Precious metal recoveries in secondary copper smelting with high-alumina slags

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Abstract
Waste electrical and electronic equipment (WEEE) represents a significant urban resource for precious metals. To maximize the recoveries and sustainable use of these metals, their behavior needs to be characterized in the secondary copper smelting of WEEE. The current study experimentally determined the distributions of gold, silver, platinum and palladium between copper alloy and FeOx–SiO2–Al2O3/FeOx–SiO2–Al2O3–CaO slags ($L_{Cu}$/$M$ = $[M]_{Copper}/[M]_{Slag}$) over the oxygen partial pressure range of $10^{-5}$ – $10^{-10}$ atm at 1300 °C. In addition, the equilibria of copper alloy, slag and Al–Fe spinel system are presented and discussed. The experiments were conducted employing an equilibration—drop-quenching technique followed by major element analysis with Electron Probe MicroAnalysis (EPMA) and trace element analysis with sensitive Laser Ablation Inductively Coupled Plasma Mass Spectrometry (LA-ICP-MS). The results showed that the distribution coefficient of silver increased from 10 to $10^3$ as a function of decreasing oxygen partial pressure. For gold, platinum and palladium, the distribution coefficients were at minimum $10^5$. Lime addition improved the recovery of silver notably and had a small positive effect on gold recovery into copper. Overall, the precious metals can be recovered very efficiently in copper alloy via secondary copper smelting with alumina-rich iron-silicate slags.

Keywords Sustainability · Recycling · Black copper · WEEE · LA-ICP-MS

Introduction
The most important industrial use of gold, silver and many PG-metals (platinum group) is in electrical and electronic equipment (EEE). Their quantities in EEE vary widely depending on the e-product, production era and location [1, 2]. In much of today’s sophisticated electronic equipment, the precious metal concentrations are considerable, even up to several thousands of ppmw. Thus, end-of-life EEE (especially high-grade) provides much richer urban resources than the current primary ores for multiple precious and rare metals.

While the high demand for new electronics in the developing world has increased global sales remarkably, the rapid pace of innovations is making their lifespans shorter and disposal rates constantly higher. Therefore, WEEE presents one of the fastest-growing waste streams globally, with a growth rate of 3–5% per year [3]. The presence of many precious and rare metals in WEEE makes it an attractive resource for recycling and secondary production. The production of precious metals from secondary resources has a lower environmental impact compared to primary production, and it implements sustainability targets with improved resource efficiency and securing the availability of precious metals.

The recovery of precious metals from the WEE is currently performed in copper-making circuits including copper smelters, converters and secondary (black) copper smelters. Black copper smelting is typically implemented for low-grade copper scrap, which may be highly oxidized and includes a wide range of impurities, typically lead, tin, zinc, iron, nickel and aluminum [4, 5]. Aluminum concentration in the copper scrap fraction varies from 1 up to 10 wt% [6, 7] depending on the initial WEEE composition and the shredding-sorting practices. The residual aluminum in secondary copper feed entering the furnace will have an influence on the slag properties and, therefore, on the success of the
entire copper smelting process. The black copper smelting process includes reduction and oxidation smelting stages carried out in either order at the same furnace or consecutive furnaces, such as Ausmelt TSL (top submerged lance) [8, 9]. This study focused on alumina-containing iron-silicate slags and shows how high-alumina (20 wt%) and lime (5 wt%) concentrations in slag affect the precious metal behavior from reducing ($pO_2 = 10^{-10}$ atm) to oxidizing atmospheres ($pO_2 = 10^{-7}$ atm).

In general, research concentrating on minor/trace element behavior in WEEE smelting is relatively sparse, as it is a new field of science, with no published studies of precious metal distributions between copper and high alumina-containing iron-silicate slag. We have recently published papers investigating rare elements In, Ga and Ge [10] and the PG-metal Ir [11] in secondary copper smelting with high-alumina slags. A review paper by Shuva et al. [12] collected the existing literature on minor element distributions in primary and secondary copper smelting. It shows that precious metal distributions in primary smelting with copper matte-slag systems have been investigated more broadly [13–18] than in secondary smelting between metallic copper and slag.

