

# Antimony Recovery from End-of-Life Products and Industrial Process Residues: A Critical Review

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**Abstract** Antimony has become an increasingly critical element in recent years, due to a surge in industrial demand and the Chinese domination of primary production. Antimony is produced from stibnite ore ( $\text{Sb}_2\text{O}_3$ ) which is processed into antimony metal and antimony oxide ( $\text{Sb}_2\text{O}_3$ ). The industrial importance of antimony is mainly derived from its use as flame retardant in plastics, coatings, and electronics, but also as decolourizing agent in glass, alloys in lead-acid batteries, and catalysts for the production of PET polymers. In 2014, the European Commission highlighted antimony in its critical raw materials report, as the element with the largest expected supply–demand gap over the period 2015–2020. This has sparked efforts to find secondary sources of antimony either through the recycling of end-of-life products or by recovering antimony from industrial process residues. Valuable residues are obtained by processing of gold, copper, and lead ores with high

contents of antimony. Most of these residues are currently discarded or stockpiled, causing environmental concerns. There is a clear need to move to a more circular economy, where waste is considered as a resource and zero-waste valorization schemes become the norm, especially for rare elements such as antimony. This paper gives a critical overview of the existing attempts to recover antimony from secondary sources. The paper also discusses the possibility of waste valorization schemes to guarantee a more sustainable life cycle for antimony.

**Keywords** Antimony · Recovery · Recycling · Secondary sources

## Introduction

The history of antimony goes back to as early as 3100 BC, when it was used by the Egyptians as cosmetic in the form of black  $\text{Sb}_2\text{S}_3$  [1]. Antimony was later used by alchemists in the form of antimony oxychloride  $\text{SbOCl}$ , which acted as a powerful emetic and thus became known as “mercury of life” or “algarot” [2]. The etymology of the current name “antimony” is still subject of discussion. Some believe it comes from the Greek word *antimonos* (“never found alone”), others have suggested it is derived from the greek word *anthos* (“flower”), due to the petal-like appearance of stibnite ore, and some even argued that it is derived from the Greek word *anti-monachos* or the French word *anti-moine* (“against monks”), which could refer to the poisonous properties of antimony and the fact that many early alchemists were monks [1, 2]. Despite its long history, antimony metal was isolated and identified as late as the 16th century. Today, antimony is used in a wide variety of products and processes. It is mainly produced from stibnite

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ore ( $\text{Sb}_2\text{S}_3$ ), but also occurs as oxide ( $\text{Sb}_2\text{O}_3$ ), and as antimonides and sulphoantimonides of metals like lead, copper, zinc, silver, and gold [3]. Antimony is mainly used in the form of  $\text{Sb}_2\text{O}_3$ , as flame retardant in plastics, coatings, and electronics, due to its synergetic effect with halogenated flame retardants, which minimizes the amount of halogenated flame retardant required [4]. This application takes up the majority of the world's antimony production [4, 5]. Antimony is also used in catalysts for the production of polyethylene terephthalate (PET) polymers and as additive in glass in the form of sodium antimonite, which acts as a decolourizing agent for optical glass in cameras, photocopiers, binoculars, and fluorescent light tubes [3, 6]. Antimony metal is used as a hardener in lead alloys such as the lead electrodes in lead-acid (LA) batteries [1, 7]. The average antimony content of automotive battery alloys has declined from 7 to 1.6 % in recent years, as Ca, Al, and Sn alloys have been used as replacements. However, the increasing car sales partly compensate for the declining use of antimony in LA batteries [5, 7]. Other minor uses include paint pigments ( $\text{Sb}_2\text{O}_3$ ,  $\text{Sb}_2\text{S}_3$ ,  $\text{Sb}_2\text{S}_5$ ), semiconductors (e.g., AsSb, GaSb, InSb), IR-reflecting camouflage paints ( $\text{Sb}_2\text{S}_3$ ), matchboxes ( $\text{Sb}_2\text{S}_3$ ), and vulcanizing agents in the production of red rubber ( $\text{Sb}_2\text{S}_5$ ) [2, 7]. Antimony compounds are also used in certain phosphors (for instance the halophosphate lamp phosphors), pesticides, ammunition, and medicine [7]. Overall, it is estimated that the global antimony consumption is distributed to flame retardants 52 %, lead alloys and lead-acid batteries 38 %, catalysts for the production of polyethylene terephthalate (PET) 6 %, and chemicals, ceramics, and glass less than 3 %, down from 13 % in 2000 [3]. More detailed overviews of the antimony market and applications can be found in industry reports and government reports [3, 5].

Antimony mining is currently dominated by China ( $\approx 78$  % of global production), which also holds the largest reserves (Table 1) [1]. The dependence on China, combined with the strong industrial demand for antimony, has raised concerns over the supply security, especially since the abundance of antimony in the earth's crust is quite low

(0.2 ppm) and current reserves (1,800,000 t) allow for only 10–11 more years of production at the current speed (Table 1) [8].

Figure 1 shows the distribution of primary and secondary production as well as the portion of illegal (non-reported) primary mining based on a 2011 consulting report [3]. Despite the presence of some deposits, no antimony is currently mined in Europe or in the U.S.A. [1, 7].

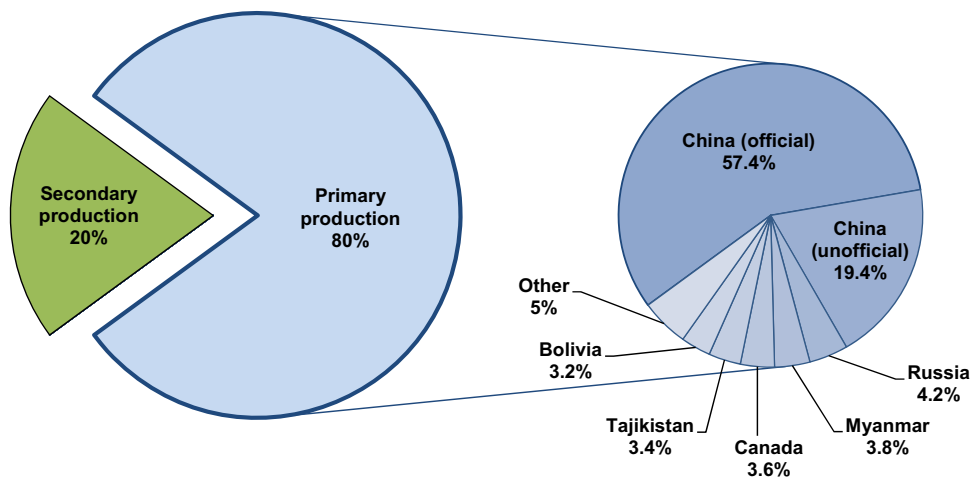
These observations have led the European Commission to highlight antimony as a critical raw material in 2014, with an expected supply–demand gap exceeding 10 % over the period 2015–2020, which is the highest amongst all critical metals [9]. The processing of antimony ores and the production of antimony metal is also concentrated in China, due to its high smelting capacity. Most of the antimony industry in Europe and the U.S.A. is therefore dependent on the import of Chinese antimony metal and is focused on the production of high-purity products and applications.

The aim of this paper is to give an overview of antimony-containing waste streams and to analyze their potential as secondary sources of antimony. Furthermore, the existing valorization methods are discussed as well as the environmental hazard associated with antimony. Finally, an overview is given of the existing water purification and air treatment methods to recover traces of antimony and avoid unwanted release of antimony into the environment. Secondary supply, through recycling and valorization of industrial residues could be a solution to ensure a more secure long-term supply of antimony [9]. Secondary production of antimony could also make Europe and the U.S.A. less reliable on Chinese antimony, thus lowering geopolitical risks. Roskill Consulting estimated that in 2010 around 20 % of the global antimony supply originated from secondary production [3]. Today, secondary production of antimony is mainly restricted to the recycling of antimony-containing lead alloys from lead-acid battery recycling plants [3]. However, interesting future secondary sources could include industrial residues (e.g., mine tailings, process residues, manufacturing scrap) from the production of lead, copper, gold, and antimony [7]. The large production volumes, especially for copper and lead, mean that these residues have the potential to replace a large portion of the primary antimony production [7]. Secondary antimony sources also include end-of-life products such as lead-acid batteries, plastics with antimony-containing flame retardants, antimony-containing glass, and phosphor powders from spent fluorescent lamps [7]. At this moment, only lead-acid batteries are being recycled on a large scale [10, 11]. Antimony could also be recycled from spent fluid catalytic cracking (FCC) catalysts or from catalysts used for the production of PET polymers [12]. Finally, efforts have been made to recover antimony

**Table 1** Antimony world mine production and reserves (tons) in 2014 [8]

| Country    | Mine production 2014 | Reserves (recoverable) |
|------------|----------------------|------------------------|
| China      | 125,000              | 950,000                |
| Burma      | 9000                 | NA                     |
| Russia     | 7000                 | 350,000                |
| Bolivia    | 5000                 | 310,000                |
| Tajikistan | 4700                 | 50,000                 |
| Other      | 8300                 | 177,000                |
| Total      | 160,000              | 1,800,000              |

**Fig. 1** Distribution of antimony ore production in 2010 (196,484 t), using data compiled by Roskill [3]



from landfills and municipal solid waste incineration (MSWI) ashes [13–15]. A schematic overview of the antimony life cycle, with its waste streams and possible recycling routes, is depicted in Fig. 2.

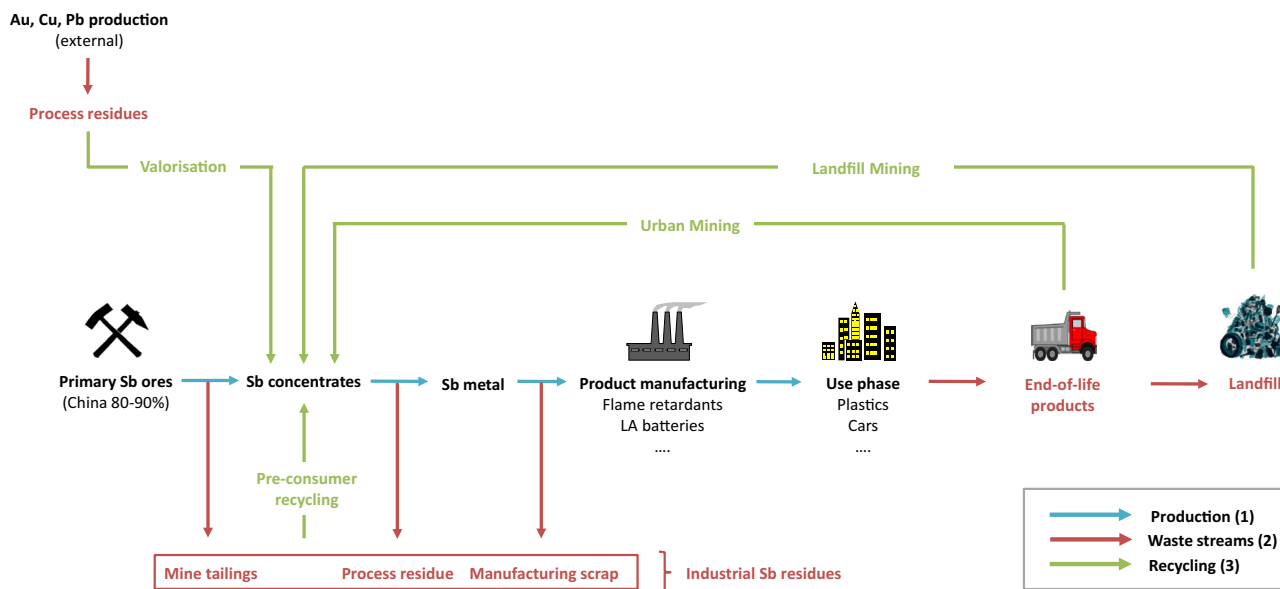
### Recovery from Industrial Residues

#### Antimony Production

The primary production and metallurgy of antimony has been reviewed by Anderson (2012) [1]. However, in order to discuss the secondary production of antimony from industrial residues, a short overview must be given here of

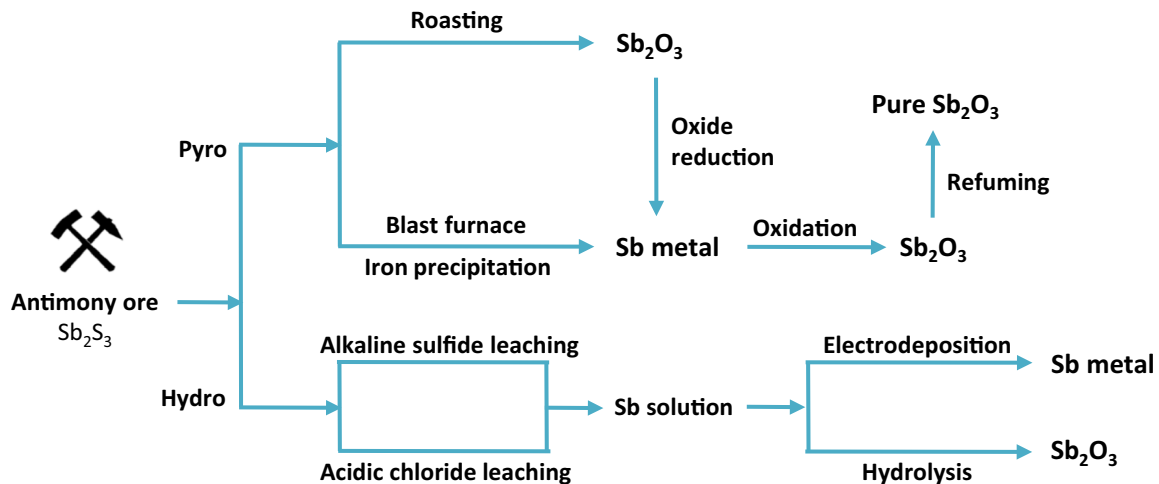
the main production routes, process residues, and metal-containing waste streams. The main pyro- and hydrometallurgical pathways to produce antimony metal and antimony oxide ( $Sb_2O_3$ ), are shown in Fig. 3.

