- 1 Characterising the within-field scale spatial variation of nitrogen in a
- 2 grassland soil to inform the efficient design of in-situ nitrogen sensor
- 3 networks for precision agriculture
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#### ABSTRACT

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The use of in-situ sensors capable of real-time monitoring of soil nitrogen (N) may facilitate improvements in agricultural N-use efficiency (NUE) through better fertiliser management. The optimal design of such sensor networks, consisting of clusters of sensors each attached to a data logger, depends upon of the spatial variation of soil N and the relative cost of the data loggers and sensors. The primary objective of this study was to demonstrate how in-situ networks of N sensors could be optimally designed to enable the cost-efficient monitoring of soil N within a grassland field (1.9 ha). In the summer of 2014, two nested sampling campaigns (June & July) were undertaken to assess spatial variation in soil amino acids, ammonium (NH<sub>4</sub><sup>+</sup>) and nitrate (NO<sub>3</sub><sup>-</sup>) at a range of scales that represented the within (less than 2 m) and between (greater than 2 m) data logger/sensor cluster variability. Variance at short range (less than 2 m) was found to be dominant for all N forms. Variation at larger scales (greater than 2 m) was not as large but was still considered an important spatial component for all N forms, especially NO<sub>3</sub>. The variance components derived from the nested sampling were used to inform the efficient design of theoretical in-situ networks of NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> sensors based on the costs of a commercially available data logger and ionselective electrodes (ISEs). Based on the spatial variance observed in the June nested sampling, and given a budget of £5000, the NO<sub>3</sub> field mean could be estimated with a 95% confidence interval width of 1.70 µg N g<sup>-1</sup> using 2 randomly positioned data loggers each with 5 sensors. Further investigation into "aggregate-scale" (less than 1 cm) spatial variance revealed further large variation at the sub 1-cm scale for all N forms. Sensors, for which the measurement represents an integration over a sensor-soil contact area of diameter less than 1 cm, would therefore, be subject to further spatial variability and local replication at scales less than 1 cm would be needed to maintain the precision of the resulting field mean estimation. Adoption of in-situ sensor networks will depend upon the development of suitable low-cost

- 37 sensors, demonstration of the cost-benefit and the construction of a decision support system
- that utilises the generated data to improve the NUE of fertiliser N management.

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- 40 Keywords: Fertilizer management; soil heterogeneity; dissolved organic nitrogen; precision
- 41 agriculture; nitrogen-use efficiency; nutrient cycling

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#### 1. Introduction

Improving nitrogen (N) use efficiency (NUE) remains one of the key challenges for global agriculture (Cassman et al., 2002; Robertson and Vitousek, 2009) and is essential for the success of sustainable intensification (Tilman et al., 2011). The deleterious environmental effects and economic costs of diffuse N pollution from farmland in Europe, where N has been applied in excess of crop requirement, are well documented (Sutton et al., 2011).

One often-cited approach to reduce N losses and improve NUE, is to ensure synchronicity of N supply with crop demand (Shanahan et al., 2008; Robertson and Vitousek, 2009), although, achieving this in practice is challenging due to the complex nature of the soil-plant system. Precision agriculture (PA) attempts to address this issue by reducing uncertainties surrounding the measurement of key variables to determine optimum N fertiliser management (Pierce and Nowak, 1999; Dobermann et al., 2004). Temporal variations in growing conditions, both within and between seasons may lead to considerable differences in optimum N fertiliser requirement and hence, inefficiencies in N fertiliser-use if temporal variations are not considered (Lark and Wheeler, 2003; McBratney et al., 2005; Shahandeh et al., 2005; Shanahan et al., 2008; Deen et al., 2014). However, conventional soil sampling techniques, coupled with laboratory analysis is expensive, labour-intensive, and time-consuming and cannot provide real-time data of sufficient resolution to accurately inform PA management (Sylvester-Bradley et al., 1999; Kim et al., 2009).