The purpose of the present study was to investigate the distribution behavior of gold, silver, platinum and palladium between molten copper alloy and alumina-containing iron-silicate slags in typical secondary copper smelting conditions, $pO_2 = 10^{-5} – 10^{-10}$ atm and $T = 1300 \, ^\circ$C. The advanced experimental method consisted of equilibration and quenching followed by EPMA and LA-ICP-MS analyses. In our previous papers [13, 19, 20], we have shown that the LA-ICP-MS technique can be utilized successfully for the analysis of the trace element contents in metallurgical slags based on true chemical dissolutions.

**Experimental section**

This study employed a static experimental technique, where data was collected after the system reached the true equilibrium. Technique consisted of three main stages: sample preparation, equilibration—drop-quenching (experiment) at controlled gas atmosphere and sample analyses.

**Sample preparation**

The copper master alloy was prepared by melting copper cathode (Boliden Harjavalta, 99.999%) with approximately 1 wt% of each precious metal, silver (Alfa Aesar, 99.95%), gold (Alfa Aesar, 99.95%), platinum (Alfa Aesar, 99.98%) and palladium (Alfa Aesar, 99.9%), in an alumina crucible at 1400 °C and under 99% Ar—1% H₂ gas atmosphere for 12 h. The CaO-free slag mixture was prepared from pure commercial powders of Fe₂O₃ (Alfa Aesar, 99.99%), SiO₂ (Umicore, 99.99%) and Al₂O₃ (Sigma-Aldrich, 99.99%). The components were weighed out to an initial composition of 52% of Fe₂O₃, 28% of SiO₂ and 20% of Al₂O₃ (Fe/SiO₂ = 1.30). The slag forming components were thoroughly ground and mixed in an agate mortar to obtain a homogeneous slag powder mixture. The CaO-containing slag was prepared by adding 5 wt% CaO (Sigma-Aldrich, 99.9%) to the prepared Fe₂O₃-SiO₂-Al₂O₃ powder mixture. Each equilibration experiment consisted initially equal masses of ~250 mg copper master alloy and slag.

**Equilibration time**

The required experimental time was determined with a pre-experimental time series consisting of 1 h, 2 h, 4 h, 8 h and 16 h equilibration experiments. The equilibrium was approached from different directions using two slags: a pre-reacted iron-silicate slag (Fe₂O₃-SiO₂-20 wt% Al₂O₃ pellet was heated for 1 h at N₂ atmosphere and 1100 °C) and a powder mixture of Fe₂O₃-SiO₂-Al₂O₃. Furthermore, two types of copper alloys (incl. 1 wt% of Ag, Au, Pd and Pt) were employed: a powder mixture and the pre-smelted copper master alloy. Equal weights (250 mg) of copper and slag were put in alumina crucible and equilibrated at 1300 °C and $pO_2 = 10^{-7}$ atm. Samples were analyzed with a Scanning Electron Microscope (SEM) LEO 1450 (Carl Zeiss Microscopy, Germany) equipped with an Oxford Instrument X-Max 50 mm² + INCA Energy Dispersive X-ray Spectrometer (EDS, Oxford Instruments, UK) using acceleration voltage of 15 kV. Figure 1 presents the results for Fe, Si and Cu in slag, as well as Fe and Al concentrations in discrete spinels.

As can be seen from Fig. 1, slag and Al–Fe spinel reached equilibrium in 8 h. For shorter times (1 h, 2 h and 4 h) the compositions of slag and spinel were not entirely homogeneous, and thus produced higher standard deviations than in the completely equilibrated samples. The equilibrium time was also determined for the Fe₂O₃-SiO₂-Al₂O₃-CaO slag and the results confirmed that 8 h was needed to reach equilibrium. However, the experimental time was set to 16 h to absolutely ensure equilibrium conditions in the final experiments at all conditions investigated. The experimental apparatus and critical steps for equilibration—drop-quenching experiments have been presented more detailed elsewhere [11, 19, 20].

