Recycling of tellurium via copper smelting processes

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Abstract
The modern world continuously demands more raw materials for manufacturing all kinds of products. Nowadays, the lifetime of a single product can be very short, as is the case with electronic appliances. Waste electrical and electronic equipment (WEEE) is one of the fastest growing waste categories, and one of the most promising recycling routes for WEEE is to use it as a feed material in pyrometallurgical copper smelting. This article presents new experimental observations regarding the behavior of tellurium in secondary copper smelting process, and compares the results to primary smelting experiments. In secondary smelting conditions, most of tellurium distributed into the copper phase, and the distribution coefficient between copper and slag decreased with increasing oxygen partial pressure. In the primary smelting experiments, most of tellurium was vaporized into flue dusts, and the distribution coefficient between copper matte and slag increased with increasing oxygen pressure, i.e. increasing matte grade.

Keywords  Copper · Trace element · Slag · Distribution · WEEE

1 Introduction
Various copper smelting processes are efficient and widely used ways of treating different kinds of copper-bearing wastes, such as waste electrical and electronic equipment (WEEE). With the reducing grade of primary copper ores and the increasing amount and complexity of electronic scrap, new elements are introduced into the smelting circuits. For the most efficient recovery of these elements, and finding optimal process parameters, fundamental knowledge about the behavior of different elements is required. This article presents experimental results on the distribution of tellurium between copper and slag in simulated laboratory-scale secondary copper smelting conditions. In addition to the iron silicate base slag, alumina and potassium oxide were introduced in the slag to investigate their effect on the process, i.e. slag composition and trace element behavior. In industrial operations, alumina originates from WEEE as well as the sand used for fluxing. This sand may, depending on the location, also contain up to 5 wt% potassium oxide, the effects of which are not well known.

Tellurium is found in primary copper ores, and most of the tellurium produced in the world comes from processing of copper anode slime, which is a by-product formed during electrolytic refining of copper. Traditionally, tellurium has been mostly used as an additive in copper, iron, steel and lead alloys [1, 2]. In recent years, the demand of tellurium has increased significantly due to the large-scale production of CdTe photovoltaic panels, where approximately 40% of the tellurium produced annually is currently used [2, 3]. The amount of tellurium in one photovoltaic panel is low, around 0.1% of the total mass of the panel [4]. However, as the demand for solar energy increases and

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most likely CdTe thin film panels will continue to be one of the main technologies in the near future, recycling of the tellurium values should be improved [4–6]. Tellurium, in different forms, is currently also used in thermoelectric devices, semiconductors, optical disks, phase change memory chips, glass optical fibres and as a vulcanizing agent [1, 3, 6].

Regarding the future uses of tellurium, several articles dealing with new chemistries for providing alternatives to lithium-ion batteries have been published recently. Zhang et al. [7] investigated tellurium–aluminium batteries, Li et al. [8] studied liquid metal batteries with tellurium–tin based electrodes, Koketsu et al. [9] and Li et al. [10] published research about lithium–tellurium batteries, just to name a few. These technologies are still years away from being in commercial use; nevertheless, it is important to establish the most efficient recycling processes before the End-of-Life (EoL) products are entering the recycling and material recovery stage.

Some previous studies have been conducted regarding the behavior of tellurium in different geological/metallurgical systems. Brenan [11] conducted high-pressure studies on the distribution of tellurium between molten and solid sulfides and silicates at 1200–1300 °C. He found that the distribution coefficient is strongly dependent on the FeO concentration in the silicate. Avarmaa et al. [12] studied the behavior of tellurium in Doré smelting, i.e. between metallic silver and different types of SiO2–Na2O slags at 1000–1300 °C. Swinbourne et al. [13] investigated the distribution of tellurium in a similar system consisting of silver–copper selenide matte and SiO2–Na2O slag.

