

# GEMAS: Geochemical background and mineral potential of emerging tech-critical elements in Europe revealed from low-sampling density geochemical mapping

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1	GEMAS: Geochemical background and mineral potential of
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3	sampling density geochemical mapping
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## 32 Abstract

33

The demand for 'high-tech' element resources (e.g., rare earth elements, lithium, platinum group elements) has increased with their continued consumption in developed countries and the emergence of developing economies. To provide a sound knowledge base for future generations, it is necessary to identify the spatial distribution of critical elements at a broadscale, and to delineate areas for follow-up surveys. Subsequently, this knowledge can be used to study possible environmental consequences of the increased use of these resources.

40 In this paper, three critical industrial elements (Sb, W, Li) from low-sampling density 41 geochemical mapping at the continental-scale are presented. The geochemical distribution and 42 spatial patterns have been obtained from agricultural soil samples (Ap-horizon, 0-20 cm; N=2108 samples) collected at a density of 1 site per 2500 km<sup>2</sup> and analysed by ICP-MS after a 43 44 hot aqua regia digestion as part of the GEMAS (GEochemical Mapping of Agricultural and 45 grazing land Soil) soil-mapping project in 33 European countries. Most of the geochemical 46 maps show exclusively natural background element concentrations with minor, or without, 47 anthropogenic influence. The maximum extent of the last glaciation is marked as a discrete 48 element concentration break, and a distinct difference occurs in element concentration levels 49 between the soil of northern and southern Europe, most likely an effect of soil genesis, age and 50 weathering. The Sb, W and Li concentrations in soil provide a general overview of element 51 spatial distribution in relation to complexity of the underlying bedrock and element mobility in 52 the surface environment at the continental-scale. The chemical composition of agricultural soil 53 represents largely the primary mineralogy of the source bedrock, the effects of pre- and post-54 depositional chemical weathering, formation of secondary products, such as clays, and element 55 mobility, either by leaching or mineral sorting. Observed geochemical patterns of Li, W and Sb 56 can be often linked with known mineralisation as recorded in the ProMine Mineral Database, 57 where elements in question occur either as main or secondary resources. Anthropogenic impact 58 has only been identified locally, predominantly in the vicinity of large urban agglomerations. 59 Unexplained high element concentrations may potentially indicate new sources for high-tech 60 elements and should be investigated at a more detailed scale.

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Keywords: agricultural soil, weathering, geochemistry, critical elements, antimony, tungsten,
 lithium

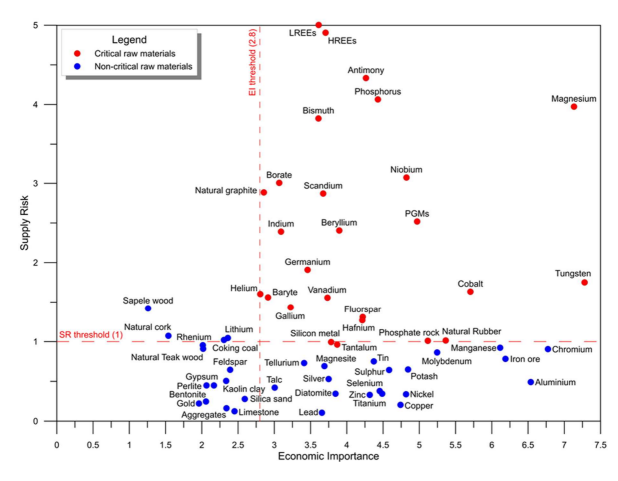
## 67 **1. Introduction**

68 The demands for a variety of mineral and/or element resources (rare earth elements, platinum 69 group elements, cobalt, beryllium, lithium, iodine, etc.) are strongly increasing (Buchert et al., 70 2009; Bertrand et al., 2016; Meinert et al., 2016). These elements are essential for maintaining 71 and improving the present and future quality of life, including many high-technology yet low-72 carbon industries. Two factors have been used by the National Research Council (NRC) to rank 73 criticality: (a) the degree to which a commodity is essential and (b) the risk of supply disruption 74 for the commodity (Ali et al., 2017). Further, the demand for energy-related minerals has 75 increased, as global energy production diversifies beyond carbon and nuclear-based sources 76 (Teske, 2019). Many of these elements are crucial to a variety of manufacturing, high-tech and 77 military applications like photovoltaic solar cells (Ga, Ge, In, Se, Ag, Te), high-strength 78 permanent magnets and magnetic and/or optical components for lighting (mostly rare earth 79 elements and Co), for wind turbines and hybrid automobiles, high performance batteries (Li, 80 La), catalysts (Pt, Pd, Ce), advanced turbines (Re), and advanced energy reactor designs (He). 81 A critical mineral and/or element is both essential in use, subject to the risk of supply restriction 82 and the induction of environmental impacts. Based on information about occurrences, resources 83 and reserves, extraction, processing, utilisation, and recycling, good candidates for the 84 designation of 'energy-critical' elements can be determined. The list of critical raw materials 85 (Table 1) is updated by the European Union Commission every few years with the last update 86 in 2017 (European Commission, 2017).

The overall results of the 2017 criticality assessment are shown in Fig. 1 where the critical raw materials are denoted by the red dots (European Commission, 2017). Of the 61 candidate raw materials assessed (58 individual and 3 grouped materials, biotic and abiotic), the following 26 raw materials and groups of raw materials were identified as critical (Table 1). It is noted that Li discussed here is not yet a critical raw material, according to the 2017 European Commission classification, but its criticality is being discussed in view of its
importance as a critical element for future sustainable technologies according to UNEP (2009).
Lithium is an essential, in fact a critical, element in the battery industry and in technical
evolution towards low-carbon society.

97 Table 1. List of Critical Raw Materials, updated in 2017 by the European Commission
98 (2017).

2017 Critical Raw Materials					
Antimony	Cobalt	Helium	Natural graphite	Phosphorus	
Baryte	Fluorspar	HREEs	Natural Rubber	Scandium	
Beryllium	Gallium	Indium	Niobium	Silicon metal	
Bismuth	Germanium	LREEs	PGMs	Tantalum	
Borate	Hafnium	Magnesium	Phosphate rock	Tungsten	
				Vanadium	



103 Figure 1. Criticality assessment of economically important raw materials for the European

104 Union in 2017 (redrawn from European Commission, 2017, Fig. 6, p.39).

106 Geoscientists have a prominent role in the exploration for, management of, and 107 environmentally safe handling of critical mineral and/or element resources considering that 108 these needs will become even more important as the world's population and standards of living 109 continue to increase (Nickless et al., 2015; Ali et al., 2017). It is worth noting that the concept 110 of criticality is variable (and to large degree political) and depends principally on the context. 111 What is critical for a state, a manufacturer or a product may not be critical for another, and what is critical for national defence may be different when compared to what is necessary to make 112 113 an electronic apparatus cheaper. Therefore, prioritisation of critical minerals and/or elements 114 are crucial for modern societies and has important implications for our economic prosperity 115 (Bringezu et al., 2016).

116 To provide a solid resource base for the future, but also to timely identify any possible 117 new environmental impact of the increased use of these elements, it is necessary to identify 118 their natural spatial distribution with a wider view, from the order of a country or a continental 119 block, e.g., European Union (Lambert et al., 2013). Any significant change in their cycle at the 120 Earth's surface, due to their increased use in a variety of technological applications, needs 121 robust preliminary knowledge. The growing consumption of critical elements generates the 122 demand for new production, both from primary (ore deposits) and secondary (mineral-based 123 and non-mineral-based wastes) sources. At some point, they may become new contaminants 124 and their identification in various environmental compartments is an urgent issue for future 125 pollution and waste management.

The chemical elements, which are now key components for the development of new technologies, are according to the present understanding of their concentrations, transformation and transport in the different environmental compartments, poorly studied (Reimann et al., 2010). This paper presents the GEMAS (Geochemical Mapping of Agricultural and grazing land Soil) results for Li, W and Sb, a project carried out by the Geochemistry Expert Group of

131 EuroGeoSurveys in cooperation with Eurometaux (European Association of Metals). The 132 GEMAS project aims were to detect and to map the element variation in productive soil at the 133 European scale. Soil samples were collected from 33 European countries, covering an area of 134 5.6 million km<sup>2</sup> (Reimann et al., 2012a; 2014a, b). Reimann et al. (2016) argue for low sampling 135 densities (1 site/100 to 1 site/18 000 km<sup>2</sup>) rather than the costly and time-consuming very high 136 sampling densities (100s to 1000s of samples/km<sup>2</sup>) employed for geochemical exploration, 137 giving the same conclusions as those obtained in previous studies of geochemical mapping in 138 continental- and regional-scale surveys (Smith and Reimann, 2008; Garrett et al., 2008; 139 Cicchella et al., 2013; Birke et al., 2015). Reimann et al. (2007a, 2016) stated that this low-140 sampling density geochemical survey approach has also proved efficient in the early stages of 141 mineral exploration programmes through identification of more suitable regions for detailed 142 mineral exploration.

143 As part of GEMAS valorisation, a series of peer-reviewed publications, where detailed 144 interpretation of the continental-scale distribution of single elements or related groups of 145 elements, have been published, e.g., Reimann et al. (2012b - Pb); Ottesen et al. (2013 - Hg); 146 Tarvainen et al. (2013 - As); Scheib (2012 - Nb); Sadeghi et al. (2013 - Ce, La, Y); Birke et al. 147 (2014a, 2016, 2017 – Cd); Poňavič and Scheib (2014 – Se); Cichella et al. (2014 – U, Th, K); 148 Ladenberger et al. (2015 - In); Albanese et al. (2015 - Cr, Ni, Co, Cu); Négrel et al. (2016 - Ge; 149 2018a – U, Th; 2018b - Rb, Ga, Cs); Jordan et al. (2018 – Ni). Among these publications, the 150 elements In, Nb, Ge and the rare earth elements, although defined as technology-critical 151 elements, have been interpreted in the frame of weathering patterns related to climate and 152 inheritance from processes controlling element binding to mineral phases.

Even if we are not facing an imminent absolute shortage of energy critical elements (e.g., see the Hubbert's Peak scenario in Hubbert, 1982), market-driven shortages are possible, and no country can become independent from energy critical elements, which are nowadays

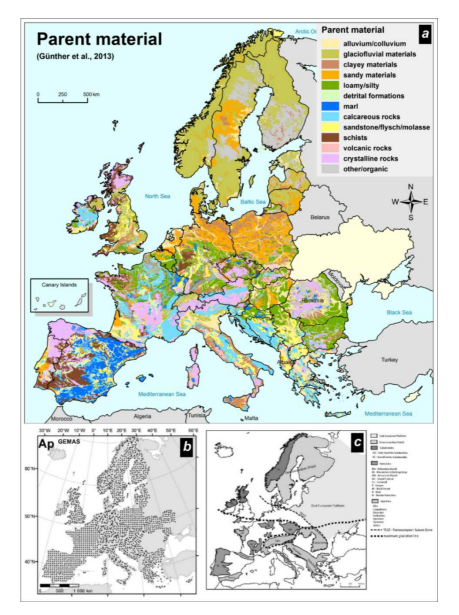
156 produced often in politically unstable countries. Therefore, securing energy critical elements 157 requires an advanced research plan including geological, metallurgical and materials science, 158 the impact of which on knowledge and, therefore, the capacity to exploit resources is crucial 159 for future development (Nickless et al., 2015; Ali et al., 2017). In this paper, we investigate the 160 geochemical distribution and behaviour of three critical elements Sb, W and Li using low-161 sampling density geochemical mapping at the continental-scale and interpret the geochemical 162 patterns in relation to their mineral potential. We need to point out the difference between a 163 single mineralisation event, which cannot be located by using the GEMAS density of 1 sample/2500 km<sup>2</sup> and a mineralising process which affects a much larger area, a mineral 164 165 province that can be identified with this data set (Bölviken et al., 1990). The GEMAS data set 166 make it possible to define, therefore, potential areas, where follow-up and detailed 167 geochemical exploration survey can be carried out in order to better define the mineralised 168 zones (Reimann et al., 2007b).

## 169 **2. Materials and methods**

170 The GEMAS project (Reimann et al., 2014a, b) was carried out by the Geochemistry Expert 171 Group of EuroGeoSurveys in cooperation with Eurometaux (European Association of Metals) 172 and managed by the Geological Survey of Norway (NGU). It was the second large geochemical 173 mapping survey at the European continental-scale following the FOREGS project (Salminen, 174 Tarvainen et al., 1998; Salminen et al., 2005; De Vos, Tarvainen et al., 2006). Agricultural and 175 grazing land soil samples were collected in 33 European countries, covering an area of about 5.6 million km<sup>2</sup> (Reimann et al., 2012a; 2014a, b). The survey area is shown in Figure 2. The 176 177 main objective of the project was to detect and to map the element variation in agricultural and 178 grazing land soil and to present their geochemical baseline variation for Europe. The soil 179 samples were not taken at known contaminated sites, in the immediate vicinity of industry or 180 power plants, cities, railway lines or major roads, directly below high power electric lines or

near to pylons and wooden fences. The GEMAS aqua regia and XRF results are provided in a
two-volume geochemical atlas (Reimann et al., 2014a, b).

183 For the GEMAS project two types of soil samples from two land use categories have been collected at an average density of 1 site per 2500 km<sup>2</sup> each (Fig. 2b). Grazing land soil 184 (Gr; N=2024 samples) has been defined as 'land under permanent grass cover' and a sample 185 186 depth of 0-10 cm was used, according to the REACH regulation (Registration, Evaluation, 187 Authorisation Restriction of chemicals) guidance document requirements and 188 (https://echa.europa.eu/). Agricultural soil (Ap; N=2108 samples) was collected from the 189 ploughing layer of an agricultural arable field from a depth of 0-20 cm. Each sample (ca 3.5 190 kg) corresponds to a composite of five sub-samples taken from the corners and centre of a 191 10×10 m square. Field duplicate samples were collected from both land use categories at a rate 192 of 1 in 20 routine samples.



195 Figure 2. Modified from Négrel et al. (2015, Fig. 1, p.3). (a) Map of parent materials in Europe 196 showing the distribution of various lithologies across the continent (modified from Gunther et 197 al., 2013 and adapted from Négrel et al., 2015, Fig. 1, p.3). (b) Sample locations (dots) of the 198 ploughed agricultural soil (Ap-samples; N=2108). Map projection: Lambert Azimuthal Equal 199 Area (ETRS\_1989\_LAEA), with central meridian at 10°. (c) Generalised geological map of 200 Europe with major lithotectonic units, Variscan and Alpine belts, Transeuropean Suture Zone 201 (TESZ) and the extension of maximum glaciation (modified from Reimann et al., 2012b, Fig. 1, 202 *p.533*).

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A single facility (State Geological Institute of Dionyz Stur, Slovakia) prepared all samples for analysis. Soil samples were air-dried, sieved to <2 mm using a nylon screen, homogenised and finally split into 10 sub-samples. Two reference materials, e.g., standards, prepared in the same laboratory, were inserted at a rate of 1 in 20 samples together with field and analytical replicates to monitor analytical performance during the project. Samples were
analysed by total X-ray fluorescence (XRF) and partial extraction methods like aqua regia (AR)
and mobile metal ion (MMI<sup>®</sup>) (Reimann et al., 2012a; 2014a, b), and here the results of the AR
method will be discussed.

The aqua regia extraction applied to the soil samples prior to analysis used a 15 g aliquot of the unmilled <2 mm fraction, which was leached in 90 ml of aqua regia (95°C, 1 hour) and then made up to a final volume of 300 ml with 5% HCl. The solutions were analysed using a combination of inductively coupled plasma-atomic emission spectrometry (ICP-AES) and inductively coupled plasma-mass spectrometry (ICP-MS) at ACME laboratories in Vancouver, Canada (now Bureau Veritas Mineral Laboratories). A total of 53 elements was determined on the over 5000 soil samples using matrix matched standards and reference materials.