A number of different approaches have been used to address this issue. Crop canopy sensing techniques, for determination of plant N status, are now in commercial use and can be used to inform variable rate fertiliser application (e.g. wheat, maize; Raun and Johnson, 1999; Diacono et al., 2013). Whilst the advantages of this approach in some situations have been evidenced (Diacono et al., 2013), plant N status and yield is the product of many variables and may not always correlate with soil mineral N status. On-the-go soil sampling

for NO<sub>3</sub>, using electrochemical sensor platforms attached to agricultural vehicles have been developed (Adsett et al., 1999) and, for the case of pH, commercialised (Adamchuk et al., 1999). The results have been used to develop field nitrate maps (Sibley et al., 2009) which could be used to define within-field management zones and to calculate variable fertiliser application rates. On-the-go sampling is generally more spatially intensive than manual field sampling, allowing better spatial resolution, although key information on how soil mineral N varies over small spatial scales may not be obtained. This can lead to increased uncertainties of interpolative predictions, especially if the sample volume is small (Schirrmann and Domsch, 2011). Furthermore, increasing the temporal resolution of this approach requires additional economic costs and as both these approaches rely on reactive management, crucial changes in soil mineral N status may be missed.

One approach, which has yet to be explored, is the use of in-situ sensors capable of monitoring soil mineral N in real time. At the time of writing, there are no such quasipermanent field sensors in use commercially. However, potential for the development and deployment of such sensors exists (Shaw et al., 2014). For example, ion-selective electrodes (ISEs) have many characteristics suitable for soil sensing networks. They are relatively cheap, simple to use, require no mains electrical power supply and the concentration of the target ion can be easily calculated via a pre-calibration. Nitrate (NO<sub>3</sub><sup>-</sup>) ISEs have previously been successfully deployed for monitoring streams and agricultural drainage ditches (Le Goff et al., 2002; Le Goff et al., 2003) as well as for on-the-go soil sampling of agricultural soils (Sinfield et al., 2010) and on-farm rapid tests for soil NO<sub>3</sub><sup>-</sup> (Shaw et al., 2013). Similarly, ammonium (NH<sub>4</sub><sup>+</sup>) ISEs have been used for water monitoring in a variety of situations (Schwarz et al., 2000; Müller et al. 2003). Direct soil measurement, which is essential for the success of in-situ monitoring, has been shown to be possible (Ito et al., 1996; Adamchuk et al., 2005), although improvements in accuracy and robustness of the sensing membrane are

required. Increasing use of nano technologies for the construction of electrochemical sensors may result in significant advances in sensor performance (Arrigan, 2004; Atmeh and Alcock-Earley, 2011).

Using in-situ sensor networks may enable a step away from predetermined fertiliser N recommendations (Defra, 2010) to a more dynamic system that responds in real-time to changes in growing conditions. It potentially has many benefits compared to on-the-go soil sampling and crop canopy sensing. The data provided by in-situ sensors will be of significantly higher temporal resolution, negating the need for repeated sampling surveys throughout the year, which represent an economic cost to the farmer. Furthermore, this may enable more accurate timing of fertiliser application, reducing the risk of yield penalties caused by N-nutrition deficiencies, and the risk of N transfers to water and air as a result of excessive fertiliser N applications. It is also likely that the data generated at a high temporal resolution by an in-situ sensor networks will increase knowledge of the controls of soil N processes and thus enable development of models which allow for a proactive approach to fertiliser N management. However, there is a trade-off to be made. The increase in temporal resolution gained from in-situ sensor networks may be offset by the costs of achieving sufficient spatial resolution.

It is, therefore, important that consideration is made as to how such sensor networks could be optimally designed to enable sufficiently precise estimates of mean field or management zone (MZ) soil N at minimum cost. Two factors complicate this. First, each sensor must attach to a data logger, over a relatively short distance. As data loggers cost more per unit than sensors, and one logger can support several sensors, then feasible networks will comprise sensor clusters, each associated with a logger. As such, sensor networks can be regarded as multi-scale sampling schemes with data loggers (primary units) and sensors (secondary units) randomly placed in an area around each data logger. Second,

soil N is variable, at multiple scales. Efficiently designed sensor networks will have sufficient replication at the most variable scales, achieving this within the constraints of feasible clustered designs. As shown by de Gruijter et al. (2006), the optimum configuration of such a sampling scheme depends on the relative costs of additional primary and secondary units and the within- and between-primary unit variability.

