

## Article

# Mineralogical and Geochemical Characterization of Gold Mining Tailings and Their Potential to Generate Acid Mine Drainage (Minas Gerais, Brazil)

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**Abstract:** For more than 30 years, sulfide gold ores were treated in metallurgic plants located in Nova Lima, Minas Gerais, Brazil, and accumulated in the Cocoruto tailings dam. Both flotation and leaching tailings from a deactivated circuit, as well as roasted and leaching tailings from an ongoing plant, were studied for their acid mine drainage potential and elements' mobility. Detailed characterization of both tailings types indicates the presence of fine-grain size material hosting substantial amounts of sulfides that exhibit distinct geochemical and mineralogical characteristics. The samples from the ongoing plant show high grades of Fe in the form of oxides, cyanide, and sulfates. Differently, samples from the old circuit have higher average concentrations of Al (0.88%), Ca (2.4%), Mg (0.96%), and Mn (0.17%), present as silicates and carbonates. These samples also show relics of preserved sulfides, such as pyrite and pyrrhotite. Concentrations of Zn, Cu, Au, and As are higher in the tailings of the ongoing circuit, while Cr and Hg stand out in the tailings of the deactivated circuit. Although the obtained results show that the sulfide wastes do not tend to generate acid mine drainage, leaching tests indicate the possibility of mobilization of toxic elements, namely As and Mn in the old circuit, and Sb, As, Fe, Ni, and Se in the tailings of the plant that still works. This work highlights the need for proper management and control of tailing dams even in alkaline drainage environments such as the one of the Cocoruto dam. Furthermore, strong knowledge of the tailings' dynamics in terms of geochemistry and mineralogy would be pivotal to support long-term decisions on wastes management and disposal.

**Keywords:** geochemistry and environmental mineralogy; tailings; mobility of toxic elements; acid mine drainage; Minas Gerais—Brazil



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## 1. Introduction

Mining of metals throughout the world produces high volumes of wastes represented by different types of materials [1]. Although highly variable as a function of the characteristics of the ore deposit and the beneficiation processes, as an example [2] estimated around 50,000 Mt of wastes, with 33% of them in tailing dams. In plants for Au metallurgy, the ratio tailings/concentrate can achieve 200:1 [3]. In South Africa, one of the largest Au producing countries, [2] reported a production of  $7.4 \times 10^5$  t of tailings from 1997 to 2006.

These waste materials are discharged in tailing dams and/or ditches in conditions that could be dangerous to the environment. Even though the tailings treatment and control processes use alkaline products [4], the main sources of Au are often associated with sulfides, such as pyrite ( $\text{FeS}_2$ ), arsenopyrite ( $\text{FeAsS}$ ), and pyrrhotite ( $\text{Fe}_{1-x}\text{S}$ ). Deposition of these waste materials in subaerial storage facilities allows the oxidation of sulfides, which may promote the generation of acid mine drainage (AMD) [5–8] with the consequent mobilization of potentially toxic elements (PTE). Unregulated mining also deserves a special mention for its contribution to AMD pollution in regions with great influence of artisanal and small-scale mining operations [9,10].

In order to assure safe closure conditions, it is important to evaluate the AMD potential in the long term and understand the factors that may control the mobility of PTE. Several parameters, in addition to the presence of sulfides, can contribute to generation of AMD and contamination of the surrounding environment [11–15]. Tailings can have a wide variety of chemical and mineralogical compositions and can suffer geochemical evolution controlled not only by the original ore paragenesis, but also depending on the mineral processing techniques and weathering conditions [15]. Supergenic evolution of the tailings results in a secondary paragenesis with high diversity of mineral phases, including soluble sulfates, oxides and iron oxyhydroxides, and arsenates, among others. These secondary minerals, occurring as efflorescences, crusts and other iron-rich precipitates, play important roles in the dynamics of the sulfide oxidation and in mobility of PTE in surface and groundwater [16–18]. Another important factor is the dimension of the tailing particles, as the relationship between size and exposure to weathering can directly influence the oxidation rate of sulfides [7,19].

In general, AMD develops when the neutralization capacity of carbonate minerals or alkaline reagents used in the mineral processing plants (as lime) is depleted due to ongoing sulfide-mineral oxidation. In addition to carbonates, or when they are absent, silicate minerals can provide some level of acid buffering [19–22]. Nevertheless, major and trace elements, such as Se and As can be leached from the tailing dams, contributing to the contamination of water bodies, even if the global geochemical balance is neutral or even alkaline [8]. Furthermore, [23] presents a review of the geochemistry of these elements, emphasizing their sources and mobility in naturally contaminated rocks, namely due to desorption from iron-rich minerals.

Once the production of AMD starts, it is very complex to stop the process. In addition, the remediation techniques based on passive or active processes [24,25] are often inefficient or expensive as they require long-term application of the treatment strategies. Moreover, as mentioned by [26], these classical approaches demand an adequate control and disposal of the resulted PTE-enriched sludge.

Thus, for the management and control of the tailings, it is crucial to assess the AMD potential as well as the mobility of toxic elements such as As, Ni or Se. In general, a good understanding of the environmental behavior of the tailings involves a set of procedures, including acid base accounting and leaching tests (e.g., batch, column and/or in situ) [3,14,15,27–35].