219 A rigorous quality control (QC) procedure was part of the analytical protocol (Reimann 220 et al., 2009; Demetriades et al., 2014). Samples were analysed in batches consisting of 20 221 samples (Reimann et al., 2009). In each batch, one field duplicate, one analytical replicate of 222 the field duplicate and a project standard were inserted. The practical detection limit (PDL) was 223 estimated from results of the project replicate samples by calculating regression line coefficients 224 with the 'reduced major axis line' procedure (Demetriades, 2011). This PDL in the aqua regia 225 extraction by ICP-MS is 0.003 mg/kg for Sb, 0.01 mg/kg for W and 0.04 mg/kg for Li. Details 226 on analytical procedures and quality control are provided in Reimann et al. (2009; 2011; 2012c), 227 Birke et al. (2014b) and Demetriades et al. (2014).

Geochemical data are compositional data, element concentrations reported in wt % or mg/kg sum up to a constant and are thus not free to vary (Reimann et al., 2012d). Compositional data do not plot in the Euclidean space, but rather on the Aitchison simplex (Aitchison, 1986; Buccianti et al., 2006; Pawlowsky-Glahn and Buccianti, 2011). Thus, only order statistics should be used in statistical data processing. The colour surface maps were produced by kriging, based on a careful variogram analysis (Filzmoser et al., 2014). Kriging was used to interpolate
values from the irregularly distributed sampling sites to a regular grid and into unsampled space.
Class boundaries for the colour surface maps are based on percentiles (5, 25, 50, 75, 90 and 95).
Geochemical maps displayed hereafter (Figs. 4-6) were produced in ArcGIS.

### 237 **3. Results and discussion**

#### 238 **3.1. Element concentrations in agricultural soil**

The concentrations of Sb, W and Li generated by the GEMAS project constitute a homogeneous
data set and can be compared, but are not compatible, with other continental data sets, e.g., from
Europe (FOREGS; Salminen et al., 2005; De Vos, Tarvainen et al., 2006), U.S.A. (Smith et al.,
2013, 2014), Australia (Caritat and Cooper (2011a, b, c, d) and China (Xie et al., 2012) as
tabulated in Table 2.

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#### 245 3.1.1. Sb concentrations in agricultural soil

246 The median for Sb in the Ap and Gr soil samples in Europe in the hot aqua regia 247 extraction are very close, 0.234 and 0.275 mg/kg, respectively. The combined plot histogram -248 density trace one - dimensional scattergram - boxplot shows the univariate Sb data distribution 249 (Fig. 3a for Ap, and 3b for Gr samples). The existence of outliers in the Sb distribution is shown 250 by one-dimensional scattergram and boxplot. The density trace displays a smooth multimodal 251 distribution, but is overall symmetrical about the median, a feature also indicated by the very 252 symmetrical boxplot (log-scale). Compared to the upper continental crust (UCC) estimated Sb 253 abundance of 0.4 mg/kg (Hu and Gao, 2008), the aqua regia extractable Sb (median Ap 0.234 254 and Gr 0.275 mg/kg) is lower by a factor of 1.74 in the Ap samples, and 1.45 in the Gr soil 255 samples. Due to high detection limits of GEMAS XRF results, it is not possible to directly relate 256 GEMAS total concentrations to the UCC values. However, estimated extractability of Sb (from

- the comparison of aqua regia and XRF results) is 9 and 13% for the Ap and Gr samples,
- 258 respectively (Reimann et al., 2011).

Table 2. Median values of Sb, W and Li (in mg/kg) from continental-scale geochemical surveys.

D	Analytical	Sb median	W median	Li median	9
Provenance	information	(Sb range)	(W range)	(Li range)	Source
World soil	<2 mm, total, estimate	1 (<0.2 – 10)	1.5 (0.5 - 83)	25 (3 - 350)	Bowen (1979)
FOREGS <sup>1</sup> -Atlas, EU, topsoil (0 - 25	<2 mm, total (ICP-MS <sup>2</sup> ,	0.6	<5.0 <sup>3</sup>	20.8	Salminen et al.
cm)	N=845)	(<0.02 – 31.1)	$(<5.0-14)^3$	$(0.28-271)^4$	(2005)
GEMAS <sup>5</sup> , EU, agricultural soil (0	<2 mm, AR <sup>4</sup> (ICP-MS <sup>2</sup> , N =	0.23	0.073	11.4	Reimann et
-20 cm)	2108)	(<0.02 – 17)	(<0.01 – 5.2)	(0.161 – 136)	al. (2014a)
GEMAS, EU, agricultural soil (0	<2 mm, total XRF	<5	<5	n.d.	Reimann et
– 20 cm)	(N = 2108)	(<0.5 – 36)	(5 – 25)		al. (2014a)
GEMAS <sup>5</sup> , EU, grazing land soil (0	$<2 \text{ mm}, \text{AR}^4$ (ICP-MS <sup>2</sup> , N =	0.28	0.077	11.3	Reimann et al. (2014a)
– 10 cm)	2024)	(<0.02 - 25)	(<0.01 – 19)	(<0.1 – 153)	
GEMAS, EU, grazing land soil (0	<2 mm, total XRF (N = 2024)	<0.5	<5	n.d.	Reimann et al. (2014a)
– 10 cm)		(<0.5 – 33)	(<5 – 37)		
USA, surface soil (0 - 5 cm)	<2 mm, total (ICP-MS <sup>2</sup> , ICP-	0.57	0.8	20	Smith et al. (2014)
(0 - 5 cm)	AES, N = 4857)	(<0.05 - 482)	(<0.1 – 1150)	(<1-300)	(2014)
USA, soil A-	<2 mm, total (ICP-MS <sup>2</sup> , ICP-	0.57	0.8	20	Smith et al.
horizon	$AES^{6}, N = 4857)$	(<0.05 - 630)	(<0.1 – 299)	(<1-514)	(2014)
Australia, NGSA <sup>7</sup> , TOS <sup>8</sup> (0 - 10 cm)	<2 mm, AR <sup>4</sup> (ICP-MS <sup>2</sup> , N =	0.12	N/A <sup>9</sup>	N/A <sup>9</sup>	Caritat and Cooper
103 (0 - 10 cm)	1190)	(<0.02 – 18.1)	( <0.1 – 174)		(2011a, b)
Australia, NGSA <sup>7</sup> , $TOS^8 (0, -10 \text{ am})$	<0.075 mm, AR <sup>4</sup> (ICP-MS <sup>2</sup> , N =	0.15	N/A <sup>9</sup>	N/A <sup>9</sup>	Caritat and Cooper
$TOS^{8} (0 - 10 cm)$	1177)	(<0.02 – 27.1)	(<0.1 ->200)		(2011a, b)
Australia, NGSA <sup>7</sup> ,	<0.075 mm,	N/A <sup>9</sup>	1.6		Caritat and
TOS <sup>13</sup> (0 - 10 cm)	HF+HNO <sub>3</sub> (ICP- MS <sup>2</sup> , N = 1187)	(<0.4 – 28.9)	(<0.1 - 643)	N/A <sup>9</sup>	Cooper (2011a, b)

Australia, NGSA <sup>7</sup> , TOS <sup>13</sup> (0 - 10 cm)	<2mm, HF+HNO <sub>3</sub> (ICP-	N/A <sup>9</sup>	1	N/A <sup>9</sup>	Caritat and Cooper
105 (0 - 10 cm)	MS <sup>2</sup> , N = 1190)	(<0.4 – 14.5)	(<0.1 – 327)		(2011a, b)
China, RGNRP <sup>10</sup> , silt or fine sand on the drainage bottom, bank soil or dry stream sediment	<0.22 mm, total, (N = 1,568,528)	0.6	1.71	31.1	Xie et al. (2012)
		(<0.03 – 911) <sup>11</sup>	(<0.04 - 1073) <sup>12</sup>	(<0.4 - 560)	
BSS <sup>14</sup> , agricultural soil (0 - 25 cm)	<2 mm, HF (ICP- MS <sup>2</sup> , N = 747)	0.24	N/A <sup>9</sup>	N/A <sup>9</sup>	Reimann et al. (2003)
	(<0.1 – 3.2)				× /

 <sup>1</sup> FOREGS - Forum of European Geological Surveys; <sup>2</sup> ICP-MS - Inductively Coupled Plasma Mass Spectrometry;
 <sup>3</sup> XRF - X-ray Fluorescence Spectrometry; <sup>4</sup> Stream sediment; <sup>5</sup> GEMAS - Geochemical Mapping of Agricultural and Grazing Land Soil in Europe; <sup>6</sup> ICP-AES - Inductively Coupled Plasma Atomic Emission Spectrometry; <sup>7</sup> NGSA – National Geochemical Survey of Australia; <sup>8</sup> TOS - top outlet sediment; <sup>9</sup> N/A – not available/applicable;
 <sup>10</sup> RGNRP – Regional Geochemistry National Reconnaissance Programme; <sup>11</sup> HG-AFS – Hydride Generation-Non-Dispersions Atomic Fluorescence Spectrometry; <sup>12</sup> POL – Polarography; <sup>13</sup> GF-AAS - Graphite Furnace Atomic Absorption Spectrometry; <sup>14</sup> BSS - Baltic Soil Survey: Agricultural Soils in Northern Europe. A Geochemical Atlas.

270 An explanation for the difference between UCC (total concentrations) and hot aqua regia 271 results is that a major quantity of Sb occurs in minerals and phases which are not easily 272 dissolved by acids. Sulphides, major host minerals for Sb, may not be very common in average 273 soil. On the other hand, Sb can sorb onto Fe, Mn and Al oxides and hydroxides, clay minerals, 274 and organic matter (Belzile et al., 2001; Buschmann and Sigg, 2004; Johnson et al., 2005), 275 especially, when it occurs as oxyanions in low pH environments (Takahashi et al., 2010; 276 Hockmann and Schulin, 2013). Some of the Sb can also be volatilised and lost during the hot 277 acid extraction (Reimann et al., 2010).

The median Sb values and their concentration ranges in soil samples of different continental-scale surveys are provided in Table 2. The median Sb values vary between 0.12 mg/kg in Australia and 0.6 mg/kg in the European FOREGS topsoil. The Sb 'world soil' median of 1 mg/kg (with a range from <0.2 to 100 mg/kg), as provided by Bowen (1979), is about four times higher than the GEMAS hot aqua regia extractable Sb median. Kabata-Pendias and 283 Mukherjee (2007) provide better-constrained Sb concentrations with the background level of 284 Sb in topsoil samples between 0.05 and 4.0 mg/kg. Recently, the Sb data set of the LUCAS Soil 285 Survey (AR; ICP-AES; N=23,000; Cristache et al., 2014; ECS, 2010) published by Tóth et al. (2016) reports values from 0.01 to 10.9 mg/kg with a calculated mean of 0.25 mg/kg. The 286 287 GEMAS median values for agricultural and grazing land soil samples (Reimann et al., 2014a, 288 b), based on 2108 samples, are thus comparable to those reported by the LUCAS project based 289 on over 23,000 samples. An advantage of the GEMAS results is that they are based on two 290 separately analysed and statistically treated land-use sample types (Ap and Gr) evenly covering 291 the whole survey area, while in the LUCAS project various land-use materials were analysed 292 and processed together. None of these different land-use classes provided an even coverage of 293 the European continent.

294 Some European regional geochemical surveys also provide Sb background data. In the 295 Baltic Soil Survey (Reimann et al., 2003), the median Sb concentration (HF extraction) is 0.24 296 mg/kg in TOP (0-25 cm) soil, which is nearly identical with the GEMAS median and its 297 concentration ranges between <0.1 and 3.2 mg/kg, while in the BOTtom (50-75 cm) soil it is 298 lower, 0.19 mg/kg (range: <0.1 - 1.41 mg/kg). The total Sb concentrations in topsoil samples 299 (0-15 cm) of England and Wales (Rawlins et al., 2012) range from 0.1 to 55 mg/kg, with a 300 median value of 0.7 mg/kg. In Irish topsoil (0–10 cm), the median value of Sb is 0.53 mg/kg 301 and the total concentrations vary from <0.05 to 5.29 mg/kg (Fay et al., 2007).

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- 303

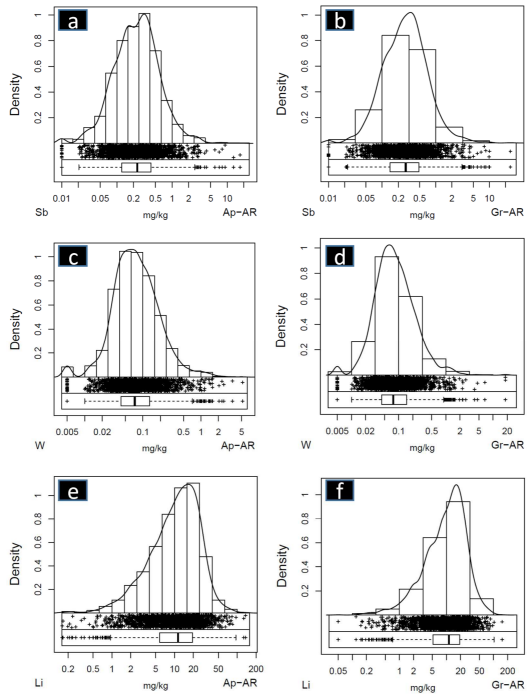


Figure 3. Combined plot of histogram, density trace, one-dimensional scattergram and boxplot
distribution in European Ap and Gr samples following a hot aqua regia extraction for the three
elements Sb (a for Ap and b for Gr samples), W (c for Ap and d for Gr samples), Li (e for Ap
and f for Gr samples).

304

310 3.1.2. W concentrations in agricultural soil

The median concentration of extractable W (aqua regia extraction) in the GEMAS European soil is 0.073 mg/kg in the Ap and 0.077 mg/kg in the Gr samples. The combined plot 313 histogram - density trace one- dimensional scattergram - boxplot shows the univariate data 314 distribution of the aqua regia extractable W (Fig. 3c for Ap and 3d for Gr samples). The one-315 dimensional scattergram and boxplot display the existence of upper and lower outliers in the W 316 distribution. The density trace, histogram and one-dimensional scattergram highlight the 317 samples below detection. The main body of the density trace and histogram exhibit an almost 318 symmetrical data distribution in the log-scale.

319 Because W does not substitute into most rock-forming minerals, its distribution in soil 320 is related largely to naturally occurring W mineralisation. Tungsten is enriched in mica (5-50 321 mg/kg), and muscovite in altered granite near W deposits may contain up to 500 mg/kg 322 (Wedepohl, 1978). A comparison of the GEMAS Ap soil W median to the upper continental 323 crust (UCC) estimated abundance of 1.9 mg/kg (Hu and Gao, 2008) shows that the hot aqua 324 regia extractable W is a factor of 27 lower in the European agricultural soil. Clearly, only a very 325 small percentage of total W in soil can be extracted by the mixture of HCl and HNO<sub>3</sub>, even after 326 heating. XRF results for W in the GEMAS project vary around the method's detection limit, 327 but the interpretation of the data range and its maximum values indicates that the data set is 328 quite homogeneous, without many outliers. The extractability of W from the comparison of 329 aqua regia to XRF results is 3 and 4% for the Ap and Gr samples, respectively (Reimann et al., 330 2011). The AR extraction is, therefore, not a sufficiently good method for extracting W from 331 soil samples, mainly because of the presence of resistant W-bearing minerals and precipitation 332 of insoluble tungstates and the polymerisation of tungstates onto soil particles (Bednar et al., 333 2010).

Apart from the GEMAS project XRF results for W (Ap samples: <5 to 25 mg/kg, with a median of <5 mg/kg; Gr samples: <5 to 37 mg/kg, with a median of <0.5 mg/kg), total W concentrations, have been reported in three continental-scale soil geochemical projects, i.e., the Geochemical Atlas of Europe (FOREGS; Salminen et al., 2005), the Geochemical and Mineralogical Atlas of the Conterminous United States; Smith et al., 2014), and the Geochemical Atlas of Australia (Caritat and Cooper, 2011a,b), all ranges and medians are tabulated in Table 1. In the United States of America, the areas of very high total W values in the Central and Southern Rocky Mountains are related to the Colorado Mineral Belt and the Butte Mineral district in Montana (Smith et al., 2014). Mineral deposits enriched in W are widespread throughout the western United States and are most often genetically related to intrusions of felsic rocks.