Hydrometallurgical methods are based on two steps: leaching (alkaline sulfide or acidic chloride system), followed by the electrodeposition of antimony metal at the cathode, or hydrolysis with NaOH or  $NH_4OH$  to produce  $Sb_2O_3$  [1]. At this moment, mainly pyrometallurgical processes are used, but hydrometallurgical processes based on alkaline sulfide leaching have been employed industrially in the former Soviet Union, China, Australia, and the United States [1]. Alkaline sulfide technology has the advantage of having a high selectivity toward antimony,



**Fig. 2** Schematic overview of the antimony life cycle, which includes (1) the main value chain between primary mining and final disposal, (2) the major waste streams, and (3) the possible secondary

antimony production routes through recycling and recovery of antimony-containing waste (Color figure online)

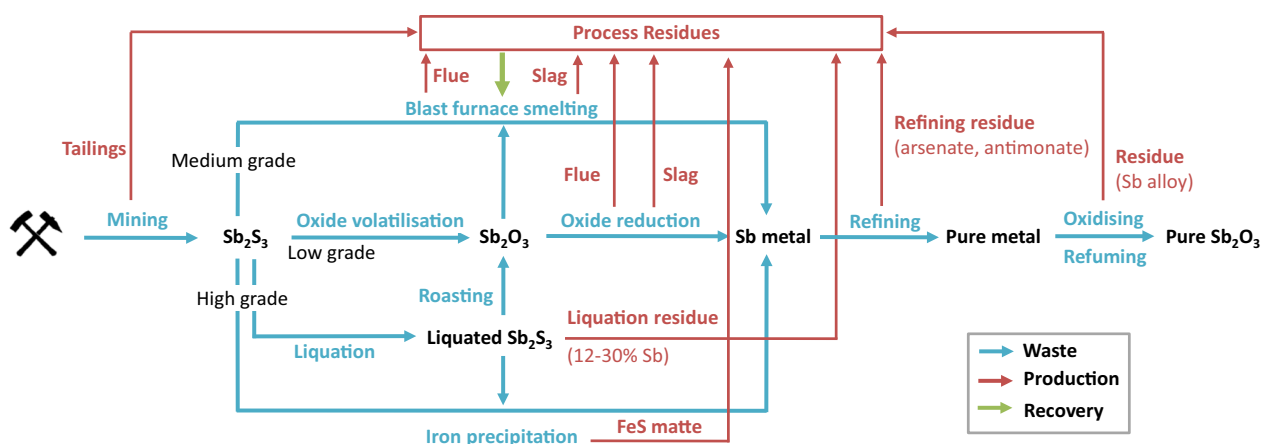


**Fig. 3** Primary production of antimony through pyrometallurgical and hydrometallurgical methods

and having less issues with corrosion. However, a considerable amount of research and pilot-scale work has been undertaken to utilize chloride-based technology [1, 16, 17]. A more detailed overview of the pyrometallurgical pathways and process residues is given in Fig. 4.

A stibnite concentrate is first produced by grinding, milling, flotation, and gravity concentration [1]. These steps are usually carried out close to the mining site, and result in Sb-containing mine tailings. Depending on the grade of antimony in the concentrate, different pyrometallurgical methods can be used to extract antimony [1]. For low-grade ores (5–25 % Sb), *oxide volatilization* is used. The ores are roasted at 1000 °C, and the volatile  $Sb_2O_3$  is recovered. Intermediate grade ores (25–40 % Sb), and Sb-rich residues, slags, mattes, and flue dusts are smelted in a blast furnace at 1300–1400 °C which produces antimony

metal,  $SO_2$  gas, and a slag. Sb-rich ores (45–60 % Sb) can be treated by *liquation* and *iron precipitation*. Liquation consists of heating the ore to 550–600 °C under a reducing atmosphere, to extract liquid  $Sb_2S_3$  from the ore. This product is called crude, liquated, or needle antimony. The liquation residue (12–30 % Sb) can be treated to further improve the recovery of antimony. The *iron precipitation* method is used to produce antimony metal from concentrated antimony ores or from the crude antimony issued from the liquation process. In this process, scrap iron is added to molten  $Sb_2S_3$  to displace the antimony in order to form antimony metal and a *matte* which contains iron sulfide. The reduction of  $Sb_2O_3$  to antimony metal is done with charcoal in a reverberatory furnace (1200 °C), with the formation of a slag. The loss of antimony by volatilization is high (12–20 %), meaning that the flue



**Fig. 4** Pyrometallurgical pathways (blue) and metal-containing waste streams (red) for the primary production of antimony from stibnite ore. The different process residues can be used as secondary antimony sources and fed back into the blast furnace (green) (Color figure online)

dusts must also be caught and then reprocessed in a blast furnace. Finally, refining steps are needed to obtain pure antimony metal (e.g., arsenic removal with NaOH). This creates arsenate- and antimonate-containing residues. Highly pure Sb<sub>2</sub>O<sub>3</sub> is usually obtained by oxidation of technical grade antimony metal (99.0–99.8 % pure), followed by a refining process (volatilization + condensation) [2]. These oxidation processes produce dusts and residues, which can also contain important quantities of antimony. It is clear from Fig. 4 that antimony production creates many residues. The composition of these residues is highly variable on the type of feed material and the process parameters, but some representative compositions are given in Table 2.

In some countries, these residues are stockpiled or landfilled, causing metal pollution in the areas around antimony mining sites and processing plants [18, 22–24]. A better approach is to try to immobilize these waste residues by applying various stabilization, solidification, and geopolymerization techniques [22]. However, a better solution is to valorize these residues, thus creating *secondary sources* of antimony. This increases the extraction efficiency of antimony from the initial ore and also eliminates or reduces the need to discard residues containing toxic metals [18]. Guo et al. [18] characterized the process residues (e.g., slags, flue dusts) from an antimony smelting plant in Xikuangshan area in China (the “antimony capital”). They found large amounts of antimony in the flue dust of the blast furnace (23.4 wt%) and in the residue from the arsenic removal process (37.8 wt%). The leaching of the different residues was studied in an effort to remove the antimony and other metals (e.g., Hg, Pb, As, Cr) from these waste residues. Fuxu et al. investigated the use of a three-phase fluidized bed reactor for arsenic removal from Sb-refining residue (20–40 % Sb, 3–5 % As) [21]. Their set-up achieved a 97 % removal rate of arsenic, leaving behind a valuable antimony concentrate. Kequiang et al. [21] prepared Sb<sub>2</sub>O<sub>3</sub> (98.50 % pure) from Sb-containing slag, using

a vacuum evaporation method. The slag was placed in the vacuum chamber and heated (893–1073 K) in order to selectively evaporate the volatile Sb<sub>2</sub>O<sub>3</sub>. Luo et al. [20] studied the recovery of gold and antimony from antimony smelting slag by direct reduction in an electric furnace. Using this method, the antimony content in the slag was reduced from 32 to <1 wt%. During the pyrometallurgical production of antimony oxide from antimony metal, waste residues are also created, which contain alloys of antimony and other base and precious metals. Anderson [7] studied the recovery of antimony from these pyrometallurgical production residues. Using alkaline sulfide leaching, 99.5 % of antimony was recovered.

At the same time, research is still ongoing to optimize the primary production of antimony from stibnite ore and to reduce the amount of (toxic) waste streams [7, 16, 25, 26]. Yang et al. [25], for example, developed a low-temperature sulfur-fixing smelting process, using a NaOH–Na<sub>2</sub>CO<sub>3</sub> flux with ZnO to fix the sulfur as ZnS. This approach reduces SO<sub>2</sub> emissions (98.61 % sulfur-fixing) and results in an antimony recovery rate of 97.07 % in the form of antimony metal. Ye et al. [26] investigated a similar system, using a eutectic Na<sub>2</sub>CO<sub>3</sub>–NaCl molten salt and ZnO as a sulfur-fixing agent. Under optimum conditions, the average recovery ratio of antimony reached 92.88 %. Also of interest, are the efforts to recover antimony from a low-grade or complex stibnite ore, as well as from secondary antimony minerals. Yang and Wu [16] developed a new hydrometallurgical process to recover antimony from complex stibnite concentrate. They dissolved antimony using a chlorination–oxidation procedure to obtain an SbCl<sub>3</sub> solution. Using electrodeposition, hydrolysis, hydrolysis–smelting, or hydrolysis–washing, they obtained cathode antimony metal (99.98 %), Sb<sub>2</sub>O<sub>3</sub> (99.9 %), crude antimony metal (99 %), or SbOCl (99.9 %), respectively. Gök (2014) published a hydrometallurgical method for the catalytic production of antimonate from stibnite concentrate [27]. Alkaline leaching was used to extract the antimony, followed by the oxidation of Sb(III) in a hydroquinone-catalyzed alkaline electrolyte, which precipitates antimony as sodium hydroxyantimonate NaSb(OH)<sub>6</sub> (with 90 % recovery).

Besides stibnite, researchers are also investigating other, complex antimony minerals. An overview of complex antimony minerals and their mineralogy has been compiled by Roper et al. [28]. Complex antimony minerals often require adapted flowsheets in order to ensure the recovery of antimony and the other main metal constituents. The last primary producer of antimony in the U.S.A. (Sunshine Mining & Refining Company) was mining the complex copper-silver-antimony sulfide: freibergite (Cu,Ag)<sub>12</sub>Sb<sub>4</sub>S<sub>13</sub> [1]. This mine produced both silver and antimony concentrates until its closure in 2001. Other interesting Sb-

**Table 2** Heavy metal content (wt%) in various process residues from the primary production of antimony

| Residue                  | Sb    | As   | Hg   | Zn   | Pb   | Cu   |
|--------------------------|-------|------|------|------|------|------|
| Smelter slag [18]        | 1.11  | 0.57 |      | 0.07 |      | 0.01 |
| Smelter flue dust [18]   | 23.4  | 0.13 | 0.93 | 0.03 | 0.05 | 0.04 |
| Gas treatment slag [18]  | 0.69  | 0.07 | 0.05 |      |      |      |
| Refining residue [18]    | 37.8  | 11.4 |      |      |      |      |
| Refining residue [19]    | 34.85 | 4.41 |      |      |      |      |
| High-Sb slag [20]        | 32.00 | 0.63 |      |      | 0.30 |      |
| High-Sb slag [21]        | 39.49 |      |      |      |      |      |
| Sb oxidation residue [7] | 63.0  |      |      |      | 18.0 | 12.0 |

The data are based on literature reports [18–20]



containing minerals are tetrahedrite  $(\text{Cu,Fe})_{12}\text{Sb}_4\text{S}_{13}$  and jamesonite  $\text{Pb}_4\text{FeSb}_6\text{S}_{14}$ . Baláž et al. [29] investigated the leachability of antimony and arsenic from tetrahedrite and jamesonite with alkaline sulfide leaching. Awe and Sandström [30] also studied the selective leaching of arsenic and antimony from a tetrahedrite-rich complex sulfide concentrate and concluded that by removing these elements, a valuable copper concentrate was obtained, which could be used as a feedstock for copper smelting. Yang et al. [31] developed a flowsheet for the recovery of antimony from a low-grade jamesonite concentrate ( $\text{Pb}_4\text{FeSb}_6\text{S}_{14}$ ). They used sodium sulfide to dissolve the antimony as sodium thioantimonate ( $\text{Na}_3\text{SbS}_4$ ) and subsequently used air oxidation to produce sodium pyroantimonate ( $\text{Na}_2\text{H}_2\text{Sb}_2\text{O}_7$ ). Duchao et al. [32] improved this process by using pressured air oxidation. Under optimum conditions, the precipitation ratio of antimony was 99.38 %. Gold, copper, and lead production are also associated with large amounts of antimony impurities, which end up in the process residues.

