2$O$_3$, Fe$_3$O$_4$, SiO$_2$, garnet, and others from the silicate group of minerals that, under favorable conditions, undergo oxidation and reduction processes to release metastable thiosulphate ions (S$_2$O$_3^{2-}$) and bisulphide ions (HS$^-$) to trigger the dissolution of Au, Ag, and other metals within the hydrothermal systems [52,54].
These identified elemental species Si, Ag, Fe, Al, N, O, Hg, and Ti can act as pathfinder elements in the host minerals of the Kubi Au ore and can lead to exploration success in this mining area. Despite similarities in geology, mineralogy, and the structural setting of the Kubi Gold project with that of Northern belts of Ghana, there exist differences in pathfinder elements due to weathering, erosion, chemical, and landscape processes leading to differences in regolith (loose materials covering solid rock material) [46,47]. In the Kubi Gold area, the association of Ti and Fe with Au is unique and differs from what exists in northern Ghana. The saprolitic and ferruginous clay that have been formed by weathering profile are dominated by garnet (that contains TiO$_2$), Fe-oxides, and iron sulfide minerals through secondary processes and are capable of hosting Au mineralization.

The most interesting part of this study is the scope of economic benefits to be considered from the 1.90 kg artisanal small-scale sediments (soil samples or mining wastes), which produced 1.0 g of Au bulk nugget (22 carats measured with a digital electronic Au purity Analyzer DH 300 K from VTSYIQ), 0.85 g of coarse-grained, and 1.20 g of fine-grained concentrate samples. These samples are found to contain a high percentage of Au as well as metallic and semi-metallic elements such as Ag, Fe, Al, and Ti that are important mineral commodities of economic value.

These new findings from the near surface sediments call for further investigations into the ore-body of deep drilled holes where materials have been subjected to high temperatures and pressures. We anticipate that there will be depth-dependent composition with a lot of deformations and defects due to hydrothermal and physio-chemical processes, among others, in the ore body.

4. Conclusions

By combining core-level and valence band X-ray photoelectron spectroscopy with EDX electronic structure calculations, we investigated the electronic structure and chemical bonding in various Au powder samples that remain after Au panning, in comparison to bulk Au metal and Au film. For the impure Au powder (fine-grained and coarse-grained concentrate) samples, there is a chemical shift of 0.14 eV for the Au $4f_{7/2}$ and $4f_{5/2}$ peak positions towards higher energies as compared to the metal values of 83.95 and 87.60 eV due to a screening effect and charge transfer, indicating the bonding of Au atoms with the surrounding ligands in the indicator minerals.

This study demonstrates that indicator and pathfinder methods have a broader application, which includes the Kubi Gold project, a fact that is not commonly known for artisanal small-scale mining wastes (alluvia small-scale mining sampling). The indicator minerals in this alluvia small-scale mining are part of a large group of indicator minerals that can be used to explore a broad range of Au deposits and other commodities in this mining area.

From the study, we identified that Si, Ag, Fe, Al, N, O, and Ti act as pathfinder elements and can be used to infer indicator minerals such as quartz, hematite, pyrite (marcasite), garnet, and other occasional silicate minerals such as biotite and hornblende.

The results also demonstrate that the mobility of Au and pathfinder elements are very high within the oxide zones (near surface) and these elements bond to the indicator minerals hosting the Au in the mining site.

Quantitatively, we found that the fine-grained Au concentrate has a relatively high amount of pure Au that forms electrum with a significant amount of Ag; this finding makes mining waste an economically viable option to be considered. The significant percentage of oxygen in the powder samples are due to contaminations from the fresh alluvia samples, while the C is attributed to the carbonic or graphitic alterations in the Kubi mining site.

Finally, we conclude that fine-grained sediments considered to be “mining waste” could be of economic importance to miners and investors, and are important materials for exploration. Mining geologists may also apply XPS, EDX, geochemistry analysis, and other techniques in order to elucidate the indicator elements associated with Au host minerals.
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