The oxygen partial pressure in the furnace was controlled with CO (purity ≥ 99.997%, AGA-Linde) and CO₂ (purity ≥ 99.9992%, AGA-Linde) gases at the experimental temperature (1300 °C) by mass-flow controllers (Aalborg DFC26, USA; accuracy ± 1% of full scale). The equilibrium reaction considered for gas atmosphere is

$$CO + 1/2O_2 = CO_2. \quad (1)$$

The calculated CO₂/CO ratios for each oxygen partial pressure using $K$ value of 71.985.23 (HSC 6.12) are shown
in Table 1. The $\text{CO}_2/\text{CO}$ ratio was set for a total gas flow of 300 ml/min (STP) during the experiments.

**Sample analyses**

The samples were analyzed with a Cameca SX100 (Cameca, SAS, France) EPMA employing five wavelength dispersive spectrometers (WDS), and an Excite 193 nm ArF laser ablation system (Teledyne CETAC Technologies, Omaha, USA) with a Nu AttoM single collector ICP-MS (Nu Instruments Ltd., Wrexham, UK).

The EPMA analyses were performed for copper, slag and (solid) Al–Fe spinel phases. The acceleration voltage for EPMA analyses was 20 kV and the electron beam current 40 nA. The beam diameter varied depending on the analyzed phase. For copper, the beam diameter was set to 100 µm, for spinel 1 µm and for slag between 10 and 100 µm. The average elemental detection limits obtained are presented in Table 2. The detection limits were not sufficient to determine the precious metal concentrations in slag and Al–Fe spinels reliably.

The LA-ICP-MS analyses were performed for accurate and sensitive determination of precious metal concentrations in slags. Both slag types were investigated with 40 µm and 110 µm ablation spot sizes, and some of the CaO-free slag samples were also analyzed with a spot size of 155 µm. The analysis parameters and standards used are presented in Table 3.

The analyses were conducted in time-resolved analysis (TRA) mode, with each analysis consisting of 20 s of background measurement prior to switching the laser on for 40–60 s of signal measurement. Baseline reduction and quantification of the trace element data was performed using the Glitter software package (GEMOC Laser ICP-MS Total Trace Element Reduction; Macquarie University, Australia) [21]. The instrumental uncertainty of the LA-ICP-MS technique was ±5–10% based on analyses of NIST 610 [22], BHVO-2G and BCR-2G [23] reference materials carried out during the analytical sessions.

The detection limits of LA-ICP-MS with 40 µm spot size were not sufficiently low to detect gold, platinum and palladium in slags at experimental conditions $pO_2 < 10^{-5}$ atm. The detection limits for 40 µm spot were approximately 30 ppbw for Ag and Pd, and 5 ppbw for Pt and Au. Eight of the samples, covering both slags and the entire $pO_2$ range, were re-analyzed with larger laser spots (110 and 155 µm), which improved the accuracy and lowered the detection limits. The trace element concentrations of Al–Fe spinels formed within the liquid slag domain

![Fig. 1 Experimental results for equilibration time determination at 1300 °C and $pO_2 = 10^{-7}$ atm. Black marks refer to pre-reacted slag and white marks to slag powder mixture](image-url)

**Table 1** Calculated $\text{CO}_2/\text{CO}$ flow ratios at 1300 °C

| $pO_2$ (atm) | $\text{CO}_2/\text{CO}$ ratio |
|-------------|-----------------------------|
| $1 \times 10^{-5}$ | 227.6 |
| $1 \times 10^{-6}$ | 72.0 |
| $1 \times 10^{-7}$ | 22.8 |
| $1 \times 10^{-8}$ | 7.2 |
| $1 \times 10^{-9}$ | 2.3 |
| $1 \times 10^{-10}$ | 0.7 |

**Table 2** Elemental detection limits of EPMA (ppmw)

| O | Si | Cu | Al | Fe | Ag | Au | Pd | Pt |
|---|----|----|----|----|----|----|----|----|
| 1315 | 223 | 239 | 240 | 230 | 402 | 1753 | 401 | 1255 |

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The spot size used was limited to 5 µm due to the small size of the spinel phase. Detection limits for Ag, Au and Pt in Al–Fe spinels were approximately 3, 0.2 and 0.5 ppmw, respectively.

The detection limits of $^{107}$Ag, $^{197}$Au, $^{108}$Pd and $^{195}$Pt in slags obtained using different laser spot sizes are presented in Fig. 2. Considerably lower detection limits were obtained with larger spots. The results correspond well with the observations by Chen et al. [24] in the corresponding spot size range below 80 µm.