### Gold Production

Gold-bearing sulfide ores often contain important amounts of antimony (Table 3). Kyle et al. [38, 39] made a series of reports on the occurrence and deportment of trace elements (Sb, Bi, Se, Te, As, Hg, Cd, Pb) in gold processing. They estimated that in Australia alone, total emissions of antimony (air, water, land) reach as much as 3.8 t/year during mining and 8.7 t/year during metal manufacturing. This represents a significant loss of valuable antimony and it is a challenge to find better processing technologies which can minimize the loss of antimony by valorizing the different gold-mining residues and waste streams.

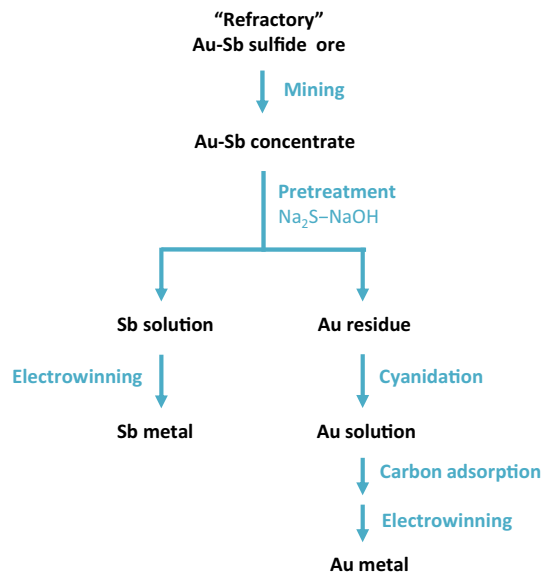
These “refractory” gold ores are resistant against the standard gold recovery method of cyanidation and carbon adsorption, because the sulfide mineral traps the gold particles, making it difficult for the cyanide leach solution to complex with the gold [2, 40]. Pretreatment is therefore required to make the cyanidation effective. Different pretreatment options exist: chemical leaching, roasting, bacterial oxidation, pressure oxidation, and ultrafine grinding [2, 40]. The choice of pretreatment process depends on the

type of ore and the impurities present in the ore and may be preceded by a flotation concentration step. Depending on the concentrations of gold and antimony, different categories of Au-Sb-containing sulfide ores can be distinguished: (1) ores in which gold is the main valuable component and antimony is a harmful impurity, (2) gold-bearing ores in which the amount of antimony is considerable and for which it is economically viable to recover antimony as a commercial byproduct, (3) antimony ores in which gold is a minor accompanying element, the recovery of which could increase the economic efficiency of the process [38, 41]. For the ores of group (1) and (3), standard gold-processing and antimony-processing technology is applied, respectively. The recovery of traces of gold and antimony from these ores is only interesting for waste products, as this does not cause losses in the recovery of the main metal. The processing of the second group of ores (2), usually requires changes in the flowsheet in order to ensure the recovery of both the gold and antimony fraction. For this type of ores, alkaline sulfide leaching is a preferred pretreatment method as it isolates antimony and simultaneously increases the recovery rate of gold in the consequent cyanidation step (Fig. 5).

Ubal dini et al. [40] used alkaline sulfide leaching as a pretreatment step, to remove antimony from a gold-bearing stibnite ore. Antimony was then recovered by electrodeposition. Results showed that this pretreatment improved the recovery of gold in the consequent cyanidation process from 30 to 75 %. Celep et al. [35, 42] studied similar antimonial refractory gold and silver ores. Their findings confirmed that alkaline NaOH and  $\text{Na}_2\text{S}$  leaching were appropriate pretreatment methods to recover antimony prior to the conventional cyanidation for the recovery of the noble metals. Using this approach, 85 % of antimony could be leached out and during the subsequent cyanidation step, silver recovery rates increased from less than 18 % up to 90 % and gold extraction was also enhanced by 20–30 %. Solozhenkin et al. (2010) investigated the complex antimony ores and gold-antimony concentrates, found in the Sarylakhsky and Sentachansky deposits in Russia [34, 43]. They developed different hydrometallurgical processes to treat these ores and produce both gold and antimony concentrates with antimony recovery rates of

**Table 3** Literature examples of Au-Sb-rich ores and waste from which both gold and antimony can be valorized into concentrates

| Antimony concentrate        | Au (g/t) | Ag (g/t) | Sb (wt%) | As (wt%) |
|-----------------------------|----------|----------|----------|----------|
| Refractory Sb-Au ore 1 [33] | 3.6      |          | 0.3      | 0.4      |
| Refractory Sb-Au ore 2 [34] | 7.4      |          | 16.73    |          |
| Refractory Sb-Au ore 3 [34] | 42.2     |          | 28.7     |          |
| Refractory Sb-Au ore 4 [35] | 20       | 220      | 1.6      | 0.03     |
| Refractory Sb-Au ore 5 [36] | 10.5     | 2.1      | 0.22     | 1.67     |
| Electrorefining Slime [37]  | 210      |          | 0.31     |          |



**Fig. 5** Schematic visualization of the pretreatment of refractory Au-Sb sulfide ore with alkaline sulfide leaching. The pretreatment method improves the cyanidation of gold and allows the recovery of antimony [40]

95–98 %. The studied methods include antimonite flotation, sodium dimethyldithiocarbamate treatment, and biotreatment of the ores. The authors also introduced a processing plant for Sb-Au-bearing alloys and demonstrated the successful electrolytic refining and production of cathode antimony and noble metal slurry [39, 43]. Kanarskii et al. [38] investigated the concentration and isolation of antimony and arsenic in Sb-As-bearing gold ores by flotation. They demonstrated the flotation separation of antimonite and arsenopyrite into separate products, thus increasing process efficiency and profit by improving the gold recovery rate and reducing the consumption of cyanide. Karimi et al. [36] investigated the influence of different pretreatment methods ( $H_2O_2$ /air oxidation, roasting, and  $HNO_3/HCl$  leaching) on the cyanidation of Sb-rich gold ores and the movement of antimony and impurities. Saleh et al. [37], examined the recovery of Au, Sb, and Sn from gold electrorefining slime. The solid waste was dissolved in HCl (2 M), before extracting the metal ions with the quaternary ammonium chloride extractant Aliquat<sup>®</sup> 336.

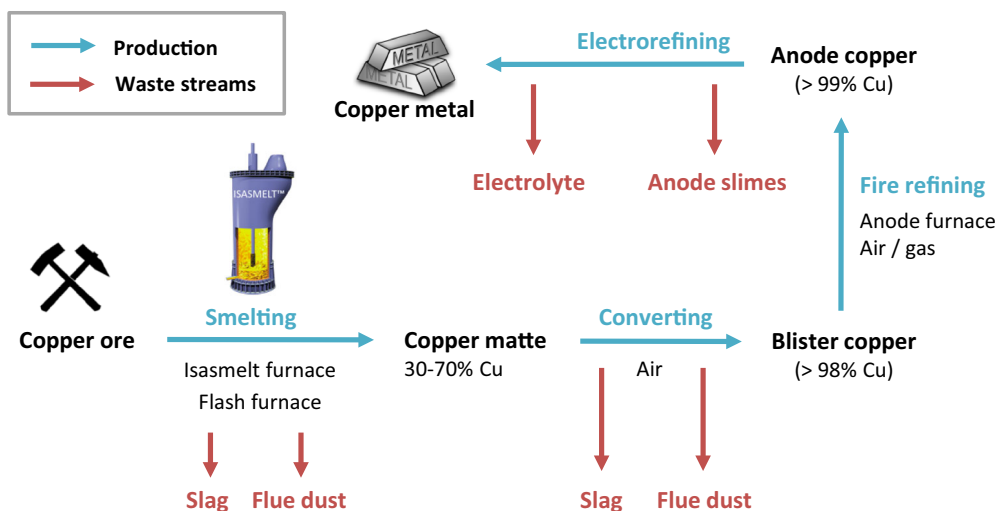
## Copper Production

Copper is mainly produced from copper ores such as chalcopyrite,  $CuFeS_2$ , which often contain antimony, arsenic, and bismuth as impurities [2]. These impurities must be removed and therefore end up in various residues. The formation of the residues is discussed here, as well as the recovery of antimony from these residues. In the past,

the copper concentrate (20–40 % Cu) was first roasted and then introduced in blast furnaces or reverberatory furnaces, but nowadays direct smelting is favored using for example Outokumpu Flash furnaces or Isasmelt furnaces [2, 44]. During smelting, silica is added as a flux to remove iron as an iron-silicate *slag*, which floats on top and which can be used as building material. A *copper matte* is also formed which is a mixture of copper, iron, and sulfur that is enriched in copper. The copper matte (30–70 % Cu) produced in the smelter is then introduced in a converter, where air is blown into the matte to remove the sulfur as  $SO_2$  gas and to form *blister copper* (>98 % Cu) and an iron-silicate “Fayalite” slag ( $Fe_2SiO_4$ ). The blister copper is then put into an anode furnace (fire refining), where it is purified to *anode-grade copper* by removing most of the remaining sulfur, oxygen, and iron. Oxygen is usually removed by blowing natural gas through the melt (*poling*). The resulting *anode copper* (>99 % Cu) is further purified by electrorefining. The copper anodes are placed in an electrolysis set-up filled with an aqueous solution of  $CuSO_4$  and  $H_2SO_4$ . By applying a voltage, copper and the less noble metals (e.g., Fe, Ni, Co, Zn, Pb, Sb, Bi, As) dissolve at the anode, while the more noble metals (e.g., Ag, Au, Se, Te) settle at the bottom of the cell as *anode slime*. The  $Cu(II)$  ions migrate through the electrolyte to the cathode, and copper metal is deposited. Part of the less noble metal ions remain in solution as soluble sulfates (e.g., Fe, Ni, Co, Zn), while others precipitate as insoluble sulfates (Pb) or hydrolyzed species (As, Sb, Sn, Bi) and are therefore found in the anode slime on the bottom of the cell, together with the noble metals. The electrolyte solution and anode slimes are valuable by-products of copper electrorefining and can be further treated. Metals can also accumulate in the *flue dust* of the smelter or converter. A detailed overview of the existing copper production processes has been compiled by Moskalyk and Alfantazi [45]. Here, a condensed (schematic) overview of the copper production process and its residues is given (Fig. 6).

The composition of the process residues is very dependent on the type of ore and the process that is used to extract the copper. However, some representative examples can be found in the literature (Table 4).

Antimony-containing residues are an increasingly big issue in copper processing due to the deteriorating quality of primary copper ores. However, no recycling activities are currently carried out on industrial scale although there is a strong commitment to develop these in the future. Antimony is found in different intermediates and residues of the copper production process, and a wide variety of processes has been developed on laboratory scale to recover antimony from these waste streams. Awe et al. [53] developed a flowsheet to remove antimony from copper concentrates which feed the copper industry. Alkaline