345 In Europe, total W contents in the topsoil of England and Wales vary at the regional 346 scale from <0.6 to 70 mg/kg, with a median of 2.1 mg/kg (Rawlins et al., 2012). The anomalous 347 W concentrations in southern Devon and Cornwall are associated with mining and extraction 348 processes and occur around the granite intrusions of Dartmoor, Bodmin Moor and St. Austell. 349 In the Soil Geochemical Atlas of Ireland, W concentrations range between <0.1 and 7.72 mg/kg, 350 with a median of 0.59 mg/kg (Fay et al., 2007). High and elevated levels of total W are found 351 in soil developed on igneous rocks as well as on greywacke, black shale and fine-grained 352 sandstone (Fay et al., 2007).

353

#### 354 3.1.3. Li concentrations in agricultural soil

355 The median concentration of Li in an aqua regia extraction in the GEMAS European 356 soil is 11.4 mg/kg in Ap and 11.3 mg/kg in Gr samples. The combined plot histogram - density 357 trace one - dimensional scattergram - boxplot displays the Li univariate data distribution in Ap 358 and Gr soil samples (Fig. 3e for Ap and 3f for Gr samples). One-dimensional scattergram and 359 boxplot highlight the existence of many outliers at the lower end of the Li distribution and very 360 few at the upper end. Though the density trace, histogram and boxplot suggest a slight skew, 361 the data distribution is still rather symmetrical in the log-scale, for both Ap and Gr types of soil. 362 During weathering processes, water-rock interaction and erosion, the geochemical behaviour of Li is similar to that of Na, despite the difference in the abundance of both elements. However, Li is more mobile in the surficial environment and during water-rock interactions. Owing to its weaker binding than Na in ion exchange reactions, Li is preferentially leached during weathering of silicate rocks. Lithium is very mobile during hypergenetic processes, as well as in the initial stage of soil formation (Kabata-Pendias and Mukherjee, 2007). It may become more stable due to its adsorption to clay minerals, Fe and Mn hydroxides, and organic matter (Millot et al., 2010).

The comparison of Li to the estimated upper continental crust (UCC ) abundance of 24 mg/kg (Hu and Gao, 2008) shows that the hot aqua regia extractable Li is almost a factor of 2 lower in the European agricultural soil samples. Large differences in the median values of Li in GEMAS and in continental-scale geochemical soil surveys (Table 2) from China, U.S.A. and Australia indicate that natural factors such as underlying geology, and technical aspects like number of samples, sample size and methodology play a crucial role in data processing and interpretation.

377

#### 378 **3.2.** Spatial distribution of Sb, W and Li in European soil and their source geology

379 As very few differences are observed for the three elements between Ap (ploughing layer 380 of agricultural arable fields) and Gr (land under permanent grass cover) types of soil only the 381 data obtained for the Ap soil samples will be considered hereafter.

At first glance, geochemical maps for Sb, W and Li (Figs. 4b, 5b and 6b) reflect natural sources of these elements and their spatial distribution seems to be governed by weathering type and degree, Quaternary history, the underlying bedrock and mineralisation. The chemical composition of soil represents largely the primary mineralogy of the source parent materials, the effects of pre- and post-depositional chemical weathering, formation of secondary products such as clays, and element mobility, either by leaching or mineral sorting. Geographical distribution (based on ProMine Mineral Database, Cassard et al., 2012, 2015) of the mineralised
areas are shown in Figures 4a, 5a and 6a for Sb, W and Li, respectively, and the maps of Sb, W
and Li in the hot aqua regia extraction of agricultural soil samples are correspondingly displayed
in Figures 4b, 5b and 6b, respectively.

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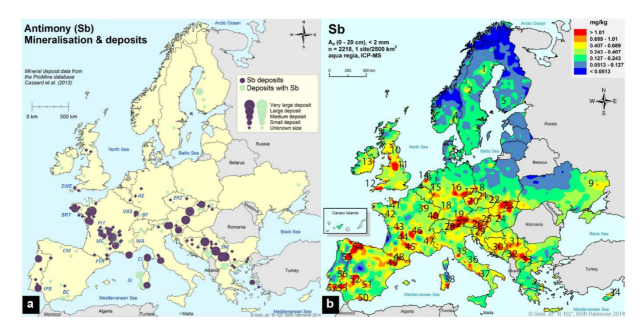
393 3.2.1. Spatial distribution of Sb

394 The Ap map of Sb in the hot aqua regia extraction show one of the most striking 395 differences in element concentration between the Ap soil of all analysed elements/parameters 396 (Fig. 4a) between northern (median 0.11 mg/kg) and southern Europe (median 0.35 mg/kg). It 397 is worth noting the difference in Sb concentrations between the upper continental crust and 398 world soil, 0.75 and 0.9 mg/kg, respectively, compared to values in Ap soil. This pronounced 399 disparity of Sb concentrations is most likely an effect of glaciation, soil age and weathering. 400 Soil pH is generally lower in northern than in southern Europe facilitating thus the removal and 401 accumulation of many elements including Sb from soil. Low pH and/or high content of organic 402 matter are two main factors controlling the fate of Sb in soil. Interestingly, Ap soil samples 403 collected in the far north over Archaean bedrock in Fennoscandia are characterised by the 404 lowest Sb contents in Europe, a feature also known from various local studies and national 405 geochemical surveys, e.g., Andersson et al. (2014). It seems, that soil developed on old cratonic 406 areas, such as the Fennoscandian Shield and East European Platform with the oldest bedrock in 407 Europe, are highly depleted in Sb in comparison with soil formed from parent materials 408 composed of younger bedrock in western and southern Europe. Even though Sb mineralisation 409 is usually located within granitic bodies, the pristine granitic massifs at several locations in 410 Europe show very low Sb contents in overlying soil, e.g., in NE Sardinia, northern Portugal and 411 western-central Spain (Fig. 4b). The enrichment in Sb occurs in soil developed on sedimentary 412 rocks such as shale, mudstone and young argillaceous rocks. Soil within the Carpathian Belt,

where flysch-type rocks dominate, has elevated Sb contents. More rarely, Sb can substitute Fe
and occurs as trace element in minerals typical of mafic rocks (e.g., in olivine) and, therefore,
certain Sb enrichment may occur in areas where mafic to intermediate volcanic rocks occur,
e.g., in Italy, France (Massif Central) and Czech Republic (part of the Bohemian Massif) exist.
A couple of Sb anomalies in Italy are associated with the Roman and Neapolitan alkaline
volcanic provinces (Nos. 36 & 37, Fig. 4b).

In the hypergene zone, Sb is often associated with Fe and Al oxides/hydroxides, and can be sorbed onto silicate grains, representing a potentially bioavailable and extractable fraction (Gal et al., 2007). Such a process may be possibly the cause of Sb enrichment in soil in warm

422 Mediterranean climates.



423

424 Fig. 4. (a) Distribution of Sb-bearing ore deposits in Europe (based on the ProMine Mineral 425 Database; Cassard et al., 2012, 2015; Demetriades and Reimann, 2014). Abbreviations of 426 mineralised districts: AE: Ardennes-Eifel; BC: Betic Cordillera; BF: Black Forest; BRT: 427 Brittany; CI: Corsica; CM: Cantabrian Mts.; ERZ: Erzgebirge (Ore Mts.); IPB: Iberian Pyrite 428 Belt; MC: Massif Central; PIT: Poitou; PYR: Pyrenees; SI: Sardinia; SM: Serbo-Macedonian 429 ore district; SWE: South-west England ore district; VGS: Vosges; WA: Western Alps; (b) Soil 430 geochemical map for hot aqua regia extractable Sb concentrations in ploughed agricultural 431 soil (Ap, N=2108). Numbered anomalies are listed in Table 3.

- 432
- 433 3.2.2. Spatial distribution of W

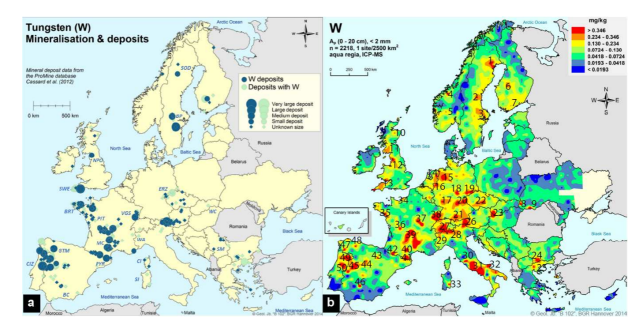
434 The geochemical map of W shows an interesting distribution pattern, which is different 435 from that of Sb (Fig. 5b). Though the northern European Ap soil displays somewhat lower W 436 values (median 0.058 mg/kg) than those from southern Europe (median 0.083 mg/kg), the 437 boundary does not exactly follow the limit of the last glaciation. This is due to the many high 438 values observed in German soil and, in general, in central Europe. In Germany, several 439 anomalies (from Hannover to the border of The Netherlands) coincide with the occurrence of 440 sedimentary iron ore deposits. Poland and Ukraine are in contrast marked by very low W 441 concentrations in Ap soil. This can be explained by the presence of aeolian deposits, coversands 442 and loess, deposited prior to, and just after, the last glaciation maximum and are of Weichselian 443 age (Zeeberg, 1998). These deposits are rich in Hf and Zr (Scheib et al., 2012) but depleted in 444 fine-grained fraction W, and the primary W minerals, such a scheelite and wolframite, are not 445 easily dissolved by traditional hot acid extractions. In fact, most of the elevated W background 446 in Europe is observed in areas underlain by crystalline bedrock.

447 Many granitic intrusions (see Fig. 5b; Cornwall in south-west England (No. 13), 448 northern Portugal (Nos. 49 & 50), Massif Central (No. 39) in France, for example) are clearly 449 marked by W anomalies, as are the Roman and Neapolitan alkaline volcanic rocks in Italy (No. 450 31). The Vosges (No. 38) and the Alpine region (especially the western part, Nos. 27, 28 & 29) 451 are marked by enhanced W concentrations. This is most likely related to the crystalline massifs 452 and to the extensive tectonic lines with associated magmatic rocks (Rhône-Simplon-Insubria-453 Periadriatic Line) and metamorphic rocks (trace W concentrations are common in mica-rich 454 rocks). In Sweden and Finland, the predominance of granitic bedrock (Fennoscandian Shield) 455 over large parts of these countries is well reflected in somewhat enhanced W concentrations. 456 The higher nappes in the Scandinavian Caledonides (mainly calcareous sedimentary rocks) and 457 the Archaean bedrock in the north are characterised by low W soil concentrations. In eastern 458 Finland, W depletion delineates the location of so called greenstone belts. Generally, low W

459 contents in soil occur in areas with glacial drift (central Europe) and with transported calcareous460 materials, e.g., in Denmark.

It is worth noting that the geochemical mobility of W is different from that of Sb; W is mobile under oxidising conditions in alkaline waters and tends to adsorb to Mn-oxides, clay minerals and binds to organic matter. Primary W exists almost exclusively in the form of tungstate minerals (scheelite CaWO<sub>4</sub>, wolframite [Fe/Mn]WO<sub>4</sub>), which are resistant to weathering. Under specific conditions, the formation of soluble W complexes, with many inorganic and organic ligands, enhance its mobility in the surface and subsurface aquatic environment (Koutsospyros et al., 2006).

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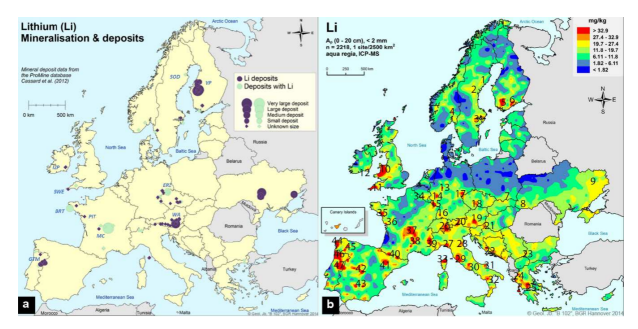
470 Fig. 5. (a) Distribution of W-bearing ore deposits in Europe (based on the ProMine Mineral 471 Database; Cassard et al., 2012, 2015; Demetriades and Reimann, 2014). Abbreviations of 472 mineral districts: BC: Betic Cordillera; BP: Bergslagen ore province; BRT: Brittany; CI: 473 Corsica; CIZ: Central Iberian ore zone; ERZ: Erzgebirge (Ore Mts.); GTM: Galicia-Trás-os-474 Montes zone; MC: Massif Central; NPO: North Pennine orefield; PIT: Poitou; PYR: Pyrenees; 475 SI: Sardinia; SM: Serbo-Macedonian ore district; SOD: Skellefte ore district; SWE: South-west 476 England ore district; VGS: Vosges; WA: Western Alps; WC:Western Carpathians (b) Soil 477 geochemical map for hot aqua regia extractable W concentrations in ploughed agricultural soil 478 (Ap, N=2108). Numbered anomalies are listed in Table 4.

479

480

#### 481 3.2.3. Spatial distribution of Li

482 The aqua regia extractable Li map displays typical GEMAS spatial distribution patterns 483 (observed for many elements), with predominantly low concentrations in Ap soil in northern 484 Europe (median 6.4 mg/kg Li) and evidently higher values in southern Europe (median 15 485 mg/kg Li; Fig. 6a). The maximum extent of the last glaciation is visible as a discrete 486 concentration break on the Ap geochemical map of Li (Fig. 6b). The principal Li anomalies are 487 spatially associated with granitic rocks throughout Europe, e.g., northern Portugal (Nos. 46, 488 47), Massif Central (Nos. 37 & 38) in France, Wales (No. 10), south-west England (No. 11). 489 While in Fennoscandia the Central Scandinavian Clay Belt (Ladenberger et al., 2014) is clearly 490 visible (Nos. 3 & 5), and this feature emphasises the tendency of Li to bind to clay (although 491 developed on the crystalline, mainly granitic bedrock). The Alpine Region (especially the 492 western part) is marked by enhanced Li concentrations (Nos. 26 & 30). This is most likely 493 related to the crystalline massifs and to the extensive tectonic lines with associated magmatic 494 rocks (Rhône-Simplon-Insubria-Periadriatic Line). In northern Portugal, the extensive Li 495 anomaly (Nos. 46 & 47, Fig. 6b) is related to the occurrence of Li-pegmatites. The anomaly in 496 the Wolfsberg area in Austria (No. 26) is also associated with Li-pegmatites with no known 497 relation to any granite. High Li values occurring over karst limestone areas in southern Europe 498 (Croatia, No. 22) are possibly due to secondary Li enrichment attributed to climatic conditions, 499 and to phosphorite and bauxite occurrences (Hellas, Nos. 24 & 25). In arid areas, Li can be 500 enriched via precipitation (together with other evaporite minerals). Lithium's tendency to bind 501 to clay leads to enrichment in fine-grained sedimentary rocks, both young and unaltered and 502 those metamorphosed into schist. Additionally, Li occurs in trace amounts in mica and 503 amphibole minerals, and soil with mica-rich parent materials (such as mica schist) can have 504 high Li concentrations.

