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- Title: Revisiting herbage sample collection and preparation procedures to minimise risks of
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# Title: Revisiting herbage sample collection and preparation procedures to minimise risks of trace element contamination

34

#### 35 Abstract

A renewed interest in trace elements (TE), as micronutrients as well as potentially toxic 36 elements, and new options for multi-element analysis has led to an increased number of 37 scientists engaging in TE studies. Accreditation, certification and quality control of TE 38 39 analyses often applies only to the last step in the sample chain when prepared samples are sent 40 to the laboratory for digestion/extraction and subsequent analysis. However, all stages of the chain from initial sampling to final analysis require an understanding of the specific 41 challenges involved in TE studies and an awareness of the contamination risks as well as 42 approaches to limit these. Contamination can potentially be introduced during all stages of 43 handling and preparation of plant samples, e.g. through dust and the materials that make up 44 the different work surfaces, tools and containers used. Milling devices originally used during 45 preparation of two sets of archived herbage samples were tested to indicate the degree of 46 47 contamination that can arise from milling. For example, some of the milling devices tested showed effects on several TE concentrations while also increasing the variability between 48 samples. A titanium knife mill which was included for comparison gave the best results, 49 showing no measurable contamination by TE of primary interest, while it allowed a high 50 throughput of samples. To enhance the quality of data on TE in bulky plant material such as 51 herbage and to ensure future usability of newly archived samples, we suggest that field 52 handbooks and sample preparation protocols (where needed) are revised to include 53

precautions against TE contamination in all handling steps. This will ensure reliable data on
concentrations of micronutrients and potential toxic TE in plant material.

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57 Keywords: micronutrient, plant sample, sample drying, sample milling, sample storage

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#### 59 Introduction

60 Over recent years, there has been a renewed interest in trace elements (TE) from various perspectives including: agronomic requirements, feed/food quality, and the environmental 61 impacts of potentially toxic elements (e.g. Alloway, 2008; Stein, 2010; Cooper et al., 2011; 62 Tidemann-Andersen et al., 2011). Technical advances in analytical equipment and preparation 63 procedures have opened up new possibilities for including comprehensive multi-element 64 analyses of soil and plant samples. Such analyses are frequently carried out on newly 65 collected samples, but there is also interest in re-analysing archived samples, e.g. samples 66 67 from surveys and monitoring studies as well as long-term field experiments. These samples 68 may have been originally collected for a specific purpose, but they now have a key role in the study of time trends for a range of elements or to pursue new research questions beyond the 69 scope of the original sampling programme. This change in emphasis has led to an increase in 70 71 the number of researchers involved in TE studies. In the past, specialists with their own rigorous procedures and analytical equipment determined TE in studies specifically designed 72 to secure high quality data. More recently, however, it is less common to find the same people 73 responsible for the whole chain from sampling through to analysis. The new generation of 74 scientists often have a primary focus other than TE per se and may not be familiar with 75 76 practical aspects relating to TE research, especially contamination risks.

Trace element analyses of plant material pose specific demands with regard to sampling,
sample preparation and pre-treatment. There are various potential sources of contamination
which include soil and the equipment used for the different processing stages. Various aspects
of uncertainties and errors along the whole sampling, sample preparation and analysis
sequence were discussed during a workshop on 'Improvements of trace element in plant
matrices' held in Brussels in May 1994 (Quevauviller, 1995).

If samples from studies that were originally designed and undertaken with a different focus 83 are to be reused for contemporary TE studies, the potential risks for TE contamination must 84 be evaluated and the consequences this might pose for archived material assessed. It is 85 therefore appropriate to revisit some of the issues associated with such TE studies, 86 87 particularly for the benefit of researchers who are relatively new to the research subject. This is supported by the fact that out of the ten most recent papers on micronutrients or TE in 88 herbage found during a search of Web of Knowledge only one paper clearly stated 89 90 precautions against contamination (Smith et al., 2009), whereas in the remaining nine papers either no reference was made to this or the described methods indicate that contamination was 91 likely. 92

The overall aim of this paper is to provide an overview of risk of contamination from sources 93 associated with herbage sample collection and preparation. The overview is based on a 94 95 literature review complemented by examples from our own laboratories to demonstrate the 96 issues. Literature searches of peer-reviewed publications, and other sources such as conference publications, reports and field protocols were thus undertaken with keywords that 97 included, but were not limited to, sample collection, sample storage, sample preservation, 98 99 sample preparation, milling and TE contamination. References were reviewed with a special 100 focus on herbage samples.