**Fig. 6** Schematic overview of primary copper production from copper ores (Color figure online)

**Table 4** Literature examples of copper processing residues and their metal contents (wt%) [7, 46–51]

| Residue            | Sb   | Cu   | Ni  | Se  | Te  | As   | Bi  | Pb   | Zn  | Ag   | Sn   | Fe   |
|--------------------|------|------|-----|-----|-----|------|-----|------|-----|------|------|------|
| Slag [46]          |      | 0.4  |     |     |     |      |     |      |     |      |      | 49.3 |
| Flue dust [47]     | 3.1  | 5.3  |     |     |     | 2.8  | 2.8 | 28   | 8.9 |      | 12.0 | 1.2  |
| Anode slime 1 [48] | 3.0  | 25.4 |     | 8.2 | 1.3 | 3.4  | 0.8 | 2.9  |     | 14.5 |      |      |
| Anode slime 2 [49] | 24.6 | 1.44 | 2.4 |     |     |      |     | 0.5  |     |      | 17.7 | 2.4  |
| Anode slime 3 [7]  | 8.3  | 1.0  | 1.0 | 0.2 | 0.2 | 2.0  | 2.0 | 35.7 |     | 18.2 |      |      |
| Anode slime 4 [50] | 5.09 | 11.9 |     | 5.2 | 0.6 | 4.1  |     | 16.2 |     | 10.5 | 1.0  |      |
| Electrolyte [51]   | 1.0  | 90.3 |     |     |     | 7.51 | 1.2 | 0.1  |     |      |      |      |
| Electrolyte [52]   | 1.0  | 97.5 |     |     |     |      | 0.6 |      |     |      |      | 1.0  |

sulfide leaching was used ( $\text{Na}_2\text{S}/\text{NaOH}$ ) to extract antimony from the ore, which was then electrodeposited as antimony metal. Using this approach, the antimony content in the concentrate was reduced from 1.7 % to less than 0.1 % Sb, which is desirable for copper. Zhang et al. [54] studied the recovery of antimony and bismuth from pressure-leached flue dust created during the smelting of copper. They used a three-step process consisting of a kerosene desulfurization step, a chloride leaching step, and a final hydrolysis step to recover 95.80 % of Bi and 90.8 % of Sb. Vircikova et al. [47] investigated the removal of arsenic from converter flue dust as well as the behavior of antimony and bismuth. Arsenic was leached using a  $\text{Na}_2\text{S}$  solution and then precipitated using different methods. Recovery rates up to 99.9 % were achieved for arsenic but only 6.6 % for antimony. Fernández et al. [48] investigated the leaching of antimony and arsenic from anode slimes in the electrorefining of copper. The oxidized arsenic and antimony compounds in the anode slimes were selectively and almost completely dissolved in 0.4 M KOH at 80 °C. Meng et al. [49] studied the recovery of antimony(V) from the chloride leachate of copper anode slimes. Antimony

was recovered by hydrolysis in the form of  $\text{Sb}_2\text{O}_5$  with a recovery rate of 97 %. Anderson [7] reported the selective leaching of antimony and arsenic from copper electrorefining slimes, using alkaline sulfide leaching ( $\text{Na}_2\text{S}-\text{NaOH}$ ). Recovery rates were 99.3 % for Sb and 99.5 % for As. Li et al. [50] investigated alkaline fusion leaching as a method to recover valuable metals from copper anode slimes. The slime was fused with NaOH and  $\text{NaNO}_3$ , followed by water leaching. Se, As, Sn, and Pb were leached out, leaving behind a valuable concentrate of Cu, Sb, Te, and precious metals (Ag, Au, Pt).

Much research has also been carried out to remove impurities (e.g., Sb, Bi, As, Pb) from copper electrolyte solutions with sorbents and solvent extraction methods [55]. Some metals such as antimony can cause passivation of the electrodes during electrorefining and is therefore important to remove these from the electrolyte solution [56]. Ando and Tsuchida [52] studied the recovery of trace impurities (Sb and Bi) from copper electrolyte solutions, using adsorbents with aminophosphonic acid functional groups. High recovery rates of 99.5 % for Bi and 100 % for Sb were achieved. Deorkar and Tavlarides [57]



investigated the use of a silica gel ceramic support, functionalized with covalently bonded pyrogallol groups as a sorbent to separate antimony from copper and lead. The bed was regenerated by desorbing Sb(III) with HCl (4 M) and potassium hydrogen tartrate (0.05 M), resulting in a concentration factor of 25–30. Navarro and Alguacil [58] reported the adsorption of antimony and arsenic from a copper electrorefining solution onto activated carbon. Wang et al. [51] developed a new adsorbent based on Sb(V) and BaSO<sub>4</sub> as a carrier to recover impurities (e.g., Sb, Bi) from a copper electrolyte solution. Sb(V) can combine with Bi(III) and Sb(III) to precipitate these ions as Bi(SbO<sub>4</sub>) and Sb(SbO<sub>4</sub>). The sorbent was able to remove 90 % of Bi and 80 % of Sb from the acidic copper electrolyte solution. Xiao et al. [59] conducted studies on the removal of antimony and bismuth impurities from synthetic copper electrolyte solution. They used As(III) ions as collector to precipitate the impurities, but low removal rates were reported: 53 and 52 % for Sb and Bi, respectively. Xiao et al. [60] also studied the removal of Sb, As and Bi impurities using Sb(III) ions as collector to precipitate these impurities from a synthetic copper electrolyte solution, but with low recovery rates: 48.0 % for Sb and 38.4 % for Bi.

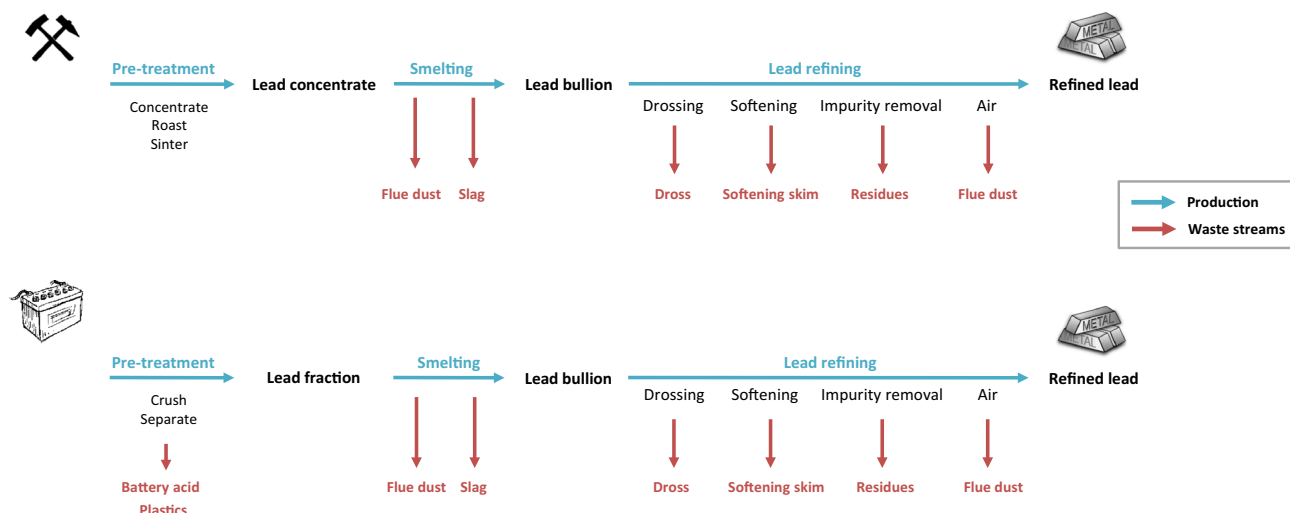
Solvent extraction has also been studied as a way to purify the copper electrolyte solutions and to extract valuable metals such as antimony or bismuth [61]. Szymanowski [61] discussed the use of phosphor-based extractants (e.g., TBP, DEHPA, Cyanex<sup>®</sup> 923), hydroxamic acids, and (poly)alcohols as a way to recover Sb, As, and Bi from copper electrolytes. Navarro et al. [62] studied the extraction of antimony from copper electrorefining solutions with the hydroxamic acid extractant LIX 1104SM and HCl as a stripping agent. They found that the order of extraction was Sb(III) > As(V) > Fe(III) ≫ Cu(II), with a distribution factor *D* for Sb of 299. Sarkar and Dhadke [63] studied the solvent extraction separation of Sb(III) and Bi(III) with bis(2,4,4-trimethylpentyl) monothiophosphinic acid (Cyanex<sup>®</sup> 302) from H<sub>2</sub>SO<sub>4</sub> and HCl solutions. They reported quantitative extraction of antimony and bismuth and separated the metals by first stripping with HNO<sub>3</sub> (2 M) to remove bismuth and then H<sub>2</sub>SO<sub>4</sub> (8.5 M) to strip antimony. Iyer and Dhadke [64] reported the quantitative extraction and separation of Sb(III) and Bi(III) from aqueous H<sub>2</sub>SO<sub>4</sub> and HCl solutions with the trialkyl phosphine oxide extractant Cyanex<sup>®</sup> 925. Sb(III) was stripped from the organic phase with H<sub>2</sub>SO<sub>4</sub> (8 M) and Bi(III) with HNO<sub>3</sub> (2–3 M). Other relevant studies include the work of Fuyii et al. [65] on the extraction behavior of Sb(III) in the TRUEX<sup>®</sup> system and a study of the co-extraction of Sb(III) by the Cu(II) extractant Acorga<sup>®</sup> CLX50 performed by Lin [66]. This pyridine-3,5-dicarboxylate ester extractant is used in the CUPREX<sup>®</sup> process to produce cathode-grade

copper with a hydrometallurgical process in which sulfide ore concentrates are leached in chloride medium [67]. The extraction of Sb(III) and Sb(V) ions with amine extractants in different acid solutions and the separation from other relevant metal ions (e.g., Bi(III), Sn(II), Cd(II), Te(IV), Se(IV), Pb(II), Cu(II), Au(III), Fe(III), and Zn(II)), have been studied extensively by Alian and Sanad [68] and Sargar et al. [69]. Facon et al. [70] also investigated the removal of Sb(III) from Bi(III), Pb(II), and Sn(IV) with Cyanex<sup>®</sup> 301 from chloride solutions.

## Lead Production

Antimony is also connected to primary and secondary lead production because it occurs in primary lead ores (e.g., galena PbS) as well as in the lead alloys used in lead-acid (LA) batteries, which use up the majority of the world's lead (76.7 % in 2006) [10]. Due to the large production volumes of lead (11,000,000 t/year in 2014), compared to antimony (160,000 t/year in 2014), lead production residues are considered as important secondary feedstocks of antimony [71]. Stringent environmental regulations and the exhaustion of high-quality primary lead deposits have caused an accelerated shift toward lead recycling as the main supply source [10]. In 2006, 60 % of the world's annual lead consumption was met by recycling of LA batteries, lead pipes, etc. [10]. In the U.S.A., secondary supply meets more than 80 % of the demand for lead and in countries such as Austria, the Netherlands, Spain, and Belgium the entire lead production is based on secondary raw materials from lead recycling [72]. Here, an overview is given of primary and secondary lead production with an emphasis on the formation of the different Sb-containing residues (e.g., slags, drosses, speiss, slimes) and the existing technologies to recover antimony from these residues (Fig. 7). Almost all antimony in secondary lead is currently being recycled [73]. For example, antimonial drosses are used to produce antimonial lead [73]. Efforts have also started to upgrade the process in order to produce pure Sb<sub>2</sub>O<sub>3</sub> from these secondary lead residues [74–76]. Residues with low antimony contents such as the slags and matte are less interesting when it comes to antimony recycling, but can have other applications [73]. Lead refiners are highly interested in valorizing the valuable Sb-containing waste streams both from an economic and environmental standpoint. The composition of the different process residues is dependent on the feed, equipment, and process, but some representative examples are shown in Table 5.

*Primary lead production* from lead sulfide ores (Galena: PbS) starts with the concentration of the ore, followed by a roasting and sintering step to form sinter (Fig. 7) [81, 82]. The lead concentrate is then fed into a blast furnace



**Fig. 7** Schematic overview of primary lead production from lead ores, and secondary lead production from spent lead-acid (LA) batteries. The different metal-containing waste streams and residues are indicated in red [73] (Color figure online)

**Table 5** Literature examples of lead processing residues and their metal contents (wt%) [7, 76–80]

| Residue            | Sb   | Pb   | Cu   | As   | Fe   | Ni  | Zn   | Ag  | Sn  | Bi  |
|--------------------|------|------|------|------|------|-----|------|-----|-----|-----|
| Slag [77]          |      | 2.8  | 0.1  |      | 37.1 |     | 11.2 |     |     |     |
| Harris dross [78]  | 8.2  | 67.0 |      |      | 0.8  |     | 0.7  |     |     |     |
| Speiss [7]         | 3.3  | 14.8 | 43.5 | 12.2 | 1.7  | 1.4 | 0.9  | 0.9 | 0.5 |     |
| Matte [80]         | 0.9  | 41.1 | 45.3 | 0.6  | 0.3  | 0.3 | 0.4  |     |     |     |
| Softening skim [7] | 31.7 | 52.9 |      | 3.3  |      |     |      |     |     |     |
| Sb dust [76]       | 42.4 | 7.5  | 0.12 | 10.4 |      |     |      |     |     | 0.6 |
| Slime [79]         | 63.6 | 12.6 | 1.47 | 4.0  |      |     |      | 1.1 |     | 3.3 |

together with limestone (flux) and coke in order to reduce the oxides to the metal [81]. New reactor designs also allow the roasting and smelting in a single reactor (e.g., Isasmelt furnace) [2, 82]. The non-metallic fraction (e.g., sulfides, silicates) forms a slag with the fluxing materials. During the smelting, a lighter phase is formed which is known as *speiss*. *Speiss* consist mainly of iron arsenides and antimonides, and other elements such as Pb, Cu, Ni, Sn, and significant levels of precious metals (Ag 9 kg/t and Au 45 g/t) [7]. It rises to the top of the melt and can be skimmed off. Furthermore, a *matte* layer consisting of copper and other metal sulfide impurities is formed. The *speiss* and *matte* can be sold to copper smelters, where they are refined for copper recovery. The lead coming from the smelting furnace, called lead bullion, still contains many impurities (e.g., Cu, As, Sb, Sn, Bi, Zn, Ag, Au), and needs to be refined. First, copper is removed in a drossing process [81]. The melt is cooled to decrease the solubility of copper, and a *copper dross* is formed on the surface [81]. Sulfur can also be added to remove the last traces of copper. Dross is usually skimmed off and sent to a dross