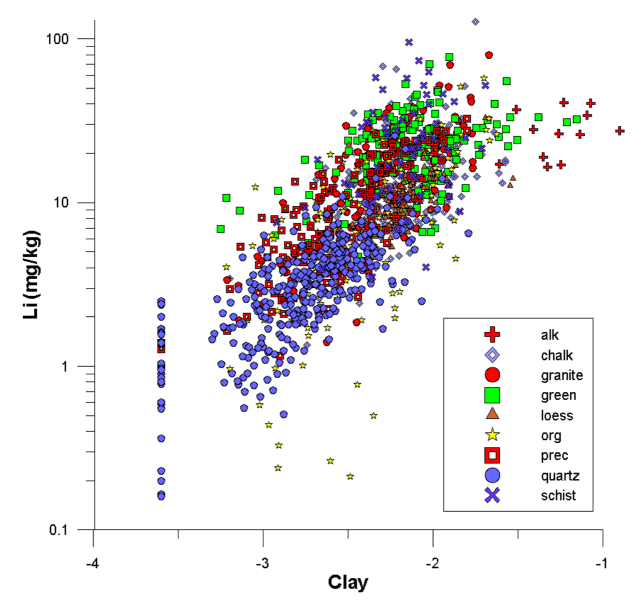


505

Fig. 6. (a) Distribution of Li-bearing ore deposits in Europe (based on the ProMine Mineral
Database; Cassard et al., 2012, 2015; Demetriades and Reimann, 2014). Abbreviations of
mineral districts: BRT: Brittany; ERZ: Erzgebirge (Ore Mts.); GTM: Galicia-Trás-os-Montes
zone; IZP: Irish Zn-Pb ore district; MC: Massif Central; PIT: Poitou; SOD: Skellefte ore
district; SWE: South-west England ore district; VP: Vihanti-Pyhäsalmi ore district; WA:
Western Alps. (b) Soil geochemical maps for hot aqua regia extractable Li concentrations in
ploughed agricultural soils (Ap, N=2108). Numbered anomalies are listed in Table 5.

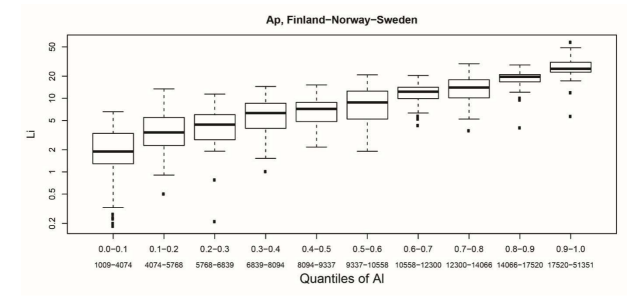
514 The Li concentration is compared with the clay content in Fig. 7. The classical 515 representation of Li concentration (mg/kg) vs. clay content (%) evidenced the existence of 516 heteroscedasticity in the data. To overcome this, the Li concentrations are plotted with 517 logarithmic scaling and the clay contents are expressed as a binary logit function, i.e., a logistic 518 transformation according to  $\log[P/(100-P)]$  where P is the clay content. A clear trend is 519 observed in Figure 7, less visible, however, for soil developed on coarse-grained sandy deposits 520 ('quartz') and calcareous rock ('chalk'), since these parental materials do not produce clay 521 minerals under weathering processes. Better correlation occurs in soil developed on greenstone 522 ('green') and Precambrian bedrock (granitic gneiss - 'prec') - these parent materials are sources 523 to thick clay overburden as a result of weathering, especially in warm and humid climate.

524 The correlation between Li and Fe and Al was earlier reported by Reimann et al. 525 (2012a). The strong Li-Al correlation can be explained by the fact that kaolinite, the most aluminous mineral, easily fixes large amounts of Li compared to other minerals such as illite or
chlorite (Tardy et al., 1972). Using a selection of regional data from the GEMAS project (e.g.,
from Nordic countries), this very strong relationship between Li and Al is shown (Fig. 8).



529

Fig. 7. Plot of clay (%) and Li (mg/kg) contents in European Ap soil samples. Data are classified according to the ten geological parent material subgroups (Reimann, 2012a, b), comprising alk: alkaline rock; chalk: calcareous rock; granite: granitic bedrock; green: greenstone, basalt, mafic bedrocks; loess: loess; org: organic soil; other: unclassified bedrock; prec: predominantly Precambrian granitic-gneiss; quartz: soil developed on coarse-grained sandy deposits; schist: schist.; the category 'other' in the original classification has not been plotted on the graph.



538

Fig. 8. Multiple boxplots showing relationship between Li concentrations (mg/kg) and Al
quantiles in Ap GEMAS samples from Sweden, Finland and Norway (Ladenberger et al., 2013).

#### 541 **3.3. Emerging critical elements in soil as mineral potential indicators**

542 Nowadays, high-tech elements are rarely mined as a main ore. Most often, they are co-products 543 from the principle ore (e.g., Zn, Fe, Cu). In order to identify economic resources of high-tech 544 materials, Sb, W and Li from agricultural soil (Ap) are used as proxies, and it is here discussed 545 whether the geochemical anomalies in soil can be utilised for mineral exploration purposes. It 546 is vitally important to be able to discriminate 'real' anomalies, high element concentrations 547 related to potential economic mineralisation, from secondary enrichment caused by weathering, 548 climate, contamination, etc. Historical and inactive mining regions can be of interest since there 549 are a strong political and environmental reasons to utilise mining wastes, such as tailings and 550 waste heaps.

The majority of Sb, W and Li GEMAS geochemical anomalies in Ap soil can be directly related to known mineralisation and ore deposits (Figs. 4 to 6 and Tables 3 to 5). Except for deposits in the Fennoscandian Shield, most of the ore deposits in Europe are located within Palaeozoic to Caenozoic tectonic units and are related to respective orogenic events, e.g., in the Ordovician (Caledonian orogeny), Carboniferous (Variscan orogeny) and Tertiary (Alpine 556 Orogeny). Active volcanism, regional metamorphism and hydrothermal activity often 557 accompanied these events. In western, central and southern Europe, most mineralisation is 558 located in Variscan and Alpine units, while within the Fennoscandian Shield in northern Europe 559 the majority occurs in Proterozoic (Sweden, Finland) and Caledonian (Sweden and Norway) 560 terranes.

561 In order to evaluate the mineral potential of Sb, W and Li, GEMAS maps have been 562 compared to the ProMine Mineral Database and European Minerals Knowledge Data Platform 563 (Cassard et al., 2012, 2015, which is based on the ProMine Mineral Database; 564 http://minerals4eu.brgm-rec.fr/) (Tables 3 to 5). These online databases contain compiled 565 information about mineralisation and ore deposits for Europe and are often used by companies 566 and policy makers. While most mineralisation is not reflected in the Ap soil geochemistry, there 567 are several relatively large anomalies which cannot be easily explained by the ProMine Mineral 568 Database. Such an example is Sb anomaly No. 39 in eastern France, which coincides with a Pb 569 anomaly (No. 22 in Reimann et al., 2012b) where there is no known mineralisation, no smelter, 570 no major city but correlate well with the proximity to Verdun, a small city made famous because of a fierce First World War (WW1) battle. Anomalies of Sb (No. 49) and Li (No. 41) are also 571 572 unexplained by the ProMine Mineral Database and they appear to be significant in northern 573 Spain. These geochemical anomalies can be found in the Geochemical Atlas of Spain (Locutura 574 et al., 2012) and they seem to correlate with known polymetallic mineralisation (Pb-Zn-Cu-Ag-575 Sb-Bi-As-W-Au) in the pre-Pyrenean and Pyrenean region and to the existence of clays and 576 evaporitic salts north of Zaragoza city and 100 km east in the Cardona-Suria Salt mines.

577 Interestingly, many anomalies of Sb, W and Li can be correlated to the ProMine Mineral 578 Database deposits, which do not list these particular elements as associated commodities, e.g., 579 Nordli Mo deposit in Norway and base-metal sulphide deposits in northern Sweden (Skellefte 580 district). Additionally, a difference in background concentrations between northern and

- 581 southern Europe requires a separate interpretation of Fennoscandian countries by taking into
- 582 account the local background levels.
- 583

### 584 Table 3. Explanation of Sb anomalies in Europe: comparison with ProMine Mineral

- 585 Database (PMD; Cassard et al., 2012, 2015). For anomaly numbers see Fig. 4a.
- 586

No.	Country	Explanation (PMD)	Other explanation
1	Sweden	Various sulphide deposits with	Skellefte Ore District
	(N Sweden)	Zn, Ag, Au, Cu, Pb	
2	Sweden	Zn-Cu-Pb deposits	Black shale in the Lower
	Central		Allochthon in the
	Sweden)		Caledonides
3	Sweden (SE	Associated with Zn-Pb and W	
	Sweden)	mineralisation (associated with	
		Mo and Sb)	
4	Norway	Sulphide mineralisation in	
	(S Norway)	volcanic rocks of the Oslo Rift,	
		Cu-Mo and Au deposits	
5	Finland	Au deposits associated with Ag,	
	(S Finland)	Cu, Bi and Sb	
6	Denmark, (E	Fe mineralisation in young	
	Jutland)	postglacial sediments	
7	Poland (central-	Lignite deposits (Miocene) or	
	northern	sedimentary Fe deposits	
	Poland)	(Jurassic)	
8	SW Poland/SE	Lignite deposits (active opencast	Former lignite power plants
	Germany	mines)	and ash dumps, old opencast
			lignite mines
9	Ukraine	Hg deposits associated with	
		silicified sandstone and coal	
		(Donbass region	
10	United	Various sulphide mineralisation	
	Kingdom	with Sb, associated with Cu, Mo,	
	(Scotland)	Au and Ag; Coalfield region.	
11	United	Old mining region with Pb	
	Kingdom	deposits, sphalerite, pyrite,	
	(Bradford-	galena, chalcopyrite, F and Ba.	
10	Manchester)	Coalfield region	
12	United	Sb sulphide mineralisation,	
	Kingdom	associated also with Pb, W and	
10	(Cornwall)	Sn. Fault-related vein deposits	
13	Eastern Ireland	Sulphide deposits with Zn, Au,	Possible contribution of
		Cu, Pb and Sb, e.g., in	contamination from the
		Glendalough, Wicklow and	capital city of Dublin
1.4		Avoca region)	
14	Netherlands		Anthropogenic anomaly
	(Amsterdam)		related to Amsterdam city

No.	Country	Explanation (PMD)	Other explanation
15	Germany	Sb sulphide deposits, limonite	Pb-Zn-(Ag) sulphide deposits
	(Rhenish Slate	occurrences and historical	in the Stolberger mining
	Mts, Nordrhein-	smelting	district
	Westfalen,		
	Rheinland-Pfalz		
16	Central	Northern anomaly: Cu	Southern anomaly:
	Germany	Kupferschiefer deposits; Pb	Thuringian Forest, Cu-Pb-Zn-
	(Mansfeld,	smelter	Ag mineralisation following
	Lutherstadt		the Hercynian Eichenberg-
	Eisleben,		Gotha-Saalfelder fault zone;
	Gräfenroda)		Copper shale deposits
			(abandoned Cu mines and
	~		mining heaps)
17	Germany	Polymetallic vein mineralisation	Eastern Ore Mts.: Sn
	(Erzgebirge,	Bi-Co-Ni-Ag-U, arsenides	deposits, Altenberger ore
	East Thuringian		district, Schellerhauer granite;
	Slate Mts)		Western Ore Mts.:
			Kirchberger granite, W-Mo
			mineralisation, hard coal
			deposits near Oelsnitz; Sb-(Ag-Pb-Fe-As)
			mineralisations in the Berga
			Antiform
18	SW Germany	Open pit mining with fluorite	Anthonin
10	(Karlsruhe-	associated by Bi and Cu in vein	
	Stuttgart region;	and breccia deposits	
	Pforzheim in		
	Schwarzwald)		
19	S Germany - W	Leogang (inactive plant) with Fe,	
	Austria	Mn, Co, Ag, Cd, Hg, Ga, In, Ge,	
	(Insbruck	Sn, Sb	
	region)		
20	Czech Republic	Coal and lignite deposits with Ge	
	(Plzen)	(Plzen and Radnice basin); Sb	
		(antimonite)-Zn-Pb deposits	
21	Czech Republic/	Fe deposits with Sb; coal	
	Poland	deposits	
22	Czech Republic	Pb mineralisation; minor coal	
	(Opava)	deposits	
23	Central	Numerous Sb deposits (with Sb,	Few localities with larger
	Slovakia	Pb, Zn, Ag and Au)	findings of Sb minerals,
			together with Au and Ag in
			hydrothermally altered
			Miocene andesites -
0.1			www.mindat.org)
24	Hungary (N of	Pb-Zn mineralisation	Anthropogenic anomaly –
	Budapest,		Budapest City
	Visegrad)		

No.	Country	Explanation (PMD)	Other explanation
25	E Austria/W	Sb-Hg mineralisation; lignite	
	Hungary	deposits	
26	S Austria	Carbonate-hosted Pb-Zn deposits	
		(MVT) with Ga, Ge, Pb, Zn; Cu	
		VMS deposits; inactive plants	
		(mainly surface storage)	
27	W Austria	Pb-Zn deposits; skarn deposits	
	(Insbruck)	with W; Cu deposits; inactive	
		plants (mainly surface storage)	
28	W Austria/N	Minor Zn mineralisation (with	
	Italy	Ga and In); inactive plants	
		(mainly surface storage)	
29	Slovenia	Sb mineralisation (vein and	
		disseminated); Pb-Zn carbonate-	
		hosted deposits (MVT)	
30	Bosnia &	Sb mineralisation; Pb-Zn	
	Herzegovina	mineralisation; open pit mining	
	(Sarajevo	(Pb-Zn with Sb)	
	region)		
31	Serbia	Cu-Au VMS deposits; Sb vein	
		deposit; Sediment-hosted	
		precious metal deposits	
32	S Serbia	Sediment-hosted precious metal	
		deposits with Sb; Pb-Zn Sedex	
		deposits; coal deposits	
33	SE Serbia/	Sb deposits; Pb-Zn +Ag skarn	
	North	mineralisation; Sb deposits with	
	Macedonia	stibnite	
34	N Cyprus		Western part of the Kyrenia
	• 1		Terrain (Alpine orogenic
			belt); however, it is most
			likely a transported anomaly,
			and related to the Troodos
			Ophiolite complex massive
			sulphide mineralisation; it
			correlates with anomalous
			Ag, Au, As, Hg, Pb, S, V and
			Zn
35	Central Italy	Sb deposits	
36	Central Italy	Sediment-hosted Sb	Rome – anthropogenic
		mineralisation, related to shallow	anomaly; young volcanic
		intrusions	rocks in the Rome region
37	S Italy (Naples)		Naples anthropogenic
			anomaly; young volcanic
			rocks (Vesuvius) which are
			responsible for Sb enrichment
			in local soil and groundwater

No.	Country	Explanation (PMD)	Other explanation
38	SE Sardinia (Italy)	VMS deposit with Sb (antimonite); Pb-Zn deposits, Ag-Sb mineralisation, Zn-Sb carbonate-hosted stratabound and vein mineralisation (MVT)	
39	E France	anthropogenic	Verdun – WW1
40	E France (Vosges)	Sb mineralisation; Pb-Cu-Ag mineralisation	Similar anomaly is observed for Pb (Reimann et al., 2012b); Variscan orogen
41	N France	Sediment-hosted baryte-Sb deposits, vein and metasomatic polymetallic deposits with Sb, Cu, Pb	
42	NW France	Polymetallic, mainly vein types deposits with Sb, Bi, Mo, Sn, W in granitic host rocks	
43	W France	Carbonate-hosted (in karst) Pb- Zn (+Ag) deposits MVT, known since medieval time; historical Ag smelting	
44	France (Limoges, Haute-Vienne, W Massif Central)	Carbonate-hosted Pb-Zn (Ag) MVT deposits; several Sb mineralisation with stibnite and berthierine, related to numerous Au deposits	Au-bearing granitic mica schist
45	France (S Massif Central)	Pb-Zn (Ag) deposits; Sb mineralisation with stibnite, a large system of Au and sulphide- rich quartz veins	
46	France (Massif Central, S of Clermont Ferrand, and west of Lyon)	Numerous Sb mineralisation associated with Au, Ag, Pb, Zn; mainly fault-related	
47	SE France (Grenoble)	Cu-Pb-Ag + Au (±Sb) fault related deposits, sandstone ± dolomite hosted Pb-Zn + Ag +Sb deposits	
48	NE Spain/Andorra (Pyrenes)	Volcano-sedimentary Zn-Ag deposits	
49	NE Spain (S Pyrenes)		Various polymetallic mineralisation (Pb-Zn-Cu- Ag-Sb-Bi-As-W-Au) in the pre-Pyrenean and Pyrenean region (Locutura et al., 2012)