101

## Sample handling and preparation of herbage samples to avoid TE contamination

Published scientific literature was generally focused on the individual steps in the sample 104 105 collection or preparation chain and also included other aspects of each step beyond risks of 106 TE contamination, e.g. procedures to ascertain collection of representative samples (Table 1). Notable exceptions were a special issue reporting on the 1994 workshop 'Improvements of 107 trace element in plant matrices' (STOTEN, 1995), and two publications from the early 1970's 108 109 (Scott et al., 1971; Scott and Ure, 1972). Protocols for sample collection and preparation for use by field staff was generally found in 'grey' literature. Sample collection and preparation 110 protocols to minimise TE contamination have, for example, been published for a range of 111 grain and tuber crops and for plantains and bananas (Stangoulis and Sison, 2008), but other 112 protocols do not always include considerations of TE (e.g. Försökshandboken, 2009). Sample 113 handling procedures to prevent accidental contamination are also mentioned as an important 114 aspect when implementing the EC Directive concerning the performance of sampling and 115 116 analysis for the official control of different substances (including some TE) in foodstuffs 117 (European Commission, 2007), although few practical directions are given.

The 1994 workshop on the state of the art of TE determinations in plant matrices summarised the most crucial aspects of plant material sampling, preparation, pre-treatment and detection (Quevauviller, 1995). However, the discussion covered all possible types of plant matrices, and as a result conclusions and recommendations were very general, pointing out the need for adjustment of procedures in relation to the aims and objectives of each individual study.

Mixed species herbage samples involve special challenges during collection and sample 123 preparation as the major part of the above-ground plant material is collected, potentially 124 giving rise to highly heterogeneous samples. The heterogeneity of herbage materials 125 126 emphasises the importance of extracting a representative sample both at the time of collection and also subsequent preparation stages, together with the need for herbage sample 127 homogenisation. In the following text, the recommendations and conclusions from the 1994 128 workshop (STOTEN, 1995) will hence be revisited and developed specifically for herbage 129 and with the aim of illustrating the need for overall quality assurance in TE studies of herbage 130 and other bulky crops. 131

Trace element studies demand rigorous protocols to avoid contamination during sample 132 133 collection and preparation. Dust evolving from soil and plant material and other incidental sources constitutes a potential contamination risk and obviously calls for a high standard of 134 hygienic maintenance of rooms and equipment used during sample preparation. It follows that 135 work areas and equipment used for plant material processing should be kept separate from 136 those used for soil processing. Work facilities should also be designed to give a minimum and 137 138 predictable level of contamination, e.g. by the use of impermeable surfaces (Hamilton, 1995). Equipment should be stored in closed containers when not in use to protect it and the test 139 materials from dust (Stangoulis and Sison, 2008). Samples may also become contaminated 140 from the surfaces of containers and tools (e.g. metals, paints, tanned leather, rubber) 141 (Lockman, 1980; Fleming et al., 1986; Stangoulis and Sison, 2008). Tools, containers and 142 procedures used throughout the various stages should therefore be chosen with care. Further 143 144 potential TE contamination sources during different stages of the sample chain are transfers from metal structures and from skin-care products via hands (Stangoulis and Sison, 2008). 145

#### 146 Sample collection

147 Factors such as sample collection strategy, plant species identification, and collection of consistent proportions between plant parts is of importance for acquiring representative 148 149 samples (e.g. Ernst, 1995; Wagner, 1995) and avoiding erroneous and highly variable results. For herbage sample collection, a standardised cutting height some distance above the soil 150 151 surface not only decreases variability in sample composition but also decreases the risk of soil contamination. Risks of contamination by soil and dust during growth or sampling have been 152 recognised for decades and recommendations issued to minimise it; including avoiding 153 sampling after high winds, heavy rains and prolonged drought, and waiting to sample until at 154 155 least two weeks after grazing (Scott et al., 1971). Soil or dust contamination is obviously most critical for elements where concentrations are much higher than the corresponding plant 156 concentrations: most notably cobalt (Co), chromium (Cr) and iron (Fe), but also copper (Cu), 157 158 zinc (Zn) and boron (B) (Fleming et al., 1986; Wyttenbach and Tobler, 1998). As part of quality assurance procedures, indicators of soil contamination (e.g. aluminium (Al), Fe, 159 160 titanium (Ti) or scandium (Sc)) should thus be observed (Scott et al., 1971; Bargagli, 1995; Wyttenbach and Tobler, 2002; Elias et al., 2008; Cook et al., 2009). 161