furnace to recover the non-lead components. The lead is then subjected to further purification steps either by thermal refining or by electrorefining [81]. In thermal refining, the lead is heated and cooled under different conditions and at different temperatures to oxidize or to remove the metal impurities from the lead. Arsenic, tin, and antimony are best removed in a softening step, which consists of heating the lead in a furnace in the presence of air, which causes the impurities with a greater affinity for oxygen than lead, to be oxidized. The *softening skims* can then be collected from the surface of the bath. An alternative is the *Harris process*, which uses a flux of molten NaOH and NaNO<sub>3</sub> to remove the impurities as sodium arsenate, stannate, and antimonate [83, 84]. Further purification steps are used to remove gold and silver (Parkes process), zinc (vacuum distillation), bismuth (Kroll–Betters process), and other impurities such as antimony (caustic refining) [2, 82, 84–86]. These refining processes are combined to obtain a lead bullion with sufficient purity, before casting it into ingots [81]. Instead of this sequence of thermal refining steps, pure lead can also be obtained by electrorefining after the

initial decoppering step [87]. Solutions of  $\text{H}_2\text{SiF}_6$  (Betts process) or  $\text{HBF}_4$  are used and the work-lead is cast into anodes and then dissolved [88]. The metals that are more noble than lead (e.g., As, Sb, Bi, Au, Ag) do not dissolve and are collected in the *anode slimes* [2]. The advantage of electrorefining is that unlike thermal refining all the impurities are removed in one step and 99.999 % pure lead is obtained [2]. In the sintering, smelting, and refining stages, fumes and dusts may contain between 10 and 60 % Pb and a wide range of other metals (e.g., Cu, Zn, As, Sb, Sn, Cd, Ag, Au, Bi). Plants are normally designed to capture these *flue dusts* in filters in order to discard them safely or resmelt them in certain cases [81].

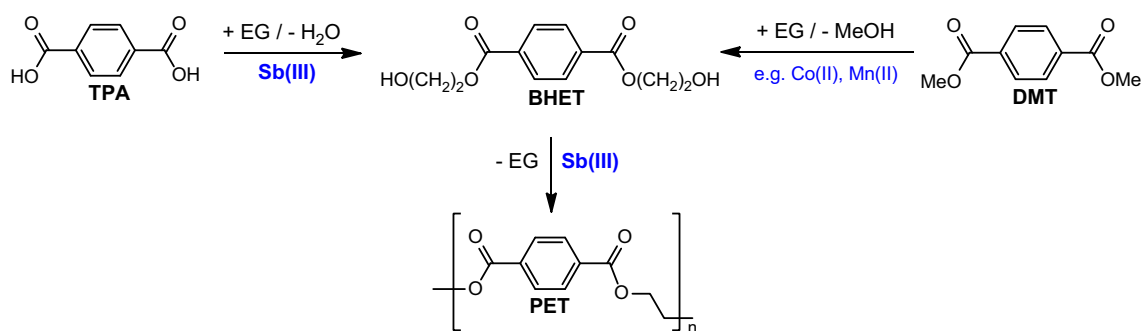
*Secondary lead production* is almost entirely dependent on the recycling of spent lead-acid batteries (Fig. 7) [2, 73]. After removing the acid, the batteries are crushed to separate lead from the plastic. The lead can be separated in a lead metal fraction and a lead oxide fraction by a mechanical process [2]. The lead metal is then simply remelted, whereas the lead oxide is reduced in several types of furnaces (e.g., rotary furnaces or Isasmelt furnaces) [2, 11]. Besser et al. reviewed the different types of furnace designs and processes for battery recycling as well as their efficiency, and ability to produce pure lead and lead-antimony alloys for battery grid manufacturers and other lead alloy applications [10, 89]. The lead is refined using similar processes as the primary lead production and the same type of residues (e.g., skims, drosses, flue dust) are obtained [73].

The recovery of antimony from lead production residues has been investigated for some time [73]. Anderson [7], studied the use of alkaline sulfide leaching as a way to selectively remove antimony (and arsenic) from lead smelting residues. A mixture of  $\text{Na}_2\text{S}$  and  $\text{NaOH}$  was used to solubilize antimony as thioantimonite  $\text{Na}_3\text{SbS}_3$ . High antimony recovery rates were reported for lead smelter speiss (99.4 %), skims (89.0 %), and flue dust (95 %) [7]. Singh [78] reported the recovery of antimony from the antimony-containing Harris dross (8.2 % Sb) of the refining section of a lead plant.  $\text{HCl}$  was used to leach antimony from the dross (95 % recovery) and  $\text{Sb}_2\text{O}_3$  was formed by hydrolysis (73.6 % recovery) [78]. Peterson and Twidwell [90] investigated the removal of arsenic and antimony from lead smelter speiss, using volatilization techniques. The removal of antimony from antimony dust captured during the direct lead smelting process has been studied by Liu et al. They proposed a pyrometallurgical process based on reduction smelting, alkaline refining, and oxidation to extract antimony from antimony dust and produce  $\text{Sb}_2\text{O}_3$  with a purity above 99.8 % [76]. Cao et al. studied the recovery of antimony from lead anode slimes, using a potential-controlled chlorination leaching ( $\text{Cl}_2$  leaching) and continuous distillation [79]. High-purity  $\text{SbCl}_3$  was

prepared, with an antimony recovery rate of more than 95 %. The process forms a closed loop and no waste is created [79]. Itoh et al. [91], proposed an alternative process to recover antimony from the anode slimes (Pb–Bi–Sb alloy) of spent lead-acid battery processing. In a method called *volatile oxide formation*, antimony is selectively oxidized due to its higher affinity for oxygen compared to lead or bismuth.  $\text{Sb}_2\text{O}_3$  is then evaporated because of its high vapor pressure. Lin and Qiu [92] also developed a process to recover antimony and arsenic from anode slimes. They used a new process called *vacuum dynamic flash reduction* to evaporate the volatile oxides of antimony and arsenic, leaving silver behind. A recovery of 93.67 % was achieved for antimony and 98.92 % for arsenic. The vacuum treatment eliminates much of the air pollution and material losses associated with other conventional treatment methods. Qui et al. also used vacuum evaporation technology to recover antimony from Sb-rich anode slimes from the lead industry [93]. The antimony recovery rate was 92 %, and a 99.7 % pure antimony oxide distillate was obtained. Binz et al. [75] developed a bottom-up process to recover antimony from lead-refining residues. They investigated a more selective oxidation during lead softening and the carbothermic reduction of the slag to produce a concentrated antimony slag and improve the recovery of antimony. Kaporal et al. [94] studied the feasibility of antimony removal from accumulator acid found in spent lead-acid batteries. They achieved 100 % removal of antimony from synthetic solutions, using electrodeposition with a copper electrode. Bergmann and Kaporal [95] continued this work and studied the electrochemical recovery of antimony from real spent accumulator acid found in lead-acid batteries. They managed to reduce antimony levels from 5 ppm to less than 0.15 ppm, while also regenerating the accumulator acid. Copper and graphite cathodes produced the best results.

### Spent Antimony Catalyst

Antimony is also used in several catalysts [96–101]. When these catalysts reach their end-of-life, antimony can be recovered and reused. The most important application is the production of polyethylene terephthalate (PET), one of the most common polyester thermoplastic polymers. This polymer is mostly used in textile fibers and as container for food or liquids. PET is produced by the Sb(III)-catalyzed polycondensation of bis(hydroxyethyl)terephthalate (BHET) (Fig. 8). This precursor is either synthesized by the transesterification of dimethyl terephthalate (DMT) or the esterification of terephthalic acid (TPA). Titanium-based catalysts have been proposed, but today more than 90 % of the world's polyester production still runs on Sb-based catalysts (150–300 ppm) [96, 102].



**Fig. 8** Two pathways for the production of PET (EG ethylene glycol)

The most common forms are  $\text{Sb}_2\text{O}_3$ ,  $\text{Sb}(\text{OCH}_2\text{CH}_2\text{O})_3$ , and  $\text{Sb}(\text{OAc})_3$  [103]. High-purity compounds are required, especially in regard to arsenic since the catalyst will be present in the final PET polymer used for making plastic bottles. Despite the efficiency of antimony catalysts, there are drawbacks such as the decomposition, which deposits antimony metal particles [103]. This causes a grayish discoloration in the polymer, especially at concentrations  $>250$  ppm Sb [96]. The mechanism behind the formation of antimony metal during PET polymerization and the discoloration was studied in detail by Aharoni [104]. Although antimony is present in the PET polymer (100–300 ppm), it has been demonstrated that water stored in PET bottles contains less than 1 ppb of Sb, which is far below the EEC limit of 20 ppb [96, 105]. The mass recycling of PET plastic could therefore be an opportunity to recover antimony, although antimony contents are typically lower than in plastics containing antimony-based flame retardants. Therefore, most studies have focused on the recovery of antimony from spent catalysts so far. Dougherty and Garska [12] developed a process to recover antimony from spent ethylene glycol residues resulting from the manufacture of PET polymers. Their method consists of combusting the spent glycol residues to produce an ash, from which antimony can be recovered. Several patents have also been published on the recovery of antimony from halocarbon solutions, because Sb-containing catalysts (e.g.,  $\text{HF}/\text{SbCl}_5$ ) are used in the fluorination of chlorinated hydrocarbons [99, 106, 107]. Fernschild et al. [106] described a method for the recovery of an Sb-containing catalyst by chlorination, followed by the distillation of  $\text{SbCl}_5$ . Hyatt [107] developed an alternative process based on the reduction of  $\text{SbCl}_5$  to  $\text{SbCl}_3$ , followed by extraction of  $\text{SbCl}_3$  from the halocarbon solution to an acidic aqueous phase. Finally, Anderson [7] also studied the recovery of antimony from a spent catalyst used in the production of acrolein from propylene. Antimony was successfully leached out (99.5 %) using HCl, leaving the silica substrate intact.

## End-of-Life Products and Municipal Waste

Antimony can be found in a variety of products, which eventually end up as waste. It is therefore important to understand the presence of antimony in different items, in order to assure that no antimony is released into the environment. The high antimony content in certain products also makes them very interesting as secondary sources of antimony either through direct recycling or through municipal waste incineration. Table 6 gives an overview of the different antimony products and their approximate growth rates [3].

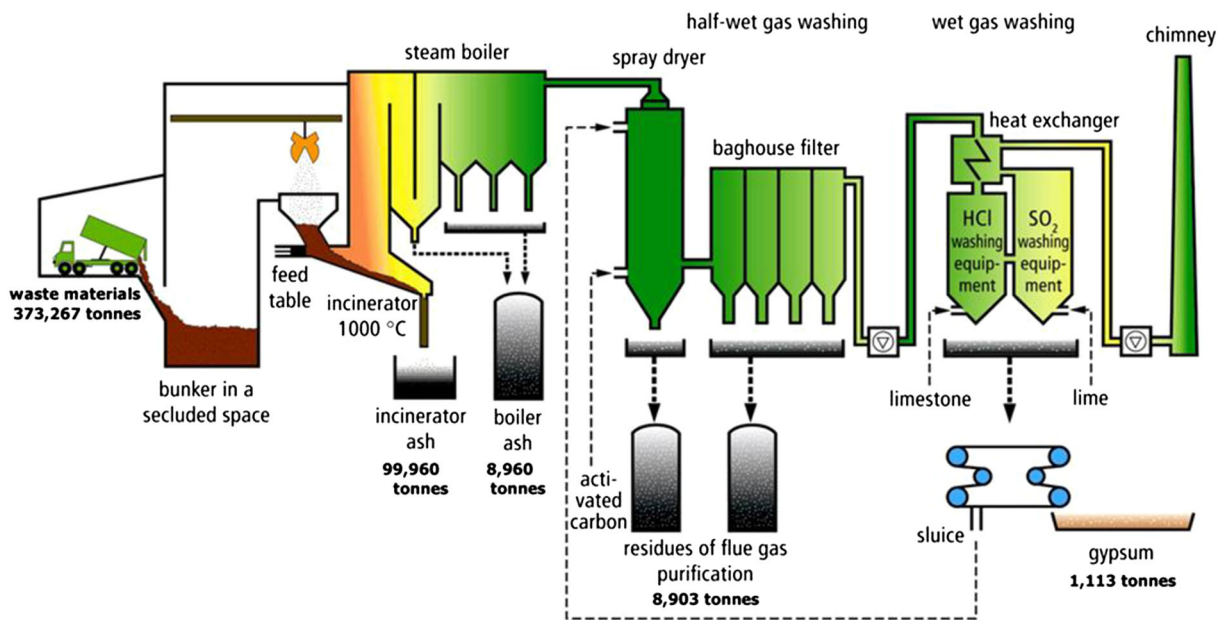
## Municipal Solid Waste Incineration

Municipal solid waste incineration (MSWI) is a vital part of many waste management systems. The residues formed during the incineration contain many valuable elements and have been investigated as secondary source of ferrous and non-ferrous metals. In a MSWI plant, the waste is incinerated at a temperature of 850–1000 °C, forming a solid bottom ash fraction and flue gas, which is captured and cooled to form boiler ash, fly ash, and air pollution control (ACP) residues. A schematic overview of a MSWI plant is shown in Fig. 9.