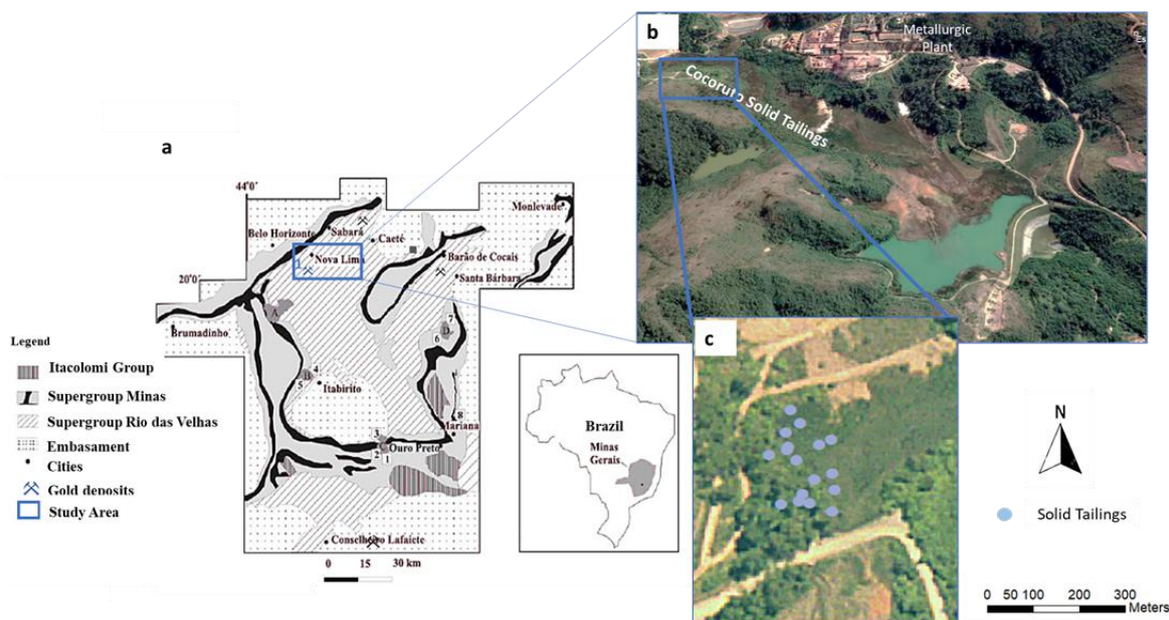
In this work, results of chemical, mineralogical and environmental evaluation of two different set of tailings from Au beneficiation are presented and discussed. The tailings represent different scenarios regarding mineralogy and geochemistry as well as mineral processing techniques and temporal evolution. Therefore, there are tailings from an old circuit (already deactivated) that are accumulated in an abandoned dam (Cocoruto dam) and tailings from an active plant, both located in Minas Gerais, Brazil. These two types represent distinctive beneficiation processes. The old circuit was a sulfide flotation plant and direct leaching. The ongoing circuit includes stages of calcination of sulfide concentrates and leaching.

The general objective is to compare the two types of tailings, using detailed geochemical and mineralogical characterization, namely for determining the AMD potential and the ability to mobilize PTE. This comparison intends to (i) understand the supergenic evolution

of different tailings and (ii) support monitoring and identify possible opportunities for environmental improvements and management of both types of tailings.

## 2. Location and Characterization of the Study Area

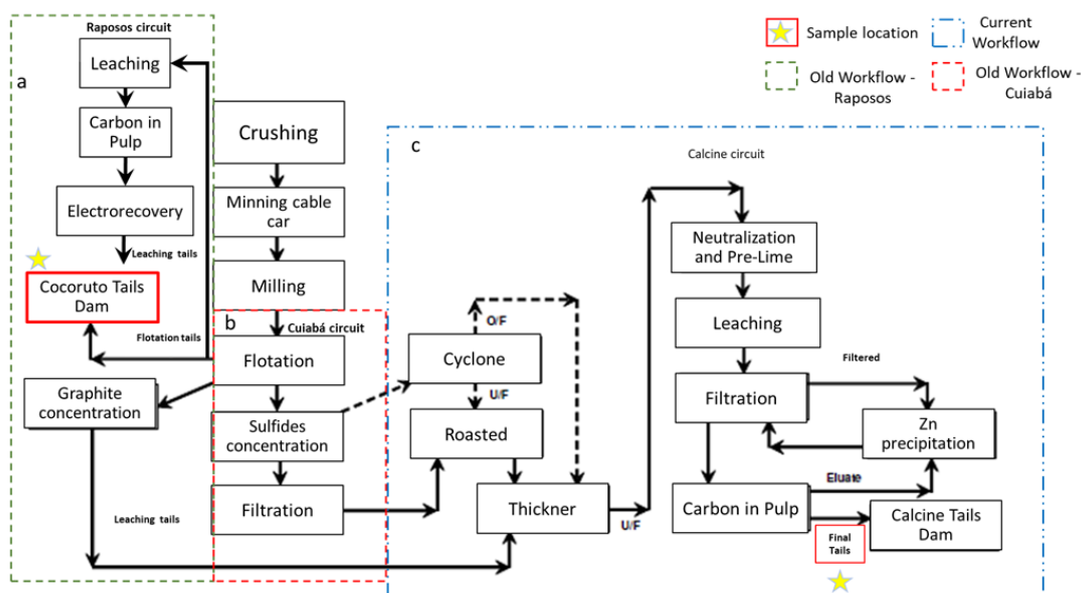
The study area is in the Iron Quadrangle (QF), one of the most gold-producing regions in Brazil [36,37]. The gold world-class deposits are part of the Rio das Velhas Greenstone Belt, largely located in State of Minas Gerais (Figure 1a), with an estimated 4.5% of the world's ore reserves, almost half, about 936 tonnes [38–40]. The orebodies, hosted in Archaean rocks, are structurally associated and controlled by hydrothermal alteration. The mineralization is related to different typologies such as iron-banded formation, clastic and turbiditic metasedimentary rocks, and quartz veins enriched with sulfides, mainly pyrite, arsenopyrite, and pyrrhotite [38,40–42].



**Figure 1.** Location of the study area and Iron Quadrangle map (a) [43,44]; (b,c) Images of the Cocoruto Dam sampled area (SIRGAS2000—10-09-2019).

These sulfide Au ores have been treated for over 30 years in metallurgic plants located in Nova Lima, in the northern part of the QF, Minas Gerais, 25 km away from its capital, Belo Horizonte (Figure 1a). The region has a warm and temperate climate, according to the Cfa classification (humid subtropical climate). The average rainfall is around 1390 mm per year, with the month of August being the driest and the month of December having the highest rainfall, with an average of 302 mm. The average temperature is 23.3 °C, with January the hottest month of the year while the lowest annual temperature is in June (average temperature of 17.6 °C) [45].

The circuits of the Nova Lima plant are and were fed by ores from the Raposos (Raposos circuits (Figure 2a)) and Cuiaba mines (Cuiaba circuit (Figure 2b) and ongoing roasted plant (Figure 2c)). The Raposos circuit (Figure 2a) is a plant that treated non-refractory sulfide (pyrite, pyrrhotite and arsenopyrite) ore. The circuit reached 90% of Au recovery and was divided into grinding, gravity concentration, conventional leaching and CIP (carbon in leach), elution and electrorecovery. This part of the plant was deactivated in 1998 with the deactivation of the Raposos underground mining works, [46].