Procedures for counteracting sample contamination by soil and dust through picking, 162 brushing, and washing of samples have been developed (Porter, 1986; Markert, 1992; Aboal 163 164 et al., 2008; Elias et al., 2008) and can, to some extent, counteract differences over the year in the magnitude of contamination by dust. Apart from this, it has been shown that variation 165 between repeated samplings may be decreased by sampling under similar weather conditions, 166 167 as well as using similar storage times and storage conditions before sample cleaning (Fernández et al., 2010). Washing of plant material may lead to losses of TE from inside the 168 cells though, the magnitude increasing if unfavourable ratios between solvent and plant 169 material or long washing times are applied (Markert, 1992; Rossini Oliva and Raitio, 2003). A 170

summary of different washing techniques and recommendations for when to apply them is 171 172 given by Rossini Oliva and Raitio (2003). However, cleaning of samples is not always recommended. For example, where the aim is to study the contribution of atmospheric derived 173 174 'contamination' or actual intake by livestock, then a direct analysis of unwashed material would be required. Sampling of lodged herbage should be avoided, though, if at all possible. 175 Plant concentrations of TE are influenced by soil factors, hydrological conditions, plant 176 177 species, phenological stage and plant part, and ley/pasture management (Mayland and Sneva, 178 1983; Anke et al., 1994; Belesky et al., 2000; Fystro and Bakken, 2005; Sinclair and Edwards, 2008; Roche et al., 2009). Hence it is important to use the same sampling protocol on every 179 180 occasion and, unless corresponding soil samples are collected, at least notes of the soil and 181 hydrological conditions, farm management and signs of herbivory and pathogen infestation, should be taken. Examples of such protocols are given by Ernst (1995) and Hamilton (1995) 182 and may be adapted to suit herbage sampling. 183

184

#### 185 Sample drying and storage

Herbage is generally bulky and heterogenous and large samples are needed to attain a
representative sample. Hence the freeze-drying procedures recommended by Hamilton (1995)
for preparing plant material prior to TE analysis are generally only applicable when large
capacity freeze driers are available. Instead forced ventilation drying ovens are frequently
used. It is important that driers (and surroundings) are cleaned thoroughly before use and that
separate driers are used for soil and plant material. Also sample bags (e.g. perforated plastic
bags) should be clean.

If herbage samples are stored prior to further preparation or stored after milling, containers for 193 194 storage should similarly be clean (e.g. new or acid washed) and samples stored in a dry and clean environment. The composition of storage containers is also important. Glass containers 195 196 work well in many ways, but may contaminate samples with B from the glass or other elements from the closures. Some TE are further used in colouring of e.g. plastics and are also 197 found as likely traces from the manufacturing process (Waheed et al., in press) and may be 198 released into the samples. Details on drying and storage of samples are given by Lockman 199 200 (1980), Houba et al. (1995), Quevauviller (1995) and Stangoulis and Sison (2008).

201

#### 202 Sample milling

Creation of a representative subsample is a crucial step in all analytical work and the 203 204 homogenisation frequently needed for this can be the most risky step with regard to 205 contamination, in particular if the plant material contains mineral particles which are likely to 206 abrade grinding equipment (Hamilton, 1995). To avoid the risk of contamination from the mill, samples of e.g. grains in some laboratories are not milled but digested as whole grain 207 208 (e.g. Öborn et al., 1995; Wångstrand et al., 2007). Where whole grains are used it is important 209 to ensure representativeness by using larger sample weights and digestion volumes compared with standard procedures. For bulky samples of heterogeneous material such as whole 210 herbage samples, however, it is not possible to avoid particle size reduction by milling or 211 212 grinding prior to homogenisation and extraction of a representative subsample. A number of mill types made from materials low in TE (Hamilton, 1995; Markert, 1995) are used for 213 grinding small sample sizes, but larger samples still present difficulties as many agate and 214 ceramic mortar mills are suitable only for smaller sample sizes. Mills generally used for the 215 preparation of larger, fibrous samples of varying hardness are cutting mills and hammer mills. 216