### Results

#### Equilibrium compositions in copper–slag–Al–Fe spinel system

The microstructures of the slags and Al–Fe spinels at two different oxygen partial pressures are shown in Figs. 3, 4. In general, the slags were completely homogeneous, without any signs of crystallization, in the oxygen partial pressure range of $10^{-7} - 10^{-10}$ atm. The samples at $pO_2 = 10^{-6}$ atm had some ‘copper-snowflake’ formations in the slag structure, whereas the slags at $pO_2 = 10^{-5}$ atm were relatively heterogeneous with different kinds of brighter copper segregations. The copper content in the slag increased as the oxygen partial pressure increases, and thus with $pO_2 = 10^{-5}$ and $10^{-6}$, the slag was not capable of holding the dissolved copper during quenching. As the segregated copper in these high $pO_2$ experiments was part of the slag at the equilibration temperature, they were included in slag analyses.

The equilibrated system was complex as it included two liquid phases, slag and copper, and two solid phases, alumina crucible and Al–Fe spinel. The Al–Fe spinels were formed at the slag-crucible interface as a continuous spinel layer as well as within the slag as discrete spinels (see...
Figs. 3, 4). Thus, the copper-slag system was in direct Al–Fe spinel saturation. The following sections present all the results separately for each phase in the system (excluding pure alumina crucible).

**Copper alloy**

Figure 5 presents the analysis results of copper and iron in the copper alloy phase. When the oxygen partial pressure increased, the copper concentration stayed constant and iron concentration decreased. The EPMA detection limit for iron was 230 ppmw, which explains the large standard deviations in the analyses of the lower oxygen partial pressure experiments. As expected, lime addition in slag did not have an impact on the copper phase composition.

The concentrations of precious metals in the copper alloy decrease somewhat as a function of decreasing oxygen partial pressure. Interestingly, as can be seen in Fig. 6, Au and Ag show similar trends between each other, likewise palladium and platinum concentrations. In the highest oxygen partial pressure(s), the volume of the copper alloy phase decreased somewhat as part of copper dissolved into the slag phase. Thus, the concentrations of precious metals, which highly favored copper, in every experimental condition, show small increase towards the higher oxygen partial pressures.

**Slag**

The composition of the slags can be defined by Fe/SiO2 ratio, Cu, Al2O3 and CaO concentrations in slag. The concentration of aluminum was constant around 9–10 wt% in slag (17–20 wt% as Al2O3). The concentration of calcium in lime-containing slags was between 3.3–4.3 wt% (4.6–6 wt% as CaO). Figure 7 shows the obtained Fe/SiO2 ratios as well as the copper loss in the slag as a function of oxygen partial pressure.

The iron-to-silica ratio increases somewhat with the decreasing oxygen partial pressure. As the SiO2 level in slag stayed relatively constant at 30–33 wt% (except at the highest pO2 = 10−5 atm, where SiO2 decreased to ~25 wt%), iron behavior in slag was mostly responsible for the changes in the Fe/SiO2 ratio. The behavior of iron was further related to the spinels, where iron concentration also changed along with the oxygen partial pressure. The copper loss in slags was highly dependent on the oxygen partial pressure and
the slag composition, as shown in Fig. 7. Similar copper solubilities in slags were determined by Avarmaa et al. [10], Nishijima and Yamaguchi [25] and Hidayat et al. [15]. The copper loss increased drastically with the increased oxygen pressure, approximately with a slope of 0.28–0.29 indicating to oxide form of CuO\(_{0.5}\) (Cu\(^+\)) in the slag. The copper loss in the slag was decreased by approximately 20–30% with an addition of 5 wt% CaO. This positive influence of lime has also been observed in other studies [10, 26, 27].
The silver concentration in slags was reliably determined from all the samples employing the LA-ICP-MS technique, even with the small analysis spot of 40 μm (Fig. 8). Silver concentration in slag was clearly dependent on the oxygen partial pressure and the slag composition. Lime addition decreased its concentration in slag. Other precious metals (Au, Pt and Pd) were not detected in slags with the smallest spot size employed, except gold at the highest oxygen partial pressure [19].