**Figure 2.** Old and ongoing workflow of Nova Lima's plant (a,b). Old circuit sources of Cocoruto Dam. (c) Ongoing workflow [46]. The yellow stars represent sampling points.

The Cuiaba circuit (Figure 2b) treats refractory ore (mainly Au enclosed in pyrite) with 92% of Au recovery and was divided into grinding, gravity concentration, sulfide flotation, roasting, neutralization, calcined leaching, CIP, elution and electrodeposition [46]. Currently at the Nova Lima plant, the material is treated from the roasting stage (Figure 2c), as the previous steps are carried out directly at the plant located in the Cuiaba mine in Sabará, Minas Gerais. The tailings from the Raposos and the Cuiaba flotation circuits were deposited in a dam, already deactivated, named by Cocoruto (Figure 2a,b). The tailings from the ongoing circuit (roasting and leaching) are available in a dam known as calcined (Figure 2c).

Both structures are downstream type, monitored and declared safe according to the mining national agency [47]. In this study, the tailings samples representing the old circuits were collected directly at the Cocoruto dam, while the tailings from the ongoing circuit were collected directly at the calcined dam discharge. Therefore, the Nova Lima facility provides the opportunity to examine the potential environmental impact promoted by leaching and mobilization of PTE [48].

### 3. Materials and Methods

The sampling campaign was performed in the spring, a season (September 2019) in which weather conditions vary greatly. Typically, during this month it is common to alternate between dry and humid periods, with temperatures between 19 °C and 25 °C [45]. A total of 15 sites were selected in the Cocoruto dam (CoT samples) to represent the old circuit. Sampling, using percussion and auger methods, was performed at a maximum of 10 m. Another set of fresh tailings samples (CaT samples) was collected over 28 days in September 2019 during the production stage, representing a total of 40 samples from the active plant (Figure 2c). All samples were immediately sealed and refrigerated until analysis. Additional material was transferred to polypropylene bags and frozen until analysis. Refrigerated and frozen samples were packaged and shipped to the chemical laboratory for analysis.

The chemical composition was analyzed by inductively coupled plasma mass spectrometry (ICP-MS, PerkinElmer SCIEX, Waltham, MA, USA—detection limit 0.001 mg/kg) after acid digestion (nitric acid, hydrogen peroxide and hydrochloric acid) in accordance with Method 3050B published by the U.S. Environmental Protection Agency (USEPA) [49]. In addition, Au analyses were performed by atomic absorption spectroscopy (AAS, Varian,

Palo Alto, CA, USA—detection limit 0.05 mg/kg), using the fire assay method. All metal solutions were prepared from concentrated stock solutions (Sigma Aldrich). High-purity water produced with a Millipore Milli-Q system was used throughout the analytical process. Quality assurance and quality control (QC/QA) procedures included duplicates, blanks and standard reference materials for ICP-MS. Certified Reference Materials Si81 (Rocklabs) for solid tails were selected to represent a wide range of total elemental concentrations. Results of blanks were always below detection limits. Values for precision (expressed as RSD %) were less than 5% for all elements.

The particle size distribution (PSD) was obtained by vibratory sieving between 212–38 µm. In addition to the geochemical and PSD data, polished sections were prepared in composed samples for mineralogical characterization. The mineralogical study was carried out through optical microscopy and scanning electron microscopy (SEM—Field Electron and Ion Company—FEI; Hillsboro, OR, USA) at the UFMG, Belo Horizonte. The samples were analyzed in an FEI electronic microscope, Quanta 600 FEG, high vacuum mode, coupled to the automated analyzer software (MLA—mode GXMAP and SPL-DZ) and the EDS Spirit Bruker (20 Kve) microanalysis system.

The evaluation of acid mine drainage potential consisted of the comparison of the Modified Acid-Base Accounting (MABA) and Net Acid Generation Test (NAG-test) [50,51]. The acid base accounting (ABA) test is conducted to assess the static potential of generating acidic drainage from a material. It was conducted in accordance with the MEND guide [50] and aims to determine the balance between acid production and consumption from the mineral components of a given sample. Acid production and consumption were measured, respectively, using the parameters acidity potential (AP) and neutralization potential (NP). The NAG-test was carried out in accordance with the MEND guide [51], in which hydrogen peroxide was used to oxidize an aliquot (2.5 g) of comminuted sample (<75 µm). Final pH values (NAG-pH), and electrical conductivity were noted, and the liquor, filtered, was titrated with NaOH solution to defined pH values (4.5 and 7.0) in order to obtain indications on the global amount of non-neutralized acidity, per sample weight. The results of the NAG and MABA tests are compared with each other, allowing a refinement of the interpretation of the latter, and a more improved classification (according to both criteria) regarding the sample's AMD potential [50,51]. In addition, leaching test derived (SPLP) was carried out in accordance with the USEPA 1312 method [52].

## 4. Results

### 4.1. Chemical Composition

Table 1 and Figure 2 show the composition of the tailings. The elements were selected primarily based on the environmental relevance and inclusion in the Brazilian legal framework as well as potential economic interest. The samples from the ongoing process (CaT) comprise the following major elements, according to the decreasing order of their average concentration: Fe > Ca > Al > Mg > K > Mn, Na. The samples from the old Cocoruto dam (CoT) contain higher levels of Mn than CaT tailings: Fe > Ca > Mg > Al > Mn > K, Na (Table 1 and Figure 2). A relative depletion of the iron concentrations could be verified for the tailings deposited historically in the Cocoruto Dam. The elements Na and, especially K, on the other hand, had an enrichment of the order of 3× and 10×, respectively (Table 1 and Figure 3).

The average concentrations of sulfur and cyanide in the ongoing process samples were also an order of magnitude higher than that of the CoT samples (Figure 3). Based only on the sulfur contents, this result could suggest that the CaT samples may have greater acid generation potential, compared to the samples from the Cocoruto Dam.

The trace elements with environmental interest [40] presented the following decreasing order of their average concentrations for CaT: (As > Zn) 10 mg/kg > (Cu > Ni > Pb > Co) 102 mg/kg > (Sb, Cr, Ba, Cd) 101 mg/kg and CoT (As) 103 mg/kg > (Zn) 102 mg/kg > (Cr > Cu > Ni > Pb > Hg) 101 mg/kg. The As followed by Zn remained as the main trace elements of environmental interest in the samples of tailings collected both in mineral

processing and in the Cocoruto Dam (Table 1 and Figure 4). The element Hg was not detected in the CaT samples, but it had an average content of 21 mg/kg in the samples from the Cocoruto Dam.