These are most often made from steel with TE as major constituents or as minor components and thus likely to introduce these elements into the samples through wear. Use of reference materials is of little help in the quality control of this step as these are generally already milled (or otherwise fine powder) and thus will not be milled or ground in the laboratory along with the material to be analysed. Reference materials consequently constitute only a quality control for onward steps in the analysis and not for all stages during sample preparation.

223 The European Commission (2007) regulation for the methods of sampling and analysis for the 224 official control of the levels of some TE in foodstuffs, states that the analyst should ensure that samples do not become contaminated during sample preparation. According to their 225 226 recommendations devices should be of inert materials such as polypropylene or 227 polytetrafluoroethylene, but high quality stainless steel is (surprisingly) permitted for cutting 228 edges. However, Cubadda et al. (2001) tested a range of milling devices (glass and porcelain 229 mortars, and four steel mills) and revealed significant contamination by all the tested devices 230 with one or several TE. Statistical differences with respect to the control were thus detected for ten TE (Al, cadmium (Cd), Co, Cr, Cu, Fe, manganese (Mn), molybdenum (Mo), nickel 231 232 (Ni), and lead (Pb)). The contamination was found to be higher with hard durum wheat than when softer wheat was milled, indicating that the scope of the contamination risk may differ 233 depending on the hardness of the material to be milled. On the other hand, Sager and 234 Mittendorfer (1997) did not find any significant difference between continuously and 235 discontinuously operating milling devices, nor did they find any significant differences in the 236 efficiency of different cleaning methods (washing, blowing, brushing and discarding of the 237 238 first milled portion).

### A practical test of different cutting and milling devices for preparation of herbage samples

241 Materials and methods

242

243 our institutions (SLU, SAC) were tested in two experiments comprising a) a steel hammer 244 mill, a steel hammer mill followed by ball mill, and a Ti knife mill using a plexi-glass knife as a control (Test 1), and b) a steel cutting mill using plastic scissors as a control (Test 2) (Table 245 246 2). The plant material used for Test 1 was mature mixed hay consisting of perennial ryegrass (Lolium perenne L.), timothy (Phleum pratense L.) and white clover (Trifolium repens L.), 247 248 and for Test 2 timothy harvested at the emerging ear stage. For each experiment, the plant material was split into equivalent weight subsamples which were randomised with five 249 replicates being processed by each cutting or milling device. 250 Digestion of the plant material was carried out according to the procedures developed and 251 routinely used in the laboratory of the Department of Soil and Environment, Swedish 252 University of Agricultural Sciences. 253

Mills that had been used during preparation of potentially useful archived samples available in

Day 1: One gram plant sample was weighed into acid-washed Tecator glass tubes (Höganäs,
Sweden). Ten ml conc. (15.6 M) HNO<sub>3</sub> (Merck suprapure) was added and the sample, covered
with a glass pear, incubated in the Tecator blocks at 30 °C for 9.5 h, followed by 100 °C for 1
h, and 135 °C for 1.5 h.

258 Day 2: When cooled to approx. 70 °C, the tubes were removed from the Tecator blocks, and 259 another 5 ml conc. HNO<sub>3</sub> was added, where-after incubation was resumed at 135 °C for 260 another 2.5 h.

Filter papers (Munktell 00H, Ø185 mm) were washed twice with 10% (1.56 M) HNO<sub>3</sub>. The

digests were diluted to a total volume of 100 ml with ultrapure water (maximum  $0.055\mu S$  cm<sup>-</sup>

<sup>1</sup>) and then filtered directly into acid washed plastic bottles.

