

1 **Local versus field scale soil heterogeneity characterization - a challenge for**  
2 **representative sampling in pollution studies**

3  
4  
5  
6

7 Zahra Kardanpour<sup>1,2</sup>, Ole Stig Jacobsen<sup>1</sup>, Kim H. Esbensen<sup>1,2</sup>

8

9 1. Geological Survey of Denmark and Greenland (GEUS)

10 2. ACABS research group, University of Aalborg, campus Esbjerg (AAUE)

11

12

13

14

15

16 Abstract

17 This study is a contribution to development of a heterogeneity characterisation facility for  
18 'next generation' soil sampling for example aimed at more realistic and controllable pesticide  
19 variability in laboratory pots in experimental environmental contaminant assessment. The  
20 role of soil heterogeneity on quantification of a set of exemplar parameters is described,  
21 including a brief background on how heterogeneity affects sampling/monitoring procedures  
22 in environmental pollutant studies. The Theory of Sampling (TOS) and variographic analysis  
23 has been applied to develop a more general fit-for-purpose soil heterogeneity  
24 characterization approach. All parameters were assessed in large-scale transect (1-100 m) vs.  
25 small-scale (0.1 -0.5 m) replication sampling point variability. Variographic profiles of  
26 experimental analytical results from a specific well mixed soil type concludes that it is  
27 essential to sample at locations with less than a 2.5 meter distance interval to benefit from  
28 spatial auto-correlation and thereby avoid unnecessary, inflated compositional variation in  
29 experimental pots; this range is an inherent characteristic of the soil heterogeneity and will  
30 differ among other soils types. This study has a significant carrying-over potential for related  
31 research areas e.g. soil science, contamination studies, and environmental monitoring and  
32 environmental chemistry.

33

34 Keywords: Heterogeneity characterisation, soil, variogram, large-scale, representative  
35 sampling, Theory of Sampling (TOS), MCPA, biomass, CFU

36 1. Introduction

37 All parameters for realistic, effective integration of variability over different scales are directly  
38 related to soil heterogeneity. There is a growing need for an integrated understanding of  
39 contaminant behaviour in soil pollution studies (Arias-Estévez et al. 2008; Crespín et al.  
40 2001; Johnsen et al. 2013; Li et al. 2006; Rodríguez-Cruz et al. 2006; Sørensen et al. 2006;  
41 Torstensson and Stark 1975; Rasmussen et al. 2005). In this context there is a missing link in  
42 the form of soil heterogeneity and its effective characterization, a feature often overlooked.  
43 Heterogeneity characterisation is the first, and in some cases the most important step, in soil  
44 contaminant studies, with relationships to various other aspects of environmental research  
45 and monitoring. A result of introducing more valid soil heterogeneity characterisation will be  
46 improved soil sampling procedures (Kardanpour et al. 2014; Kardanpour et al. 2015a,b),  
47 which in turn will contribute towards improved environmental fate study reliability  
48 (Boudreault et al. 2012; Chappell and Viscarra Rossel 2013; de Zorzi et al. 2008; Lin et al.  
49 2013; Mulder et al 2013; Totaro et al. 2013).

50 Even in simple systems, the variability and risk for misinterpretation may have strong effect  
51 on parameterisation of processes relating to compound fate studies. These latter issues are  
52 being increasingly more recognised, as is the lack of appropriate methods to ensure  
53 documented representativity of the experimental batch volumes/masses with respect to the  
54 surrounding geology and biotic/abiotic soil characteristics. There is an urgent need for  
55 scientifically based experimental approaches, scale-up procedures and attendant principles  
56 for parameterisation of variability in these types of natural systems(Kardanpour et al. 2014;  
57 Adamchuk et al. 2011; Chappell and Viscarra Rossel 2013; de Zorzi et al. 2008).

58 Of particular interest will be a newly developed facility for empirical variability  
59 characterisation, which allows heterogeneity to be mapped at problem-dependent scale  
60 hierarchies. Based on this, it is possible to devise optimised sampling strategies that will allow  
61 fit-for-purpose representativity with respect to laboratory experiments depending of similar  
62 (or at least comparable) soil samples (pots). For this purpose the Theory of Sampling (TOS)  
63 delivers benchmarks measures expressing acceptable maximum heterogeneity limits and in  
64 the case of violations/transgressions furthers a complete understanding of how to identify  
65 and eliminate the detrimental sampling errors and provides tools for unambiguous mixing  
66 effectiveness. Combining these tools with specific knowledge on the relevant contaminant  
67 processes and compound properties, it will be possible to address the critical scale-dependent  
68 variability with increased confidence based on more realistic environmental parameter  
69 delineation.

70 We here introduce the variographic approach mainly for the cases of 1-D as a means of  
71 characterising the heterogeneity in one transect direction. Compared to the typical major  
72 variability in the Z-direction of soil depth profiles (soil horizons, layers, geological  
73 formations), the linear (1-D) or 2-D heterogeneity *within* soil horizons is significantly smaller,  
74 although this is exactly the kind of heterogeneity the present study aims at controlling.

75 Contrary to depth profile zonation a.o. the within-horizon 1-D and 2-D heterogeneity complies  
76 with the requirements of both TOS and geostatistics, i.e. spatial heterogeneity can be modelled  
77 variographically w.r.t. a physically meaningful *average level* (the inherent *stationarity*  
78 *assumption* in geostatistics). It is not meaningful to apply variographic characterisation on  
79 measurement series which contain discontinuous shifts, upsets or other disrupt, level  
80 changes, as is the prime characteristic of soil depth zonations. The geostatistical tradition of  
81 modelling 2-D patterns based on projection onto a 1-D transect is also not free from debatable  
82 issues<sup>1</sup>. In the present context all *isotropic* 2-D heterogeneity patterns can be characterised  
83 comprehensively by a *randomly selected* 1-D direction (transect). In all sampling operations  
84 there should preferentially always be some sort of random selection involved, unless  
85 compelling geo-science reasons exists for choosing a direction related to the genesis of the  
86 specific heterogeneity met with, e.g. choosing a 1-D transect either along a dominant plow  
87 direction.

88 This study focuses on development of the necessary heterogeneity characterisation for  
89 sampling/monitoring and multi-parameter modelling practices, allowing implementation of  
90 realistic pesticide variability in experimental environmental contaminant assessment studies.  
91 The study has a significant carrying-over potential for related research areas e.g. soil science,  
92 contamination studies, and environmental monitoring.

93 We here focus on characterization of soil heterogeneity in terms of soil moisture, organic  
94 matter (LOI), biomass, microbiology, MCPA sorption and mineralization. The measured  
95 parameters are here used to illustrate effective management of heterogeneity; this particular  
96 location has been studied before in its own right. Following two earlier complementary  
97 studies, the focus below is on the necessary representativity demands when facing compound  
98 fate and mineralization studies (Kardanpour et al. 2014; Kardanpour et al. 2015). Field  
99 observation indicates a very well mixed sandy soil with almost no visual heterogeneity  
100 features. But the main issue is: does this apparent uniformity extend to all fate compounds?  
101 How is it possible to document that small sample masses, as typically used in pot experiments,  
102 are representative of their entire parent field, or to which sub-field scale? In other words, how  
103 can results and conclusions from laboratory experiments be reliably scaled-up and  
104 generalized to larger scales?

## 105 2. Materials and Methods

### 106 2.1. Location and sampling pattern

---

<sup>1</sup> The present authors do not wish to reject the 2-D geostatistical tradition with this statement, but in relation to the present matters this issue is better deferred to another occasion in which the 2-D modelling issue can be presented and discussed in full - this issue is a legitimate and interesting area for a fruitful debate. Entering into a 3-D geostatistical modelling realm, there are also here issues that in need of further discussion, e.g. the required minimum number of samples (measurements) needed for meaningful, and stable variogram calculation, The present foray only aims at presenting the power of a simple 1-D variogram characterisation operator based on TOS, upon which several versions of potential follow-up generalisations to 2-D and 3-D cases may be entertained.