No.	Country	Explanation (PMD)	Other explanation
50	S coast of Spain	Fault-related vein and breccia	
	(Granada	deposits with Ag, Zn (galena);	
	region)	vein and disseminated Sb	
		deposit; carbonate-hosted Pb-Zn	
	~	deposits with Ag	
51	Central Spain	Sb vein deposits; fault-related Pb-Zn-Ag deposits	
52	Central Spain	Pb-Zn-Ag deposits (with galena,	
02	(N of Cordoba)	sphalerite)	
53	SW Spain	Iberian Pyrite Belt: Zn-Pb-Cu	
		(+Ag) VMS deposits	
54	NW Spain	Coal deposits; numerous Au	
	(Galicia, Oviedo	deposits (quartz veins,	
	region)	conglomerate and alluvial	
		placer); Sb (stibnite) deposits	
		with cinnabar	
55	NE Portugal	Au-As mesothermal deposits	
		(Au, Ag, As, Zn, Pb); Sn-W	
		deposits (cassiterite, wolframite,	
		scheelite, silver, pyrite,	
		arsenopyrite, marcasite) in	
		granitic veins and stockworks	
		(greisen)	
56	Central Portugal	Sb-W fault-related deposits; Au-	
		Sb mineralisation	
57	S Portugal	Sb fault-related deposits	
		(antimonite, malachite, baryte,	
		silver); Zn-Pb-Cu (+Ag) VMS	
		deposits	

#### 588 3.3.1. Antimony in soil as a mineral potential indicator

589

590 By studying the Sb maps in more detail (Figs. 4a, b), it can be observed that most of the 591 identified anomalies correspond with known mineral belts or ore deposits (Table 3). The highest 592 Sb contents in Ap soil occur in Portugal where several known Sb occurrences are located, for 593 example, within the so called Iberian Pyrite Belt, in south-eastern Portugal. In Austria, an Sb 594 anomalous region stretches in a form of a belt in the Alps and coincides with many known 595 mineralised occurrences and historical mines. The Massif Central in France with its extensive 596 hydrothermal mineralisation stands out as an anomaly, as does the Iberian Pyrite Belt (southern 597 Spain and Portugal), the Vosges (French and German border), Ardennes-Eifel (Germany), the 598 Ore Mountains (German and Czech border), Bohemian Forest (Czech Republic), Carpathians 599 (Slovakia), and the Pennine and south-west England ore-fields in the United Kingdom. High 600 Sb contents in Ap soil occur also in the Balkan countries, Bosnia and Herzegovina, Serbia (with 601 Kosovo), North Macedonia and Hellas, and they all are related to various types of 602 mineralisation, mainly polymetallic with Pb-Zn-Cu, the Serbo-Macedonian mineralised zone.

603 Due to strong contrast in Sb concentrations between northern and southern Europe, the 604 Sb concentrations in the soil of Fennoscandia, Estonia, Latvia, Lithuania, northern Poland, 605 Denmark and northern Germany should be displayed as a separate data set (Fig. 9). In 606 Fennoscandia, relatively high Sb concentrations are mainly related to polymetallic 607 mineralisation, e.g., Pb-Zn-As deposits in northern Sweden (Skellefte district), Sb, Bi, Au and 608 As mineralisation north-west of Luleå, and in central Sweden (Bergslagen district). Known Zn-609 Pb mineralised occurrences in SW Scania in southern Sweden are indicated by a large Sb 610 geochemical anomaly in grazing land soil. High Sb concentrations in two Ap soil samples are 611 unexplained; one in the northernmost Sweden within the Archaean bedrock and the other in SE 612 Sweden (Småland, near Oskarshamn). The remarkable enrichment in Sb in the fine-grained 613 sedimentary rocks (mainly in black shale, e.g., in the Lower Allochthon of the Caledonides), 614 although appearing very promising, has potentially no economic significance mainly because 615 Sb is dispersed at low levels and does not form economic ore-bodies, as its mining is difficult 616 and requires processing of large volumes of rocks. In Norway, large Sb anomalies are closely 617 related to mineralisation within the Oslo Rift and in SW Norway with some sulphide 618 polymetallic deposits (Fig. 9).

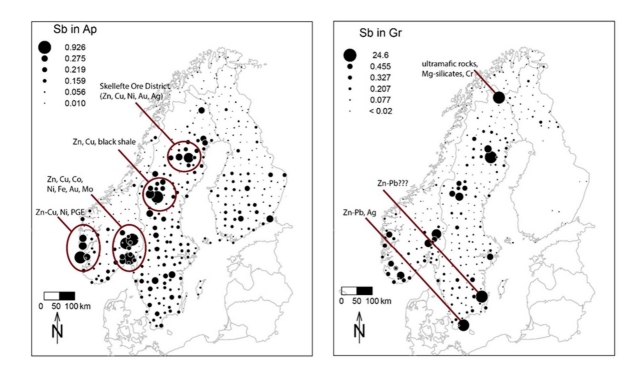


Fig. 9. Sb geochemical maps (mg/kg) in Ap (n=453) and Gr (n=350) soil of Norway,
Sweden and Finland. Regional Fennoscandian percentile calculation allows clear detection of
anomalies. Most of them can be explained by mineralisation or lithology.

Economic occurrences of Sb in Europe are mainly associated with sulphide deposits, and Sb can be used as a pathfinder element for sulphide mineral deposits (Boyle, 1974; Boyle and Jonasson, 1984). Another example is from deposits in Austria that were economic in their time related to their mono-mineral characteristic (stibnite). Antimony is also used together with As and Bi as a pathfinder element for Au mineralisation (Boyle, 1974; Boyle and Jonasson, 1984; Plant et al., 1989). In other words, using Sb, sulphide ores can be classified according to their potential for hosting by-products of economic importance.

Because many of the Sb-hosting deposits have been mined historically (e.g., in France), it is difficult to evaluate to what extent Sb anomalies are natural anomalies (due to the occurrence of the deposits themselves) or whether the extent of mining and smelting activities has accentuated originally very local anomalies. To answer the question 'natural or anthropogenic' in these areas, detailed studies would be needed in every single case. On the other hand, the size of an anomaly can be used as pathfinder for potential future recovery at the 637 source (e.g., from past tailings). Otherwise, typical anthropogenic Sb anomalies related to 638 urbanisation, although rare, they have been recognised near Amsterdam, Budapest, Rome and 639 Naples. Additionally, interesting historical Sb impacts can be observed, e.g., in Naples region 640 (Italy). The rather large Sb anomaly, apart from possibly being anthropogenic (Naples), 641 coincides with the location of Europe's largest volcano – Vesuvius. Antimony occurs naturally 642 in groundwater close to volcanoes, and Pompeii's proximity to Mount Vesuvius could have 643 provided Sb concentrations in the water that were even higher than in a typical Roman city at 644 the time. Antimony poisoning is known in this area from historical times. The Pb pipes in 645 Pompeii, studied by archaeologists, contain Sb which was likely a cause of health problems 646 among the residents (Charlier et al., 2017).

647

648 Table 4. Explanation of W anomalies in Europe: comparison with ProMine Mineral Database

649	(PMD; Cassard et al.,	, 2012, 2015).	For anomaly numbers	see Fig. 5.
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No.	Country	Explanation (PMD)	Other explanation
1	N Sweden	Cu-Ag mineralisation with	W mineralisation in skarn;
		scheelite occurrences	associated Cu, Zn, Au, Ag
			mineralisation (SGU database)
2	Central Sweden	Au-Cu+Ag mineralisation	Many small W mineralised
			occurrences and historical W
			(+Pb+Cu) mining (SGU database)
3	Central Sweden	Scheelite mineralisation	
	(Bergslagen)	(+Mo, Fe-Cu-Ag) and	
		historical mining sites	
4	W Norway		Mineralisation in quartz-rich
	(Molde-		lithologies, V-Fe-Ti
	Ålesund)		mineralisation in western Norway
			(FODD database)
5	S Norway		Small granitic intrusions, Zn-Cu-
	(Bergen, Odda)		Au mineralisation (source: FODD
			database); Odda - a major Zn
			smelter in Norway
6	Central Finland	Mineralisation in pegmatites,	
		numerous gold occurrences	
7	S Finland	Scheelite-Mo (As+Co) skarn	
		mineralisation, associated Au	
		mineralisation	
8	SW Ukraine		Epithermal polymetallic deposits
			in Transcarpathian region. Vein-

No.	Country	Explanation (PMD)	Other explanation
			type Au-base metal deposits hosted by volcano-sedimentary sequences of the Pannonian basin in the Carpathian Neogene volcanic belt
9	SW Ukraine		Au-Ag-Zn-Pb-Cu mineralisation and epithermal deposits in Transcarpathian region
10	Scotland	Wolframite-scheelite mineralisation in Devonian granites (associated with cassiterite, molybdenite, sphalerite, chalcopyrite, topaz)	
11	Scotland W (Southern Uplands)	Au-Pb-Cu-As-Ag mineralisation	
12	NW England (southern part of Lake District and Pennines)		W mineralisation in the Lake District (Cumbria)
13	United Kingdom (SE Wales, Cornwall)	Wolframite-cassiterite mineralisation (Cs+Li) in Cornubian orefield	W mineralisation in Cornubian batholith
14	NE Netherlands	unexplained	
15	NW Germany (Lower Saxony)	Fe-Ag-Pb-Zn mineralisation in felsic alkaline plutonic rocks; metasomatic carbonate-hosted haematite- magnesite deposits; Fe smelter	Abandoned iron ore mines
16	W Germany (North Rhine- Westphalia)	Fe smelter	Hard coal deposits and mining, inactive hard coal power plants
17	Rhineland- Palatinate	Minor Ag-Cu mineralisation	Donnersberg rhyolite massif, historical Cu mines
18	Central Germany (Thuringian Forest)	Cu-Pb-Zn-Ag mineralisations	Cu-Pb-Zn-Ag vein mineralisation following the Hercynian Eichenberg-Gotha-Saalfeld fault zone; Cu shale deposit
19	SE Germany (Erzgebirge, Ore Mts)	Sn-W (+Ag+Zn+Ge) mineralisation in greisen	W-Mo mineralisation in the area of Kirchberger granite
20	S Germany (W and NE Nürnberg) / W Czech Republic	U-W (+Sn) mineralisation	Pb-Zn mineralisation, Baryte-fluorspar-haematite mineralisation

No.	Country	Explanation (PMD)	Other explanation
21	S Germany	Minor Pb-Zn mineralisation	Pitch coal occurrences (strong
	(Molasse basin)		pressed lignite) in the south
22	Central Czech	W and Pb-Ag mineralisation	
	Republic		
23	SW Slovakia-	Au-Sb-Ag-Zn-Pb	
	NW Hungary	mineralisation	
24	SW Bulgaria	polymetallic W-Bi-Mo	
	(Western	deposits; W-Pb-Mo-Cu	
	Rhodopes Mts)	mineralisation in granite and	
		skarn	
25	Hellas (Central	Sb-W vein mineralisation;	
	Macedonia)	Cu-Ag-Au porphyry deposits	
26	W Austria	Numerous W occurrences	
	(Insbruck, Hohe	(scheelite-Mo skarn)	
	Tauern)		
27	S Switzerland	Au, REE, Pb-Ag-Zn	
20		mineralisation	
28	N Italy	Zn-Ag carbonate-hosted	
20		mineralisation	
29	NW Italy	Cu-Ag mineralisation	V-lessie entrie
30	Central Italy	Minor Sb (Hg, Cu, Zn, Pb) mineralisation	Volcanic origin
31	Control Itoly	mineralisation	Valaania arigin
51	Central Italy		Volcanic origin
	(Rome-Naples region)		
32	Italy (Gargano		Plutonic and volcanic alkaline
52	Peninsula)		rocks (gabbro and syenite,
	i chinisulu)		anorogenic lamprophyres); skarn
			mineralisation at the contact with
			limestone
33	SE Sardinia	W-Zn-Pb mineralisation;	
		Mo-W mineralisation in	
		greisen; Sb-W VMS	
		deposits; Au mineralisation;	
		Ag-Sb-Pb-Zn polymetallic	
		veins	
34	N France	unexplained	
35	NW France	W (scheelite) – Mo skarn	
		mineralisation; wolframite in	
		greisen	
36	W central	Numerous W (+Mo, Sn, Zn,	
	France (Poitu,	Cu) mineralised occurrences	
	Limousin,	in relation to granitic	
	Haute-Vienne)	intrusions and in greisen	
37	Central France	W (+Sn, Mo, Py)	
		mineralisation in granite and	
		greisen	

No.	Country	Explanation (PMD)	Other explanation
38	E France	Numerous W mineralisation;	
	(Vosges)	W-Mo skarn; wolframite and	
		scheelite in greisen	
39	Central France	Numerous W mineralisation	
	(Massif Central)	(+Bi, Sn, As, Mo, Cu)	
40	S France	Numerous W mineralisation,	
	(Provence-	carbonate hosted Pb-Zn	
	Alpes-Côte	MVT deposits; W-Mo skarn	
	d'Azur region)	deposits in greisen (with Sn)	
41	S France	W-Mo skarn mineralisation,	
	(Pyrenees)	associated with Au and Ag	
42	S France	W-Mo skarn mineralisation,	
	(Pyrenees)	associated with Ag, Zn and	
		Pb	
43	N Spain	Minor Ag occurrences	
44	Central Spain,	W-Mo in skarn; W-Sn	
	near Madrid	mineralisations	
45	W Spain	W mineralisation	
46	S central Spain	W mineralisation (with	
	(Cordoba)	associated As, Ag,	
		tourmaline)	
47	NW Spain	W-Fe-Sn mineralisation	
	(Galicia)		
48	NW Spain	Au and W mineralisations in	
	(Galicia)	granite-greisen-quartz veins	
49	N Portugal	W mineralisation in	
		pegmatites (scheelite,	
		wolframite); W-Sn	
		mineralisation with	
		associated Au (Ag)	
50	N Portugal	W-Sn-Cu-Zn mineralisation	
		in granite and greisen	

650

651 3.3.2. Tungsten in soil as a mineral potential indicator

652

Tungsten anomalies in European Ap soil are most pronounced in regions with granitic bedrock and associated with various types of mineralisation, especially Au (and Ag) and base metals hosted by granitic rocks (Figs. 5a and b). This is typical geological environment where various W ore deposits can be found, e.g., granitic intrusions in northern Portugal, Massif Central in France, Bohemian Massif in Czech Republic, the Vosges Mountains in eastern France and the Alpine region (mainly in Switzerland and western Austria). Scheelite and 659 wolframite (often accompanied by cassiterite and fluorspar) are the economic sources of W, 660 and they are mainly found in quartz-rich veins, pegmatites and skarn deposits. Most W deposits 661 are of metasomatic or hydrothermal origin. Tungsten is used as a pathfinder for gold deposits. 662 High W concentrations occur in Ap soil that overlies igneous and metamorphic rocks of the 663 Fennoscandian Shield, mainly of Palaeoproterozoic age. In Scandinavia, the largest W 664 anomalies occur in the Skellefte ore district, Bergslagen and sporadically along the coast of the 665 Gulf of Bothnia. These anomalies reflect known W mineralisation occurring in granite, gneiss 666 and quartz veins (together with U, Au, Ag and Li). In Lapland, anomalies occur within the so 667 called Gold Line and they follow Sn, Mo, Cu, Fe, U and precious metal ore deposits with Au 668 and Ag. The Bergslagen mining district comprises the most important W (scheelite) ore deposits 669 in Sweden, which are located mainly in skarn-limestone, commonly in contact with 670 Svecokarelian granitic rocks (Andersson et al., 2014).