264	The digests were analysed for Cd, Co, Cr, Cu, Fe, Mn, Mo, Ni, Pb and Zn by inductively
265	coupled plasma mass spectrometry (Elan 6100 ICP-MS; Perkin Elmer SCIEX instruments).
266	Dry matter content in the plant material was determined and metal concentrations expressed
267	in mg kg <sup>-1</sup> plant material dry weight (dw).

Certified reference material (NIST Wheat Flour, National Institute of Standards and
Technology, Gaitersburg, MD, USA) was included in all batches. There were no values for Cr
or Ni concentrations provided with the certified reference material and therefore the in-house
average of the NIST material was used as test values for these two elements. Detection and
analytical limits were calculated from the composition of 10 blanks with the detection limit
set to 3×standard deviation and analytical limits to 10×standard deviation for each element.

Differences due to the milling devices were evaluated through ANOVA using JPM 8.0.1
software (SAS, Cary, NC, USA) using ln transformed data to get a normal distribution of
residuals where appropriate (Zn in Test 1, and Cr in Test 2). Where ANOVA indicated
significant differences (P<0.05) between means, the effects of individual devices were tested</li>
by Tukey's HSD.

279 Results and conclusions

In the first of the current tests of mills the pattern of contamination relative to controls for 10
elements fell into three broad groups. In the group which included Cd, Co, Cu, Fe, Mn and
Mo, the mills tested generally showed a difference of <20% from the control (Tables 3, 4, Fig.</li>
1a). Nickel and Zn approximately doubled in samples milled in the hammer mill and/or ball
mill, and an increasing, massive, contamination of Cr and Pb was caused by the hammer mill
and subsequent ball mill (Table 3, Fig. 1b,c). An increase in variability of Pb and Zn (Fig. 1b)
was apparent in samples that had been hammer milled, and this was accentuated by the ball

mill that also increased the Cr concentrations (Fig. 1c). This was not the case with the other 287 288 elements, or was only expressed as a trend, presumably because the contamination that arouse during milling contributed less to the total concentrations in the analysed plant material, and 289 that the inherent variability within the original material was large. On the other hand with the 290 Ti mill, there was no significant difference in element concentrations or variability as 291 292 compared to the control. Titanium is a very hard, strong and corrosion resistant metal and thus suitable for construction of cutting and milling devices. However, it can also include some 293 impurities; an example of the TE composition (21 elements) in Ti used to construct cutting 294 blades and bearings for processing other biologically derived materials showed that it 295 contained Fe 130, Sn 100, Cu 24 and Cr 4 mg kg<sup>-1</sup> as impurities (Shand et al., 1983). 296 In the second test there were no significant differences between the Cd, Co and Pb in the steel 297 milled samples compared with the control samples whereas the steel mill significantly 298 299 increased concentrations of all other elements (Table 4, Fig. 1a-c). The variability, however, was not affected by milling with the steel mill, except for Cr (Fig. 1a-c). 300

The test of mills used for archived samples demonstrated contamination with a range of TE. 301 302 For some elements the milling introduced an error much greater than that suggested by Markert (1995), indicating that the samples could only be used for studies on a few of the 303 304 tested elements. Furthermore, two of the mills increased the variability of some element concentrations, contrary to the objective of milling, which is carried out to increase 305 homogeneity and reduce variability in samples. On the other hand, the Ti knife mill did not 306 significantly contaminate the processed plant material with any of the focus elements (Cd, Co, 307 308 Cr, Cu, Fe, Mn, Mo, Ni, Pb, and Zn), at least not detectable with the analytical protocol and sensitivity of the instruments used. One drawback of using a Ti cutting mill for sample 309 preparation is that the Ti concentration cannot be used as indicator of soil contamination of 310 the plant samples but other elements such as Al or Sc may be used instead (see above). 311

#### **Reanalysis of archived samples to answer new research questions**

The use of historical data and sample archives potentially has great value in improving our 314 315 understanding of TE dynamics, e.g. in different ecosystems and the food chain. However, an appreciation of the potential contamination issues surrounding TE studies, some of which are 316 317 outlined in the present paper, for each set of data or archived sample will be of key importance in reducing the risk of data misinterpretation and inaccuracy in calculations. In 318 319 order to assess the data or sample quality, there is a need to know what equipment and procedures were used during sample collection and preparation and these must subsequently 320 be tested for potential contamination. Some elements are more likely to be introduced via 321 322 contamination and the prospect of using existing samples from earlier studies may be limited. Other elements, as suggested by this study, may less often be introduced via contamination. 323 Research questions concerning these elements may well benefit from investigating the large 324 325 amounts of samples stored in archives at different institutions.