The waste incineration residues can undergo different treatments (e.g., dry, wet, thermal, mechanical) aimed at removing ferrous and non-ferrous metals and/or valorizing it as a granulate for the construction industry, e.g., in road foundation. A lot of different elements are present in these ashes, but the discussion is restricted here to the recovery of antimony. Jung et al. [109] and Allegrini et al. [13] compiled an overview of the composition of MSWI bottom ash, fly ashes, and other waste incineration residues, and compared this with typical ore concentrations to determine the recovery potential (Table 7) [13, 109]. The typical ore concentration given in Table 7 reflects the average grade of the deposits, but does not take into account the ore's abundance. The widely available incinerator residues could

**Table 6** Antimony consumption (tons) in 2000 and 2010 per product category and its main market drivers (Source: Roskill consulting) [3]

| Product             | 2000 (t) | 2010 (t) | Annual growth rate (%) | Market driver |
|---------------------|----------|----------|------------------------|---------------|
| Flame retardants    | 70,000   | 103,500  | 4.0                    | Plastics      |
| Lead-acid batteries | 40,000   | 53,000   | 2.9                    | Automotive    |
| Lead alloys         | 11,000   | 23,000   | 7.7                    | Construction  |
| PET catalysts       | 6000     | 11,400   | 6.6                    | PET           |
| Heat stabilizer     | 1,400    | 2600     | 6.4                    | PVC           |
| Ceramics            | 1700     | 2500     | 3.9                    | Construction  |
| Glass               | 16,000   | 1700     | −20.1                  | CRT glass     |
| Others              | 1500     | 1840     | 2.1                    |               |
| Total               | 147,600  | 199,540  | 4.1                    |               |



**Fig. 9** Schematic representation of the MSWI plant in Doel, Belgium. This figure is reproduced with permission from KU Leuven SIM<sup>2</sup> [108]

therefore become an attractive secondary resource. Fly ashes are particularly interesting, due to the volatility of antimony which results in an important enrichment of antimony in the fly ashes. However, presorting of materials with high antimony contents (e.g., plastics with flame retardants), could drastically improve the efficiency of such an incineration process.

Nakamura et al. [111] investigated the origin of antimony, lead, and cadmium in incinerator waste. They concluded that the relatively high content of antimony in these incineration residues is mainly due to the presence of antimony in plastics, glass, and textiles (Table 8). Curtain fabrics had a particularly high antimony content (2100 ppm), due to the use of flame retardant coatings

**Table 7** Content (mg kg<sup>-1</sup>) of a selection of metals in MSWI bottom ash, boiler ash, and fly ash, compared to typical ores

| Element | Bottom ash | Boiler ash  | Fly ash     | Typical ore     |
|---------|------------|-------------|-------------|-----------------|
| Sb      | 10–400     | 200–1000    | 260–1100    | 27,000          |
| As      | 1–200      | 20–60       | 40–300      | 1000–40,000     |
| Cu      | 300–8000   | 500–1000    | 600–3200    | 5000–20,000     |
| Pb      | 100–14,000 | 1000–35,000 | 5300–26,000 | 300,000–400,000 |
| Sn      | 2–400      | 200–700     | 550–2000    | 4000            |

Note: these are representative values, but other reports may differ [109]

A range is given, based on the work of Van Gerven et al. [110] and Allegrini et al. [13, 110]



[111]. In general, Watanabe et al. [112] estimated that raw incinerator waste contains approximately 40–50 mg kg<sup>-1</sup> of antimony.

The removal of antimony from incinerator residues has been an important topic of discussion, mainly because of its toxicity and the risk for uncontrolled release of antimony. Osaka et al. [113] wrote a report on the risks associated with the presence of antimony in municipal waste incineration residues. They studied the influence of the pH on the leaching of antimony from these residues and proposed risk management measures. Paoletti et al. (2001) and Nakamura et al. (1996) studied the movement and distribution of antimony in the different incinerator residues [111, 114]. These risk management studies focused mainly on limiting the leaching of antimony from these residues. However, due to the high criticality of antimony, the recovery of antimony from these residues may become of higher interest, especially since consumer waste contains important amounts of antimony in textiles, plastics, and glass (Table 8). Miravet et al. [115] analyzed the leachability and speciation of antimony in incinerator fly ash. They used citrate solutions to leach out antimony and concluded that Sb(V) was the main Sb-containing species. The extraction of metals from incinerator residues through leaching was also studied by Van Gerven et al. [110]. They performed column and batch leaching tests with acidified water (HNO<sub>3</sub>) to estimate the leachability of a wide range of metals from MSWI bottom ash, boiler ash, fly ashes, and air pollution control (APC) residues. Van Gerven et al. [116] also studied the removal of heavy metals from MSWI bottom ashes using organic solutions of citric acid and ammonium citrate. Cornelis et al. [14, 15] investigated the influence of the pH on the leaching of antimony from carbonated and non-carbonated MSWI bottom ash. Hong et al. studied the extraction of heavy metals from MSW incinerator fly ashes, using acids (HCl), and chelating agents (NTA, EDTA, DTPA) in batch experiments. The leaching test on the residues after the treatment with chelating agents showed that the fly ashes were successfully detoxified to meet the guideline for landfilling. Okkenhaug et al. [117] showed that iron-rich sulfuric acid

**Table 8** Antimony content and contribution in crushed municipal waste

| Waste type | Sb content (mg kg <sup>-1</sup> ) | Contribution to waste (%) |
|------------|-----------------------------------|---------------------------|
| Plastics   | 180                               | 44.7                      |
| Glass      | 1008                              | 15.2                      |
| Textiles   | 111                               | 14.7                      |
| Wood       | 11                                | 9.3                       |
| Other      | 93                                | 16.1                      |

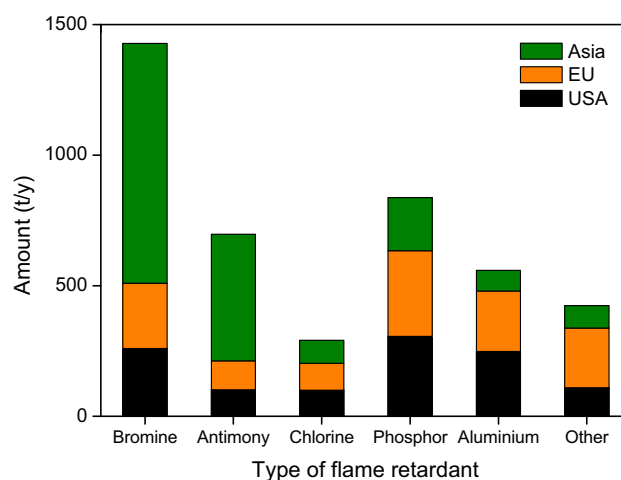
The data were adapted from Nakamura et al. [111]

waste is efficient to immobilize antimony in MSWI air pollution control residues. These technologies are all promising but the winning technology will be the one that can *selectively* remove certain elements such as antimony. Due to the mix of elements in these residues, it is crucial to develop methods which can deal with the complexity of these powders in a low-cost and efficient manner. Despite the technical challenges, there is a significant opportunity to recover valuable metals from these ashes as they are available in large quantities.

### Flame Retardants in Plastics

In 2014, the world consumption of flame retardants was more than 2.2 million tons, representing a market of approximately 5 billion EUR with a 4–5 % annual growth rate due to rising safety standards worldwide [4, 118]. Flame retardants are crucial to guarantee the safety of plastics, construction materials, electronics, textiles, and coated products [119, 120]. There are three main classes of flame retardants: (1) metal hydroxides such as Al(OH)<sub>3</sub> (ATH) and Mg(OH)<sub>2</sub> (MDH), (2) organohalogen compounds (chlorinated or brominated) in combination with antimony trioxide (Sb<sub>2</sub>O<sub>3</sub>), and (3) organophosphorus compounds [119]. An overview of the flame retardant market (2007) is given in Fig. 10.

Continuous research and regulations are driving new developments in the different categories of flame retardants. For example, research is being conducted to graft flame retardant groups in the structure of the polymer, thus avoiding the release of the flame retardants into the environment [122–124]. The most effective commercial fire retardant systems are currently based on brominated organic compounds (known as brominated flame retardants



**Fig. 10** Flame retardant sales per region and per category (2007), expressed in t/year. Adapted from SRI-Consulting [121] (Color figure online)

(BFRs)), preferably in conjunction with antimony trioxide  $\text{Sb}_2\text{O}_3$  due to the strong synergetic effect [120]. Flame retardants were responsible for 52 % of the demand for antimony in 2010 ( $\approx 103,500$  t) [3]. Camino and Costa [119], reviewed the performance and mechanisms of antimony fire retardants in polymers. They attributed the synergetic effect to the fact that metal halides are far more effective flame inhibitors, compared to the hydrogen halides formed in the absence of the metal compound [120]. In reality, a variety of different oxyhalide metal species are formed and their thermal stability and chemical reactivity have therefore been studied in great detail [125]. A discussion of the fire retardant mechanism is beyond the scope of this work, and what follows is therefore only a discussion of the possible recovery of antimony from plastics, electronics, and coatings. The antimony content in plastics is highly dependent on the type of plastic and the application. Table 9 provides an overview of the typical formulations used to fireproof different polymers [6].

Non-halogenated polymers are inherently flammable and require large amounts of halogenated flame retardants in combination with  $\text{Sb}_2\text{O}_3$  to be flame resistant [6]. Halogenated polymers (e.g., PVC, PVDC) on the other hand, are already partially flame retarded due to their chlorine content and do not require the addition of halogenated compounds. Smaller amounts of  $\text{Sb}_2\text{O}_3$  can be added to these polymers to ensure their fire resistance [6]. Due to the inherent risk for fires, polymers used in building materials, electronics, and some textiles are often made fireproof using these halogenated compounds and  $\text{Sb}_2\text{O}_3$ . Plastics used for food and beverages or plastics that do not require fire-proofing, do not contain significant quantities of antimony. PET bottles are an exception, as these contain some antimony (0.2–0.3 wt%) due to the use of an antimony catalyst for the production of PET polymers [126]. Efficient recovery of antimony from plastics requires adapted screening and sorting methods to identify plastics with high antimony contents. X-ray fluorescence

spectroscopy (XRF) could provide a fast and non-destructive screening. Other interesting techniques are X-ray transparency and laser-induced breakdown spectroscopy (LIBS). Bellara et al. [127] also developed a method for the direct determination of antimony in solid PVC samples by graphite furnace atomic absorption spectrometry (GFAAS). They concluded that this screening technique provides sufficiently reliable results in a reasonably short time, but it is difficult to implement in a continuous process. Once sorted, the antimony-containing plastics can be placed in special ovens for the pyrolysis of the polymer, where antimony is caught in the residues (bottom ashes or fly ashes). A patent of Nippon Electric Co (1994) described a method to decompose a resin, containing flame retardants, and capture the bromine and antimony in a gas mixture (e.g.,  $\text{HBr}$ ,  $\text{SbBr}_3$ ,  $\text{CO}_2$ ,  $\text{H}_2\text{O}$ ) [128]. By washing the gas mixture with an alkaline aqueous solution (e.g.,  $\text{NaOH}$ ),  $\text{NaBr}$  and  $\text{Sb}_2\text{O}_3$  could be obtained. The thermal decomposition could be operated under air-free conditions or by combustion. In a follow-up patent, Masatochi et al. [129] specifically described the recovery of antimony from decomposition gasses, originating from the pyrolysis of flame resistant plastics. Masatoshi and Ikuta [130] also described the pyrolysis-based recovery of antimony and bromine from molding resins used in electronics. The electronic parts consisted of an epoxy resin, a silica filler, and flame retardants consisting of brominated compounds and  $\text{Sb}_2\text{O}_3$ . This study investigated the best pyrolysis conditions to obtain a pure silica product. The exhaust gasses generated in the pyrolysis of the molding resin were treated to remove the bromine and convert the antimony bromides to  $\text{Sb}_2\text{O}_3$ . Jakab et al. [131] studied the thermal decomposition of flame-retarded high-impact polystyrene. In a detailed overview of the decomposition pathways and mechanisms, they concluded that the brominated additives themselves do not change the decomposition temperature of polystyrene, but  $\text{Sb}_2\text{O}_3$  reduces the thermal stability of the samples by acting as an initiator for the decomposition of the brominated flame retardants. Klein et al. [132] also studied the behavior of antimony during thermal treatment of antimony-rich halogenated waste. Their aim was to ensure the environmentally friendly waste incineration of antimony-rich plastic waste, by strengthening the knowledge about the fate of antimony and the potential formation of harmful species as function of temperature, residence time and flow rate of air in the furnace. Their investigations showed that under moderate oxidative conditions, the partition of antimony between the residual bottom ash ( $\approx 64$  %) and the gas phase ( $\approx 36$  %) was constant regardless of the temperature. However, they observed that at  $850$  °C antimony was mainly present in the gas phase as  $\text{Sb(III)}$ , while around  $1100$  °C,  $\text{Sb(V)}$  was favored. Wu et al. [133] studied the thermal degradation of high-impact