671 By comparison with the ProMine mineral deposit map (Table 4; Figs. 5a and b), it can 672 be observed that several strong anomalies are not reflected on the mineral deposits map, e.g., 673 anomalies in NW Germany/Eastern Netherlands, SW Slovakia, central Italy, southern and 674 central Germany, northern Sweden and central Finland. For example, an unexpectedly 675 extensive W anomaly in central Italy (No. 31, Fig. 5b) might be related to the young volcanism. 676 Other anomalies, such as No. 18 in the Thuringian Forest indicates that mining and exploration 677 activities in this polymetallic province did not focus on W, which shows elevated soil 678 concentrations. All these observations indicate that there is a high potential for discovering new 679 W resources.

680

Table 5. Explanation of Li anomalies in Europe: comparison with ProMine Mineral Database
(PMD; Cassard et al., 2012, 2015). For anomaly numbers see Fig. 6.

No.	Country	<b>Explanation (PMD)</b>	Other explanation
1	E Sweden		Occurrences of LCT
			(Lithium-Cesium-

No.	Country	Explanation (PMD)	Other explanation
			Tantalum) pegmatites of
			the Varuträsk type
2	Central Sweden		Li (Sn, Ta)
			mineralisation (FODD
			database); Sn, Li, Nb,
			Be, Ta occurrences
			(SGU database)
3	Central Sweden	Li in pegmatites	Lithium seems to be
	(Bergslagen)	(spodumene, beryl,	enriched in some Fe
		cassiterite); Li-Fe in	deposits
		magnetite skarn; REE and	
		pegmatite occurrences	
4	Norway (Oslo Rift)		Mineralisation in
			pegmatites; Mo
_			mineralisation
5	S Finland	Ta, Be, Li, Sn in pegmatites	Rapakivi granites; Li
6			mineralisation
6 7	S Finland SE Poland	unavalained	Rapakivi granite
8	SW Ukraine	unexplained	
0	(Carpathians)	unexplained	
9	E Ukraine (Donetsk	Li mineralisation	Phosphate-Be-REE-Ta-
2	region)	Li mineransation	Nb deposits; Li deposits
			in eastern Ukraine
			(petalite and spodumene
			in pegmatites)
10	United Kingdom (Wales)		Li-bearing secondary
			manganese minerals in
			hydrothermal ore
			deposits and sedimentary
			Mn deposits
11	United Kingdom	Li and Cs in pegmatites and	Cornubian batholith
	(Cornwall)	aplites (Li, Ta, Nb, Be)	(lithium-mica-albite-
			topaz granites)
12	SE Ireland (Wexford)	Li mineralisation in	
		pegmatites; LCT pegmatites	
		(Lithium-Cesium-Tantalum)	
13	W Germany (Northern	Unexplained; coal?	Slate mining (roof slate);
	Hesse, Sauerland, SE		fluorspar and baryte
	Dortmund)		mining; hard coal power
			plant in the region;
			industrial region with
			different types of
			industry (metalworking) and steel production
14	W Germany (Rhenish	Kaolin occurrences;	
14	Slate Mts., SE Bonn)	phosphorites	
15	SW Germany (Saarland)	Oolitic Fe-ores; Cu-Pb-Zn	Hard coal deposits,
10	Surfaind)	mineralisation	former hard coal power
L		minorunoution	Torritor hurd cour power

No.	Country	Explanation (PMD)	Other explanation
			plants and iron and steel works
16	SW Germany (Central	Au-Ge-Ag, galena, fluorspar	Pb-Cu-Fe and Fe
	Black Forest)	occurrences	mineralisation, Triberger biotite granite
17	SE Germany/W Czech	Numerous Li occurrences in	Li in Sn mineralisation
	Republic (West	LCT pegmatites	(cassiterite-quartz
	Erzgebirge, Western Ore Mts.)		association) and in Li- bearing micas in granites
18	S Czech Republic	unexplained	
19	S Austria	Li mineralisation in LCT pegmatites	
20	W Austria (Insbruck)/N	Li mineralisation in LCT	
	Italy	pegmatites	
21	E Slovenia	unexplained	
22	SE Croatia/S Bosnia &	Fe deposits (oolitic iron	Karst/bauxite weathering
	Herzegovina/Montenegro	deposits); bauxite deposits in	
		karst and laterite; inactive Al	
22	W Dulcomio	plant Granite-controlled Au-Cu	
23	W Bulgaria	deposits with quartz veins;	
		supergene clay deposits	
24	NW Hellas (Epirus)	Laterite related polymetallic	Karst/bauxite
		deposits (Ni, As, Au, Co, Cr,	weathering; correlation
		Fe, Mg, Mn, Pd, Pt);	to mafic and ultramafic
		Ophiolite related VMS	rocks (ophiolite) and Fe-
		deposits (Cu, Zn, Co)	Ni deposits;
			phosphorites; evaporites
25	C Hellas (Sterea Elladha)	Bauxite and Al-rich rocks	(by the coast) Karst/bauxite
23	C Hends (Stered Endula)	deposits (in karst and	weathering? Possible
		laterite); mine waste dump	correlation to mafic and
		with 'red muds' from bauxite	ultramafic rocks and Fe-
		refining	Ni deposits
26	SE Switzerland	Ta-U mineralisation in felsic	Mineralisations in quartz
		alkaline rocks	veins, pegmatites
27	NW Italy (Genoa,	Volcano-sedimentary Mn	
	Portofino)	mineralisation with jasper and chert; Cr mineralisation	
		in mafic-ultramafic rocks	
		(Cu-Zn, Co VMS deposit)	
28	Central Italy (Ravenna)		Valli di Comacchio:
			brackish lagoons with
			brines which might be
0.0			enriched in lithium
29	Central Italy (including	Li - Cs - REE mineralisation	Related to the igneous
	island of Elba)	in pegmatites; Sb (+Hg) mineralisation	rocks from Roman Alkaline Province
30	Italy	unexplained	Aikainie Province
50	Italy	unexplained	l

No.	Country	Explanation (PMD)	Other explanation
31	Italy (Gargano Peninsula)	Bauxite deposits and Al-rich rocks in karst/laterite	Possibly related to the plutonic and volcanic alkaline rocks (gabbro and syenite, anorogenic lamprophyres
32	Italy (Southern Apennines)	unexplained	
33	N Corsica (France)	Granitic-pegmatitic veins and greisens (W-Mo, Ag)	
34	S Belgium (Dinant, Ardennes)	Vein deposits with fluorspar, baryte, sphalerite, galena, etc; historical mining	Kaolinised regoliths
35	NW France (Hercynian Armorican Massif)	Peraluminous leucogranites, pegmatites and peri-granitic veins (associated with U, Sn, W, REE, Zr and beryl)	
36	NW France (Poitu)	Associated with W-baryte- kyanite deposits, carbonate- hosted veins with Ba and F deposits (MVT)	
37	S Central France (Massif Central)	Li mineralisation in pegmatites associated with beryl, cassiterite, Ta-Nb, Y and REE; numerous occurrences with Sn, W, Au, Ag	
38	S Central France (Massif Central)	Numerous occurrences (+Sn, As, Fe, W, Ag, Au) in granitic and peri-granitic veins and in greisens	
39	SE France (Grenoble)	Cu-Sn-Ag-Mo-Bi and baryte-fluorspar granite- controlled mineralisation; In- Zn, fluorspar mineralisation, fault-related; granitic and peri-granitic veins; historical mining	
40	S France/N Spain (Pyrenees)	W-Mo skarn mineralisation, associated with baryte; Ag, Zn and Pb; Zn-Ag, Co-Sn Pb (+Ga, Ge) mineralisation; historical mining	
41	N Spain (Zaragoza)	Clays and evaporitic salts deposits	This anomaly in south Pyrenean Zone correlates well with Li anomaly (partial extraction) in sediments in Geochemical Atlas of

No.	Country	Explanation (PMD)	Other explanation
			Spain (Locutura et al., 2012).
42	Central Spain (Madrid)	Sn, W, Ag mineralisation related to granite, pegmatite, greisen and quartz- and mica-rich (muscovite) veins; W-Mo skarn mineralisation	
43	S Central Spain	W, Sn, quartz, tourmaline; As, Fe, Ag, Au, Pb mineralisation in granite- pegmatite rocks	Related to Iberian Massif (Central Iberian Zone); the Almadén mercury mining district
44	NW Spain (Galicia)	Associated Au-As mineralisation in quartz-rich veins; Fe-Sn-W-Cu in granites and pegmatites	Hercynian granite; Li- rich mineral occurrences associated with aplite- pegmatite dykes and sills intruded in granitic and metasedimentary rocks of Galicia – Trás-os- Montes geotectonic zones
45	NW Spain (Galicia)	Associated with Au-Ag in quartz-rich veins; Sn-Ta-Nb, quartz, kaolin and REE mineralisation	Hercynian granite
46	N Portugal (Braga)	Lepidolite, spodumene, petalite mineralisation in pegmatite, associated with W-Sn-Au-Ta vein mineralization zone	Galicia - Trás-os-Montes Zone; aplite-pegmatite veins in a late Hercynian granite, rich in spodumene, embedded in Silurian metasediments
47	Central Portugal (Castelo Branco)	Li mineralisation in aplites and pegmatites, associated with W-Sn-Au, Ta-Nb, ilmenite, beryl mineralisation	The Central-Iberian Zone; aplite-pegmatite veins in a late Hercynian granite

683

## 684 3.3.3. Lithium in soil as a mineral potential indicator

Lithium in agricultural soil (Table 5, Figs. 6a and b) shows its high concentrations predominantly in regions with granitic parent materials. A comparison between Li-bearing ore deposits in Europe and the Ap soil geochemical map for Li reveals that many anomalies are associated with Li-bearing mineralisation, especially with occurrences of LCT (Lithium-Cesium-Tantalum) pegmatites, for example, in the Galicia-Trás-os-Montes zone (anomalies 690 No. 46 and 47 and in western Alps (anomaly No. 39). In NW Spain and northern Portugal, Li 691 anomalies are closely associated not only with known Li mineralisation but also with Sn, W, 692 Nb and Ta occurrences. These Li deposits in the Central Iberian Zone have been mined 693 intensively in recent years and Portugal became the world's sixth largest Li producer (Roda-694 Robles et al., 2016). In northern Europe, high Li concentrations in Ap soil occur in regions with 695 granitic parent materials (e.g., southern Finland), but also in till overburden enriched in 696 postglacial and marine clay, for example, in central Sweden (see also Fig. 8). In southern 697 Lapland, anomalies coincide with Au, W, Mo and scheelite mineralisation of the 'Gold Line' 698 ore province. Recently, the Bergby deposit was discovered in central Sweden, which lies 25 km 699 north of the town of Gävle with Li<sub>2</sub>O averaging 1.71% and ranging from 0.01% to 4.65% 700 (http://leadingedgematerials.com/leading-edge-materials-discovers-lithium-mineralization-in-701 outcrop-at-bergby-project-sweden/).

702 There is a clear correlation of Li with mica- and clay-rich soil (as reflected by Ga, Al, 703 Cs, Rb) but also with some Fe-rich phases, which can be a result of both weathering and 704 presence of more mafic parent materials (correlation together with Fe, Cr, Co, Sc). High 705 correlation with chalcophile elements such as Zn, In, Tl, Cu makes geochemical behaviour of 706 Li in soil more complicated and requires careful interpretation at the regional to local scale. 707 Lithium anomalies are more difficult to follow in the ProMine Mineral Database (Cassard et 708 al., 2012, 2015). There are several relatively extensive anomalies, which do not have a direct 709 explanation with respect to the ProMine Mineral Database, e.g., Li anomaly in Wales (No. 10, 710 Fig. 6) in the United Kingdom, Li anomalies in western Germany (Nos. 13 and 14), in W Hellas 711 (Nos. 24 and 25) and in NE Spain (No. 41). Locally, high Li concentrations in Ap soil occur at 712 Lisbon city limits in the delta of the Tajo river, and in the vicinity of the brackish lagoons with 713 brines near Ravenna in Italy, and they may have secondary and even anthropogenic origin.

These enhanced secondary Li concentrations can be misleading when looking for economic Lioccurrences.

716

717 3.3.4. Combined Sb, W and Li anomalies in soil as mineral potential indicators

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Table 6 summarises the co-existing anomalies for Sb, W and Li. The most straightforward observation is that in several regions (Cornwall, Massif Central, Erzgebirge, and the Alps) all three commodities can be found close to one another and, thus, polymetallic ore provinces can be defined, even if the host rock is not of the same lithology and the oreforming processes also differ.

724

725	Table 6. Summary of co-existing Sl	o, W and Li anomalies.
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	1	1	1
Region	Sb	W	Li
Cornwall (United Kingdom)	12	13	11
Massif Central (France)	44-46	39	37-38
W Pyrennes (N Spain/S	48	42	40
France)			
Erzgebirge (E Germany/N	19	17	17
Czech Republic			
W Salzburg/Tyrol (Austria)	27	26	20
Brittany (NW France)		35	35
Central Portugal		50	47
Southern Czech Republic		18	22
SE France	47		39
NE Spain	49		41
E Ukraine	9		9
W France	43	36	
E France	40	38	

726 727

## 728 **4. Conclusions**

729

730 The demand for 'high-tech' element resources stimulated efforts to find new deposits and even 731 possibilities to extract them from active and historical mines throughout the world. The 732 GEMAS data set offers European scale results for several so-called high-tech elements, and 733 almost all listed on the Critical Raw Materials list by the European Commission (2017). Here, 734 three of the high-tech elements, Sb, W and Li were used as test elements. The geochemical 735 distribution of Sb, W and Li in agricultural soil of Europe appears to be a useful starting tool 736 for identifying areas for discovering new sources of high-tech elements. Most of the patterns 737 observed on the geochemical maps of Sb, W and Li can be directly interpreted in terms of 738 known ore deposits and mineralised occurrences and most of the anomalies reflect natural 739 sources of these elements. Additionally, their spatial distribution seems to be modified by 740 weathering type and intensity, Quaternary history and climate, like, for example, the clear 741 difference in element concentrations between northern and southern Europe. The chemical 742 composition of agricultural soil represents largely the primary mineralogy of the source parent 743 material, with superimposed effects of pre- and post-depositional chemical weathering, 744 formation of secondary products such as clays, and element mobility. For Sb, the most 745 promising are regions with known sulphide mineralisation, while for Li and W, regions with 746 crystalline bedrock and evolved granitic rocks should be of interest.

Natural background concentrations of Sb, W and Li in agricultural soil define areas of interest and their relationship to mineralisation, rock types and mining centres, and provide important ground for further investigations. The application of GEMAS soil data to mineral exploration is promising, especially as most of the Sb, W and Li anomalies can be explained by known and mapped mineralisation (such as the ProMine Mineral Database and national registers). In conclusion, several unexplained anomalies require more detailed geochemical surveys, these could shed light on potential mineral resources that have to date remainedundiscovered.

755

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## 773 References<sup>1</sup>

774

- Aitchison, J., 1986. The statistical analysis of compositional data. London: Chapman & Hall, 416 pp.
- Ali, S.H., Giurco, D., Arndt, N., Nickless, E., Brown, E., Demetriades, A., Durrheim, R, Enriquez, M.A.,
- 780 Kinnaird, J., Littleboy, A., Meinert, L.D., Oberhänsli, R., Salem, J., Schodde, R., Schneider, G.,
- Vidal, O., Yakovleva, N., 2017. Mineral supply for sustainable development requires resource
  governance. Nature 543, 367–372.

<sup>Albanese, S., Sadeghi, S., Lima, A., Cicchella, D., Dinelli, E., Valera, P., Falconi, M., Demetriades, A.,
De Vivo, B. and the GEMAS Project Team, 2015. GEMAS: Chromium, Ni, Co and Cu in agricultural
and grazing land soil of Europe. Journal of Geochemical Exploration 154, 81-93.</sup> 

<sup>&</sup>lt;sup>1</sup> Note: All web links checked on the 9<sup>th</sup> March 2019

Andersson, M., Carlsson, M., Ladenberger, A., Morris, G., Sadeghi, M., Uhlbäck, J., 2014. Geokemisk
atlas över Sverige (Geochemical Atlas of Sweden), Geological Survey of Sweden, 210 pp.;
http://doi.org/10.13140/2.1.1341.6642.

