Introduction

Assays have been applied for hundreds of years and remain the standard methods for analyzing samples containing noble metals [1,2]. However, gamma-activation analysis (GAA) is expected to penetrate the market of gold-bearing ore analysis in the coming years [3-5]. The GAA method involves the irradiation of the studied samples with high-energy gamma rays (bremsstrahlung radiation) from a linear electron accelerator (LINAC) and measurement of the induced activity of the excited gold nuclei (isomers) using a gamma-ray spectrometer. The advantages of gold GAA, such as high sensitivity and accuracy of the analysis, ability to assay representative analytical samples (500g), and high expressivity and productivity, have been repeatedly discussed and are well known [3-6]. Unfortunately, the development of the GAA method for gold analysis has been limited by the higher cost of primary investment than in standard assay analysis [7]. The total cost of a GAA laboratory, including construction work, can reach $4-5 million $US, approximately 2 million $US more than the initial investment in an assay laboratory. According to economic calculations, this initial investment gap between GAA and standard assay laboratories will close after one year of operating the GAA facility; nevertheless, investors must understand the outcome of investing in a GAA facility [7].

In this work, we present the metrological characteristics and technical features of the GAA demo-testing facility, where methods for determining the concentrations of noble metals and other elements in rocks, ores, and their technological processing products are developed. Applying this installation, we can analyses samples from various mineral resources under various simulation conditions and optimize the measurement conditions of their samples. Such laboratory tests will provide clients with the real limits of detection of the elements being determined and the accuracy of their determination well in advance of their investment in their own GAA installations.
Demo-Testing Facility

The demo-testing facility is located inside a concrete biological shielding, designed to test more powerful accelerators with energies up to 10MeV and beam powers up to 10kW. Concrete shielding itself is located within an industrial hangar (Figure 1).

![Concrete shielding with a GAA demo-testing facility inside an industrial hangar.](image1)

LINAC

Industrial laboratories of gold GAA require LINACs with energies of 7-9MeV and operating powers of up to 10kW [7,10]. Such LINACs were used in our previous industrial works [7, 10-12]. The demo-testing facility is based on a 7-9MeV LINAC and a power of 2kW (Figure 2). In this accelerator, the microwave source was an МИ-456A magnetron with an average power of 5kW operated at 2856MHz. The magnetron is powered by a modulator based on insulated gate bipolar transistors and is constructed as a Marx generator circuit. Owing to the high efficiency of this modulator and the short fronts of the voltage pulses, the power consumption of this accelerator was reduced to ~12kW at a beam power of 2kW.

![LINAC of the GAA demo-testing facility.](image2)

Measuring system

To measure the spectra of gold-bearing ores, one requires precision spectrometers with high energy resolution and maximum gamma-ray detection efficiency. Under industrial conditions, we increased the detection efficiency of the radiation from excited gold nuclei with two-channel spectrometers based on three-crystal assemblies of high-purity germanium (HPGe) detectors, which achieved a gold detection limit below 0.05g/t [11,12]. The demo-testing facility uses a single-channel spectrometer based on a coaxial HPGe detector with a gamma radiation detection efficiency of 38%. The spectra are processed by a digital multichannel analyzer equipped with analytical software for calculating the peak intensities and gold contents in the measured samples.

Measurement collection

In the sample preparation room, the crushed ore is manually packed into a research cylindrical container of diameter 100mm and height 35mm (Figure 3). It should be noted that in contrast to assay analyses, which generally require micron-sized (74 μm) particles in the sample material, the GAA analyses coarse samples with particle sizes of up to 2-4mm [3-6]. The operator manually fixes the container on the holder opposite the accelerator target (Figure 4), exits the irradiation zone, and turns on the accelerator beam for 8-12 seconds. Once arranged in the irradiation position, the round sample container rotates around its axis at several revolutions per second, ensuring uniform irradiation of the container contents. Immediately after exposure to gamma rays, the sample is carried
to the spectrometer that measures the radiation (the measurement zone) by a movable rod. If necessary, the sample can be repeatedly moved between the irradiation and measurement zones during subsequent irradiation-measurement cycles. The irradiation and measurement zones are separated by shielding of the structural elements. The time of moving the sample from the irradiation zone to the measurement zone can be adjusted from 1.5 to 5 seconds.

![Figure 3: Open container packed with the sample to be analyzed.](image3)

The measured spectra are processed and stored in computer, and the measurement results are presented on the monitor screen. After carrying out the required number of exposures-measurement cycles, the operator manually replaces the sample. The time of replacing the sample and returning the accelerator to the mode of operation is approximately 5 minutes.