107 Fladerne Bæk is situated on the Karup peri-glacial outwash plain, Jutland, Denmark (56°N,  
108 9°E) South West of Karup airport. The substratum is an arable sandy soil which has been tilled  
109 and cropped for more than 100 years, mainly supporting barley and potatoes during last 30  
110 years. Thus this is a typical “very well mixed” soil type compared to the much more  
111 heterogeneity glacial clayey soil types treated in (Kardanpour et al. 2014). Soil samples were  
112 collected from the topsoil (A-horizon) in cylindrical cores; the present samples cover depth  
113 interval from 0-15 cm. The 60 m long sampling transect was roughly N-S. Each field sample  
114 included 200-300 grams of fresh soil. At the center of this transect at point 29, seven  
115 additionally samples form a roman grid (3 x 3) replication experiment with 0.3 meter  
116 equidistance.

117 The sampling rationale aimed at variographic fate characterization commensurate with a long  
118 profile at a scale length between 1m and 60 m; the roman square was intended as a basis for  
119 conventional statistical treatment (average and, standard deviation). This central sample  
120 layout serves as a small scale local ‘replication experiment’ compared with the transect  
121 dimensions (Kardanpour et al. 2014). In total 64 samples were collected, 57 samples from the  
122 long profile and nine samples of the small grid (two samples identical to two from the  
123 transect), one in between and three more in each side of transect with the same distance as  
124 the first three in the center of transect. The original fresh soil was kept frozen until use.

125 The primary sampling was specifically intended to correspond to current sampling traditions  
126 in the soil and microbiology communities. In other studies efforts have been made to optimize  
127 each individual field sample, for example with respect to the famous “Gy’s formula”, from  
128 which control over the so-called Fundamental Sampling Error is often sought. However, in the  
129 present study it is a major point to outline how the variographic approach a.o. lead to a  
130 procedure with which to characterize the magnitude of the total sampling-plus-analytical  
131 error and thus to be warned of the need to control (better) all the inherent sampling errors,  
132 see e.g. DS 3077 (2013) for a comprehensive introduction.

## 133 2.2. Theory of sampling and variographic analysis

134 The Total Analytical Error (TAE) is most often under acceptable control in the analytical  
135 laboratory as regards to both accuracy and precision. A sampling procedure must be both  
136 *correct* (ensures accuracy) and *reproducible* (ensures precision); TOS defines *representativity*  
137 in a rigid conceptual and mathematical approach. The critical issue is always, even for TOS-  
138 compliant sampling, that analytical results are but an *estimate* of the true (average) analytical  
139 grade of the lot sampled, because the aliquot is based on only a miniscule mass (0.5 – 2.0 g)  
140 compared to the entire field topsoil layer it is supposed to represent (typical mass/mass  
141 sampling ratios range 1:10<sup>3</sup> to 1:10<sup>9</sup>). The full sampling-analysis process and its  
142 characteristics is therefore the only guarantee for the relevance and reliability of the aliquot  
143 brought forth for analysis. The fundamental TOS principles need to be applied to all  
144 appropriate scales along the entire ‘field-to-aliquot’ pathway, not only to the primary  
145 sampling, but in particular also to the successive stages of mass reduction in the laboratory

146 before the ultimate analytical aliquot extraction. The only change in this multi-stage sampling  
147 chain is the operative scale (TOS principles and unit operations are scale-invariant). A  
148 comprehensive overview of all subsampling issues (laboratory mass reduction) has been  
149 published in (Petersen et al. 2004), which does not include the ‘coning-and-quartering’  
150 approach, despite the fact that this approach has enjoyed some popularity e.g. for certain field  
151 applications to soils (Gerlach et al. 2002). However the coning-and-quartering approach has  
152 been severely criticized in the professional TOS literature, e.g. most recently in (Esbensen and  
153 Wagner 2014); from a representativity point of view coning this mass reduction approach  
154 must be strongly discouraged

155 On the basis of a correct sampling and mass reduction regimen, it is possible to characterize  
156 the inherent auto-correlation between units of a process/lot or along 1-D transect (or  
157 transect). The *semi-variogram* (in this work referred to simply as the ‘variogram’) is employed  
158 to describe the variation observed between sample pairs as a function of their internal  
159 distance.

160 To calculate a variogram a sufficient number of units (increments/samples) are extracted  
161 equidistantly, spanning the process interval of interest, or the full transect length, as needed.  
162 The variogram is a function of a dimensionless, relative lag parameter,  $j$ , which is this distance  
163 between two units, the analytical results of which are compared. Full details of the  
164 variographic approach are described in (DS3077 2013; Esbensen et al. 2007; Esbensen et al.  
165 2012a; Esbensen et al. 2012b; Gy 1998; Minkkinen et al. 2012; Petersen and Esbensen 2006;  
166 Petersen et al. 2005). Variograms may have apparent different specific appearances, but three  
167 fundamental characterizing features carry all the important information related to sampling  
168 errors and the heterogeneity along the transect in any-and-all variogram: the *sill*, the *range*,  
169 and the y-axis intercept, termed the *nugget effect*. Definitions of these features are given  
170 below.

171 The Sill is the y-axis value at which the variogram levels off and becomes horizontal. The Sill  
172 represents the total variance calculated from all experimental heterogeneity values. The sill  
173 corresponds to the overall maximum variance for the data series if/when calculated *without*  
174 taking their ordering into account.

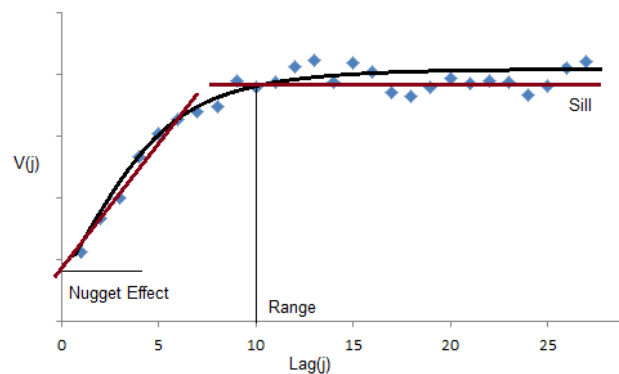
175 The Range is the lag distance beyond which the variogram  $v(j)$  levels off and reaches a stable,  
176 constant Sill. Samples taken at lags below the Range are auto-correlated to a larger and larger  
177 degree as the lags gets smaller and smaller. The range carries critical information as to the  
178 local heterogeneity with respect to the objective of the present method development.

179 The Nugget Effect indicates the amount by which the variance differs from zero when a  
180 variogram is extrapolated backwards so as to correspond to what would have been a lag = 0. A  
181 lag equal to zero has no physical meaning, but it represents the hypothetical case of two  
182 samples extracted at the same time and location (indeed from exactly the same physical  
183 volume of the lot). Thus although ‘true replicates’ from the exact same soil location (volume)  
184 are not physically possible, the nugget effect never-the-less allows to estimate the

185 corresponding discontinuous variance difference. This can be viewed as a collapse of the 1-D  
186 sampling situation (profile, transect) to a stationary sampling situation (small lots, 2-D and 3-  
187 D lots), see (DS3077 2013; Esbensen et al. 2007, 2012a, 2012b) for further descriptions.

188

189 The nugget effect has a special interest, it contains all sampling, - sample handling/processing  
190 and analytical errors combined, which makes up the total measurement uncertainty. A  
191 variogram with a high nugget effect w.r.t. the sill signifies a measurement system not in  
192 sufficient control (DS3077 2013; Esbensen and Wagner 2014).



193

194 Figure 1. A generic variogram, schematically defining nugget effect, sill, and range. The  
195 illustration depicts an *increasing variogram*, which is the most often occurring type of  
196 variogram in the case of significant auto-correlation (for lags below the range)(Kardanpour et  
197 al. 2014). The nugget effect magnitude relative to the sill in this illustration is significant of an  
198 acceptable total measurement system, < 20%.