Bednar, A.J., Jones, W.T., Chappell, M.A., Johnson, D.R., Ringelberg, D.B., 2010. A modified acid
digestion procedure for extraction of tungsten from soil. Talanta 80, 1257-1263.

- Belzile, N., Chen, Y.-W., Wang, Z., 2001. Oxidation of antimony (III) by amorphous iron and
  manganese oxyhydroxides. Chemical Geology 174, 379-387.
- Bertrand, G., Cassard, D., Arvanitidis, N., Stanley, G., EuroGeoSurveys Mineral Resources Expert
  Group, 2016. Map of critical raw material deposits in Europe. Energy Procedia 97, 44-50;
  https://doi.org/10.1016/j.egypro.2016.10.016.
- 793 Birke, M., Reimann, C., Rauch, U., De Vivo, B., Halamić, J., Klos, V., Gosar, M., Ladenberger, A.,

2014a. Distribution of Cadmium in European Agricultural and Grazing Land Soil. Chapter 5 In: C.

Reimann, M. Birke, A. Demetriades, P. Filzmoser, P.O'Connor (Editors), Chemistry of Europe's

agricultural soils – Part B: General background information and further analysis of the GEMAS data

set. Geologisches Jahrbuch (Reihe B103), Schweizerbarth, Hannover, 89-115.

- Birke, M., Reimann, C., Fabian, K., 2014b. Analytical Methods Used in the GEMAS Project. Chapter
  5 In: C. Reimann, M. Birke, A. Demetriades, P. Filzmoser, P.O'Connor (Editors), Chemistry of
  Europe's agricultural soils Part A: Methodology and interpretation of the GEMAS data set.
  Geologisches Jahrbuch (Reihe B102), Schweizerbarth, Hannover, 41-46.
- Birke, M., Rauch, U., Stummeyer, J., 2015. How robust are geochemical patterns? A comparison of low
  and high sample density geochemical mapping in Germany. Journal of Geochemical Exploration
  154, 105-128.
- Birke, M., Reimann, R., Oorts, K., Rauch, U., Demetriades, A., Dinelli, E., Ladenberger, A., Halamić,
  J. Gosar, M., Jähne-Klingberg, F., The GEMAS Project Team, 2016. Use of GEMAS data for risk
  assessment of cadmium in European agricultural and grazing land soil under the REACH Regulation.
  Applied Geochemistry, 74, 109-121.
- Birke, M., Reimann, C., Rauch, U., Ladenberger, A., Demetriades, A., Jähne-Klingberg, F., Oorts, K.,
  Gosar, M., Dinelli, E., Halamic, J., The GEMAS Project Team, 2017. GEMAS: Cadmium
  distribution and its sources in agricultural and grazing land soil of Europe original data versus clr-
- transformed data. Journal of Geochemical Exploration, 173, 13-30.
- Bölviken, B., Kullerud, G., Loucks, R.R., 1990. Geochemical and metallogenic provinces: a discussion
  initiated by results from geochemical mapping across northern Fennoscandia. Journal of
  Geochemical Exploration, 39, 49-90; https://doi.org/10.1016/0375-6742(90)90069-M.
- Bowen, H.J.M., 1979. Environmental chemistry of the elements. Academic Press, London, New York,
  333 pp.

- Boyle, R.W., 1974. Elemental associations in mineral deposits and indicator elements of interest in
  geochemical prospecting. Energy, Mines and Resources Canada, Geological Survey Paper 74-45, 40
  pp.; https://doi.org/10.4095/102553.
- Boyle, R.W., Jonasson, I.R., 1984. The geochemistry of antimony and its use as an indicator element in
  geochemical prospecting. Journal of Geochemical Exploration 20, 223-302.
- 823 Bringezu, S., Potocnik, J., Schandl, H., Lu, Y., Ramaswami, A., Swillin, M., Suh, S., 2016. Multi-Scale
- 824 Governance of Sustainable Natural Resource Use—Challenges and Opportunities for Monitoring
  825 and Institutional Development at the National and Global Level. Sustainability 2016 8, 778, 1-25.
- Buccianti, A., Pawlowsky-Glahn, V., Mateu-Figueras, G. (Editors), 2006. Compositional data analysis
  in the geosciences: from theory to practice. Geological Society, London, 224 pp.
- 828 Buchert, M., Schüler, D., Bleher, D. 2009. Critical metals for future sustainable technologies and their
- 829 recycling potential. DTI/1202/PA, UNEP DTIE, Sustainable Consumption and Production Branch,
- 830 Paris, http://www.unep.fr/shared/publications/pdf/DTIx1202xPA831 Critical%20Metals%20and%20their%20Recycling%20Potential.pdf.
- Buschmann, J., Sigg, L., 2004. Antimony(III) Binding to Humic Substances: Influence of pH and Type
  of Humic Acid. Environmental Science & Technology 38, 4535-4541.
- Caritat, P. de, Cooper, M., 2011a. National Geochemical Survey of Australia: Data Quality Assessment,
  Geoscience Australia, Record 2011/21, GeoCat # 71971 vol. 1. Australian Government, Canberra,
  pp. 1-268.
- 837 Caritat, P. de, Cooper, M., 2011b. National Geochemical Survey of Australia: Data Quality Assessment,
  838 Geoscience Australia, Record 2011/21, GeoCat # 71971 vol. 2. Australian Government, Canberra,
  839 pp. 269-478.
- Caritat, P.de, Cooper, M., 2011c. National Geochemical Survey of Australia: The Geochemical Atlas of
  Australia, Geoscience Australia, Record 2011/20, GeoCat # 71973 vol. 1. Australian Government,
  Canberra, pp. 1-268.
- Caritat, P.de, Cooper, M., 2011d. National Geochemical Survey of Australia: The Geochemical Atlas
  of Australia, Geoscience Australia, Record 2011/20, GeoCat # 71973, vol. 2. Australian Government,
  Canberra, pp. 269-557.
- Cassard, D., Bertrand, G., Maldan, F., Gaàl, G., Kaija, J., Aatos, S., Angel, J.M., Arvanitidis, N., Ballas,
  D., Billa, M., Christidis, C., Dimitrova, D., Eilu, P., Filipe, A., Gazea, E., Inverno, C., Kauniskangas,
- E., Maki, T., Matos, J., Meliani, M., Michael, C., Mladenova, V., Navas, J., Niedbal, M., Perantonis,
- 849 G., Pyra, J., Santana, H., Serafimovski, T., Serrano, J.J., Strengell, J., Tasev, G., Tornos, F., Tudor,
- 850 G., 2012. ProMine pan-European mineral deposit database: a new dataset for assessing primary
- mineral resources in Europe. Workshop on: Mineral Resources Potential Maps: A Tool for
  Discovering Future Deposits. 12<sup>th</sup>-14<sup>th</sup> March 2012, Nancy, France, Proceedings, pp. 9–13.
- Cassard, D., Bertrand, G., Billa, M., Serrano, J.J., Tourlière, B., Angel, J.M., Gaál, G., 2015. ProMine
  mineral databases: New tools to assess primary and secondary mineral resources in Europe. Chapter

- 855 2 In: Weihed, P. (Editor), 3D, 4D and Predictive Modelling of Major Mineral Belts in Europe.
  856 Mineral Resource Reviews. Springer International Publishing, 9-58; https://doi.org/10.1007/978-3857 319-17428-0\_2.
- Charlier, P., Bou Abdallah, F., Bruneau, R., Jacqueline, S., Augias, A., Bianucci R., Perciaccante A.,
  Lippi D., Appenzeller O., Rasmussen, K.L., 2017. Did the Romans die of antimony poisoning? The
  case of a Pompeii water pipe (79 CE), Toxicology Letters 281, 184-186.
- 861 Cicchella, D., Lima, A., Birke, M., Demetriades, A., Wang, X., De Vivo, B., 2013. Mapping
  862 geochemical patterns at regional to continental scales using composite samples to reduce the
  863 analytical costs. Journal of Geochemical Exploration 124, 79-91.
- Cicchella, D., Albanese, S., Birke, M., De Vivo, B., De Vos, W., Dinelli, E., Lima, A., O'Connor, P.J.,
  Salpeteur, I., Tarvainen, T., 2014. Natural Radioactive Elements U, Th and K in European
  Agricultural and Grazing Land Soil. Chapter 8 In: C. Reimann, M. Birke, A. Demetriades, P.
  Filzmoser, P. O'Connor (Editors), Chemistry of Europe's agricultural soils Part B: General
  background information and further analysis of the GEMAS data set. Geologisches Jahrbuch (Reihe
  B103), Schweizerbarth, Hannover, 145-159.
- 870 Cristache, C., Comero, S., Locoro, G., Fissiaux, I., Ruiz, A.A., Tóth, G., Gawlik, B.M., 2014.
  871 Comparative study on open system digestion vs. microwave-assisted digestion methods for trace
  872 element analysis in agricultural soils. Joint Research Centre, Institute for Environment and
  873 Sustainability, Luxembourg, 38 pp.;
- 874 http://publications.jrc.ec.europa.eu/repository/bitstream/JRC90183/lb-na-26636-en-n.pdf
- Bernetriades, A., 2011. Understanding the quality of chemical data from the urban environment Part
  2: Measurement uncertainty in the decision-making process. Chapter 6 In: C.C. Johnson, A.
- Bernetriades, J. Locutura, R.T. Ottesen (Editors), Mapping the chemical environment of urban areas.
  John Wiley & Sons Ltd., Chichester, U.K., 77-98. https://doi.org/10.1002/9780470670071.ch6.
- Bernetriades, A., Reimann, C., 2014. Mineral deposits of Europe. Chapter 3 In: Reimann, C., Birke, M.,
  Demetriades, A., Filzmoser, P., O'Connor, P. (Eds.), Geologisches Jahrbuch (Reihe B 103).
  Schweizerbarth, Hannover, 71–78.
- Demetriades, A., Reimann, C., Filzmoser, P., 2014. Evaluation of GEMAS project quality control
  results. Chapter 6 In: C. Reimann, M. Birke, A. Demetriades, P. Filzmoser, P. O'Connor (Editors),
  Chemistry of Europe's agricultural soils Part A: Methodology and interpretation of the GEMAS
  data set Geologisches Jahrbuch (Reihe B102), Schweizerbarth, Hannover, 47-60.
- Be Vos, W., Tarvainen, T., Salminen, R., Reeder, S., De Vivo, B., Demetriades, A., Pirc, S., Batista,
  M.J., Marsina, K., Ottesen, R.T., O'Connor, P., Bidovec, M., Lima, A., Siewers, U., Smith, B.,
- 888 Taylor, H., Shaw, R., Salpeteur, I., Gregorauskiene, V., Halamić, J., Slaninka, I., Lax, K., Gravesen,
- 889 P., Birke, M., Breward, N., Ander, E.L., Jordan, G., Duris, M., Klein, P., Locutura, J., Bel-lan, A.,
- 890 Pasieczna, A., Lis, J., Mazreku, A., Gilucis, A., Heitzmann, P., Klaver, G., Petersell, V., 2006.
- 891 Geochemical Atlas of Europe. Part 2 Interpretation of Geochemical Maps, Additional Tables,

- Figures, Maps, and Related Publications. Geological Survey of Finland, Espoo, 618 pp.;
  http://weppi.gtk.fi/publ/foregsatlas/.
- ECS, 2010. Sludge, treated biowaste and soil Digestion for the extraction of aqua regia soluble fraction
  of trace elements. prEN 16174, CEN/TC400 (Draft). European Committee for Standardisation,
  Brussels, 15 pp.;
  https://www.ecn.nl/docs/society/horizontal/BT\_TF151\_WI\_CSS99025B\_Aqua\_regia\_2992007.pdf
- 898
- European Commission, 2017. Study on the review of the list of Critical Raw Materials: Criticality
  assessments. Brussels, 13.9.2017, COM(2017) 490 final; 92 pp.; http://doi.org/10.2873/876644.
- Fay, D., Kramers, G., Zhang, C., McGrath, D., Grennan, E. 2007. Soil Geochemical Atlas of Ireland.
  Teagasc, Environmental Protection Agency, Wexford, 117 pp.;
  https://www.teagasc.ie/media/website/publications/2011/Soil\_Geochemical\_AtlasofIreland.pdf.

Filzmoser, P., Reimann, C., Birke, M., 2014. Univariate Data Analysis and Mapping. Chapter 8 In: C.
Reimann, M. Birke, A. Demetriades, P. Filzmoser, P. O'Connor (Editors), Chemistry of Europe's

agricultural soils – Part A: Methodology and interpretation of the GEMAS data set. Geologisches
Jahrbuch (Reihe B102), Schweizerbarth, Hannover, 67-81.

- Gál, J., Hursthouse, A., Cuthbert, S. 2007. Bioavailability of arsenic and antimony in soils from an
  abandoned mining area, Glendinning (SW Scotland). Journal of Environmental Science and Health,
- 910 Part A Toxic/Hazardous Substances and Environmental Engineering 42, 1263-1274.
- Garrett, R.G., Reimann, C., Smith, D.B., Xie, X., 2008. From geochemical prospecting to international
  geochemical mapping: a historical overview. Geochemistry: Exploration Environment Analysis 8,
  205-217.
- Günther, A., Reichenbach, P., Malet, J.-P., Van Den Eeckhaut, M., Hervás, J., Dashwood, C. and
  Guzzetti, F. 2013. Tier-based approaches for landslide susceptibility assessment in Europe
  Landslides 10, 529-546.
- Hockmann, K., Schulin, R., 2013. Leaching of Antimony from Contaminated Soils. In: Selim, H.M.
  (Ed.), Competitive Sorption and Transport of Heavy Metals in Soils and Geological Media. CRC
  Press, Boca Raton, 119-145.
- Hu, Z., Gao, S., 2008. Upper crustal abundances of trace elements: A revision and update. Chemical
  Geology 253, 205–221.
- Hubbert, M.K., 1982. Techniques of Prediction as Applied to Production of Oil and Gas. In: Gass, S.I.
  (Editor), Oil and Gas Supply. US Department of Commerce, NBS Special Publication 631, May
  1982.
- Johnson, C.A., Moench, H., Wersin, P., Kugler, P., Wenger, C., 2005. Solubility of Antimony and Other
  Elements in Samples Taken from Shooting Ranges. Journal of Environmental Quality 34, 248-254.
- 927 Jordan, G., Petrik, A., De Vivo, B., Albanese, S., Demetriades, A., Sadeghi, M., The GEMAS Project
- 928 Team, 2018. GEMAS: Spatial analysis of the Ni distribution on a continental-scale using digital

- 929 image processing techniques on European agricultural soil data. Journal of Geochemical930 Exploration186, 143-157.
- Kabata-Pendias, A., Mukherjee, A.B., 2007. Trace Elements from Soil to Human. Springer Verlag,
  Berlin, Heidelberg, 550 pp.
- Koutsospyros, A., Braida, W., Christodoulatos, C., Dermatas, D., Strigul, N. 2006. A review of tungsten:
  From environmental obscurity to scrutiny. Journal of Hazardous Materials 136, 1-19.
- 935 Ladenberger, A., Andersson, M., Reimann, C., Tarvainen, T., Filzmoser, P., Uhlbäck, J., Morris, G.,
- Sadeghi , M. 2013. Geochemical mapping of agricultural soils and grazing land (GEMAS) in
  Norway, Finland and Sweden regional report. Geological Survey of Sweden, SGU-rapport
  2012:17, 160 pp.; http://resource.sgu.se/produkter/sgurapp/s1217-rapport.pdf;
  https://www.researchgate.net/publication/260872161\_Geochemical\_mapping\_of\_agricultural\_soils
  and\_grazing\_land\_GEMAS\_in\_Norway\_Finland\_and\_Sweden\_-\_regional\_report.
- Ladenberger, A., Demetriades, A., Reimann, C., Birke, M., Sadeghi, M., Uhlback, J., Andersson, M.,
  Jonsson, E., the GEMAS Project Team. 2015. GEMAS: Indium in agricultural and grazing land soil
  of Europe its source and geochemical distribution patterns. Journal of Geochemical Exploration
- 944 154, 61–80.
- Ladenberger, A., Uhlbäck, J., Andersson, M., Reimann, C., Tarvainen, T., Morris, G., Sadeghi, M.,
  Eklund, M., Filzmoser, P., 2014. Elemental Patterns in Agricultural and Grazing Land soil in
  Norway, Finland and Sweden: What have we learned from continental-scale mapping? Chapter 14
- In: C. Reimann, M. Birke, A. Demetriades, P. Filzmoser, P. O'Connor (Editors), Chemistry of
  Europe's agricultural soils Part B: General background information and further analysis of the
  GEMAS data set. Geologisches Jahrbuch (Reihe B103), Schweizerbarth, Hannover, 235-251.
- Barbert, I., Durrheim, R., Godoy, M., Kota, M., Leahy, P., Ludden, J., Nickless, E., Oberhaensli, R.,
- Anjian, W., Williams, N. 2013. Resourcing Future Generations: A proposed new IUGS initiative.
  Episodes 36, 82-86.
- Locutura, J., Bel-lan, A., García, C.A., Martínez, S., 2012. Atlas Geoquímico de España. Instituto
  Geológico y Minero de España, Madrid, 592 pp. (A3 size);
  http://www.igme.es/actividadesIGME/lineas/CartoGeo/geoquimica/geoquimicaIng.htm.
- Meinert, L.D., Robinson, G.R., Nassar, N.T., 2016. Mineral resources: reserves, peak production and
   the future. Resources 5, 14.
- Millot R., Vigier, N., Gaillardet, J., 2010. Behaviour of lithium and its isotopes during weathering in the
   Mackenzie Basin, Canada. Geochimica et Cosmochimica Acta 74, 3897–3912.
- 961 Négrel, Ph., Sadeghi, M., Ladenberger, A., Reimann, C., Birke, M., the GEMAS Project Team. 2015.
- 962 Geochemical fingerprinting and sources discrimination in soils and sediments at continental scale.
- 963 Chemical Geology 396, 1-15.