Eight samples were re-analyzed with a larger laser spot (110 μm), and 10 analysis points were taken from each sample. Furthermore, four of the samples (CaO-free slag) were re-checked with a 155 μm spot size. Table 4 presents the analysis results, the calculated averages with the standard deviations, and shows in how many analysis points the metal was observed relative to the total number of analysis points. The detection limits are marked in brackets () for each element and sample. The average values for platinum were calculated from the determined concentrations excluding the zero-results, using isotopes 194Pt and 195Pt. Their natural abundance is close to each other, and thus they provide similar minimum detections limits. The standard deviations were calculated only if the trace element was detected in nearly every analysis point taken.

Gold was detected reliably in the samples at the highest oxygen partial pressure \( pO_2 = 10^{-5} \) atm. Reliably in this case means detected in every analysis point taken, clearly above the detection limit and with clear signals in the TRA-spectrums. Moreover, the detected concentrations were the same with all spot sizes employed. Figure 9 (left side) presents time-resolved analysis (TRA) spectrums for gold in SiO2–FeOx–Al2O3 and SiO2–FeOx–Al2O3–CaO slags. The analysis started at approximately 24 s. Both slags gave clear signals: CaO-free slag with higher Au concentration generated a larger signal (cps) than the CaO-containing slag. For the rest of the samples at \( pO_2 = 10^{-6} - 10^{-10} \) atm, gold concentrations were near or below detection limits and the TRA spectrums did not show signals distinguishable from the background (see Fig. 9 right side). Platinum concentrations were not explicitly above the detection limit in any sample (spectrums as in Fig. 9 right side), except with 155 μm spot size at \( 10^{-5} \) atm.

The palladium concentration in slags was measured with isotopes 104Pd, 105Pd, 106Pd, 108Pd and 110Pd. However, because of significant interference of palladium isotope 105Pd with \( ^{40}\text{Ar}^{65}\text{Cu}^+ \), this isotope provided false results with too high concentrations and cannot be used for determining palladium concentration in slags with high concentrations of copper. Moreover, 106Pd, 108Pd and 110Pd isotopes have isobaric interferences when using NIST 612 SRM as the primary standard material from Cd-isotopes and additionally \( ^{94}\text{Zr}^{10}\text{O}^+ \) for 110Pd [28]. In this case, the calculated Pd concentrations were lower than the true concentrations.

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Isotope 104Pd has fewer interferences, and would seem ideal to provide reliable results, but the 104Pd isotope makes up only 11% of Pd abundance, producing results mainly below the detection limit. For these reasons, corrected palladium concentrations were calculated for isotopes 106Pd and 108Pd, which should provide the most accurate results. The correction formula can be presented as:

\[
(C_x)_{\text{samp}} = \frac{[I_x/I_{111\text{Cd}}]_{\text{samp}} * (C_{111\text{Cd}})_{\text{std}}}{[I_x/I_{111\text{Cd}}]_{\text{std}} * (C_{111\text{Cd}})_{\text{std}}},
\]

where \( I \) (Intensity) refers to background-subtracted mean counts per second (cps), \( C \) to concentration (ppmw), \( x \) to isotope under investigation (106Pd or 108Pd) and Is to internal standard (29Si). Moreover, subscripts samp and std refer to values determined for the unknown sample and external NIST612 standard, respectively. Corrections were calculated employing the interference-free 111Cd isotope to obtain true \( I_x \) values of 106Pd and 108Pd, respectively:

\[
I_x = I_{106\text{Pd}} - (I_{111\text{Cd}}) * \text{NA}(106\text{Cd/111Cd}), \quad (3a)
\]

\[
I_x = I_{108\text{Pd}} - (I_{111\text{Cd}}) * \text{NA}(106\text{Cd/111Cd}), \quad (3b)
\]

where NA denotes natural abundance in at-%. These formulas corrected the isobaric interferences of 106Cd on 106Pd and 108Cd on 108Pd from NIST612 standard, which were impossible to separate with the low mass resolution settings required for sensitive laser ablation analyses. This type of correction is typical in geology [29] but is now applied for the first time in the analysis of metallurgical slags. The corrected results are presented in Table 5.

The results for the highest oxygen partial pressure were reliable for both slags. However, the results for the rest of the samples with spot size 110 μm were most likely somewhat
too high, and the results with spot size 155 µm closer to the true concentrations.