199 Variogram calculations are strongly influenced by *outliers* and/or *trends*. A valid variographic  
200 analysis often necessitates outlier deletion after proper recognition and description and  
201 occasionally also de-trending of the raw transects data if/when trends are dominant or  
202 severe. In this study the raw data transect was de-trended using a simple regression slope  
203 subtraction from the data set where needed.

### 204 2.3. Mass reduction/subsampling procedure

205 After the stored samples were thawed and accommodated for 20 °C for a week, before being  
206 processed further. The primary field sample size (200-300 gram) must be reduced to the  
207 analytical sample size (1-2 gram), not at all a trivial mass-handling issue. In order to provide  
208 representative sub-samples, TOS principles were applied scrupulously to all mass reduction  
209 steps. Thus samples were dried and macerated, or ground, where appropriate, and  
210 subsequently deployed in a longitudinal tray, forming a 1-D lot, using the soil-adapted bed-  
211 blending/cross-cut reclaiming technique described in detail in (Petersen et al. 2004) and  
212 Kardanpour et al. (2015b). These pre-blended micro-beds were cut by 10 randomly selected  
213 transverse increments along the elongated dimension which were aggregated, resulting in  
214 subsamples of 20-30 gram each. The exact same procedure was repeated in a secondary mass

215 reduction step ending up with the final analytical mass (2 gram) for the wet samples analyses.  
216 This procedure has been honed to full representativity in the course of this project specifically  
217 so as to do away with all of the post-primary-sampling errors in order better to be able to  
218 focus in the latter and the variogram deployment, *ibid*.

219 The remainders of the secondary sub-samples were air-dried for four days in lab temperature  
220 (20 °C), to be used in parallel sorption experiments. As a further scale-down iteration, a  
221 similar bed-blending/cross-cut reclaiming were used to provide analytical samples of 2 gram,  
222 also based on 10 increments each.

223 Kardanpour et al. (2015b) describe the “from-field-sampling-to-aliquot” pathway in full  
224 details, complete with an exhaustive pictorial exposé.

## 225 2.4. Analytical experiment methods

### 226 MCPA Sorption

227 The sorption experiment started in glass vials with Teflon caps containing 1 g of the  
228 respective soils, and 9 ml of Milli-Q water. The vials were kept for 24 hours and then shaken in  
229 a horizontal, angled shaker prior to addition of 1 mL<sup>14</sup>C-MCPA stock solution, with 10,000  
230 dpm in each individual vials. Sorption experiments were performed with two initial  
231 concentrations: 1 and 100 mg MCPA/L. Sorption was determined for MCPA in all off the 64  
232 soil samples, using <sup>14</sup>C-labeled MCPA.

233 After adding the stock solution, the vials were incubated in the shaker for 48 hours and then  
234 placed vertically for another 48 hours, all at 20 °C. Subsequently 2 mL of the solution were  
235 transferred to the 2 mL Eppendorf micro-centrifuge tubes and centrifuged at 14,500x g for 7  
236 min. Radioactivity in 1.5 mL supernatant was determined using a Wallac 1409 Liquid  
237 Scintillation Counter after mixing it with 10 mL OptiPhase Hisafe3 scintillation cocktail.

### 238 MCPA Mineralization

239 Mineralization experiments were carried out in 100 mL glass jar with air tight lid. Two gram  
240 soil (wet weight) was placed in small plastic vials before adding 0.5 mL of <sup>14</sup>C -labeled MCPA  
241 (5 mg MCPA kg<sup>-1</sup> soil) with a radioactivity of 2,000 dpm. In the glass jar a LSC vial was also  
242 placed containing 2 mL 0.2 M of NaOH as a CO<sub>2</sub> trap. The jars were incubated at 20°C for 14  
243 days. Mineralization encountered as %-evolved <sup>14</sup>CO<sub>2</sub> was measured at day 3, 7 and 14. The  
244 CO<sub>2</sub>-traps were changed and replaced with a fresh trap at each sampling date. <sup>14</sup>C in the NaOH  
245 was measured as described in the sorption experiment by Liquid Scintillation Counting.

### 246 Biomass; substrate induced respiration (SIR)

247 The same set up as used for MCPA was used for the glucose mineralization with adding 0.5  
248 mL<sup>14</sup>C -labeled glucose with 5000dpm to the 2 gram of soil. All other set up details, equipment  
249 and experimental design were identical. Alkaline traps were replaced with fresh alkaline traps  
250 and measured after 4 and 24 hours considering the rapid respiration of the glucose and <sup>14</sup>C

251 measured as described in the sorption experiment by Liquid Scintillation Counting.  
252 Conversion into biomass were according to ( Dictor et al 1998; Tate et al. 1988).

253 Microbiology, Bacteria Colony Formation Units (CFU)

254 A suspension was made with 2 gram of soil into 200 mL sterile water and after shaking for 15  
255 minutes, diluted with sterilized water ended in two different dilutions for each sample; with  
256 three and four order of magnitude To measure the soil microbiology, 1 mL of each sample  
257 were placed on a Petrifilm® (3M, Saint Paul, Minnesota, USA) sheet and CFU was counted after 3  
258 and 7 days of incubation at 20°C.

259 3. Result

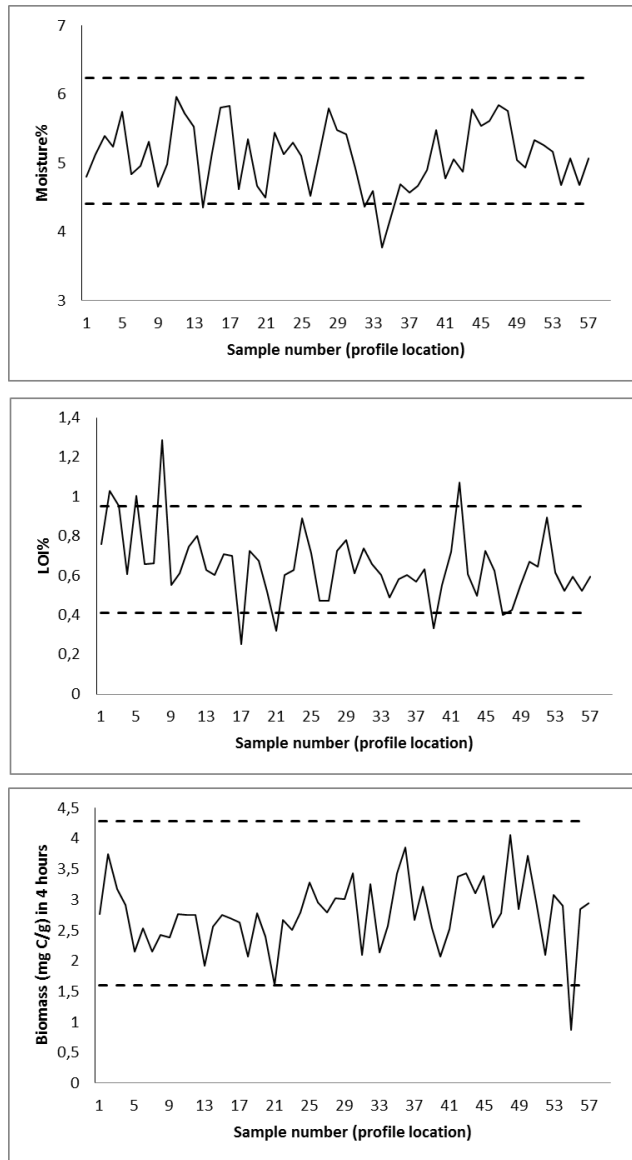
260 3.1. Geochemical profiling

261 In order to show the natural soil heterogeneity in a comparable format, Figures 2-5 illustrates  
262 the individual large-scale parameter transects; concentration vs location of the samples taken  
263 from the transect in Fladerne field. Also shown is the variation of the central *small-scale*  
264 replication samples is shown as mean concentration  $\pm$  2 SD with dashed horizontal lines in  
265 the figures. The large-scale variation of the soil moisture, loss on ignition (LOI) and the  
266 biomass content are to be compared to the small scale replication result for the same  
267 parameter in each graph, Figure 2.