326

#### 327 Concluding remarks and recommendations

There is a wealth of archived material that can potentially be used for TE studies. These include samples from field experiments, surveys and environmental monitoring programmes where research funds have been invested in maintaining experiments and collecting and archiving samples, and for which other data are already available. If uncontaminated, such samples can be used for contemporary TE studies, potentially providing added value. To enable this, the general consciousness about the risks of TE contamination in archived samples needs to be raised among non-specialists. One step towards reaching quality

assurance throughout the entire chain is to incorporate precautions against TE contamination 335 into the general protocols for e.g. field experiment maintenance and sampling, and 336 environmental monitoring. To our knowledge, such protocols either contain insufficient 337 338 information on TE issues, or none at all (e.g. Försökshandboken, 2009). Thus quality assurance with respect to TE often depends on the personal interest of individuals engaged in 339 research, advisory services or environmental monitoring. Considering the recent increase in 340 interest in TE, from nutritional as well as toxicological and environmental perspectives, it is 341 timely to raise these issues, and e.g. introduce a comprehensive approach to sample collection 342 and preparation that allows for complementary TE analysis of future archived samples. 343 344 However, this cannot be the responsibility of the individual non-TE specialists alone but needs to be a joint effort of TE specialists and non-TE specialists along the entire sample 345 chain within the fields of research, advisory service and environmental monitoring. 346

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463

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#### 468 Table 1. Potential sources of sample contamination and otherwise erroneous data in the collection-analysis chain of herbage samples. Bold lettering indicates

469 potential contamination sources, normal lettering other sources of error. Literature references for contamination sources are given below.

Should be given	in Field handbooks, S	tandard Operating Procedure	s, ISO etc.			Included in Accreditation	Schemes	
Sampling		Sample preparation		Storage	Sub-sampling	Extraction/ digestion	Analysis	
					for analysis			
Sub-sample	Cutting, handling	Drying & sub-sampling	Milling					
from field or	and transportation	from bullt commis						
plot (sampling		from bulk sample						
design)								
Area	Soil <sup>1,2,3, 4, 5, 6, 7</sup>	Washing/cleaning <sup>9</sup>	Device or technique;	Container <sup>4,11</sup>	Subsampling	Chemical agents	Working below	
No of samples	Equipment/	Surfaces/	glass, metal,	Chemical &	method	Water	detection/analytical limits	
Sampling	surfaces <sup>4, 8, 9</sup>	containers <sup>4,11</sup>	porcelain; cutting,	biological	(affecting	Vessels	Instability in analytical	
pattern (grid,	Weather	Subsampling method	grinding <sup>4,12,13,14,15,16,17</sup>	effects of	size/quality	Chance contamination	performance (e.g.	
random etc)	conditions <sup>10</sup>	Temperature ( <b>freeze</b> <sup>12</sup> ,	Surfaces <sup>4</sup>	unsuitable	separation)	Digestion	quality/purity of gas and	
	Phenological	dry etc)	Type of plant material	storage	Surfaces/	Lack of GLP – blanks	chemicals, electricity,	
	stage		Cleaning procedures <sup>16</sup>	conditions	devices	etc	temperature, humidity etc).	
	Part of plant			Size/quality		Lack of reference	Analysing and reporting	
				separation		samples (cross-lab	elements not planned for in	
						tests etc)	previous stage	

• Dust<sup>4</sup>

• Skin care products on bare hands<sup>4</sup>

Human variation

470	<sup>1</sup> Bargagli (1995); <sup>2</sup>	<sup>2</sup> Calder and Voss (1957);	<sup>3</sup> Cook et al. (2009); <sup>4</sup>	<sup>4</sup> Stangoulis & Sison (2008); <sup>5</sup>	<sup>5</sup> Wolterbeek (1995);	<sup>6</sup> Wyttenbach and Tobler
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- 471 (1998); <sup>7</sup> Wyttenbach and Tobler (2002); <sup>8</sup> Fleming et al. (1986); <sup>9</sup> Lockman (1980); <sup>10</sup> Fernández et al. (2010); <sup>11</sup> Waheed et al. (in press); <sup>12</sup>
- 472 Hamilton (1995); <sup>13</sup> Allan et al. (1999); <sup>14</sup> Cubadda et al. (2001); <sup>15</sup> Markert (1995); <sup>16</sup> Sager and Mittendorfer (1997); <sup>17</sup> Santos et al. (2008).