Figure 10 presents 108Pd spectrums at the oxygen partial pressure of 10⁻⁵ atm with two spot sizes (110 and 155 µm). The spectrums show reasonably clear signals with both spot sizes, but especially with 155 µm (see the point where the sample measurement starts). Also, the sample F25 (CaO-free at 10⁻⁶ atm) showed a clear signal when analyzed with the largest spot size. In the rest of the samples, the analysis signals were not distinguishable from the background.

**Spinel**

The main elements in the spinel phase were iron, aluminum and oxygen. In addition, low concentrations of copper (0.03–3.5 wt%), silicon (0.03–0.09 wt%) and calcium in CaO-containing slag (0.03–0.07 wt%) were detected. Figure 11 shows the obtained iron and aluminum concentrations, as well as copper concentration in spinel with their standard deviations, as a function of oxygen partial pressure. The EPMA measurements were performed from the discrete spinels in the bulk slag.

The composition of spinels changed as the oxygen partial pressure changed; iron concentration decreased whereas aluminum concentration increased with a decreasing oxygen partial pressure. In addition, as the oxygen pressure increased, the spinel phase dissolved increasingly more copper. The general spinel structure can be presented as A²⁺B³⁺O₄, which in this case is (Fe²⁺,Cu²⁺)(Fe³⁺,Al³⁺)₂O₄ [30]. As the oxygen pressure increased, so did the relative proportions of Fe³⁺ and Cu²⁺, while Al³⁺ and Fe²⁺ decreased.

Precious metals in spinels were checked from four samples using LA-ICP-MS technique. These samples were F12, F16, F19 and F25. According to the laser TRA-spectrums (cps) and calculated detection limits (MDL), no precious metals were detected in the analyzed spinels.
metals (Au, Ag, Pd and Pt) were detected in the spinels. The adopted spot size was very small (5 µm) due to the small size of the spinel phase, and thus, the detection limits were quite high. For silver, gold and platinum, the concentrations in spinel phase were less than 3, 0.2 and 0.5 ppmw, respectively (the detection limits of LA-ICP-MS for these elements). Palladium concentration has not been determined in the external standard GSE-1G, and thus no detection limits or concentration values for this metal were obtained from the samples. However, based on the TRA-spectrums, there was no palladium dissolved in the spinel. The detection limit for Pd can be estimated to be in the same order of magnitude as for Au and Pt.

### Discussion

When examining the minor/trace element behavior in copper smelting, one of the most important parameters to be defined is the distribution coefficient. In general, the distribution equilibria of minor element M between copper metal and slag can be presented with the following equation:

\[
L^{Cu/s}[M] = \frac{[\text{wt}\% M]_{\text{in copper}}}{([\text{wt}\% M]_{\text{in slag}})}.
\]  

where, in this study, [-]-brackets refer to copper alloy phase and ()-brackets to slag phase. This parameter is independent on the size of the system i.e. amounts of the phases and minor elements (following Henry’s law). Thus, the determined coefficients are compatible with industrial secondary copper smelting operating at corresponding conditions with high-alumina slags. Here, the distribution coefficients were determined from the EPMA and LA-ICP-MS analyses for the alloy and slag, respectively. The ‘zero-results’ from LA-ICP-MS were not included in the calculations of average values. The uncertainties for the distribution coefficients were calculated similarly as in our previous papers [10, 31].

In general, the distribution reaction for minor element M between metal and slag can be presented with the following equation:

\[
[M] + n/2O_2(g) = (MO_n).
\]  

Using the equilibrium constant equation for reaction (5), the thermodynamic interpretation for the distribution coefficient can be further modified and rewritten in logarithmic form as:

\[
lgL^{Cu/s}[M] = -n/2lgpO_2 + lgC
\]  

where the constant C includes the activity coefficients (γ), the equilibrium constant (K), and the total number of (cationic) moles (n) in slag and copper. This thermodynamic
approach can be applied for determining the oxidation state of the minor element in the slag. The obtained distribution coefficients for all precious metals were collected in Fig. 12.