**Table 1.** Statistical summary of the major and trace elements. CoT—old circuit; CaT—ongoing circuit; N—number of samples.

Element	Source	N	Average	Standard Deviation	Minimum	Maximum
Ag(mg/kg)	CoT	30	0.5000	0.0000	0.5000	0.5000
	CaT	40	3.260	0.5478	2.970	4.960
Al (%)	CoT	30	0.8790	0.4150	0.5670	1.4820
	CaT	40	0.622	0.0569	0.516	0.799
Sb (mg/kg)	CoT	30	0.5000	0.0000	0.5000	0.5000
	CaT	40	49.060	10.4800	29.600	70.300
As (mg/kg)	CoT	30	1799	171.2	1610	2018
	CaT	40	7905	878.0	3553	9275
Ba (mg/kg)	CoT	30	17.1500	7.3800	12.0000	28.1000
	CaT	40	23.427	2.4770	18.600	30.300
Cd (mg/kg)	CoT	30	2.9325	0.0670	2.8700	3.0000
	CaT	40	14.083	1.2010	11.000	17.000
Ca (%)	CoT	30	2.3850	0.4850	1.9190	2.8720
	CaT	40	1.672	0.1177	1.404	1.903
Pb (mg/kg)	CoT	30	21.1000	4.4800	15.9000	26.7000
	CaT	40	258.480	32.0600	215.000	343.000
Cyanide (mg/kg)	CoT	30	2.9850	1.2920	1.6200	4.2600
	CaT	40	51.890	21.1700	13.900	87.600
Co (mg/kg)	CoT	30	4.0000	0.0000	4.0000	4.0000
	CaT	40	116.330	11.7400	95.300	144.000
Cu (mg/kg)	CoT	30	78.0500	15.4100	56.6000	93.0000
	CaT	40	670.520	56.4700	583.000	824.000
Cr (mg/kg)	CoT	30	85.2000	20.5000	59.0000	109.0000
	CaT	40	35.020	8.4700	26.900	81.400
Fe (%)	CoT	30	6.7600	0.5520	6.3970	7.5660
	CaT	40	24.859	2.0270	21.557	28.974
P (%)	CoT	30	0.0002	0.0000	0.0002	0.0002
	CaT	40	0.047	0.0075	0.027	0.061
Mg (%)	CoT	30	0.9570	0.2720	0.6340	1.2040
	CaT	40	0.450	0.0424	0.317	0.520
Mn (%)	CoT	30	0.1728	0.0307	0.1396	0.2119
	CaT	40	0.079	0.0072	0.059	0.088
Hg (mg/kg)	CoT	30	0.0688	0.0338	0.0250	0.1000
	CaT	40	0.036	0.0274	0.025	0.160
Ni (mg/kg)	CoT	30	72.2000	15.2200	52.9000	90.1000
	CaT	40	392.630	51.7200	269.000	506.000
K (%)	CoT	30	0.0411	0.0178	0.0226	0.0644
	CaT	40	0.377	0.0310	0.326	0.471
Na (%)	CoT	30	0.0091	0.0010	0.0086	0.0106
	CaT	40	0.049	0.0050	0.041	0.062
Sulfates (%)	CoT	30	0.2175	0.1763	0.0400	0.4400
	CaT	40	1.505	0.1333	1.230	1.810
Zn (mg/kg)	CoT	30	170.5	38.3	136	0.5000
	CaT	40	2724	318.8	1941	1.430
Au (mg/kg)	CoT	30	1.055	0.6620	0.4800	2.010
	CaT	40	1.510	0.3731	0.3100	2.430

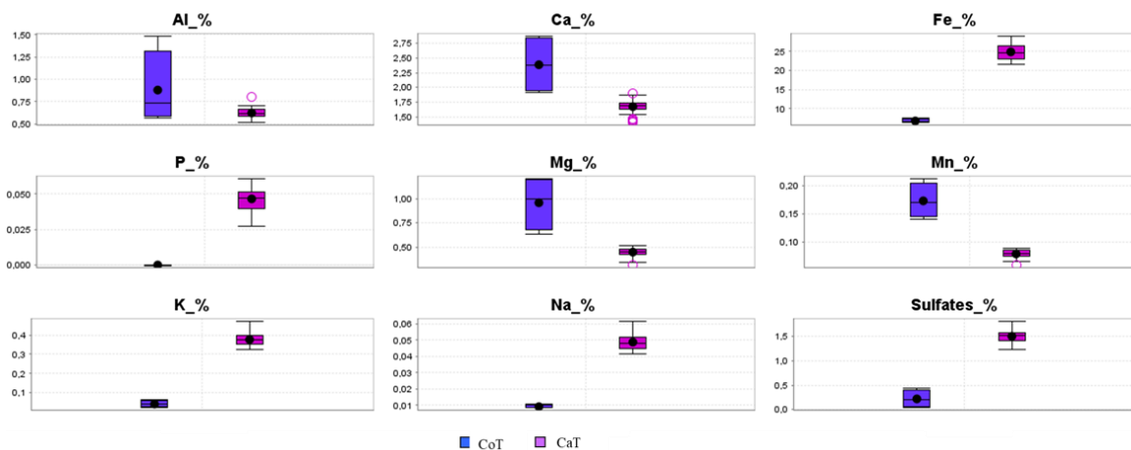


Figure 3. Basic statistics and comparison of major element (%log) for the two sets of tailings samples.