268 The same comparison graph illustrated for the MCPA sorption in Figure 3 for two different  
269 initial MCPA concentrations, as it is clear, the soil sorption behavior show different variation  
270 with different concentrations. The results of the MCPA mineralization of the soil in Figure 4  
271 also show different variability with different mineralization steps. The transect of the MCPA  
272 mineralization is illustrated for different mineralization steps: first three days, four to seven  
273 days and eight to fourteen days. The two latter periods shows rather a similar variation  
274 because these two periods are in the final part of the mineralization development, Figure 6.

275 The soil microbiology (Log (CFU/g soil)) transect after seven days of incubation is also  
276 illustrated in Figure 5.

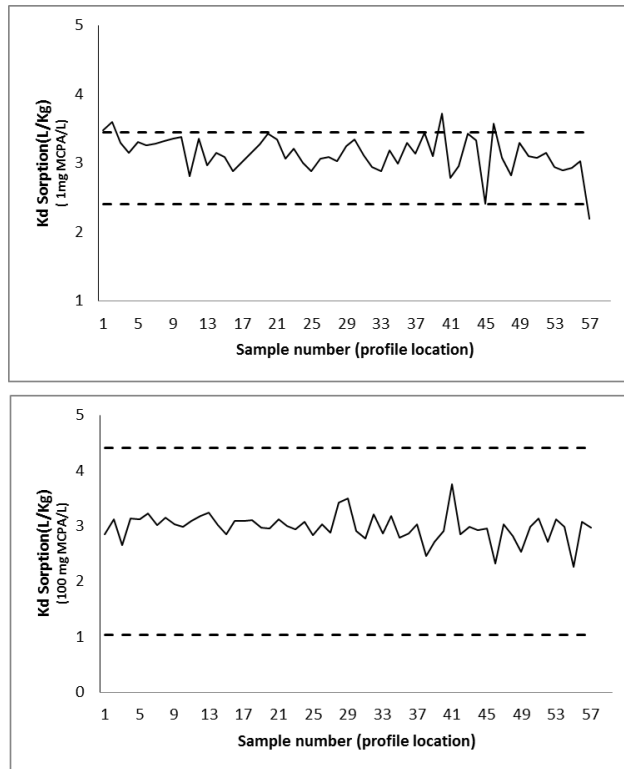




277

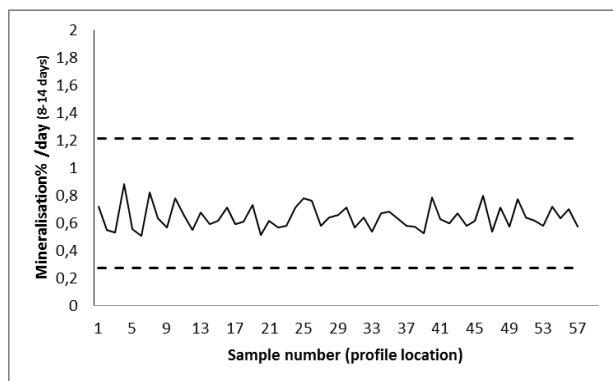
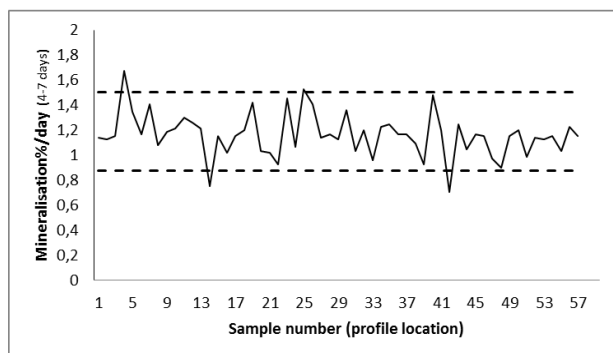
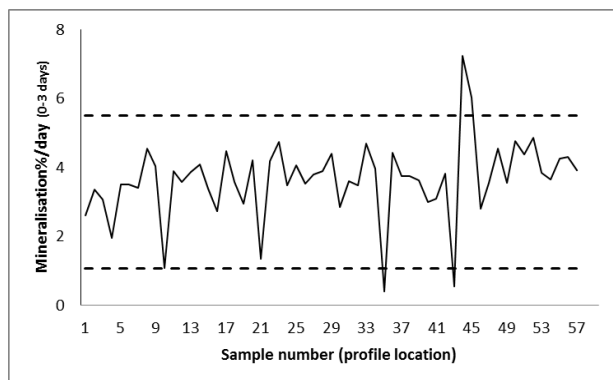
278 Figure 2. Fladerne Bæk, transects of soil moisture (%), LOI, and biomass (mg C/g); soil  
 279 biomass vs. sample number (transect location). Dashed lines represent mean  $\pm$  2 SD of the  
 280 small-scale replication experiment.

281



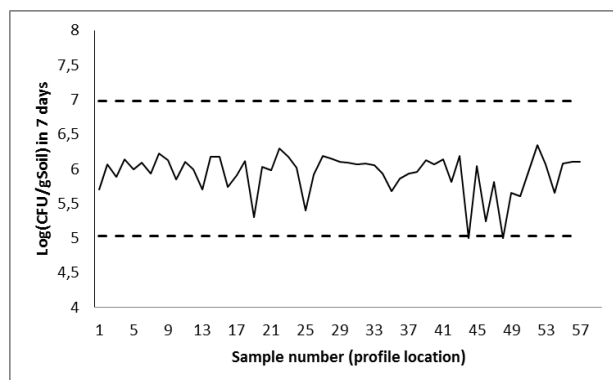
282

283 Figure 3. Fladerne Bæk, transects of  $K_d$  MCPA sorption vs sample number (transect location),  
 284  $K_{d,1}$ : MCPA (1 mg/ L),  $K_{d,100}$ : MCPA (100 mg/ L). Dashed lines represent mean  $\pm$  2 SD of the  
 285 small-scale replication experiment.



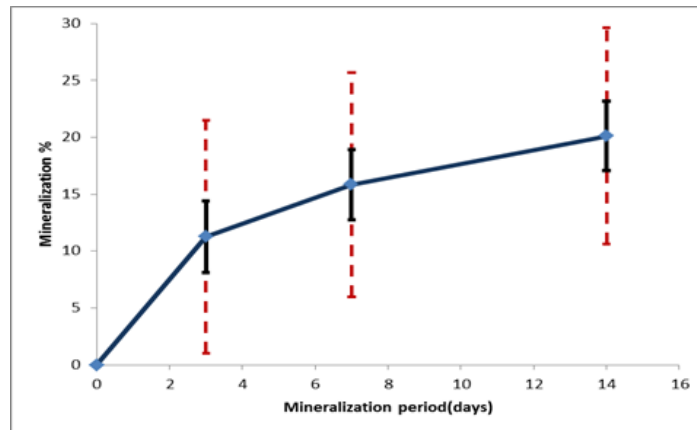
286

287 Figure 4. Fladerne Bæk, transects of MCPA mineralization in three different periods: 0-3days,  
 288 4-7days, 8-14 days vs. sample number (transect location). Dashed lines represent mean  $\pm$  2  
 289 SD of the small-scale replication experiment.