**Table 9** Required  $\text{Sb}_2\text{O}_3$  to make polymers flame resistant [6]

| Plastic | Flammability | $\text{Sb}_2\text{O}_3$ (wt%) | BFR required? |
|---------|--------------|-------------------------------|---------------|
| PE      | High         | 8–16                          | Yes           |
| PP      | High         | 5–15                          | Yes           |
| PS      | High         | 10                            | Yes           |
| PU      | High         | 10                            | Yes           |
| SBR     | High         | 5–30                          | Yes           |
| ABS     | High         | 5–12                          | Yes           |
| PVC     | Low          | 1–10                          | No            |

Polymer abbreviations: *PE* polyethylene, *PP* polypropylene, *PS* polystyrene, *PU* polyurethane, *SBR* styrene-butadiene rubber, *ABS* acrylonitrile–butadiene–styrene, *PVC* polyvinylchloride, *BFR* brominated flame retardant

polystyrene, containing brominated flame retardants and  $\text{Sb}_2\text{O}_3$ . Bauxite residue (red mud) was used as an additive to catch the bromine and antimony in a residue, while producing a valuable pyrolysis oil. Mitan et al. [134] investigated the controlled pyrolysis of polyethylene/polypropylene/polystyrene mixed with brominated high-impact polystyrene. Their work describes the effect of the brominated flame retardants and  $\text{Sb}_2\text{O}_3$  on the pyrolysis process and the distribution of pyrolysis products. Brebu et al. [135] also looked at the effect of flame retardants and  $\text{Sb}_2\text{O}_3$  synergists on the thermal decomposition of high-impact polystyrene and on its debromination by ammonia treatment. More than 90 wt% of initial bromine could be recovered as inorganic  $\text{NH}_4\text{Br}$ . Hall et al. [136] developed a method for the co-pyrolysis of flame-retarded high-impact polystyrene and polyolefins in a fixed bed reactor. They looked at the effect that the different types of brominated aryl compounds and  $\text{Sb}_2\text{O}_3$  have on the pyrolysis products.

Besides pyrolysis, other methods have also been proposed. Omwudili and Williams [137] attempted the alkaline reforming of high-impact polystyrene and traced the movement of bromine and antimony. The waste polymer was reacted in supercritical water and NaOH in a pressure reactor in order to neutralize the corrosive inorganic bromine species released during the reactions. They observed that the presence of the alkaline supercritical water led to 97 wt% debromination of the product oil, producing a valuable bromine-free oil feedstock. Furthermore, a 98 % removal rate was achieved for antimony. The same method was also used for acrylonitrile–butadiene–styrene polymers [138]. Another unusual approach was tested by Lateef et al. [139]. They used imidazolium- and pyridinium-based ionic liquids to extract brominated flame retardants and  $\text{Sb}_2\text{O}_3$  from high-impact polystyrene. They identified 1-hexylpyridinium bromide as the most suitable for solid–liquid and liquid–liquid extraction, achieving extraction percentages of 92.7 % for the brominated flame retardants and 99.9 % for  $\text{Sb}_2\text{O}_3$ . Furthermore, they demonstrated that the chain length of the polymer was not degraded by the extraction process, thus forming a polymer residue which can be recycled. Another approach could be the depolymerization of the polymers to recover  $\text{Sb}_2\text{O}_3$  and the building blocks (e.g., ethylene, HCl). This approach has not been investigated in detail so far, but some projects have looked into the possibility of separating, cleaning, and reusing bromine- and antimony-containing plastics [140].

### Plastics in Waste Electrical and Electronic Equipment

A subcategory of flame retardant plastics are the plastics used in waste electrical and electronic equipment (WEEE)

[141]. WEEE is already being recycled on a large scale, mainly to recover the precious metals (e.g., Co, In, Te, Se, Au, Ag) used in the electronic components such as printed circuit boards (PCBs) (Table 10) [142]. Besides the metallic fraction, there is also a large non-metallic fraction in WEEE [143]. Even for PCBs the non-metallic fraction is still more than 70 % [143]. This non-metallic fraction includes the silicon substrates and the plastics used in these products. The strict safety regulations for electrical and electronic equipment (EEE), require large amounts of brominated flame retardants and  $\text{Sb}_2\text{O}_3$  to be used in order to guarantee the safe use of these plastics [142]. As explained previously, approximately 52 % of antimony is consumed in flame retardants, of which 70 % is used for plastics in electrical and electronic equipment (Table 6) [3, 142].

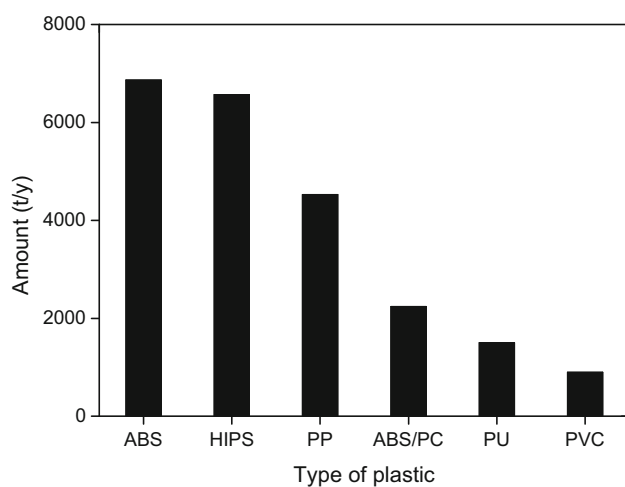
This clearly shows the potential of WEEE plastics as a secondary feedstock of antimony. Table 9 showed that certain plastics require as much as 5–15 wt% of antimony to be fire resistant, making WEEE particularly rich in antimony. The most important plastics under investigation are acrylonitrile–butadiene–styrene (ABS) and high-impact polystyrene (HIPS) as these are the most-used polymers in EEE (Fig. 11) [141]. These polymers typically contain 5–10 wt% of  $\text{Sb}_2\text{O}_3$  and an even larger amount of brominated flame retardants. [142]. Different studies have been carried out to deal with these polymers in WEEE recycling schemes. The toxic brominated compounds and antimony-containing plastics have been a source of concern for the recycling of WEEE. Regulations for the emission of brominated compounds are very strict and therefore solutions have been investigated to capture the bromine efficiently during the pyrolysis of the WEEE [142]. The fact that antimony is also present in the plastic fraction, means that bromine and antimony are intrinsically connected in WEEE recycling.

De Marco et al. and Moltó et al. reviewed the pyrolysis of electrical and electronic wastes but did not discuss in detail the recovery of antimony or bromine flame retardants [144, 145]. Guo et al. [143] reviewed the recycling of non-metallic fractions (NMFs) from waste printed circuit boards (PCBs) and demonstrated the importance of finding a solution for the recycling of this important fraction of WEEE. The non-metallic fraction was usually treated by combustion or landfilling in the past. However, uncontrolled combustion causes the formation of highly toxic polybrominated dibenzodioxins and dibenzofurans, while landfilling of the NMFs will lead to secondary pollution caused by heavy metals and brominated flame retardants (BFRs) leaching to the groundwater. This had been previously demonstrated by the work of Gullet et al. [146], who characterized the emissions and residual ash from open air burning of electronic wastes during simulated rudimentary

**Table 10** Metal consumption for electrical and electronic equipment EEE (t/year)

|    | Yearly demand for EEE (t/year) | Percentage of total production (%) |
|----|--------------------------------|------------------------------------|
| Cu | 4,500,000                      | 30                                 |
| Sn | 90,000                         | 33                                 |
| Sb | 65,000                         | 50                                 |
| Co | 11,000                         | 19                                 |
| Ag | 6000                           | 30                                 |
| Bi | 900                            | 16                                 |
| In | 380                            | 79                                 |
| Au | 300                            | 12                                 |
| Se | 240                            | 17                                 |

The data are obtained from the work of Buekens and Yang [142]



**Fig. 11** Flows of the major plastic types in Swiss WEEE products for the year 2007 [141]. *ABS* acrylonitrile–butadiene–styrene, *HIPS* high-impact polystyrene, *PP* polypropylene, *PC* polycarbonate, *PU* polyurethane, *VC* polyvinylchloride

recycling operations. They analyzed the fly ash and particulate matter and characterized the organic and metallic components in the air emissions. Therefore, Guo et al. [143] concluded that more advanced processes are required such as pyrolysis, gasification, depolymerization, or hydrogen degradation in order to convert the non-metallic fraction to chemical feedstocks and fuels. These methods have the advantage of eliminating hazardous substances such as brominated organic compounds from this waste fraction and allow the recovery of antimony from the plastic waste. In later work, Guo et al. [147] analyzed the pyrolysis of scrap printed circuit board plastic particles in a fluidized bed reactor. The gas products, liquid products, and solid residues were analyzed and it was found that the liquid yields increased with an increase in pyrolysis temperature. The main compositions of liquid products were

aromatic compounds including substituted benzenes, whereas the solid products mainly contained char and fiberglass. Morf et al. [148] also tracked the movement of brominated flame retardants and antimony in WEEE recycling plants. Hall et al. [149] investigated the fast pyrolysis of plastics recovered from computer waste. The computer case was made from PVC, but the computer monitor cover consisted of brominated acrylonitrile–butadiene–styrene and had an antimony content between 3.9 and 4.5 wt%. The pyrolysis was carried out in a fluidized bed reactor at 500 °C, and the composition products were studied. They concluded that the high bromine and chlorine content was a nuisance and that in-process sorption of the halogens was required to obtain a high-quality oil. In later work, Hall et al. [150] used zeolite catalysts and fluidized catalytic cracker (FCC) catalysts to destroy the toxic organobromines during pyrolysis of brominated high-impact polystyrene (4.6 wt% Sb) and acrylonitrile–butadiene–styrene (3.2 wt% Sb), found in WEEE. The movement of antimony in the oil and char residues was also tracked. Tostar et al. [151] specifically investigated the leaching of antimony in plastics from WEEE with various acids and gamma irradiation. The most efficient leaching medium was a heated solution of sodium hydrogen tartrate in dimethyl sulphoxide, which leached approximately 50 % of the antimony from the acrylonitrile–butadiene–styrene plastic. Gamma irradiation did not significantly change the amount of leached antimony. A patent by Zhenming et al. [152] also describes vacuum distillation as a way to recover antimony from printed circuit board waste. Yang et al. [153] reviewed the pyrolysis and dehalogenation of plastics from waste electrical and electronic equipment. The conclusion from these works is that there are technologies available to valorize the plastic components of WEEE, but that additional research needs to be done to improve the efficiency of these processes before upscaling becomes economically feasible. These plastics have to be processed appropriately due to their high content of toxic brominated compounds. The focus up till now has been on limiting the emission of brominated compounds and producing useful fuel oils. However, the high antimony content present in these same plastics offers a unique opportunity to combine the necessary treatment of these brominated plastics with the recovery of valuable antimony. Valorizing antimony from WEEE plastics could improve the economics of the total WEEE recycling process and offset the additional cost of de-brominating during the processing of WEEE plastics.