Johnson et al. [14, 15] studied the distribution of tellurium between copper matte and iron silicate slags at 1250 °C. They did not report distribution coefficients, only the equilibrium solubilities of tellurium into the slags due to tellurium-saturation of the experimental system, which lead to inconsistent Te concentrations in the matte phase. However, their conclusion was that slag additives Al2O3, CaO and MgO had a negligible effect on the behavior of tellurium. Choi and Cho [16] determined the distribution coefficient of Te between nickel–copper matte and iron silicate slags. They did not control the partial pressure of sulfur, only the partial pressure of oxygen in the system. Johnston et al. [17] investigated the effect of slag basicity on the behavior of tellurium in metallic copper–magnesia saturated calcium iron silicate slag systems, and noticed a non-linear dependency between the distribution coefficient and slag basicity. Kojo et al. [18] studied the thermodynamics of tellurium removal from copper by using sodium carbonate slags, and found a maximum in the distribution coefficient of tellurium at approximately 10−2 Pa oxygen partial pressure. Schlitt and Richards [19] measured the distribution coefficient of tellurium between copper matte and metallic copper, and found that tellurium was preferentially distributed into the matte phase, where the concentration was approximately sevenfold compared to the metal phase, independently of the iron concentration in matte and the prevailing oxygen partial pressure. The behavior of several trace elements, including tellurium, in copper smelting conditions have also been modelled to some extent [20, 21].

In this article, the distribution behavior of tellurium between metallic copper and three types of iron silicate slags at iron-alumina spinel saturation was investigated and compared to data obtained from earlier studies. The results indicated a completely different behavior for tellurium in primary and secondary copper smelting processes. The influence of potassium oxide addition into the slag was discussed with the help of constrained quasiternary phase diagrams constructed with the MTOX database [22].

2 Experimental

The experiments were conducted in a vertical tube furnace Nabetherm RHTV 120–150/18 (Nabetherm, Germany). The experimental temperature, set to 1300 °C for every experiment, was monitored throughout the experiments with a calibrated Pt/Pt10Rh thermocouple connected to a Keithley 2010 DMM multimeter (Keithley, USA), and the cold junction temperature was measured with a Pt100 resistance thermometer connected to a Keithley 2000 DMM multimeter. The desired gas atmospheres were obtained using a CO–CO2 gas mixture. The most important experimental parameters have been collected in Table 1. A more detailed description of the experimental set-up has been presented earlier [23].

The starting metal and oxide (slag) mixtures were prepared from pure powders. The copper metal mixture was mixed of copper (99.999%, Alfa Aesar) and trace elements Ni (99.996%, Alfa Aesar), Sn (99.85%, Alfa Aesar), Sb (99.9%, Alfa Aesar) and Te (99.99%, Alfa Aesar), all 1 wt% of the mass of copper. The slag, with an initial Fe/SiO2 ratio of 1.3, was mixed of SiO2 (99.99%, Umicore) and Fe2O3 (99.99%, Alfa Aesar). Additions of approximately 3 and 5 wt% of K2O (initially as K2CO3, 99.5%, Sigma-Aldrich) were used for investigating the effect of basic oxides on the distribution coefficient of tellurium, as well as the slag boundaries of the equilibrium system. The parameters for the primary copper smelting experiments, the results of which have been compared to the secondary smelting results in the Sect. 3, have been presented in another article [24].

The total sample mass was approximately 0.4 grams and equilibrations were performed in small cylinder-shaped alumina crucibles (> 99.5% Al2O3, Degussit AL23, Friatec NA, Germany; ID 8 mm and H 15 mm). The
distribution behavior of nickel, tin and antimony in the system have been presented in our previous publications [23, 25]. After equilibration, the quenched samples were cast in epoxy resin, followed by grinding, polishing and carbon coating. The microstructures and phase homogeneity were analyzed with SEM–EDS (scanning electron microscope: LEO 1450, Carl Zeiss AG, Germany—energy dispersive spectrometer: X-Max 50 mm², Oxford Instruments, UK). Based on the phase composition results from EDS, shown in “Appendix”, 16 h was deduced to be a sufficient equilibration time for the copper—slag system. Duplicate experiments were conducted for the K2O-free and 3 wt% K2O-containing samples to ensure the reliability and reproducibility of the results.