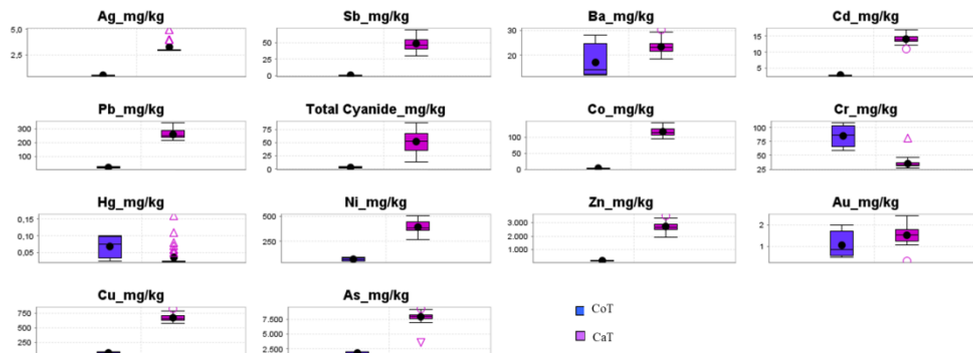


Figure 4. Basic statistics and comparison of trace element (mg/kg log) for the two sets of tailings samples.

#### 4.2. Size Particle Distribution

In general, the grain size distribution (Figure 5) is similar to other dams that store tailings from Au exploitation [7,8,27,53]. The tailings presented 80% of their particles classified as silt-sized (Figure 5). The CoT samples are shown to have a thicker distribution than the ongoing process tailings (CaT). The difference can be explained by changes in the grinding process and different features of the ore sources and their variability [30]. In addition, the samples that represent CoT originate from older circuits, which have low grinding efficiency when compared to the ongoing one (CaT).

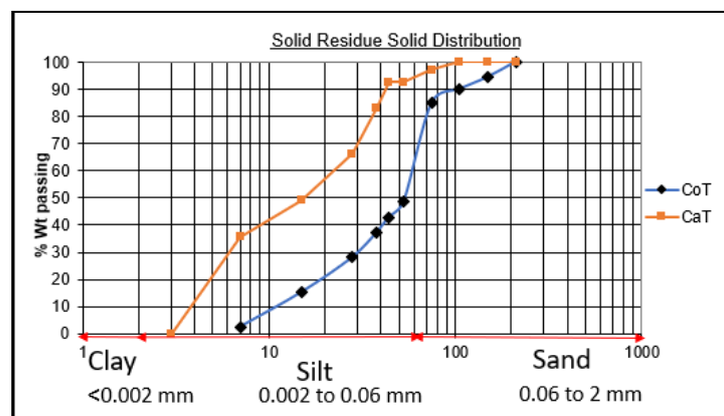


Figure 5. Size distribution (Krumbein phi scale) of solid tailings.

#### 4.3. Mineralogy

Table 2 shows the mineralogical composition of both tailings. The samples from the Cocoruto dam (old circuit) consist mainly of quartz, carbonates, iron oxides and phyllosilicates such as muscovite and chlorite (Figure 6). This is in accordance with higher Al, Ca and Mg concentrations. Sulfides are present, but they look well-preserved, mainly in the form of iron sulfides with sphalerite, covellite and arsenopyrite subordinated.

**Table 2.** Mineralogical Composition of CaT and CoT.

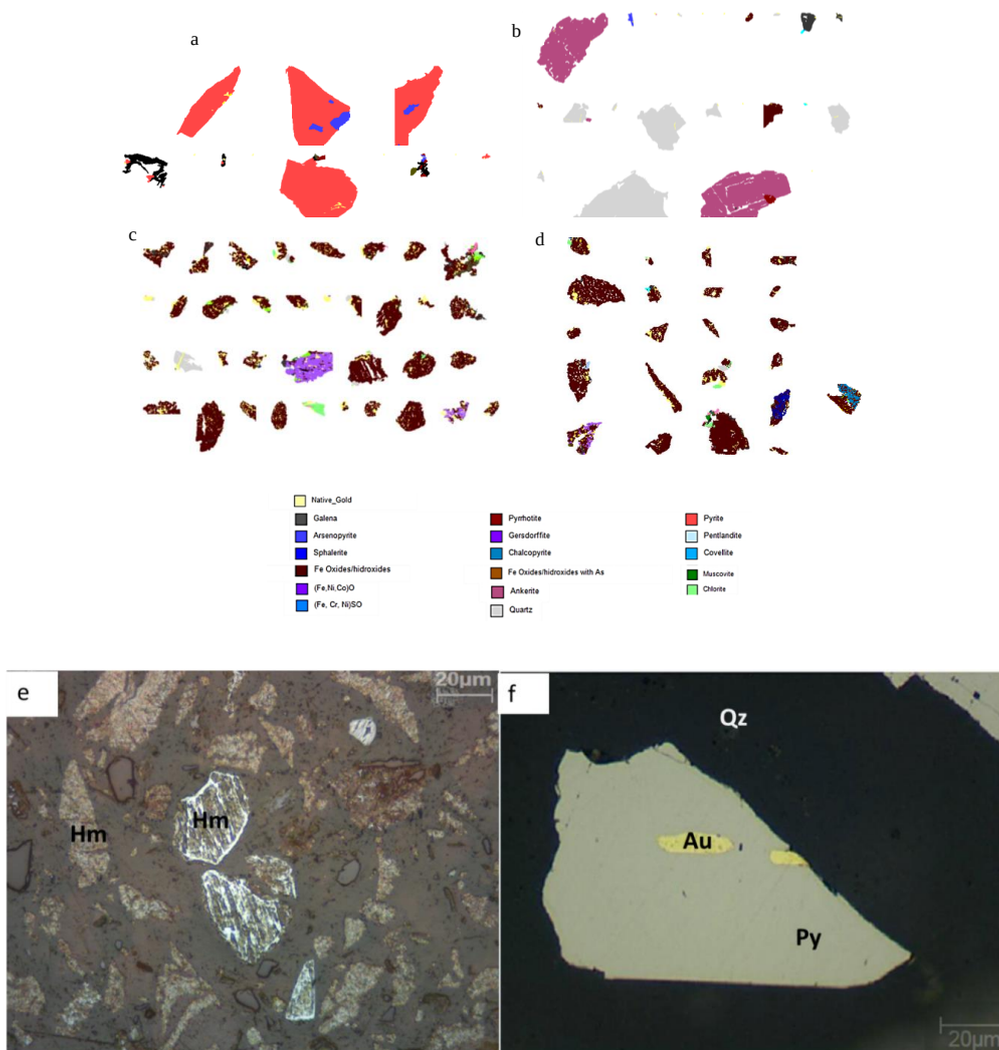
Minerals	Chemical Formula	CaT (Wt%)	CoT (Wt%)
<b>Quartz</b>	SiO <sub>2</sub>	15.6	55.8
<b>Feldspar Group</b>			
Albite	NaAlSi <sub>3</sub> O <sub>8</sub>	1.50	0.370
Anorthite	CaAl <sub>2</sub> Si <sub>2</sub> O <sub>8</sub>	0.053	0.01
K feldspar	KAlSi <sub>3</sub> O <sub>8</sub>		0.390
<b>Phyllosilicates</b>			
Biotite	KMg <sub>2.5</sub> Fe <sub>0.5</sub> AlSi <sub>3</sub> O <sub>10</sub> (OH) <sub>1.75</sub> F <sub>0.25</sub>	1.00	0.16
Smectite	(Si,Al)(Mg,Fe)O(OH)NaH <sub>2</sub> O.	1.80	0.13
Muscovite Group	KAl <sub>3</sub> Si <sub>3</sub> O <sub>10</sub> (OH) <sub>1.9</sub> F <sub>0.1</sub>	11.0	5.56
Chlorite	(Mg,Fe) <sub>3</sub> (Si,Al) <sub>4</sub> O <sub>10</sub> (OH) <sub>2</sub> ·(Mg,Fe) <sub>3</sub> (OH) <sub>6</sub>	3.30	6.12
<b>Oxides</b>			
Iron	Fe <sub>2</sub> O <sub>3</sub> /FeOOH	56.8	8.86
Oxides/Hydroxides			
Rutile/Anathase	TiO <sub>2</sub>	0.599	0.49
<b>Carbonates</b>			
Ankerite	Ca(Fe,Mg,Mn)(CO <sub>3</sub> )	1.00	11.2
Siderite	FeCO <sub>3</sub>	-	7.25
Calcite	CaCO <sub>3</sub>	0.200	2.25
<b>Sulfates</b>			
Gypsum	CaSO <sub>4</sub> 2H <sub>2</sub> O	7.00	0.030
<b>Sulfides</b>			
Pyrite	FeS <sub>2</sub>	0.002	0.500
Pyrrhotite	Fe <sub>(1-x)</sub> S	0.004	0.790
Arsenopyrite	FeAsS	0.056	0.240
Gesdorffite	NiAsS	0.010	-
Covellite	CuS	0.100	0.070
Sphalerite	ZnS	-	0.010
<b>Gold Minerals</b>			
Native Gold	Au > 80%, Ag, Cu, Hg	(526)	(364)
Electrum	Au = 80%, Ag = 20%	(42)	(10)