### Lamp Phosphor Waste

Antimony is used in the halophosphate (HALO) lamp phosphors, found in fluorescent lamps [154, 155]. These



lamp phosphors are coated on the inside of the glass and are responsible for the emission of visible light [156–159]. The HALO phosphor was invented in 1942 and large amounts of this phosphor were produced as it found widespread use in fluorescent tubes due to its emission of intense white light [156–159]. This phosphor usually consists of a fluoro-chloro apatite doped with Mn(II) and Sb(III) and has the general formula  $(\text{Sr,Ca})_5(\text{PO}_4)_3(\text{Cl,F})\text{Sb}^{3+}, \text{Mn}^{2+}$  [156–158]. Antimony is added as a blue dopant to adjust the shade of white [156–158, 160]. Today, lamp phosphor waste contains around 50 wt% of HALO, making it the largest component [154, 155, 161]. This corresponds to approximately 0.5–1 wt% of antimony which is similar to a low-grade stibnite ore [1, 155, 157, 161, 162]. With annual sales of CFLs topping  $2.5 \times 10^9$  units/y (2007), the volume of lamp phosphor waste is expected to continue growing at a rapid pace as fluorescent lamps reach their end-of-life [163, 164]. Due to their mercury content, fluorescent lamps are considered as hazardous waste and are therefore collected separately in most countries [165, 166]. The interest in lamp phosphor powders as a secondary resource has been growing recently due to their high rare-earth content (in tri-band phosphor), and the recovery of these valuable elements has been studied extensively [165–175]. Currently, a recycling process is operated on industrial scale ( $>1000$  t/y) by Solvay in France [161]. So far, the Sb-containing HALO phosphor is often still discarded as a non-valuable residue due to the absence of rare earths [165, 166, 172]. However, the HALO content in these powders is too high to ignore (50 wt%) and could serve as an interesting secondary source of antimony [176]. Our group therefore recently developed a zero-waste valorization method to recover antimony from HALO and to valorize the remaining as a calcium phosphate (apatite) product, which is a feedstock for the fertilizer industry (Fig. 12) [176]. The valorization of HALO and the recovery of antimony can be integrated in rare-earth recovery schemes and in the broader effort to recycle these lamp phosphor powders [175, 176].

## Antimony Emissions and Recovery

### Antimony Emissions: Land, Water, Air

Antimony is also lost through emissions in the air, in the water, and on the land [177]. Total antimony emissions are difficult to estimate, but it has been shown that in Australia alone, total emissions of antimony (air, water, land) reach as much as 3.8 t/year during mining and 8.7 t/year during metal manufacturing. A better understanding of these losses can help limit the unwanted emissions of antimony and increase the available supply of antimony at the same time.

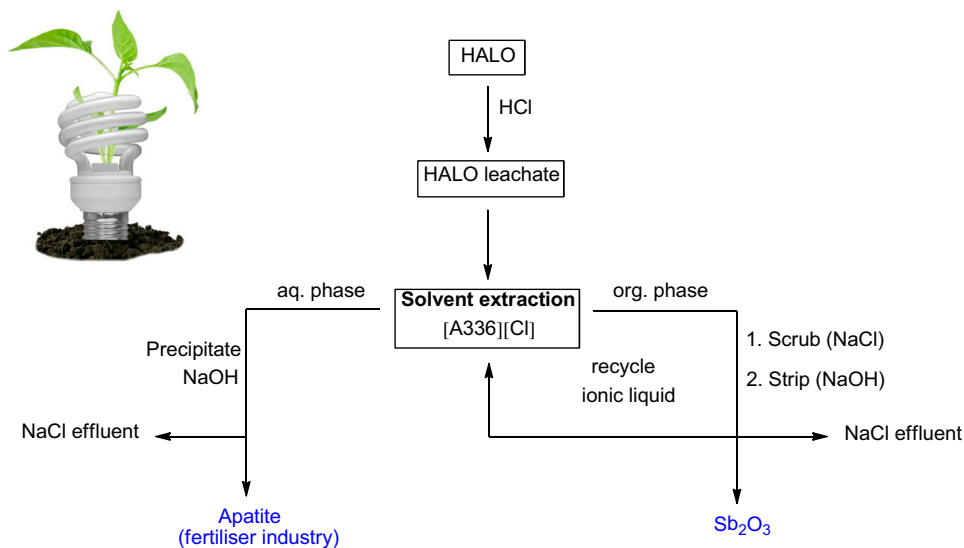
Land emissions are important, especially around antimony mining and processing sites. An overview of antimony emissions in the soil and decontamination procedures has been compiled by Wilson et al. [178] and since this antimony is often very localized and difficult to access, it is not discussed here in detail. Air emissions on the other hand are very relevant to the topic of secondary sources of antimony. We previously discussed how air pollution control residues, flue dusts, and fly ashes in lead, copper, gold, and antimony processing can contain important amounts of antimony. An overview was also given of the different valorization methods and antimony recovery processes for these residues. However, the capture of antimony air emissions still needs to be improved. Tian et al. [177] made a study of global yearly air emissions and calculated that in 2005 global air emissions of antimony reached 2232 t, and then gradually declined to about 1904 t in 2010. Atmospheric emissions are mostly caused by fuel combustion (42 %), waste incineration (24 %), brake wear (17 %), and metal production (17 %) [177]. China is the largest emitter of atmospheric antimony (649 t in 2010) due to the large-scale combustion of coal and the production of metals, as China currently has 114 antimony mines dispersed over 18 provinces [177]. The emission of antimony from coal power plants, waste incinerators, and metal production sites, can be lowered by the implementation of better filters and air pollution control systems, from which the antimony can then be recovered and reused. However, the emissions of antimony through the wear of vehicle brake linings and brake pads has proven to be a larger concern as antimony emissions are not localized and no simple solutions are available.  $\text{Sb}_2\text{S}_3$  is used as a lubricant in disc brake pads and drum brake pads, with concentrations as high as 46 g/kg, depending on the type of brake [179, 180]. Uexküll et al. [180] and Varrica et al. [179] studied the speciation of these brake wear particles and concluded that while the brake linings are mainly composed of  $\text{Sb}_2\text{S}_3$ , this compound is decomposed during the abrasion process leading to a mix of  $\text{Sb}_2\text{O}_3$  and  $\text{Sb}_2\text{O}_5$  in the airborne particulate. Finally, there is the emission of antimony to surface water, which has been reviewed by Filella et al. [181]. These emissions can also be important secondary sources of antimony if proper filtration systems are set in place. In concentrated industrial flows and mine drainage this is simple to implement, but even for very dilute water streams, decontamination is mandatory, so these large volumes can result overall in a significant amount of recovered antimony.

### Antimony Removal from Waste Water

Antimony removal from waste water was recently reviewed by Ungureanu et al. [182]. This extensive review



**Fig. 12** Zero-waste valorization process for the recovery of antimony from the halophosphate fraction of lamp phosphor waste. The phosphor is first dissolved in HCl, followed by the selective extraction of Sb(III) with Aliquat® 336. Pure  $\text{Sb}_2\text{O}_3$  is then obtained after a scrubbing (NaCl) and stripping step (NaOH). The remaining leachate is precipitated as apatite (fertilizer precursor). Reproduced with permission from Ref. [176]. Copyright Royal Society of Chemistry (2016) [176]



gives an overview of the analytic techniques for antimony speciation and the existing removal techniques [182]. Different removal techniques have been proposed such as ion-exchange, adsorption, membrane separation, electrochemical methods, bio-removal methods, or coagulation/flocculation followed by filtration or sedimentation [182]. The very large number of papers written on this topic, clearly demonstrated the importance of antimony removal from waste water [182]. In these studies, the focus is systematically on the removal of antimony to obtain clean water. This field of study is different from the previous chapters in this review, as it deals with very dilute antimony waste streams. However, the large volumes represent a sizeable amount of antimony, which should not go to waste. Antimony removal from wastewater is therefore an opportunity to collect antimony efficiently, using automatic (continuous) technologies such as column filtration with specialized sorbents. These techniques lead to a strong concentration of the antimony, which can then be used as a source of antimony, when renewing or regenerating the sorbent or filter.

The most common treatment techniques for antimony removal are flocculation–coagulation followed by filtration or sedimentation, or the use of adsorbents [182]. Flocculation–coagulation is one of the most employed treatments to produce drinking water, and has been investigated for antimony removal as well, with ferric chloride being the most suitable coagulant [182–185]. The drawback of this technique is that it is difficult to automate, it creates toxic sludge and it is not 100 % effective at removing antimony. Researchers have therefore increasingly turned to adsorbents due to the convenient automatization in a continuous set-up and the generally higher antimony-removal efficiencies [182]. The use of sorbents requires only a periodic

regeneration, which is an opportunity to recover (pure) antimony. A large variety of sorbents have been investigated to this effect, and a full list is not included here since these have already been reviewed by Ungureanu et al. [182]. The investigated materials include chars, activated carbons, graphene, alumina, metal oxides, minerals, resins, biosorbents, and sorbents based on waste materials and residues. The benchmark was set by Xu et al. [186] who achieved the highest adsorption capacity (214 mg/g) using a binary iron-manganese oxide, whose adsorption capacity significantly outperformed the individual oxides [182, 186]. However, it should be stated that this result is an outlier, with most adsorbents having an adsorption capacity around 50–100 mg/g [182]. The second best material was prepared by Biswas et al. [187], who used saponified orange waste, loaded with Zr(IV) and Fe(III), as a sorbent for the adsorptive removal of Sb(III) and Sb(V) from waste water. They reported adsorption capacities around 114–145 mg/g. However, the review by Ungureanu et al. [182] does not yet include the latest results and an overview of some additional recent examples is given here, which boast high adsorption capacities (>100 mg/g). Wang et al. prepared iron-modified aerobic granules to remove Sb(V) from waste water and investigated the adsorption mechanism in detail [188]. They reported a maximal adsorption capacity of 125 mg/g. Dou et al. [189] investigated hierarchical macro- and mesoporous amorphous alumina as a sorbent for Sb(V). The results were compared with five different commercial alumina adsorbents and showed that this material had the highest adsorption capacity (118 mg/g) and fast adsorption kinetics. Dong et al. [190] made nanocomposites from graphene oxide and schwertmannite (iron oxyhydroxysulphate mineral) to adsorb Sb(V) from aqueous solutions. A synergetic effect

on Sb(V) uptake was observed for this composite adsorbent, which had a maximum Sb(V) adsorption capacity of 158.6 mg/g. Sb(V) in spiked tap water (100 µg/L), simulated river water (6400 µg/L), and acid mine drainage (50,000 µg/L) was adsorbed by this sorbent to levels well below the regulation levels for these waters. An interesting subcategory of water purification is the removal of antimony from mine drainage. These flows are often much more concentrated and more acidic and therefore demand different solutions. Zhu et al. used electrocoagulation to partially remove antimony from mine flotation waste water [191]. The remaining concentration was still 1 mg/L, which far exceeds the drinking water limit, but meets the emission standards established by the state department of environmental protection of China [191].

## Conclusion and Outlook

Antimony is one of the most critical elements, but has remained largely out of the spotlights. However, the gap between supply and demand is expected to exceed 10 % in the coming years, making antimony supply far more critical than for example the rare-earth elements. Similar to the rare earths, antimony production is dominated by China and as the grades of primary ores diminishes, it becomes increasingly difficult to meet the demand for antimony. Antimony is an important industrial commodity and is used in flame retardants for plastics and electronics, but also in catalysts for the production of PET polymers and in lead-acid batteries. To meet demand, it is crucial that attention shifts to secondary sources such as industrial process residues and end-of-life consumer items. This review gives an overview of existing waste streams which could be of importance as secondary sources of antimony. It also encourages researchers to look at unusual sources such as air pollution control residues, low-grade ores, and unusual products such as lamp phosphor waste. Sufficient technologies are available to make antimony recycling a success and current recycling rates of approximately 20 % are expected to rise as China struggles to increase its output. However, one of the remaining obstacles for the implementation of antimony recycling in the industry is the upscaling from laboratory methods to industrial processes. Further work needs to be carried out on pilot scale to see which methods are sufficiently robust and flexible. Another challenge is the changing composition of various residues depending on the ore or product that is being processed. Changes in composition of the input material can greatly affect the performance of the recovery processes. Finally, economic feasibility studies and life cycle assessments need to be carried out to determine which techniques are the most promising for industrial upscaling. We expect

copper producers to invest in antimony recovery techniques in coming years as this is a growing problem that needs to be addressed in the copper production flowsheets. Other very interesting sources include incineration ashes from concentrated fractions such as textiles or flame retardant plastics. Recycling of antimony through the recycling of lead-acid batteries is currently carried on industrial scale and should continue to provide a steady stream of secondary antimony. Current primary antimony production is unsustainable as current reserves account for only 10–11 years of production at current levels which could be aggravated by rising demand from existing and future antimony applications such as liquid–metal batteries and thermoelectric materials. Moreover, antimony is also associated with environmental problems, so a better management of antimony resources is mandatory and the antimony life cycle must become more circular in the coming years.

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## Compliance with Ethical Standards

**Conflict of interest** The authors declare no competing financial interest.

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