The primary objective of this study was to address the above problem and to investigate how the design of a theoretical network of in-situ soil N sensors could be optimised on a cost-precision basis, to enable monitoring of soil N concentrations in a grassland field. As seen in the discussion above, the feasibility and optimal design of sensor networks depends on the variability of the target properties at different within-field scales. An effective way to collect such information is by spatially nested sampling, which has been used previously to characterise the spatial variation in a range of soil-related variables (Lark, 2011). In nested spatial sampling, sample sites are arranged in a nested hierarchical design which allows the partition of the variance of the measured variable into components associated with a set of pre-determined scales. At the highest level of the hierarchy sample, points are arranged in clusters associated with "mainstations" which may be at randomly-located sites or on nodes of a grid or transect. Within a mainstation, sample points may be divided between two or three stations at level 2 which are separated from each other by some fixed distance. Within each level-2 station, sample points may be ordered at further nested spatial scales.

As such, spatially nested sampling was performed at a range of scales to characterise the within-field spatial variability of amino acids,  $NH_4^+$  and  $NO_3^-$ . These results were then used to explore the optimisation of a  $NH_4^+$  and  $NO_3^-$  in-situ sensor network design on the basis of both cost and precision. Finally, the potential and challenges of implementing this approach within a PA framework are discussed.

#### 2. Materials and methods

#### 2.1. Field site and soil characteristics

The field used for this study is located within the Henfaes Research Station Abergwyngregyn, Wales, UK (53°14′N 4°01′W). The site has a temperate, oceanic climate, receives an average annual rainfall of 1250 mm and has a mean annual soil temperature at 10 cm depth of 11 °C. The field is roughly rectangular with a perimeter of 559 m and an area of 1.91 ha. It has an average altitude of 12.1 m asl with a slope of 1.5% in a northerly aspect. It is a semi-permanent sheep-grazed grassland, dominated by *Lolium perenne* L. The current ley was established by direct drill in April 2009 using a perennial and hybrid ryegrass mix. The field has been used for both all year round grazing and silage production since 2009, receiving an annual inorganic fertiliser input of between 100 – 130 kg N ha<sup>-1</sup> in addition to potassium (K), phosphate (P) and sulphur (S) at recommended rates. Lime has also been applied when necessary to restore the pH to a target value of 6.5. In 2014, inorganic fertiliser was applied on 12/5/14 and 11/7/14 at a rate of N:P:K 50:10:10 and 60:4:0 kg ha<sup>-1</sup>, respectively. The field was grazed until 9/6/14 and the field remained sheep free until the 2/9/14. The soil is a free draining Eutric Cambisol with a sandy clay loam texture and a fine crumb structure.

To assess the chemical characteristics of the soil, replicate samples (n=4) were collected from 4 randomly located points within the field. For each sample, the vegetation was removed from an area approximating  $30 \times 30$  cm and soil was collected to a depth of 10 cm, representing the Ahp horizon (an Ahp horizon is an Ah horizon which has been subjected to cultivation). This sampling design is not related to the spatially nested design described in section 2.2. The soil was placed in gas-permeable polyethylene bags and transported to the laboratory in a refrigerated box. All of the following procedures were performed on the same

day as field sampling. Soil pH and electrical conductivity were determined in a 1:2.5 (w/v) soil:distilled water suspension using standard electrodes. Moisture content was determined by drying for 24 h at 105 °C. Total C and N were determined with a TruSpec CN analyser (Leco Corp., St Joseph, MI, USA). Dissolved organic carbon (DOC) and dissolved organic nitrogen (DON) were measured in soil extracts (0.5 M K<sub>2</sub>SO<sub>4</sub>, 1:5 w:v) using an Analytik Jena Multi N/C 2100S (AnalytikJena, Jena, Germany). Chloroform fumigation and incubation (t = 7days) of 2 g (n = 4) of fresh soil was performed to determine microbial biomass C and N according to Voroney et al. (2008) ( $K_{EC} = 0.35 \ K_{EN} = 0.5$ ). Exchangeable cations were extracted using 0.5 M acetic acid (Sparks, 1996) and the filtered extracts analyzed using flame emission spectroscopy (Sherwood 410 flame photometer; Sherwood Scientific, Cambridge, UK). Extractable phosphorus (P) was determined by extraction with 0.5 M acetic acid with subsequent colorimetric analysis using the molybdate blue method of Murphy and Riley (1962). Basal soil respiration was determined in the laboratory at 20 °C using an SR1 automated multichannel soil respirometer (PP Systems Ltd., Hitchin, UK) and steady state CO<sub>2</sub> production rates recorded after 24 h. Potentially mineralisable N was determined using an anaerobic incubation method based on Keeney (1982). Briefly, 5 g field moist soil was place in a 50 ml centrifuge tube, which was then filled to the top with de-ionized H<sub>2</sub>O and the tubes sealed. Soils were subsequently incubated in the dark at 40 °C for 7 d. The difference in  $NH_4^+$  content between t = 0 and t = 7 d was attributed to N mineralization.