290

291 Figure 5. Fladerne Bæk, transects of log (CFU/g soil) vs sample number (transect location)



292

293 Figure 6. Average mineralisation rate for all 57 samples: Error bars are based on the standard  
 294 deviation (solid bars) and the range of the whole sample set (stippled bars)

295 The Fladerne case represents an inherently very well mixed soil type, which has been under  
 296 the plow for up to 100 years<sup>2</sup>. This case was selected to represent the one (almost extreme)  
 297 end of a spectrum (only little inherent heterogeneity) from which to compare a whole  
 298 spectrum of increasingly more heterogeneous soil types, horizon and geological formations.  
 299 Our own studies went a fair distance in this direction as possible with the (Kardanpour et al.  
 300 2015), but obviously many, even more heterogeneous cases exist and are on record in the  
 301 literature.

### 302 3.2. Experimental variograms

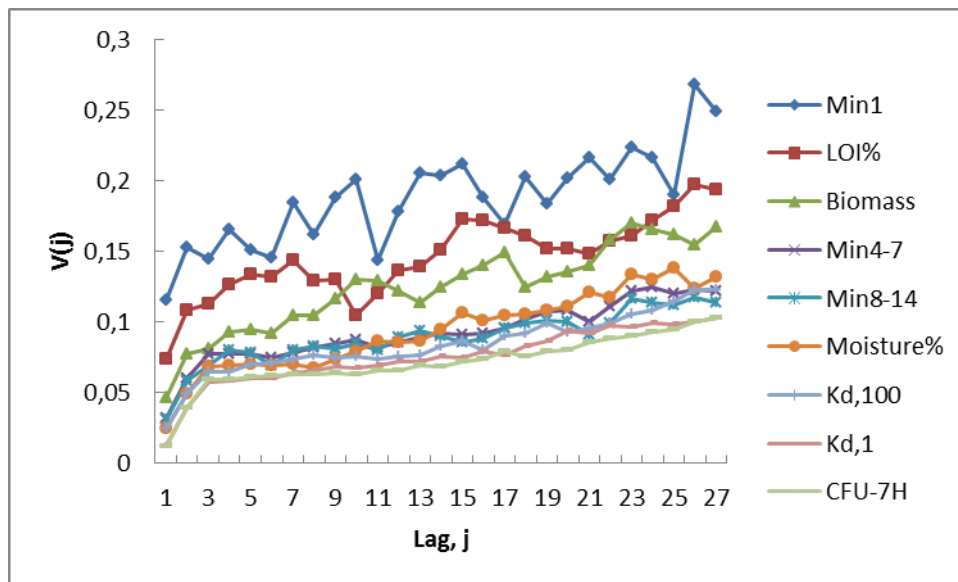
303 Prior to variogram calculation, all parameters have been checked for outliers and trends,  
 304 Figures 2-5. Variograms have been calculated with using large scale experimental transects  
 305 without model fitting of the variogram parameters. This is common in geostatistics, but not  
 306 used here as 'TOS' variogram approach is not used for kriging but solely for heterogeneity  
 307 characterization and interpretation.

308 Two different behaviors can be observed as displayed by two parameters groupings, the  
 309 increasing Min1, LOI and Biomass variograms at the top, versus the reminder of parameters,  
 310 which show a strongly similar form and behavior, Figure 7. As the sill levels represent the  
 311 maximum parameter variation along the transect, parameters Min1, LOI and Biomass clearly  
 312 display the highest transect variability. All variograms are of the increasing type with a  
 313 distinct nugget effect. Following (DS3077 2013), the %-age nugget effect in relation to the sill,  
 314 termed  $RSV_{1-dim}$ , is an expression of the total measurement uncertainty MU including TSE

<sup>2</sup> The consequence of taking care of this, low-heterogeneity end of the spectrum, is that there is a limit to the degree of transect heterogeneity to be expected, as indeed witnessed in Figs 2-5, where concentrations only comparatively rarely deviate outside the +/- 2 STD of the central Roman square design employed. This *specific* soil- and tilling history feature must not lead to untoward confusion and illegitimate generalizations however. It is the general applicability of the variographic approach which is illustrated here, as it happens, on a very well-mixed substratum. Our parallel study showcases the approach on a significantly more heterogeneous case, in which the central Roman square does not bracket most of the transect concentration manifestations.

315 (Esbensen and Wagner 2014). In the present study this  $MU_{total}$  quality index ranges from 15%  
 316 ( $K_d, 100$ ) to 75% (Min1). There is thus an appreciable difference concerning the possibility to  
 317 measure and characterize soil heterogeneity along the transect, ranging from very good to  
 318 very poor. This facility for total measurement uncertainty validation is a powerful TOS  
 319 benefit, with a wide carrying-over potential to many other sciences and application fields.  
 320 This feature was is described in full in (Esbensen and Romanach, 2015) ((Kardanpour et al.  
 321 2015) in which, by the way, the 1-D transect of the present study appears in the form of a 1-D  
 322 industrial process measurement series, illustrating the surprising generality of the variogram  
 323 approach - modeling and interpretation of the variogram from such disparate data types are  
 324 identical, showing the way for application also to natural process in the geo-science and  
 325 environmental science realms.

326



327

328 Figure 7. Synoptic variogram of all parameters in the present study comparing nugget effect,  
 329 sill and range levels

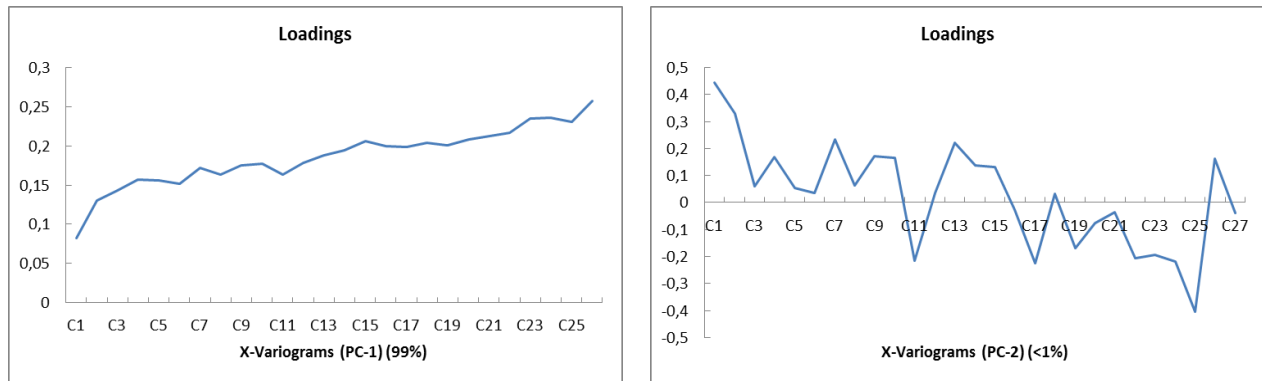
330

331 Applying the multivariate data analysis approach developed in the former studies  
 332 (Kardanpour et al. 2014; Kardanpour et al. 2015), i.e. using the variograms as the input (X-  
 333 matrix) to a Principal Component Analysis (PCA) with no centering and no scaling (see further  
 334 below), the first component is found to represent 99% of the total variogram variance over all  
 335 parameters, making it easy to find the average range characterizing the heterogeneity of the  
 336 Fladerne transect, ca. 5 meter. Figure 8 shows the loadings for PC components 1 and 2,  
 337 displayed in a fashion that mimics a *spectrum*. As expected the PC-1 loadings delineates a  
 338 general variogram shape, in fact presenting the *average* of all variograms in Figure. 7. The PC-  
 339 2 loadings accounts for deviations herefrom, as caused by the individual variograms (mainly  
 340 expressing a higher or lower average slope), a general feature, markedly overprinted by  
 341 random deviations. This component models the set of different slopes of the individual

342 variograms, and it accounts for less than 1% total variance, but never-the-less lends itself  
343 easily to be interpreted as the well-known spectroscopic 'tilting' signature, (Martens & Næs  
344 1991).