#### 473 Table 2. Milling/cutting devices tested in the two experiments and the plant material used for

the tests. 474

Experiment	Device	Туре	Device material	Plant
				material
1	Glen Creston Stanmore (bench top, swing tooth	Hammer mill	Stainless steel	
	hammers)			Mature
1	Glen Creston Stanmore (bench top, swing tooth	Hammer mill +	Stainless steel +	herbage
	hammers) + Retsch Mixer Mill MM200	Ball mill	Stainless steel	from mixed
1	Retsch Grindomix GM 200	Knife mill	Ti knives, plastic bowl	stand <sup>a</sup>
1	Plexi-glass knife, dept workshop	Knife	Plexi-glass	
2	Retch 2000	Cutting mill	Stainless steel	Vegetative
2	Plastic scissors, Kärnan AB	Scissors	Polystyrene resin	timothy

475

<sup>a</sup> Perennial ryegrass, timothy and white clover.

476 Table 3. Test of hammer mill, hammer mill+ball mill, Ti knife mill, with plexi-glass knife as

- 477 control and mature mixed herbage as test material (n=5); concentrations of elements after
- sample cutting or milling with the respective devices. Numbers in a column that are followed
- 479 by a different letter are significantly different.

Device	Cd	Co	Cr	Cu	Fe	Mn	Мо	Ni	Pb	Zn
					mg kg <sup>-1</sup>					
Plexi-glass	0.0070	0.0176	0.008a*	1.96	19.5a	61.0	1.52	0.131a	0.076a	7.95a
Hammer	0.0055	0.0174	0.172b	2.20	22.6ab	65.4	1.64	0.198b	0.127b	14.4b
Hammer+ball	0.0068	0.0166	0.586c	2.16	26.7b	62.9	1.67	0.231b	0.217b	14.3b
Ti knife mill	0.0069	0.0166	0.008a	2.13	18.6a	58.1	1.50	0.151a	0.069a	8.83a
p value	ns	ns	<0.0001	ns	0.0022	ns	ns	<0.0001	<0.0001	<0.0001

480 \*Two samples below the detection limit.

482 Table 4. Test of steel knife mill with plastic scissors as control and young timothy herbage as

test material (n=5); concentrations of elements after sample cutting or milling with the

484 respective devices.

Device	Cd	Co	Cr	Cu	Fe	Mn	Мо	Ni	Pb	Zn
					mg k	g <sup>-1</sup>				
Plastic	0.0078*	0.0204	0.060*	4.060	33.6	33.0	1.09	0.664	0.063	15.4
Steel knife mill	0.0071*	0.0226	0.414	4.478	36.8	36.9	1.41	0.812	0.063	16.3
p value	ns	ns	0.0006	0.0017	0.0114	0.0269	0.0012	0.0020	ns	0.03

485 \*Several samples below the detection limit.

#### 487 **Figure captions**

- 488 **Fig.1.** Relative concentrations of a) Cu, b) Zn, and c) Cr of two herbage materials after
- sample milling/cutting, expressed as a percentage of the average concentration in control
- 490 samples cut by plexi-glass knife (left) or plastic scissors (right).



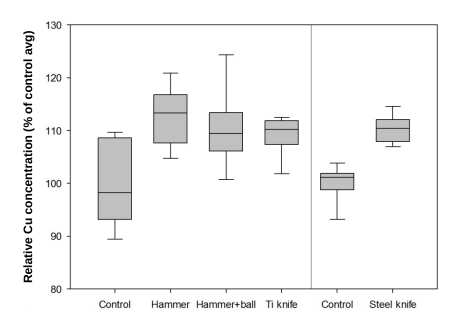


Fig. 1b