Above ground biomass was sampled on 26/6/2014. Replicate  $1 \times 1$  m blocks (n = 4) were chosen at random from within the field. The vegetation was cut to ground level, stored in paper bags and subsequently oven-dried at 80 °C to determine dry matter content. A summary of the results is shown in Table 1.

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### 2.2. Sampling design and protocol

Nested sampling for spatial variability: The aim of the sampling was to characterize the variability of plant-available N forms – amino acid-N,  $NH_4^+$  and  $NO_3^-$  – at a range of spatial scales relevant to planning the design of an in-situ sensor network. In particular, it was necessary to examine the relative importance of variance between and within local regions each of which might be represented by a cluster of soil N sensors deployed around a single data logger such that the maximum distance between any two sensors is about 2 m. In a grassland environment it was expected that one of the main sources of variation in soil N would be the uneven and relatively random distribution of urine patches of linear dimensions about 40 cm (Bogaert et al., 2000; Selbie et al., 2015). Variation at larger scales may also be important due to preferential use of certain areas of the field such as tracks, areas of shade and around drinking troughs (Bogaert et al., 2000), which may be reflected in local gradients in soil chemistry. The study field is broadly homogenous in terms of its topography and soil type. Furthermore, a visual inspection of the field revealed no obvious large-scale gradients in vegetation condition which is likely to reflect the broadly homogenous nature of the soil. Previously, the field has received uniform management in terms of its fertiliser and lime inputs and grazing regime. Because of these factors, it was decided to treat the field as singular management unit with a singular mean rather than subdivide the field into separate management zones.

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Given these considerations, a nested sampling protocol was designed with length scales within each mainstation of 1 cm, 10 cm (intermediate between the fine scale and urine patch scale), 50 cm (urine patch scale) and 2 m (upper bound on the "within-region served by a sensor cluster" scale). To assess spatial variation at larger scales, mainstations were distributed by stratified random sampling with the target field divided into four quarters (strata) of equal area. Four mainstations were established at independently and randomly-selected locations within each quarter (stratum), giving a total of 16 mainstations. The design

of the sampling scheme within each mainstation, was obtained by the optimization procedure of Lark (2011) on the assumption of a fractal or quasi-fractal process in which the variance is proportional to the log of the spatial scale. The objective function was the mean estimation variance of the variance components. With 12 samples per mainstation the total sample size was 192. The sample sites were then selected at each mainstation by randomizing the direction of the vectors between the substations at each level of the design shown in Figure 1, while keeping the lengths of the vectors fixed. For practical purposes, sampling was split over 2 successive days, with 2 strata sampled on day 1 and two on day 2, giving a total of 8 mainstations and 96 samples per day. No duplicate sampling took place as each sample site was visited only once over the 2-day period.

An initial nested sampling campaign was performed over 2 days on the 4<sup>th</sup> and 5<sup>th</sup> June, 2014. Following this, all sheep were removed and the field remained ungrazed until 2<sup>nd</sup> September, 2015. A further nested sampling campaign was performed on the 31<sup>st</sup> July and 1<sup>st</sup> August, 2014, 3 weeks after the field received a N fertiliser input of 60 kg N ha<sup>-1</sup>. These are subsequently referred to as the June nested sampling and the July nested sampling respectively. Sample site locations were set up the day before sampling took place. At each sampling location a soil corer, of diameter 1 cm, was used to sample soil. A 5 cm soil core from between depths of 5 -10 cm was sampled and placed in gas-permeable plastic bags, and stored in a refrigerated box. This depth was chosen as it represents the middle of the rooting zone and would make installation of any in-situ sensor a straight forward process. Following the sampling event, the samples were transferred immediately to the laboratory where they were refrigerated at 4 °C. Extraction of soluble N from soil was performed on the soil cores on the same day as sampling as described below. During the second nested sampling event, duplicate sub-sampling and chemical analysis were performed on 4 out of the 12 samples

from each mainstation in order to make an assessment of the error variance attributable to subsampling and analytical error.