The distribution coefficient of silver, with uncertainties, as a function of oxygen partial pressure is presented in Fig. 12a. The (logarithmic) distribution coefficients for both slags increase linearly with decreasing oxygen partial pressure. The slope of the plot is approximately 1/3, which suggests a silver oxide dissolution form of Ag\textsubscript{3}O\textsubscript{2} (monocationic form AgO\textsubscript{2/3}). The distribution coefficient of silver was approximately doubled with the addition of 5 wt% CaO. Most previous studies for a different type of slags have suggested dissolution as Ag\textsuperscript{+} ions (AgO\textsubscript{0.5}) [26, 32–34]. In general, the distribution coefficient of silver in our study for alumina-containing iron-silicate slag was lower than our results. On the other hand, \(L_{\text{Cu}}\) results in copper-slag-tridymite system (1300 °C, 10\textsuperscript{-4}–10\textsuperscript{-8} atm) by Sukhomlinov and Taskinen [35] were very close to our results with CaO-free slag. Moreover, the results by Chen et al. [24] in the matte-copper-slag-tridymite system (at 1200 °C and matte grade of 55–80 wt% Cu) were in the same range (\(L_{\text{Cu}}[\text{Ag}] = 140–520\)) as ours.

For gold, platinum and palladium, the distribution coefficients were calculated using the results obtained with a spot size of 110 µm for eight samples and 155 µm for four samples. The distribution coefficient of gold as a function of oxygen partial pressure is presented in Fig. 12b. The standard deviations (± σ) were included only for the reliable result points at higher oxygen pressures. At the highest oxygen partial pressure, the distribution coefficients of gold were 2.5 × 10\textsuperscript{5} and 4 × 10\textsuperscript{5} for CaO-free and CaO-containing slags, respectively. The slopes with spot size 110 µm between 10\textsuperscript{-5} and 10\textsuperscript{-9} atm are around 1 for both slags, referring to Au\textsuperscript{4+} (AuO\textsubscript{2}) as the dissolving species. However, for spot size...
In lower oxygen partial pressures, the distribution coefficients increased above $10^6$ and close to $10^7$. Lime addition seemed to improve the gold recovery into metallic copper. 

In general, most of the previous experimental studies [36-39] have shown much higher gold concentrations in iron-silicate slags. One reason for this may be the mechanical separation of phases followed by bulk chemical analysis, which may lead to entrained copper droplets within the analyzed slag. Obviously, it is also dependent on the initial concentration of gold in the system. In recent studies, gold distribution coefficients between copper and slag in copper—silica-saturated iron-silicate slag system (1300 °C, $pO_2 = 10^{-4} - 10^{-8}$ atm) were measured as $10^5.3 - 10^7$ [35] and in matte-copper-slag-tridymite system (1200 °C) between $10^5.3$ and $10^6$ [24]. These are in excellent agreement with our results.

Distribution coefficients of platinum between copper and slag obtained with the laser spot sizes of 110 and 155 µm are shown in Fig. 12c. The presented platinum distribution coefficients with spot size of 110 µm represent lower estimates only, as platinum was not detected in most of the analysis points in slags. Only with a spot size of 155 µm at $pO_2 = 10^{-5}$ atm the results for platinum can be considered well quantified. Previous experimental studies on the platinum distribution coefficients ($L_{Cu/Pt}$) measured much lower values of $10^2 - 10^3$ [25, 40]. Moreover, studies on platinum solubilities or concentrations in different types of slags suggest values higher in every case, e.g. 3–7 ppmw [18], 20–800 ppmw [41] and 10–100 ppmw [42], than those measured in the present study.

The distribution coefficients of palladium for both studied slags as a function of oxygen partial pressure are presented in Fig. 12d. For the slags, averages of the corrected palladium concentrations ($^{106}$Pd and $^{108}$Pd) were used. Palladium distribution coefficients increased from approximately $10^5$ to $10^6$ as the oxygen partial pressure decreased. Results at $10^{-5}$ atm can be considered quantified, whereas at lower $pO_2$, the distribution coefficients were minimum values, especially for the results with 110 µm spot size. The determined distribution coefficients were clearly higher than in
the previous studies reporting \( L_{\text{Cu}/s}[Pd] = 10–10^3 \) \[25, 40, 43\].