The elemental compositions of each phase in the equilibrated samples were analyzed directly with EPMA (electron probe microanalysis: SX-100 equipped with five wavelength dispersive spectrometers, Cameca SAS, France). Eight analysis points were taken from each phase. A defocused, 50–100 µm electron beam was used for the copper alloy and slag phases. The accelerating voltage was 20 kV and the beam current 60 nA in every analysis point. The employed standards and analyzed X-ray lines were hematite (O Kα and Fe Kα), quartz (Si Kα), sanidine (K Kα), alumina (Al Kβ), copper (Cu Kα) and antimony telluride (Te Lα). PAP-ZAF matrix correction [26] was used for raw data processing. The totals of the individual analysis points were between 98 and 100 wt%. The detection limits of each element have been collected in Table 1.

The concentrations of tellurium in slags were below the detection limits of EPMA, and therefore the slags were analyzed with LA-ICP-MS (laser ablation—inductively coupled plasma—mass spectrometry) as well. A Photon Machines Analyte Excite 193 nm ArF laser ablation device (Teledyne CETAC Technologies, USA) coupled to a Nu Atomm SC-ICP-MS (Nu Instruments Ltd, UK) was used. The laser spot size was set to 85 µm and laser energy adjusted to 30% of 5.0 mJ, resulting in a fluence of 2.5 J/cm² on the surface of the samples. The laser was fired at a 10 Hz frequency with each pulse lasting 4 ns. The mass spectrometer was operated in FastScan mode using low resolution (M/ΔM = 300) to maximize sensitivity. Eight pits were ablated from the slag phase of each sample.

NIST 612 standard reference material [27] was used as the external, and Si (measured with 29Si in the MS) as the internal standard, respectively. NIST 610 as well as BHVO-2G and BCR-2G basaltic glasses were analyzed as unknowns. The Glitter software package [28] was applied for processing the obtained time-resolved analysis (TRA) signals. The concentrations of tellurium in slags were calculated as averages of three isotopes, see Table 1.

### Table 1  Experimental parameters as well as EPMA and LA-ICP-MS detection limits in copper alloy and slag

| log₁₀ pO₂ (Pa) | CO (mL/min) | CO₂ (mL/min) | Slags used                          |
|---------------|-------------|--------------|-------------------------------------|
| -5            | 174.4       | 125.6        | ISA / ISA + 3%K / ISA + 5%K          |
| -4            | 91.5        | 208.5        |                                     |
| -3            | 36.6        | 263.4        |                                     |
| -2            | 12.6        | 287.4        |                                     |
| -1            | 4.1         | 295.9        |                                     |
| 0             | 1.3         | 298.7        |                                     |

| EPMA detection limits (ppmw) |
|-----------------------------|
| O  | Al | Fe | Si | K  | Cu | Te |
| 1215 | 4497 | 179 | 273 | 116 | 348 | 183 |
| 1443 | 3258 | 197 | 202 | 94  | 278 | 123 |

| LA-ICP-MS detection limits (ppmw) |
|-----------------------------------|
| Te                               |
| 125Te: 0.033 128Te: 0.110 130Te: 0.145 |

ISA = iron-alumina spinel-saturated iron silicate slag, K = potassium oxide

3 Results and discussion

3.1 Microstructure and phase compositions

Figure 1 illustrates a typical microstructure of the equilibrated-quenched sample. The equilibrium assembly consisted of three phases: copper alloy, slag and iron-aluminous spinel. In reducing conditions, the spinel only formed on the interface of the crucible and slag, whereas in oxidizing conditions individual spinel grains were observed floating within the slag, as seen in Fig. 1. All the alumina dissolved in the slag and spinel originated from
the alumina crucible. The formation and composition of the spinels have been discussed in detail previously [25].