345

346



347

348 Figure 8. PCA ( $X_{\text{variogram}}$ ) loading plot for PC-1 (left) and PC-2 (right). The  $X_{\text{variogram}}$  matrix has  
349 not been subjected to pre-treatment before PCA (no centering, no scaling). The range of the  
350 average variogram shape as represented by the PC1 loadings is ca. 5meters.

351 In our earlier studies, (Kardanpour et al. 2014), can be found a discussion *pro et contra* pre-  
352 treatment of an X-matrix made up of variograms. When basing variograms on heterogeneity  
353 contributions (a one-to-one transformation of the original analytical concentrations), this  
354 issue becomes moot, as this transformation is already performing what amounts to scaling. In  
355 the present paper we therefore did not apply centering, opting for the easily interpreted and  
356 useful appearance of the average variogram shape, Figure 8 (left).

#### 357 4. Discussion

358 Aiming for a general approach to soil heterogeneity characterisation, a set of naturally  
359 occurring organic, anthropogenic and biota parameters were studied at scales from 1 to 60  
360 (100) m to be compared with other, for example minerogenic parameters (see further below).  
361 The first step is always inspection of the raw data set with respect to potential outliers and/or  
362 trends. In the present study the geochemical parameter transects show no outliers and no  
363 strong trends, Figures 2-5.

364 The experimental design allows comparison of the small-scale replication variability (classic  
365 statistics) and large-scale variability. All transects can for example be directly compared with  
366 the level and variation at the small-scale experiment (less than 1 meter), by the pertinent  
367 mean  $\pm 2$  SD. In figures 2-5 the variation of the parameters in any selected small scale window  
368 cannot be overestimated to the large scale, indeed it cannot be also obtained from a small  
369 scale replication study deviation estimate. This is just for visual orientation however and not  
370 to be confused with the nugget effect, a much more general characterisation of the small(est)  
371 scale variability pertaining to below lag = 1, summing up and averaging this information for

372 all the sample pairs in the transect.

373 Any short interval on a transect Figures 2-5 can be considered as a small scale study in its own  
374 right. In this context there is a clear difference between the empirical variability in different  
375 segments *along* each transect: the local variability does not necessarily extend to larger scales.  
376 This has an important practical conclusion: any local small-scale sample collection cannot be  
377 generalised to larger scales. Unwitting or un-reflected scaling-up of small scale experimental  
378 organic, anthropogenic and biota fate and mineralization results will bring an inflated  
379 uncertainty outside experimental control. The mineralisation parameters which show  
380 different variation behaviour in the different mineralisation steps send an important message  
381 regarding studies concerning time-dependent characterisations. A similar difference is  
382 observed for MCPA sorption with different concentrations, i.e. when studies are concerned  
383 with concentration-dependent phenomena.

384 The *general* local variability behaviour is however well captured as the below-range part of  
385 the general variogram loading spectrum for PC1. The variogram is able to generalise the  
386 common local scale behaviour. With TOS, there is synoptic information residing in the range,  
387 sill and nugget effect for each individual parameter. Whenever heterogeneity variograms  
388 display a range, this relates to the ease and risk associated with attempting to secure field  
389 samples with minimum variability: Sampling with smaller inter-increment lag distances than  
390 the range makes it possible to use the inherent auto-correlation between samples in a  
391 beneficial fashion.

392 From the earlier studies (Kardanpour et al. 2014; Kardanpour et al. 2015) the overall  
393 conclusion was only to employ *composite sampling*. In the present context this means that,  
394 wherever practically possible, increments should only be collected with a maximum of half  
395 the observed range as a means to avoid unnecessary compositional variability effects due to  
396 the inherent soil scale heterogeneity. It follows that in order to minimize the total sampling  
397 error, increments must be sampled with a maximum lag of  $0.5 \times \text{range}$ , *preferentially* smaller. In  
398 the present soil variograms a general range of 5 meters is observed for multivariate  
399 variographic approach of the parameters, Figure 8. It is evident that a thorough mixing of the  
400 selected set of increments is mandatory to sample locations with less than 2.5 meters distance  
401 in between; for other soil types/analytes other numerical magnitudes apply.

402 The variograms show different behaviour with respect to mineralisation stages. This is  
403 expected from the slower rate of the mineralisation in the latter stages, Figure 6. The later  
404 stages display a flat variogram that only represent little auto-correlation between sample  
405 locations, Figure 7, and the low sill level representing low variation along the transect. As it is  
406 common in environmental studies, results of the mineralisation are mostly reported in terms  
407 of the accumulated mineralisation rate (see Figure 6 as an example), i.e. results that are  
408 mostly affected by the first stages of the mineralisation.

409 Most of the variograms level off quickly after only a few lags (range ca. 5 meters) followed by  
410 a flat (or slightly increasing) trend, while first step of MCPA mineralisation, biomass and LOI

411 show more markedly increasing variograms, Figure 7.

412 The CFU sill level is lower than natural organic and anthropogenic compounds indicating  
413 lower variability of soil microbiology at the large scale(s). This can be compared with results  
414 from a series of other large-scale studies on different microbial communities for different  
415 anthropogenic and natural compound mineralization, which also showed that microbial  
416 biomass seem to be stable intrinsic parameter of longer periods. (Sørensen et al. 2003;  
417 Bending et al. 2001; Bending et al. 2003; Walker et al. 2001).

418 It is always a matter for discussion when theoretically anticipated correlations between the  
419 physiochemical/microbial activities fail to appear in specific real-world case studies. The  
420 more complex compounds have shown a more irregular, patchy fashion of decaying due to  
421 more specific microbial communities (but still generally isotropic in nature). Analysis of soil  
422 parameters rarely gives a clear pattern; this seems to be associated to a number of not-  
423 included or unknown parameters, resulting, in some cases in a high degradation potential, but  
424 low elsewhere (Sørensen et al. 2003; Rasmussen et al. 2005; Bending et al. 2001; Walker et al.  
425 2001). Upon reflection this is no mystery however, but simply a result of local soil  
426 heterogeneity, which cannot be formulated or predicted based on the physiochemical  
427 biological or microbial correlation of the properties of soil in large scale studies. A  
428 variographic heterogeneity characterization at all scales is thus a beneficial pilot experiment  
429 able to focus on the relevant heterogeneities characterizing individual, or group of parameters  
430 in their proper scale-dependent relationships.

431 Summing up the results of all measured parameters studied here, for environmental purposes  
432 and objectives related to soil parameters at field scale, it is advantageous to employ a  
433 variographic heterogeneity characterisation as a pilot study. Results here from will lead to a  
434 comprehensive understanding of the spatial variability and auto-correlation of the  
435 parameters in the field.

436 The results from the present study show that for well-mixed sandy soil it is recommended to  
437 sample locations with less than 2.5 meters inter-distance in between, preferentially smaller. It  
438 is necessary to conduct a similar variographic pilot experiment in order to outline the  
439 relevant scale-heterogeneity characteristics for other soil types, which unavoidably will tend  
440 to show more irregular spatial heterogeneity patterns – each principal soil type will in  
441 principle be characterised by a specific range, but there is a further caveat. Each analyte may  
442 in fact display its own, more or less specific range, as witnessed above, as well as by a plethora  
443 of studies in the literature. When controlling the spatial heterogeneity is of the essence, the  
444 logical solution is to design the sampling according the analyte with the *smallest* range, i.e. the  
445 most heterogeneously distributed analyte – this will by necessity also take care of all other  
446 analytes with higher ranges. If emphasis is on sampling costs (a not totally unlikely alternative  
447 scenario that may, or may not clash with other requirements of which only one really matters  
448 though: representativity) it is a comforting thought that all analytes are measured on the  
449 same final aliquot. By carefully optimising the primary field sampling according to the  
450 principles presented here, all analytes will be measured with the same, optimal relevance,



451 indeed w.r.t. the same representativity. If sampling is done right from the start, there are no  
452 extra costs – while the opposite is a very different case, as should be abundant clear.