This study shows that if secondary copper feed includes high amounts of aluminum, precious metals can be recovered efficiently under black copper smelting conditions. Especially gold, platinum and palladium will be distributed very effectively (\( L_{\text{Cu}/s} > 10^5 \)) on the copper’s side, where they can be further recovered from anode slime \[5\]. Silver behavior was shown to be dependent on the smelting atmosphere, \( pO_2 \), and under reductive atmosphere, which typically is the first smelting stage, silver is recovered efficiently in the copper phase. The differences in results between our and previous studies can be due to different experimental conditions (atmosphere, temperature, slag composition). Moreover, the experimental technique varied between studies, and with the older techniques, quenching has not necessarily been rapid enough for achieving a homogenous phase structure compared to the modern technique \[44\]. Furthermore, the analyses have not been conducted with in-situ direct methods like EPMA and/or with highly sensitive techniques as LA-ICP-MS. This study provided concentration and distribution coefficient results based on the true chemical dissolution of precious metals in each phase, without risk of entrained phases distorting the results.

WEEE and copper scrap can be treated and fed into different smelting processes and process stages \[5\]. When compared to matte-smelting conditions, the recovery possibilities of precious metals (Au, Pd and Pt) are better in black copper smelting (or converting) as the distribution coefficients between copper and alumina-containing slags are higher than between copper matte and iron-silicate slag \[13\]. In case of silver, the distribution coefficients in both of these smelting processes were shown to be in the same range. As a consequence of high distribution coefficients, the precious metals can be considered to be lost in copper processes mainly through mechanical copper/matte droplet entrainment or operational reasons/errors. Solids in slags, such as spinels, increase droplet entrainment \[45, 46\]. Figure 13 shows how copper droplets were attached to or trapped during equilibration between spinels in the slags, and adhered to the rim spinel in the present study. In copper processing conditions, spinel and other solid phase formation should be, therefore, avoided for maximizing copper and precious metal recoveries into the copper phase.

### Conclusions

This investigation provided valuable new insight and results in the field of secondary copper smelting and copper manufacturing technologies. The main focus was to experimentally define the equilibrium distributions of gold, silver, platinum and palladium between copper and \( \text{FeO}_x-\text{SiO}_2-\text{Al}_2\text{O}_3-(\text{–CaO}) \) slags in simulated black copper smelting conditions in the oxygen partial pressure range \( 10^{-5} – 10^{-10} \text{ atm} \) at \( 1300 \text{ °C} \). The equilibrium compositions of major elements in the phases of the investigated system were also presented. An advanced and very sensitive experimental method, a combination of equilibration-rapid quenching-EPMA/LA-ICP-MS direct phase analysis, was applied. The silver distribution coefficient increased from \( 10 \) to \( 10^3 \) as a function of decreasing oxygen partial pressure. Gold distribution coefficients at \( pO_2 = 10^{-5} \text{ atm} \) was \( 2 \times 10^5 \) and \( 4 \times 10^5 \) for CaO-free and CaO-containing slags, respectively. Below \( pO_2 = 10^{-6} \text{ atm} \), \( L_{\text{Cu}/s}[\text{Au}] \) was essentially constant with values in the range of \( 10^5–10^7 \). Platinum concentration in the slag was mainly below the detection limits of LA-ICP-MS for all laser spot sizes applied giving a minimum \( L_{\text{Cu}/s}[\text{Pt}] \) of \( 10^6 \). Palladium was observed to be very difficult to analyze reliably at such
low concentrations in Cu-rich systems by LA-ICP-MS. With corrected 106Pd and 108Pd slag results, the distribution coefficients were calculated as 10^5–10^6. Initially, the copper alloy included 1 wt% of each precious metal, from which the resulting chemical solute concentrations in slags were roughly 10–50 ppmw for Ag, < 4 ppbw (25–45 ppbw at pO_2 = 10^-5 atm) for Au, < 6 ppbw for Pt and < 40 ppbw (50–80 ppbw at pO_2 = 10^-5 atm) for Pd. Lime addition improved silver recovery notably and had also a minor positive effect on the gold recovery into the copper alloy phase. Overall, all investigated precious metals can be recovered very efficiently in copper phase through secondary copper smelting and with high alumina-containing slags. The focus in smelting, to maximize the recoveries of precious metals, should be put in proper settling and phase separation.

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