The samples from the ongoing process (CaT) are characterized by high concentrations of iron oxides, in addition to sulfates and silicates (mainly quartz). The high concentration of oxides justifies higher Fe contents than the samples of Cocoruto. Sulfides are reliquaries, associated with oxidized phases, and are rare. In Figure 6b, is identified, in false images, the presence of As, Cu, Ni, and Zn with oxidized phases, in addition to Fe. In both types of samples there are still fine Au grains. These particles are associated with iron oxides in the CaT and with sulfides in CoT samples. (Figure 6a–d). The optical microscopy images (Figure 6e–f) support these observations, showing the dominant hematite in CaT samples and the presence of Au in association with pyrite in CoT samples, respectively.

#### 4.4. AMD Potential

The results of the MABA tests are presented graphically in Figure 7. Calcined samples collected in mineral processing showed relatively small variability in acidity (AP) and neutralization potentials (NP): In the first case, it ranged between 1.5 and 2.5 kg/CaCO<sub>3</sub> t and, in the second, it varied between 8.3 and 15.5 kg/CaCO<sub>3</sub> t. All these samples were

classified as potentially non-acid-forming (PNA) by the MABA assay criterion, as they presented NP/AP ratio (NPR) > 2. The average NPR value for CaT corresponded to 6.1 kg/CaCO<sub>3</sub> t.

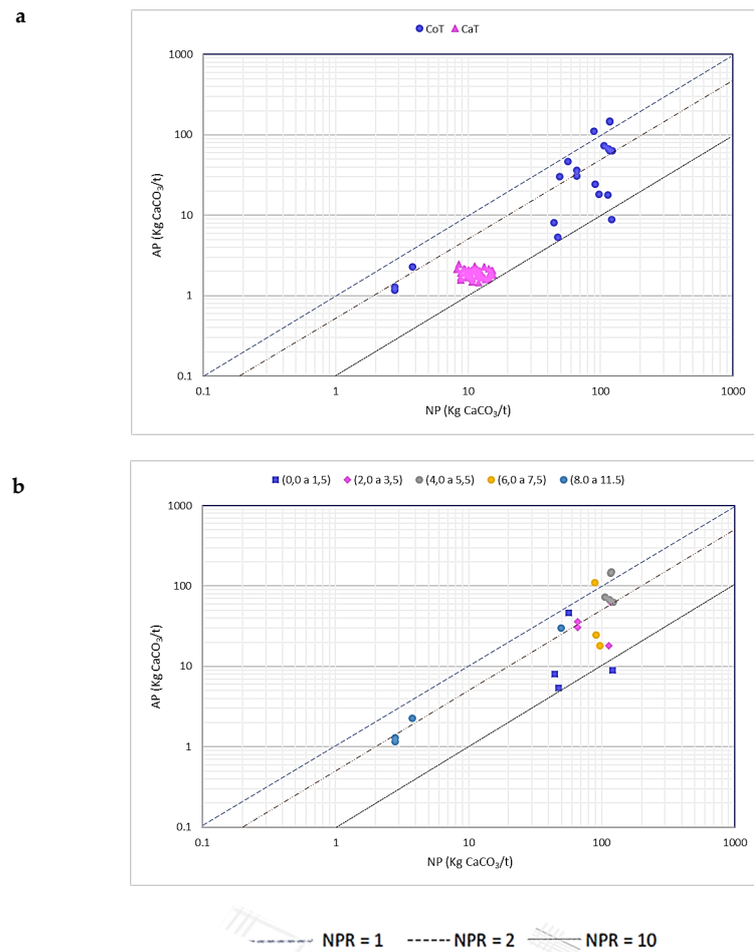


**Figure 6.** False Images (a) sulfides associated with iron oxide and Au particles—CoT, (b) quartz, carbonates and Fe oxides particles—CoT, (c,d) Fe oxides associated with Au, silicates and Fe oxides with As, Ni, Co, Al, Zn, Pb—CaT samples. Photomicrographs taken under reflected light and uncrossed nico (e) hematite (Ht) from CaT samples and (f) Au associated with pyrite (Py)—CoT samples.