453 Results from a parallel study on the *minerogenic* compounds for the same Fladerne field  
454 (Kardanpour et al. 2014) show a similar soil heterogeneity compared to the present  
455 *anthropogenic* compounds. The nugget effect for most of the minerogenic compounds are of  
456 the same order of magnitude as those for the anthropogenic compounds, i.e. the total  
457 measurement system and procedures (sampling/handling/processing/analysis) pass all the  
458 quality criteria for representative sampling established in the recent sampling standard  
459 (DS3077 2013).

460 In cases where the next step in studies might be assessment of the main factors driving the  
461 spatial heterogeneity of soil contamination analytes for example, the 1-D (or 2-D X-Y)  
462 approach advocated here, will only serve as a basis for proper selection of experimental  
463 material to be taken to the laboratory - upon which further considerations will focus on, say,  
464 the potential factors involved in contaminant input and transport a.o. Note that these latter  
465 processes manifest themselves primarily in the Z-direction, where it is by no means a given  
466 that application of the same variographic approach (or geostatistical modelling) will  
467 necessary give meaningful results (see earlier footnote).

## 468 5. Conclusions

469 A pilot experiment aimed at an intrinsic 1-D soil heterogeneity characterization is a critical  
470 success factor for laboratory studies relying on field samples to provide the experimental  
471 pots, which for replicate and comparative study objectives need to be as similar as at all  
472 possible. As a case study the variographic results for sandy soils show that the distance  
473 between two sample spot must be less than 2.5 meters for the present set of organic  
474 compounds and soil type. Specific soil types and/or other analytes will in principle display  
475 different ranges and nugget effects, and hence our call for systematic deployment of the  
476 *variographic pilot experiment*, from which can be derived all necessary information for  
477 designing an optimal sampling plan e.g. identifying the analyte with the smallest range (for  
478 significantly correlated analytes). For the case of well-mixed soil components, a general PCA-  
479 approach for modelling a whole set of variograms may be useful in addition to individual  
480 analyte consideration.

481 Without this types of information, experimental fate study work is essentially devoid a valid  
482 basis as regards interpretation, scale-up and scientific generalisation of the experimental  
483 results back to the field scale. A large-scale 1-D transect sampling can reveal the inherent  
484 heterogeneity at all scales from the smallest local sampling equidistance up to the maximum  
485 experimental length scale studied. Variographic analysis was here employed successfully to  
486 soil heterogeneity at scales between 1 and 100 meters, other scenarios may require other  
487 numerical parameters, while the general approach remains identical.

488 The TOS-guided variogram pilot study approach illustrated here has a substantial carrying-  
489 over potential to geochemistry and environmental science, as well as other application areas.

490 It is even applicable to *dynamic systems*, i.e. to natural or technological processes in these  
491 realms.

492

#### 493 Acknowledgements

494 The authors gratefully acknowledge the Danish Research Council for Ph.D. stipend funding  
495 (Stipend No. 562/06-18-10028(6)) to ZK and valuable laboratory services and assistance  
496 provided by GEUS, Dept. of Geochemistry. The authors are grateful to GEUS' management for a  
497 positive attitude with respect to development and application of TOS' principles of proper  
498 representative sampling, which are not always recognized in geosciences communities to the  
499 extent necessary in general, alas neither at GEUS specifically.

500

501 References:

- 502 Adamchuk, Viacheslav I., Raphael a. Viscarra Rossel, David B. Marx, and Ashok K. Samal. 2011.  
503 Using Targeted Sampling to Process Multivariate Soil Sensing Data. *Geoderma* 163 (1-2):  
504 63-73. "
- 505 Arias-Estévez, Manuel, Eugenio López-Periago, Elena Martínez-Carballo, Jesús Simal-Gándara,  
506 Juan-Carlos Mejuto, and Luis García-Río. 2008. The Mobility and Degradation of  
507 Pesticides in Soils and the Pollution of Groundwater Resources. *Agriculture, Ecosystems &*  
508 *Environment* 123 (4): 247–60.
- 509 Badawi, Nora, Anders R Johnsen, Jan Sørensen, and Jens Aamand. 1999. Centimeter-Scale  
510 Spatial Variability in 2-Methyl-4-Chlorophenoxyacetic Acid Mineralization Increases with  
511 Depth in Agricultural Soil. *Journal of Environmental Quality* 42 (3): 683–89.
- 512 Bending, Gary D, Suzanne D Lincoln, R Sebastian, J Alun W Morgan, Jens Aamand, Sebastian R  
513 Sørensen, and Allan Walker. 2003. In-Field Spatial Variability in the Degradation of the  
514 Phenyl-Urea Herbicide Isoproturon Is the Result of Interactions between Degradative  
515 *Sphingomonas* Spp . and Soil pH. *App. Environ. Microbiol.* 69(2): 827-834.
- 516 Bending, Gary, Eve Shaw, and Allan Walker. 2001. Spatial Heterogeneity in the Metabolism  
517 and Dynamics of Isoproturon Degrading Microbial Communities in Soil. *Biology and*  
518 *Fertility of Soils* 33 (6): 484–89.
- 519 Boudreault, Jean-Philippe, Jean-Sébastien Dubé, Mirela Sona, and Eric Hardy. 2012. Analysis of  
520 Procedures for Sampling Contaminated Soil Using Gy's Sampling Theory and Practice.  
521 *The Science of the Total Environment* 425: 199–207.
- 522 Chappell, Adrian, and Raphael a. Viscarra Rossel. 2013. The Importance of Sampling Support  
523 for Explaining Change in Soil Organic Carbon. *Geoderma* 193-194: 323-325.
- 524 Crespin, M a, M Gallego, M Valcárcel, and J L González. 2001. Study of the Degradation of the  
525 Herbicides 2,4-D and MCPA at Different Depths in Contaminated Agricultural Soil.  
526 *Environmental Science & Technology* 35 (21): 4265–4270.
- 527 De Zorzi, Paolo, Sabrina Barbizzi, Maria Belli, Ales Fajgelj, Radojko Jacimovic, Zvonka Jeran,  
528 Umberto Sansone, and Marcel van der Perk. 2008. A Soil Sampling Reference Site: The  
529 Challenge in Defining Reference Material for Sampling. *Applied Radiation and Isotopes* :  
530 66 (11): 1588–1591.
- 531 Dictor Marie- Christine, Laurent Tessier and Guy Soulas. 1998. Reassessment of the K EC  
532 Coefficient of the Fumigation ± Extraction Method in a Soil Profile. *Soil Biology and*  
533 *Biochemistry* 30 (2): 119-127.
- 534 DS3077. 2013. Representative Sampling/ Horizontal Standard. *Danish Standard Authority*  
535 44:1–38.