- 964 Négrel, Ph., Ladenberger, A., Reimann, C., Birke, M., Sadeghi, M., the GEMAS Project Team. 2016.
  965 GEMAS: source, distribution patterns and geochemical behavior of Ge in agricultural and grazing
  966 land soils at European continental scale. Applied Geochemistry 72, 113-124.
- 967 Négrel, Ph., De Vivo, B., Reimann, C., Ladenberger, A., Cicchella, D., Albanese, S., Birke, M., De Vos,
- 968 W., Dinelli, E., Lima, A., O'Connor, P.J., Salpeteur, I., Tarvainen, T., the GEMAS Project Team,
- 969 2018a. U-Th signatures of agricultural soil at the European continental scale (GEMAS): Distribution,
- 970 weathering patterns and processes controlling their concentrations. Science of the Total Environment
- 971 622-623, 1277-1293.
- 972 Négrel, Ph., Ladenberger, A., Reimann, C., Birke, M., Sadeghi, M., the GEMAS Project Team., 2018b.
  973 Distribution of Rb, Ga and Cs in agricultural land soils at European continental scale (GEMAS):
  974 Implications for weathering conditions and provenance. Chemical Geology 479, 188-203.
- 975 Nickless, E., Ali, S., Arndt, N., Brown, G., Demetriades, A., Durrheim, R., Enriquez M.A., Giurco, D.,
- Kinnaird, J., Littleboy, A., Masotti, F., Meinert, L., Nyanganyura, D., Oberhänsli, R., Salem, J.,
  Schneider, G., Yakovleva., N., 2015. Resourcing Future Generations: A Global Effort to Meet The
  World's Future Needs Head-on. International Union of Geological Sciences, 76 pp.;
- 979 http://iugs.org/uploads/RFG%20Report-sm.pdf.
- Ottesen, R.T., Birke, M., Finne, T.E., Gosar, M., Locutura, J., Reimann, C., Tarvainen, T., the GEMAS
  Project team. 2013. Mercury in European agricultural and grazing land soils. Applied Geochemistry
  33, 1-12.
- Pawlowsky-Glahn, V., Buccianti, A., 2011. Compositional data analysis: theory and applications.
  Wiley, Chichester, 378 pp.
- Plant, J.A., Breward, N., Forrest, M.D., Smith, R T., 1989. The gold pathfinder elements As, Sb and Bi:
  their distribution and significance in the south-west Highlands of Scotland. Transactions of the
  Institution of Mining & Metallurgy (Section B: Applied earth science) 98, B91-B101.
- Poňavič, M., Scheib, A., 2014. Distribution of Selenium in European Agricultural and Grazing Land
  Soil. Chapter 7 In: C. Reimann, M. Birke, A. Demetriades, P. Filzmoser, P. O'Connor (Editors),
  Chemistry of Europe's agricultural soils Part B: General background information and further
  analysis of the GEMAS data set. Geologisches Jahrbuch (Reihe B103), Schweizerbarth, Hannover,
  131-144.
- Rawlins, B.G., McGrath, S.P., Scheib, A.J., Breward, N., Cave, M., Lister, T.R., Ingham, M., Gowing,
  C., Carter, S., 2012. The advanced soil geochemical atlas of England and Wales. British Geological
- 995 Survey, Nottingham, UK, 227 pp.
- 996 Reimann, C., Siewers, U., Tarvainen, T., Bityukova, L., Eriksson, J., Giucis, A., Gregorauskienė, V.,
- 997 Lukashev, V.K., Matinian, N.N., Pasieczna, A., 2003. Agricultural Soils in Northern Europe A
- 998 Geochemical Atlas (Baltic Soil Survey), Geologisches Jahrbuch Sonderhefte Reihe D, Heft 5.
- 999 Schweizerbart Science Publishers, Stuttgart, 279 pp.

- Reimann, C., Arnoldussen, A., Englmaier, P., Filzmoser, P., Finne, T.E., Garrett, R.G., Koller, F.,
   Nordgulen, Ø., 2007a. Element concentrations and variations along a 120 km long transect in south
   Norway anthropogenic vs. geogenic vs. biogenic element sources and cycles. Applied
   Geochemistry 22, 851–871.
- Reimann, C., Melezhik, V., Niskavaara, H., 2007b. Low-density regional geochemical mapping of gold
  and palladium highlighting the exploration potential of northernmost Europe. Economic Geology,
  1006 102, 327-334.
- 1007 Reimann, C., Demetriades, A., Eggen, O.A., Filzmoser, P., The EuroGeoSurveys Geochemistry Expert 1008 Group, 2009. The EuroGeoSurveys geochemical mapping of agricultural and grazing land soils project (GEMAS) - Evaluation of quality control results of aqua regia extraction analysis. Geological 1009 1010 Survey of Norway, Trondheim, NGU report 2009.049. 94 pp. 1011 http://www.ngu.no/upload/Publikasjoner/Rapporter/2009/2009\_049.pdf.
- Reimann, C., Matschullat, J., Birke, M., Salminen, R. 2010. Antimony in the environment: Lessons from
  geochemical mapping. Applied Geochemistry 25, 175–198.
- 1014 Reimann, C., Demetriades, A., Eggen, O.A., Filzmoser, P., EuroGeoSurveys Geochemistry Working
- 1015 Group, 2011. The EuroGeoSurveys geochemical mapping of agricultural and grazing land soils
- 1016 project (GEMAS) Evaluation of quality control results of total C and S, total organic carbon (TOC),
- 1017 cation exchange capacity (CEC), XRF, pH, and particle size distribution (PSD) analysis. NGU
- 1018Report2011.043.GeologicalSurveyofNorway,Trondheim,92pp.;1019http://www.ngu.no/upload/Publikasjoner/Rapporter/2011/2011\_043.pdf.
- Reimann, C., Caritat, P. de, GEMAS Project Team, NGSA Project Team, 2012a. New soil composition
  data for Europe and Australia: Demonstrating comparability, identifying continental-scale processes
  and learning lessons for global geochemical mapping. The Science of the Total Environment 416,
  239-252.
- Reimann, C., Flem, B., Fabian, K., Birke, M., Ladenberger, A., Négrel, Ph., Demetriades, A.,
  Hoogewerff, J., the GEMAS Project Team, 2012b. Lead and lead isotopes in agricultural soils of
  Europe the continental perspective. Applied Geochemistry 27, 532–542.
- Reimann, C., Demetriades, A., Birke, M., Eggen, O. A., Filzmoser, P., Kriete, C., EuroGeoSurveys
  Geochemistry Expert Group, 2012c. The EuroGeoSurveys Geochemical Mapping of Agricultural
  and grazing land Soils project (GEMAS) Evaluation of quality control results of particle size
- 1030 estimation by MIR prediction, Pb-isotope and MMI<sup>®</sup> extraction analyses and results of the GEMAS
- 1031 ring test for the standards Ap and Gr. Geological Survey of Norway, Trondheim, NGU report
- 1032 2012.051, 136 pp. http://www.ngu.no/upload/Publikasjoner/Rapporter/2012/2012\_051.pdf.
- Reimann,C., Filzmoser, P., Fabian, K., Hron, K., Birke, M., Demetriades, A., Dinelli, E., Ladenberger,
  A., the GEMAS Project Team, 2012d. The concept of compositional data analysis in practice Total
  major element concentrations in agricultural and grazing land soils of Europe. Science of the Total
- 1036 Environment, 426, 196–210.

- 1037 Reimann, C., Birke, M., Demetriades, A., Filzmoser, P., O'Connor, P. (Editors), 2014a. Chemistry of
  1038 Europe's agricultural soils Part A: Methodology and interpretation of the GEMAS dataset.
  1039 Geologisches Jahrbuch (Reihe B), Schweizerbarth, Stuttgart, 528 pp.
- Reimann, C., Birke, M., Demetriades, A., Filzmoser, P., O'Connor, P. (Editors), 2014b. Chemistry of
   Europe's agricultural soils Part B: General background information and further analysis of the
   GEMAS dataset. Geologisches Jahrbuch (Reihe B), Schweizerbarth, Stuttgart, 352 pp.
- 1043 Reimann, C., Ladenberger, A., Birke, M., Caritat, P. de, 2016. Low density geochemical mapping and
- mineral exploration: application of the mineral system concept. Geochemistry: Exploration
   Environment Analysis 16, 48-61
- Roda-Robles, E., Pesquera, A. Gil-Crespo, P.P., Vieira, R., Lima A., Garate-Olave, I., Martins, T.,
  Torres-Ruiz, J., 2016. Geology and mineralogy of Li mineralization in the Central Iberian Zone
  (Spain and Portugal). Mineralogical Magazine 80, 103–126.
- Sadeghi, M., Petrosino, P., Ladenberger, A., Albanese, S., Andersson, M., Morris, G., Lima, A.M., De
  Vivo, B., the GEMAS Project Team, 2013. Ce, La and Y concentrations in agricultural and grazingland soils of Europe. Journal of Geochemical Exploration 133, 202-213.
- 1052 Salminen, R., Tarvainen, T., Demetriades, A., Duris, M., Fordyce, F.M., Gregorauskiene, V., Kahelin,
- 1053 H., Kivisilla, J., Klaver, G., Klein, P., Larson, J.O., Lis, J., Locutura, J., Marsina, K., Mjartanova, H.,
- 1054 Mouvet, C., O'Connor, P., Odor, L., Ottonello, G., Paukola, T., Plant, J.A., Reimann, C., Schermann,
- 1055 O., Siewers, U., Steenfelt, A., Van Der Sluys, J., Williams, L., 1998. FOREGS Geochemical
  1056 Mapping Field Manual. Geological Survey of Finland, Espoo, Guide 47, 36 pp.;
  1057 http://tupa.gtk.fi/julkaisu/opas/op 047.pdf.
- 1058 Salminen, R., Batista, M.J., Bidovec, M., Demetriades, A., De Vivo, B., De Vos, W., Ďuriš, M., Gilucis,
- 1059 A., Gregorauskienė, V., Halamić, J., Heitzmann, P., Lima, A., Jordan, G., Klaver, G., Klein, P., Lis,
- 1060 J.z., Locutura, J., Marsina, K., Mazreku, A., O'Connor, P., Olsson, S.Å., Ottesen, R.T., Petersell, V.,
- 1061 Plant, J.A., Reeder, S., Salpeteur, I., Sandström, H., Siewers, U., Steenfelt, A., Tarvainen, T., 2005.
- Geochemical atlas of Europe. Part 1 Background information, methodology and maps. Geological
   Survey of Finland, Espoo, 525 pp.; http://weppi.gtk.fi/publ/foregsatlas/.
- Scheib, A.J, Flight, D.M.A., Birke, M., Tarvainen, T., Locutura, J., GEMAS Project Team, 2012. The
  geochemistry of niobium and its distribution and relative mobility in agricultural soils of Europe.
  Geochemistry: Exploration Environment Analysis 12, 293-302.
- Smith, D.B., Reimann, C. 2008. Low-density geochemical mapping and the robustness of geochemical
   patterns. Geochemistry: Exploration Environment Analysis 8, 219–227.
- Smith, D.B., Cannon, W.F., Woodruff, L.G., Solano, F., Kilburn, J.E., Fey, D.L., 2013. Geochemical
  and Mineralogical Data for Soils of the Conterminous United States, Data Series 801. U.S.
  Geological Survey, Reston, 19 pp.

- Smith, D.B., Cannon, W.F., Woodruff, L.G., Solano, F., Ellefsen, K.J., 2014. Geochemical and
  Mineralogical Maps for Soils of the Conterminous United States, Open-File Report 2014–1082. U.S.
  Geological Survey, Denver, 386 pp.
- Takahashi, T., Shozugawa, K., Matsuo, M., 2010. Contribution of Amorphous Iron Compounds to
   Adsorptions of Pentavalent Antimony by Soils. Water, Air, and Soil Pollution 208, 165-172.
- 1077 Tarvainen, T., Albanese, S., Birke, M., Ponavic, M. Reimann, C., the GEMAS Project Team, 2013.
  1078 Arsenic in agricultural and grazing land soils of Europe. Applied Geochemistry 28, 2-10.
- 1079 Tardy, Y., Krempp, G., Trauth, N., 1972. Le lithium dans les minéraux argileux des sédiments et des
  1080 sols. Geochimica et Cosmochimica Acta 36, 397-412.
- Teske, S. (Editor), 2019. Achieving the Paris Climate Agreement Goals: Global and Regional 100%
  Renewable Energy Scenarios with Non-energy GHG Pathways for +1.5°c and +2°C. Springer Open,
  491 pp.; https://doi.org/10.1007/978-3-030-05843-2.
- Tóth, G., Hermann, T., Szatmári, G., Pásztor, L., 2016. Maps of heavy metals in the soils of the European
  Union and proposed priority areas for detailed assessment. Science of The Total Environment 565,
  1054-1062.
- UNEP, 2009. Critical metals for future sustainable technologies and their recycling potential. UNEP,
   DTI/1202/PA, Paris, 81 pp.; http://www.unep.fr/shared/publications/pdf/DTIx1202xPA Critical%20Metals%20and%20their%20Recycling%20Potential.pdf.
- Wedepohl, K.H., (Exec. Ed.), 1978. Handbook of Geochemistry. Volume II/5. Springer-Verlag, Berlin,
  Heidelberg, New York, 4400+ pp.
- Xie, X., Ren, T., Sun, H., 2012. Geochemical Atlas of China. Institute of Geophysical and Geochemical
   Exploration, Langfang, China, 135 pp.
- Zeeberg, J.J., 1998. The European sand belt in Eastern Europe and comparison of Late Glacial dune
   orientation with GCM simulation results. Boreas 27, 127–139.

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