The tailings samples collected at the Cocoruto Dam showed relevant dispersion of acidity and neutralization potentials, in accordance with the mineralogical features. The AP values oscillated mainly in the range of 5–150 kg/CaCO<sub>3</sub> t. NP values between 44 to 123 kg/CaCO<sub>3</sub> t. This dispersion resulted in the following relative potentials for the generation of acid drainage, according to the MABA test criteria: Nonacid-forming samples (NPR > 2) represent 45%, samples in the uncertainty zone (NPR 1 to 2) are 50%, and samples acid-forming (NPR < 1) are 5%. Vertically, the samples in the uncertainty zone corresponded mainly to depths of 4.0 to 5.5 m and 8.0 to 11.5 m. The non-acid-forming samples occurred mainly between 0.0 and 1.5 m and from 6.0 to 7.5 m in depth (Figure 7b).

The NAG test (net acid generation) was another method used in order to compare with the results obtained by MABA, mainly because of the S grades. Figure 8 presents the graph NAGpH versus NPR, allowing to correlate the results of the MABA and NAG-tests, and to better refine the classification of acid generation potential of the samples under

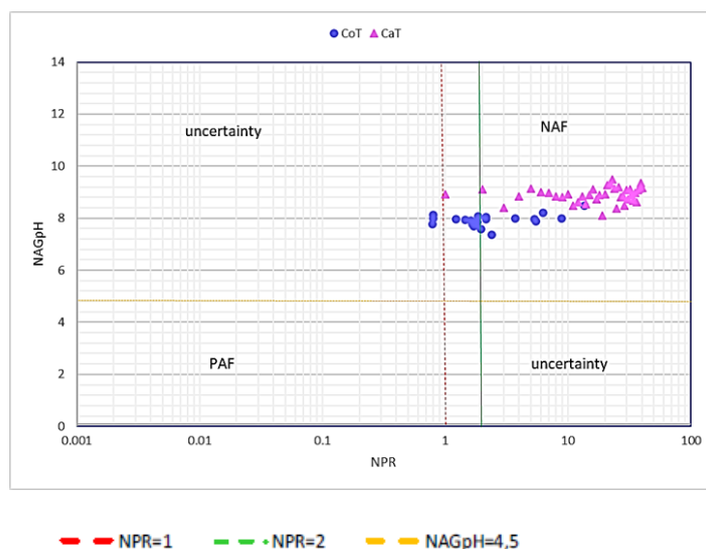
analysis. The results of the NAG test showed a pH value well above the limit of 4.5, indicating all samples as non-acid-forming, regardless of the result of the NPR. The pH fluctuated from 7.4 to 8.5 for samples from the Cocoruto Dam, and from 8.1 to 9.5 for process samples. These results indicate less potential for liquid acidity than previously indicated, only by the NPR (MABA) criterion. The summary of the assessment of the potential for acidity generation, based on the MABA and NAG, therefore, demonstrates that all process samples resulted in potentially non-acid forming (NAF). Only 45% of the samples collected at the Cocoruto Dam resulted in potentially non-acid forming, using the MABA criterion. However, this number was 100% using the NAG criterion.



**Figure 7.** Results of Modified Acid-Base Accounting (MABA) tests (a) global and (b) for CoT, by sampling depth.

#### 4.5. SPLP Leaching

In addition to the tests already presented, the SPLP leaching test (USEPA 1312 method) was carried out in order to assess the potential for leaching toxic elements. Data from leaching tests can be used as indicators of potentially contaminating substances, even though the test conditions are very limited in view of the environmental geochemical conditions. A basic criterion used to assess the magnitude of the potential impact is the comparison with benchmarks in the context of the legal framework. The Brazilian law, through CONAMA Resolution 396/2008, was used as reference for the permitted limits of the following elements: Al, Sb, As, Ba, Be, Cd, Pb, Co, Cu, Cr, Fe, Mn, Hg, Ni, Ag and Zn. Since there is no definition for the predominant industrial use, recreation was defined as the predominant use.

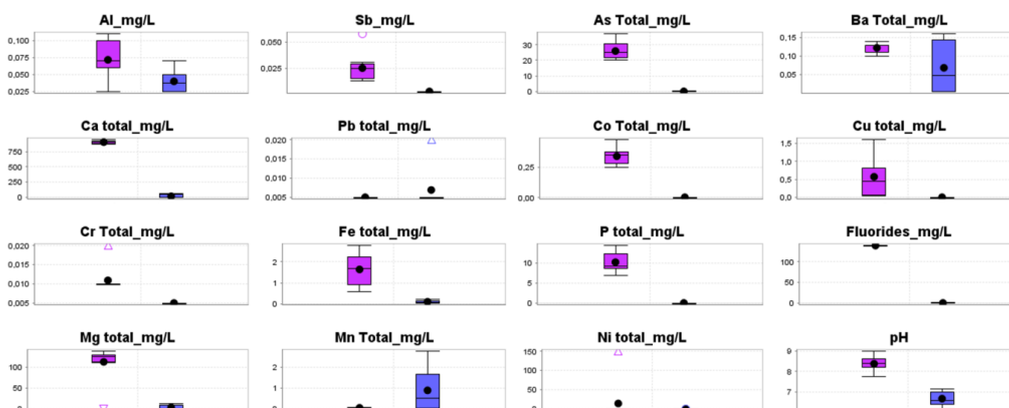


**Figure 8.** Relationship between the results of the MABA and Net Acid Generation (NAG) pH tests (NAF: region of non-acid-forming samples; PAF: region of potentially acid-forming samples).

The elements Cd, Pb, Hg, and Zn were not detected in the leaching extracts (Table 3 and Figure 9), with two exceptions: Pb equal to 0.02 mg/L, and Zn equal to 0.1 mg/L in a sample from the Cocoruto Dam. In the extracts of this dam, only the elements As and Mn showed concentrations above the respective legal reference (VMP) (i.e., 0.05 mg As/L, and 0.1 mg Mn/L). The average concentrations were around an order of magnitude higher than the VMP, in these cases (Figure 9).