- 536 Esbensen, Kim H., Hans Henrik Friis-Petersen, Lars Petersen, Jens Bo Holm-Nielsen, and Peter  
537 P. Mortensen. 2007. Representative Process Sampling — in Practice: Variographic  
538 Analysis and Estimation of Total Sampling Errors (TSE). *Chemometrics and Intelligent*  
539 *Laboratory Systems* 88 (1): 41–59.
- 540 Esbensen, Kim H., Claudia Paoletti, and Pentti Minkkinen. 2012a. “Representative Sampling of  
541 Large Kernel Lots I. Theory of Sampling and Variographic Analysis.” *TrAC Trends in*  
542 *Analytical Chemistry* 32: 154–164.
- 543 Esbensen, Kim H., Claudia Paoletti, and Pentti Minkkinen. 2012b. Representative Sampling of  
544 Large Kernel Lots III. General Considerations on Sampling Heterogeneous Foods. *TrAC*  
545 *Trends in Analytical Chemistry* 32: 178–184.
- 546 Esbensen, Kim H., and Claas Wagner. 2014. Theory of Sampling (TOS) versus Measurement  
547 Uncertainty (MU) – A Call for Integration. *TrAC Trends in Analytical Chemistry* 57: 93–  
548 106.
- 549 Esbensen, Kim H., and Rodolfo J. Romanach. (2015) Counteracting soil heterogeneity sampling  
550 for environmental studies (pesticide residues, contaminants transformation) - TOS is  
551 critical. Proceedings 7.th World Conference on Sampling and Blending (WCSB7), p.205-  
552 209.
- 553 Gerlach, Robert W., David E. Dobb, Gregory a. Raab, and John M. Nocerino. 2002. Gy Sampling  
554 Theory in Environmental Studies. 1. Assessing Soil Splitting Protocols. *Journal of*  
555 *Chemometrics* 16 (7): 321–328.
- 556 Ghafoor, Abdul, Nicholas J. Jarvis, Tomas Thierfelder, and J Stenström. 2011. Measurements  
557 and Modeling of Pesticide Persistence in Soil at the Catchment Scale. *The Science of the*  
558 *Total Environment* 409 (10) 1900-1908.
- 559 Gy, P. M. 1998. *Sampling for Analytical Purposes*. 1st ed. Chichester, West Sussex, UK: John  
560 Wily & Sons, 172pp, ISBN: 978-0-471-97956-2.
- 561 Johnsen, Anders R, Philip J Binning, Jens Aamand, Nora Badawi, and Annette E Rosenbom.  
562 2013. The Gompertz Function Can Coherently Describe Microbial Mineralization of  
563 Growth-Sustaining Pesticides. *Environmental Science & Technology* 47 (15): 8508–8514.
- 564 Johnsen, Anders R, Bjarne Styrihave, and Jens Aamand. 2014. Quantification of Small-Scale  
565 Variation in the Size and Composition of Phenanthrene-Degrader Populations and PAH  
566 Contaminants in Traffic-Impacted Topsoil. *FEMS Microbiology Ecology*, 88: 84-93.
- 567 Kardanpour, Zahra, Ole S. Jacobsen, and Kim H. Esbensen. 2014. Soil Heterogeneity  
568 Characterization Using PCA (Xvariogram) – Multivariate Analysis of Spatial Signatures for  
569 Optimal Sampling Purposes. *Chemometrics and Intelligent Laboratory Systems* 136:24–35.
- 570 Kardanpour, Z, Jakobsen, O.S. & Esbensen, K.H. 2015 Counteracting soil heterogeneity  
571 sampling for environmental studies (pesticide residues, contaminants transformation) -

- 572 TOS is critical. Proceedings 7.th World Conference on Sampling and Blending (WCSB7),  
573 p.205-209.
- 574 Li, B G, J Cao, W X Liu, W R Shen, X J Wang, and S Tao. 2006. Geostatistical Analysis and Kriging  
575 of Hexachlorocyclohexane Residues in Topsoil from Tianjin, China. *Environmental*  
576 *Pollution* 142 (3): 567–575.
- 577 Lin, Qinghuo, Hong Li, Wei Luo, Zhaomu Lin, and Baoguo Li. 2013. Optimal Soil-Sampling  
578 Design for Rubber Tree Management Based on Fuzzy Clustering. *Forest Ecology and*  
579 *Management* 308: 214–222.
- 580 Martens, Harald, and Tormod Næs. 1991. *Multivariate Calibration*. Edited by John Wiley &  
581 Sons. Chichester, West Sussex, UK, 438p, ISBN: 978-0-471-93047-1.
- 582 Minkkinen, Pentti, Kim H. Esbensen, and Claudia Paoletti. 2012. Representative Sampling of  
583 Large Kernel Lots II. Application to Soybean Sampling for GMO Control. *TrAC Trends in*  
584 *Analytical Chemistry* 32: 165–177.
- 585 Mulder, V.L., S. de Bruin, and M.E. Schaepman. 2013. Representing Major Soil Variability at  
586 Regional Scale by Constrained Latin Hypercube Sampling of Remote Sensing Data.  
587 *International Journal of Applied Earth Observation and Geoinformation* 21: 301–310.
- 588 Petersen, Lars, Casper K. Dahl, and Kim H. Esbensen. 2004. Representative Mass Reduction in  
589 Sampling—a Critical Survey of Techniques and Hardware. *Chemometrics and Intelligent*  
590 *Laboratory Systems* 74 (1): 95–114.
- 591 Petersen, Lars, and Kim H Esbensen. 2006. Representative Process Sampling for Reliable Data  
592 Analysis — a Tutorial. *Journal of Chemometrics* 19: 625–647.
- 593 Petersen, Lars, Pentti Minkkinen, and Kim H. Esbensen. 2005. Representative Sampling for  
594 Reliable Data Analysis: Theory of Sampling. *Chemometrics and Intelligent Laboratory*  
595 *Systems* 77 (1-2): 261–277.
- 596 Rasmussen, Jim, Jens Aamand, Per Rosenberg, Ole S Jacobsen, and Sebastian R Sørensen. 2005.  
597 Spatial Variability in the Mineralisation of the Phenylurea Herbicide Linuron within a  
598 Danish Agricultural Field: Multivariate Correlation to Simple Soil Parameters. *Pest*  
599 *Management Science* 61 (9): 829–837.
- 600 Rodriguez-Cruz, M. Sonia, Julie E. Jones, and Gary D. Bending. 2006. Field-Scale Study of the  
601 Variability in Pesticide Biodegradation with Soil Depth and Its Relationship with Soil  
602 Characteristics. *Soil Biology and Biochemistry* 38 (9): 2910–2918.
- 603 Rosenbom, Annette E, Philip J Binning, Jens Aamand, Arnaud Dechesne, Barth F Smets, and  
604 Anders R Johnsen. 2014. Does Microbial Centimeter-Scale Heterogeneity Impact MCPA  
605 Degradation in and Leaching from a Loamy Agricultural Soil?. *The Science of the Total*  
606 *Environment* 472:90–98.

- 607 Sørensen, Sebastian R, Gary D Bending, Carsten S Jacobsen, Allan Walker, and Jens Aamand.  
608 2003. Microbial Degradation of Isoproturon and Related Phenylurea Herbicides in and  
609 below Agricultural Fields. *FEMS Microbiology Ecology* 45 (1): 1–11.
- 610 Sørensen, Sebastian R, Anne Schultz, Ole S Jacobsen, and Jens Aamand. 2006. Sorption,  
611 Desorption and Mineralisation of the Herbicides Glyphosate and MCPA in Samples from  
612 Two Danish Soil and Subsurface Profiles. *Environmental Pollution* 141 (1): 184–194.
- 613 Tate K.R., Ross D. J. and Feltham C.W. 1988. A Direct Extraction Method to Estimate Soil  
614 Microbiology C: Effects of Experimental Variables and Some Different Calibration  
615 Procedures. *Soil Biology and Biochemistry* 20: 329–335.
- 616 Thurman, Elisabeth A Scribner. 2008. A Decade of Measuring , Monitoring , and Studying the  
617 Fate and Transport of Triazine Herbicides and Their Degradation Products in  
618 Groundwater , Surface Water , Reservoirs , and Precipitation by the US Geological Survey.  
619 *The Triazine Herbicides*, 451-475.
- 620 Torstensson, N T L, and J Stark. 1975. The Effect of Repeated Applications of 2 , 4-D and MCPA  
621 on Their Breakdown in Soil. *Weed Research* 15: 159–164.
- 622 Totaro, Sara, Paola Coratza, Caterina Durante, Giorgia Foca, Mario Li Vigni, Andrea Marchetti,  
623 Mauro Marchetti, and Marina Cocchi. 2013. Soil Sampling Planning in Traceability Studies  
624 by Means of Experimental Design Approaches. *Chemometrics and Intelligent Laboratory*  
625 *Systems* 124:14–20.
- 626 Walker, A, M Jurado-Exposito, G D Bending, and V J Smith. 2001. Spatial Variability in the  
627 Degradation Rate of Isoproturon in Soil. *Environmental Pollution* 111 (3): 407–415.
- 628