Soil properties are also likely to vary at sub-core (less than 1 cm) scales (Parkin, 1987; Stoyen et al., 2000). As such, a further sampling design and protocol was developed and performed on the 25<sup>th</sup> June, 2014 to investigate how this micro-heterogeneity affected the spatial variability of N forms at the "aggregate scale" (less than 1 cm). Two sampling locations were chosen at random within each of the 4 strata. At each location, a pair of samples were taken, using the protocol described above, with a distance of 1 cm between each sample. This resulted in a total of 16 core samples. On return to the laboratory the cores where broken apart and 4 "aggregates" of weight 60 – 80 mg were collected (diameters ca. 1-2 mm). These aggregates were then extracted for soluble N and analysed using the protocol described below.

#### 2.3. Extraction and chemical analysis of soil samples

All soil extractions were performed on the same day as sample collection, according to the following protocol. Samples were crumbled by hand, in order to prevent sieving induced N mineralisation (Jones and Willett, 2006; Inselsbacher, 2014). Large stones, roots and vegetation were removed prior to gentle mixing of the sample. To further reduce mineralisation of organic N forms, sub-samples of field-moist soil (2 g) were extracted on ice (175 rev min<sup>-1</sup>, 15 min) using cooled (5 °C) 0.5 M  $K_2SO_4$  at a soil: extractant ratio of 1:5 (w:v) (Rousk and Jones, 2010). The extracts were centrifuged (4,000 g, 15 min), and the resulting supernatant collected and frozen (-18°C) to await chemical analysis. The protocol differed slightly for the soil aggregate samples. Each aggregate, of weight 60 – 80 mg, was placed in a 1.5 ml Eppendorf® micro-centrifuge tube and crumbled gently using a metal spatula. The soil was then extracted in 500  $\mu$ l of 0.5 M  $K_2SO_4$  as described above. Total free

amino acids (referred to as amino acids) were determined by the *o*-phthaldialdehyde spectrofluorometric method of Jones et al. (2002). NH<sub>4</sub><sup>+</sup> was determined by the salicylate-nitroprusside colorimetric method of Mulvaney (1996) and NO<sub>3</sub><sup>-</sup> by the colorimetric Griess reaction of Miranda et al. (2001) using vanadate as the catalyst.

## 2.4. Statistical analysis

Nested Sampling: The aim of the statistical analysis was to compute the variance components attributable to each of the spatial-scales in order to inform the optimisation of the sensor network design. After Box-Cox transformations (see section 2.5 for details of transformations), the n data from the nested sampling may be analysed according to the following statistical model (Webster and Lark, 2013). An  $n \times 1$  vector of observations,  $\mathbf{y}$ , is regarded as a realization of a random variate,  $\mathbf{Y}$ , where

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$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{U}_{s}\boldsymbol{\eta}_{s} + \mathbf{U}_{m}\boldsymbol{\eta}_{m} + \mathbf{U}_{2}\boldsymbol{\eta}_{2} + \mathbf{U}_{0.5}\boldsymbol{\eta}_{0.5} + \mathbf{U}_{0.1}\boldsymbol{\eta}_{0.1} + \mathbf{U}_{0.01}\boldsymbol{\eta}_{0.01} + \mathbf{U}_{r}\boldsymbol{\eta}_{r}.$$
280 (1)

**X** is a  $n \times p$  design matrix which represents fixed effects in the model (e.g. p levels of a categorical factor, or p continuous covariates) and  $\beta$  is a length-p vector of fixed effects coefficients. In this analysis the fixed effects were different means for data collected on two successive days, as it was not possible logistically to sample on one day. Strata were randomly allocated to days (two strata per day) so between-stratum variation is not confounded with any temporal effect. There are 4 strata in the sampling design, and  $\mathbf{U}_s$  is a  $n \times 4$  design matrix for the strata. If the ith observation is in stratum j then  $\mathbf{U}_s$  [i, j] = 1 and all other elements in the ith row are zero. The design matrix associates each observation with one of 4 random values in the random variate  $\mathbf{\eta}_s$ . These values are assumed to be independent and identically distributed Gaussian random variables with a mean of zero and a variance  $\sigma^2_s$  which is the between-stratum variance component. Similarly,  $\mathbf{U}_m$  is a  $n \times 16$  design matrix

for the mainstations, and the variance of  $\eta_m$  is the between-mainstation variance component. The terms with subscripts 2, 0.5. 0.1 and 0.01 represent the design matrices and random effects for the components of variation associated with the 2-m, 0.5-m, 0.1-m and 0.01-m scales respectively. If duplicate material from some or all of the soil specimens is analysed then the random effect  $\eta_r$  which represents the variation due to subsampling and analytical variation can be estimated, otherwise it is a component of the variation estimated for the finest spatial scale.