The addition of potassium oxide in the iron-alumina spinel-saturated iron silicate slag increased the activity coefficients of iron oxides in the slag, therefore decreasing iron solubility in the slag and increasing the fraction of spinel in the system. This also resulted in increased solubility of silica in the slag, therefore lowering the Fe/SiO2-ratio shown in Fig. 2. The trend lines follow the oxygen partial pressures very well; at 1 Pa pO2, the Fe/SiO2-ratio is the lowest and at 10⁻³ Pa the highest. At higher oxygen partial pressures, both the increased solubility of copper in slag and the change in spinel composition contributed to the further decrease in the Fe/SiO2 ratio [25]. The alumina concentration in the slags, 17–19 wt%, remained virtually unaffected by the changes in oxygen potential. More details of the composition of the slag, spinel and copper phases have been published earlier [23, 25].

3.2 Phase diagram information

Equilibrium phase diagrams representing the influence of potassium oxide on the Al₂O₃–SiO₂–FeOₓ slag system were calculated employing MTDATA and MTOX database [22]. Figure 3 presents a constrained quasiternary phase diagram of the 50 wt% Al₂O₃-SiO₂-FeOₓ system without K₂O (black lines and labels) and with 5 wt% K₂O (pink lines) at 1573.15 K (1300 °C) and 10⁻⁴ Pa oxygen partial pressure. Figure 4 shows the influence of copper saturation (Cu-sat: black lines and labels, Cu-free: green lines) on the Al₂O₃–SiO₂–FeOₓ-3 wt% K₂O system at 1573.15 K and 0.1 Pa oxygen partial pressure. Copper dissolves increasingly into the slag as a function of increasing oxygen partial pressure, and thus a high pO2 with high equilibrium copper solubility was chosen for the calculations.

As can be seen from the figures, potassium oxide addition increases the number and complexity of equilibrium phase assemblages, especially at the low silica range and towards the alumina apex. In Fig. 3, the tridymite-liquid boundary as well as the mullite–spinel double saturation point radically move towards the silica corner with K₂O addition. When the dissolved copper is taken into consideration at high oxygen partial pressures (Fig. 4), the liquid domain is somewhat broadened by shifting the spinel-liquid phase boundary towards lower silica compositions. Our experimental results and phase diagram calculations show increasing silica concentration in slag as potassium oxide is included in the system, as well as when the oxygen partial pressure is decreased. Our experimental slag composition results are relatively close to the spinel-oxide liquid phase boundary of the calculated phase diagrams, although somewhat richer in silica. When compared to experimentally obtained results, it seems that the MTOX database exaggerates the equilibrium alumina solubility in the slag with and without K₂O addition.

3.3 Distribution behavior of tellurium

Figure 5 illustrates the equilibrium concentrations of tellurium in copper alloy (Fig. 5a) and in spinel-saturated slag (Fig. 5b) in parts per million by weight (ppmw). The exact concentrations of tellurium in the spinel phase, which were below the detection limit of EPMA (120 ppmw) in all experimental conditions, were not analyzed with LA-ICP-MS due to the small size of the individual spinels.
Fig. 3 Triangular section of 50Al₂O₃–SiO₂–FeOₓ ternary system at 10⁻⁴ Pa pO₂. The phase boundaries with 0 (black lines) and 5 wt% K₂O (pink lines) have been superimposed. OX_LIQ = slag, SPI = spinel, TRI = tridymite, MUL = mullite, HAL = halite, G = gas.

Fig. 4 Triangular section of Al₂O₃–SiO₂–FeOₓ ternary system with 3 wt% K₂O at 0.1 Pa oxygen partial pressure. The phase boundaries without copper (green lines) and at copper saturation (black lines) have been superimposed. COR = corundum, NEP = nepheline, LIQ = copper (at Cu-saturation).
Concentration values from matte–slag experiments [24] have also been plotted in Fig. 5 for comparison.