**Calibration**

![Figure 5: Dependence of gold content on the gold peak area of the reference samples.](image5)
Calibration of the demo-testing facility and preliminary assessment of the metrological characteristics of the GAA were carried out on sequentially measured reference samples provided by STC Min Standart [13]. The measurement results are displayed in Table 1. The measurement error, calculated as the relative standard deviation, did not exceed 10.1% over the entire concentration range. Figure 5 regresses the gold content against the gold peak area of the reference samples. The detection limit of the gold content under 10-fold irradiation of the sample, calculated from the spectrum of the MST Gq 158e sample (1.13g), was 0.063g/t. After multiple (8-10) irradiation cycles, the gold content was determined in the range 0.1 to 10g/t. As each sample consumed approximately 10 minutes of analysis time, the productivity of the facility after 8-10 irradiations is approximately 40 analyses in eight hours. A single ore irradiation at this facility measures the gold concentration in the range 0.75 to 100g/t, and the gold detection limit was 0.19-0.25g/t, depending on the properties of the sample matrix. The analysis time, including the time of changing the samples and switching on the accelerator, was approximately 5-6 minutes. Thus, the single-irradiation productivity of the facility is 80 analyses in eight hours.

Table 1: Measurement results of the reference samples.

| Container Number | Reference Sample | Gold Concentration, g/t | Area (peak-background)/weight | Area-Background | Weight, g | GAA, g/t | Relative Error, % |
|------------------|------------------|-------------------------|-------------------------------|-----------------|-----------|----------|-------------------|
| 3                | MST SG147f       | 0.312                   | 0.17                          | 63              | 367       | 0.34     | -10.12            |
| 4                | MST Gq157d       | 0.850                   | 0.44                          | 151             | 345       | 0.83     | 2.70              |
| 5                | MST Gq158e       | 1.130                   | 0.66                          | 251             | 378       | 1.24     | -9.59             |
| 14               | MST G-160e       | 2.870                   | 1.44                          | 478             | 331       | 2.66     | 7.45              |
| 2                | MST G173e        | 3.000                   | 1.63                          | 503             | 309       | 2.99     | 0.33              |
| 12               | MST G-144e       | 4.750                   | 2.67                          | 840             | 315       | 4.88     | -2.69             |
| 13               | MST Gq-109F      | 7.250                   | 3.96                          | 1493            | 377       | 7.23     | 0.29              |

**The First Research Results**

Real samples from two gold mining enterprises have been measured at the calibrated facility. These enterprises provided 23 samples whose gold contents had been predetermined by assay analysis. All measurements were carried out under the same conditions: sample weight 300-400g, accelerator energy 8MeV, accelerator power 2kW, 10 irradiation-measurement cycles, irradiation time 9s, sample cooling time 2s, and spectrum measurement time 15s. The gold concentrations of seven samples were within 3.5% of the assayed concentrations, but nine, five and two samples were discrepant by 15%, 30% and 60%, respectively. It is important to note that the presented samples belong to the gold-quartz type and incorporate free and large (> 0.6mm) gold particles. Figures 6 & 7 show the gamma spectra of samples with gold concentrations of 3.3 and 0.69g/t, respectively, extracted from two different mines. Almost no natural radionuclides of uranium and thorium (the main interfering elements) presented in the spectra of the gold samples from these mines.

![Gamma spectrum of an ore sample with a gold content of 3.3g/t (10 cycles, irradiation time 9s, sample cooling time 2s, spectrum measurement 15s).](image)
Figure 7: Gamma spectrum of an ore sample with a gold content of 0.69g/t (10 cycles, irradiation time 9s, sample cooling time 2s, spectrum measurement time 15s).

The clear peak of the Au-197m isomer at 279keV is the main peak in gold analysis by the GAA method. Additionally, considering the gold line at 130keV would increase the accuracy of calculating the gold concentration by another 2-3% [12], but this peak is obscured by the high background in the spectrum. The spectra also show peaks of Hf, Nb, Ir, Ba, Y and other elements. The concentrations of these elements could be calculated by appropriate measurement procedures.

Conclusion

Our projects aim is to develop and implement basic principles for the creation of industrial gamma-activation laboratories, in which clients from mining enterprises can analyses gold and related elements in samples with various compositions. The principles and technologies under development should provide a GAA method with high sensitivity, low relative error, and high productivity. Our demo-testing GAA facility, which has begun operation, serves the same purposes. By carrying out laboratory measurements of real samples at the facility, clients will understand the specific limits of gold detection, and the gold determine accuracy of their samples, which can be expected in their own GAA installations.

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