**Table 3.** VMP—maximum allowed values From Brazilian Law—CONAMA [54].

Samples	Not Detected Elements	Parameters with Values Above VMP Conama 2005 Reference
CoT	Cd, Hg, Pb, Zn	As (VMP 0.05 mg/L) Mn (VMP 0.1 mg/L)
CaT	Cd, Hg, Pb, Zn	As (VMP 0.05 mg/L) Fe (VMP 0.3 mg/L) Ni (VMP 0.1 mg/L) Se (VMP 0.01 mg/L)



**Figure 9.** Comparison of the average levels of metals, metalloids, fluorides and pH in SPLP leaching extracts. VMP Al = 0.2 mg/L; VMP Sb = NA; VMP As = 0.05 mg/L; VMP Ba = 1.0 mg/L; VMP Ca = NA; VMP Pb = 0.05 mg/L; VMP Co = NA; VMP Cu = 1.0 mg/L; VMP Cr = 0.05 mg/L; VMP Fe = 0.3 mg/L; VMP P = NA; VMP Fluorides = NA; VMP Mg = NA; VMP Mn = 0.1 mg/L; VMP Ni = 0.1 mg/L. NA—Not applied for recreation use.

On the other hand, in the calcined ongoing process the elements As, Fe, Ni, and Se showed values above the CONAMA references, in at least one of the leaching extracts.

## 5. Discussion

The differences in particle distribution could contribute to differentiate the two types of tailings regarding the speed and magnitude of AMD. It is known that the rate of oxidation of sulfides is directly related to the size of the particles and the availability of the surface for reaction [20,55]. The mineralogical characteristics confirmed the distinction between tailings that were subjected to different processes, in accordance with the geochemical signatures (Figure 1). Such geochemical and mineralogical diversity could result in different potentials for mobilizing pollutants in AMD. For example, the higher amounts of sulfides in the Cocoruto tailings (old circuit) could suggest higher potential for AMD generation. However, Figure 6 suggests that inhibition of the oxidative dissolution processes occurs, since the sulfide grains are preserved. Moreover, AMD generation could be controlled by the presence of carbonates due to the ore paragenesis and metallurgical ore treatment [7]. Neutralization conditions also occur in CaT samples due to the addition of lime.

The chemical and mineralogical study highlights the occurrence of high grades of Au in these two sets of tailings. In CoT samples, the concentration reaches 1.0 mg/kg with Au crystals associated with sulfides and rock minerals such as carbonates and silicates (Figure 6a,b). In the ongoing process, in CaT samples, the concentration reaches 1.70 mg/kg, with grains enclosed in iron oxides (Figure 6c). Both Au features portray limitations and problems in the beneficiation old and ongoing processes. However, this inefficiency of the mineral separation processes may represent an opportunity for creating profit through valorization of the mine wastes. The reuse of tailings is now an important research line, as discussed in [53,56–60]. Especially in view of the scenario experienced in Brazil after the mediatized disasters of the Mariana (2015) and Brumadinho (2019) dams [61–64], it is crucial to find sustainable solutions for the closure stage. Therefore, in the present cases, the obtained results suggest the possibility of transforming an environmental liability into an economic outcome.

The tests for prediction of AMD indicate that minimum pH is around 8, reaching a maximum of 9.6, reflecting the alkaline products used for extraction of Au with cyanide solutions. However, the levels of S present in the forms of sulfate and sulfide, thin PSD, and the presence of toxic leachable elements above permitted limits do not rule out the generation of AMD in practice if the tailings are stored improperly. These results, in addition to geochemical and mineralogical features, imply that the samples from the Cocoruto Dam are not potentially acid-forming under the conditions in which they are disposed. The net potential for acidity generation is low. However, in the case of removal of these materials, they must be disposed properly, without long-term exposure (months/years).

Under pH conditions close to neutrality, a series of trace elements can remain solubilized [8], as verified for As, Fe, Ni, Mn and Se. This behavior was also observed by [23], specifically for As and Se leached out in underground rock excavation works. The Al for the sample of the CoT tailings was an exception. Thus, the generation of effluents with metals or metalloids does not require long periods of exposure of sulfide minerals [7]. The As stood out in SPLP leaching test, with an average concentration (26 mg As/L), 3× of magnitude above the permitted limits by the Brazilian law (VMP). This could be explained by the presence of arsenopyrite in CoT samples and iron oxide in CaT. For the other parameters mentioned, the averages values cofound were up to an order of magnitude higher than the corresponding VMP.

## 6. Conclusions

An integrated characterization of tailings from different mineral beneficiation processes was performed to evaluate the environmental risk of mobilizing pollutants from the mine drainage.

The tailings presented 80% of their particles classified as silt-sized. Sulfides are present in the samples from the historic Cocoruto dam, but they appear well-preserved. The tailings consist mainly of quartz (55.8%), carbonates (20.7%), iron oxides (8.86%), muscovite (5.56%) and chlorite (6.12%). Therefore, the potential acidity associated with sulfides could be neutralized by the dissolution of carbonates, such as calcite and silicates such as chlorite, besides the reagents used during the process. This fact contributes to make it a less dangerous waste when compared to the current one (CaT).

The tailings from the ongoing process (CaT) are characterized by high contents of iron oxides (56.8%), gypsum (7%) and silicates (mainly quartz—15.6%). In both types of tailings, the chemical composition allowed to identify elements with environmental relevance by their toxicity, namely Mn in the old tailings and As, Fe, and Ni in the more recent tailings. In addition to these elements of environmental concern, it should be noted the interesting values of Au (up to 2.4 mg/kg in the recent tailings), suggesting a potential for reuse during closures of dam facilities or tailings mass-rearrangement needs.

The geochemical tests, MABA and NAG, indicate that these tailings are not potentially acid-forming, under the conditions in which they are disposed. The net potential for acidity generation is low. However, the results of the SPLP leaching tests indicated a leaching potential for As and Mn in the Cocoruto Dam. Samples that are subject to calcination indicate leaching potential for other elements, namely Sb, As, Fe, Ni, and Se. It is of note that As concentrations in these samples were two orders of magnitude higher than the ones found in samples from Cocoruto Dam. Moreover, the concentrations of S present in the forms of sulfate and sulfide, together with thin PSD and the presence of leachable PTE above permitted limits by the Brazilian law are important issues of concern. In the case of tailings mobilization or inappropriate storage, the generation of AMD and contamination by PTE is likely to occur.

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