In the copper alloy–slag system, the addition of K$_2$O increased the concentration of tellurium in the alloy in oxidizing conditions, and correspondingly slightly decreased the concentration in slag. In the matte–slag experiments, the addition of alumina and lime decreased the concentration of Te in matte, but had no clear effect on the concentrations in slags. The results in slags exhibited a large scatter, mostly due to the very low concentrations, which were often even below the detection limits of LA-ICP-MS. In the matte–slag system, by far most of the tellurium was vaporized into the gas phase, which is evident when combining the concentrations in matte and slag phases. The vaporization was far less significant in the secondary copper smelting experiments, i.e. the copper alloy–slag system. However, around 50% of tellurium was vaporized in these experiments (in reducing conditions) as well, as shown by the equilibrium concentrations in Fig. 5. The crucibles used in this study and by Sukhomlinov et al. [24] had different geometries, the latter possibly providing a better contact with the gas phase. However, the equilibration time in the present experiments was four times that used in the matte–slag equilibrations, thus eliminating the possible effects of the crucible geometry on the volatilization of tellurium.

Many researchers have stated that the vaporization of tellurium in copper matte smelting conditions is not significant, and tellurium cannot be removed either by slagging or vaporization due to its high stability in matte [19, 20]. This conclusion does not agree with the experimental observations of Sukhomlinov et al. [24] or the tellurium–sulfur phase diagram, which suggests formation of a tellurium-containing sulfide gas phase at copper matte smelting temperatures [29].

In the copper alloy–slag results obtained in our study, vaporization of tellurium seems to be more significant in reducing conditions, i.e. at low oxygen partial pressures, than in oxidizing conditions. An evident explanation for this is the behavior of oxygen as a surface active element: it enriches on the surface of copper alloy, thus preventing tellurium from evaporating [30]. The surface active property of oxygen in copper, as a function of oxygen partial pressure, is a well-documented feature in the literature, e.g. by Gallois and Lupis [31] and Nexhip et al. [32]. Hino and Itagaki [33] reported activity coefficients of tellurium in copper alloys and the low values obtained (approximately $6.5 \times 10^{-2}$ at 1200 °C) also suggest that volatilization of tellurium from liquid copper is not very significant.

The distribution coefficient $L$ of tellurium between copper and slag was calculated simply according to Eq. (1). The obtained values have been plotted in logarithmic form in Fig. 6, as a function of the logarithm of oxygen partial pressure. The dissolved form of tellurium into the slags in copper alloy–slag system can be determined from the slopes of the trend lines.

$$L_{\text{Cu–matte or alloy} / \text{slag Te}} = \frac{\text{wt}-% \text{Te in Cu–matte or Cu–alloy}}{\text{wt}-% \text{Te in slag}}$$

Based on the slope (black line in Fig. 6) [23, 25], tellurium dissolved into the slag as TeO in oxidizing conditions of secondary copper smelting, i.e. above $10^{-2}$ Pa oxygen partial pressure. In more reducing conditions, tellurium seemed to dissolve as elemental species. This conclusion regarding dissolution as neutral, elemental species should
be reliable in the sense that the concentrations of tellurium in ISA slags and copper alloy in all experimental conditions were well above the detection limits of LA-ICP-MS and EPMA, respectively. Thus, the calculated distribution coefficient values represent actual coefficients, not minimum ones, such as reported by us for platinum, for example [34]. In oxidizing conditions, 5 wt% potassium oxide in the slag notably increased the deportment of tellurium into the copper alloy, where it can be recovered in later process stages by anode slime treatment. This suggests that the dissolution mechanism of tellurium in iron silicate slags is related to the fraction of non-bridging oxygen ions. Kojo et al. [18] studied the thermodynamics of tellurium removal from liquid copper by sodium carbonate slags. They found a maximum in the distribution coefficient of tellurium at \( pO_2 \approx 10^{-2} \) Pa. Based on our results, it is also possible that the distribution coefficient starts to exhibit a slight downward trend in more reducing conditions, thus corresponding roughly to the behavior reported by Kojo et al. at \( 1200^\circ \text{C} \), even though their values are two or three orders of magnitude lower than ours. The results of Kojo et al. and the distribution coefficient value obtained by Johnston et al. [17] at \( 10^{-6} \) Pa oxygen partial pressure and \( 1300^\circ \text{C} \), between metallic copper and fayalite slag, have been marked in Fig. 6 for comparison with our results.

In primary copper smelting conditions, the dependence of the distribution coefficient on the oxygen partial pressure was much stronger than in the secondary smelting system, according to new data by Sukhomlinov et al. [24]. The trend, as a function of increasing oxygen partial pressure, i.e. increasing matte grade, was opposite compared to the metallic copper-slag results. Similar trends, although with lower slopes, have been reported before for PM's and PGM's in matte–slag systems by Avarmaa et al. [35]. The deportment of tellurium into the copper matte phase increased drastically as the matte grade (and oxygen partial pressure) increased. The addition of alumina and lime clearly decreased the deportment of tellurium into the copper matte. Avarmaa et al. [35] and Sukhomlinov et al. [24] have discussed the possible reasons for the increasing trends in distribution coefficient values as a function of increasing matte grade, attributing them to the properties of the copper matte as its sulfur-to-metal stoichiometry changes. The feature of selenium dissolution mechanism in slags as an intermetallic was also discussed recently by Sukhomlinov et al. [36].

Choi and Cho [16] studied earlier the behavior of tellurium in argon (and CO–CO\(_2\)) atmospheres (\( pSO_2 \approx 0 \)), through a wide (15–75 wt%) matte grade range with Ni–Cu mattes (Ni/Cu = 2) at 1250–1300 °C, using different slag additives. According to their findings, the distribution coefficient of tellurium was not very strongly dependent on the Ni + Cu concentration of the mattes investigated. The addition of basic oxides slightly increased the value of the distribution coefficient with all matte grades. All the aforementioned results have been plotted in Fig. 7. The differences between the results of Sukhomlinov et al. and Choi and Cho may be explained by the fact that the latter did not control the partial pressure of sulfur, and that their mattes contained more nickel than copper. The comparison with metallic copper–slag results presented in Fig. 6 is not straightforward due to the different x-axes, but in both cases, the smelting process can be considered moving forward in the chain of unit processes as the values on the x-axis increase.

4 Conclusions

As the generation of WEEE is increasing rapidly in the world, considerable efforts are being made to improve the recycling and recovery rates of different metals within. This study contributes to these efforts by determining the distribution behavior of tellurium, a common trace element in primary copper ores as well as electronic scrap, in secondary copper smelting conditions. An advanced equilibration-quenching-direct phase analyses technique was utilized, providing the true chemical solubilities of tellurium in each phase. The effect of potassium oxide on the slag boundaries was also discussed in relation to the MTOX database of MTDATA software.
According to our experimental results, the MTOX database exaggerates the equilibrium solubility of alumina in the slags investigated, as well as the effect of potassium oxide on the slag boundaries. More experimental data is required in order to optimize the database for alumina and K₂O-containing slags.

The concentration of tellurium in copper alloy was more than 50 times higher than in the slags in oxidizing smelting conditions, and more than 1000 times higher in reducing conditions. The addition of up to 5 wt% K₂O into the iron-aluminous spinel-saturated iron silicate slag increased the distribution coefficient in oxidizing conditions, thus improving tellurium deportment into the copper phase, where it can be recovered in later process stages, most commonly from anode slimes after electrolytic refining.

The vaporization of tellurium was far less significant in our experiments simulating secondary copper smelting than in recent data obtained from laboratory scale primary copper smelting experiments, and overall the recovery possibilities seem to be greater in secondary copper smelting. The results presented in this article are important for building better process models and optimizing process parameters for different feed material compositions in industrial operations.

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**Compliance with ethical standards**

**Conflict of interest**  The authors declare that they have no conflict of interest.

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**Appendix**

See Fig. 8.

**Fig. 7** Comparison of tellurium distribution coefficients between matte and slag in different studies. In the results of Sukhomlinov et al. [24], the 55 wt% Cu matte grade corresponds to an oxygen partial pressure of 10⁻³.¹ Pa, and the 75 wt% Cu matte grade to 10⁻².⁶ Pa, while pSO₂ was fixed to 10⁴ Pa. The data by Choi and Cho [16] were from argon gas atmosphere.

**Fig. 8** Concentrations of elements in slag as a function of time at 10⁻¹ Pa oxygen partial pressure. Based on the results, 16 h was deduced to be enough time for equilibration.
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