Under the linear mixed model Y has covariance matrix **H** where

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$$\mathbf{H} = \sigma_{s}^{2} \mathbf{U}_{s} \mathbf{U}_{s}^{T} + \sigma_{m}^{2} \mathbf{U}_{m} \mathbf{U}_{m}^{T} + \sigma_{2}^{2} \mathbf{U}_{2} \mathbf{U}_{2}^{T} + \sigma_{0.5}^{2} \mathbf{U}_{0.5} \mathbf{U}_{0.5}^{T} + \sigma_{0.1}^{2} \mathbf{U}_{0.1} \mathbf{U}_{0.1}^{T}$$

$$+ \sigma_{0.01}^{2} \mathbf{U}_{0.01}^{T} \mathbf{U}_{0.01}^{T} + \sigma_{r}^{2} \mathbf{U}_{r} \mathbf{U}_{r}^{T},$$
302 (2)

and the superscript T denotes the transpose of a matrix. The parameters of this matrix are therefore the variance components, and these can be estimated by residual maximum likelihood (REML), see Webster and Lark (2013). Once this has been done then the fixed effects coefficients in the model can be estimated by generalized least squares (see Lark and Cullis, 2004). Note that there is an explicit assumption that the data are a realization of a Gaussian random variable with mean  $X\beta$  dependent on the fixed effects and coefficients.

Because all sampling could not be done in one day the sampling day was randomized within strata, so as not to be confounded with the spatial variance components of interest. For this reason, it is regarded as a fixed effect in the model. The significance of the between-day effect was tested with the Wald statistic as discussed in Lark and Cullis (2004).

The significance of a random effect in the model can be tested by comparing the residual log-likelihood for a model with the term dropped  $(L^-)$  with the residual log-likelihood for the full model (all random effects, L). Any variance accounted for by a term which is dropped will contribute to variance at lower levels in the hierarchy (finer spatial

scales) for the dropped model. For this reason the ultimate component of the model ( $\eta_r$  when there are duplicate analyses and  $\eta_{0.01}$  otherwise) cannot be dropped. Dropping a term from the model will usually reduce the log-likelihood (and will not increase it). Whether the reduction in likelihood is strong enough evidence that the inclusion of the term in the full model is justified can be assessed by computing Akaike's information criterion (AIC), A, for each model:

$$A = -2L + 2P \tag{3}$$

where *P* is the number of parameters in the model. The AIC penalizes model complexity, by selecting the model with smaller AIC, one minimises the expected information loss through the selection decision (Verbeke and Molenberghs, 2000).

Aggregate scale sampling: After Box-Cox transformations (see section 2.5 for details of transformations) the n data collected to investigate variation within cores were analysed according to the following statistical model. An  $n \times 1$  vector of observations, y, is regarded as a realization of a random variate, Y, where

$$\mathbf{Y} = \mathbf{X}\boldsymbol{\beta} + \mathbf{U}_{s}\boldsymbol{\eta}_{s} + \mathbf{U}_{p}\boldsymbol{\eta}_{p} + \mathbf{U}_{c}\boldsymbol{\eta}_{c} + \mathbf{U}_{a}\boldsymbol{\eta}_{a},$$

332 (4)

As in Equation (1),  $\mathbf{X}$  is a design matrix for fixed effects and  $\boldsymbol{\beta}$  is a vector of fixed effects coefficients (here just a constant mean). Again, as in Equation (1), Us is a  $n \times 4$  design matrix for the strata and  $\eta_s$  is assumed to be an independent and identically distributed Gaussian random variate with a mean of zero and a variance  $\sigma_s^2$ . In the same way  $\mathbf{U}_p$  and  $\eta_p$  are the design matrix and the random variate for the between-pair within-stratum effect, with variance  $\sigma_p^2$ ;  $\mathbf{U}_c$  and  $\eta_c$  are the design matrix and the random variate for the between-core within-pair component, with variance  $\sigma_c^2$  and  $\mathbf{U}_a$  and  $\mathbf{\eta}_a$  are the design matrix and the random variate for the between-aggregate within-core component, with variance  $\sigma_a^2$ . This latter component is effectively the residual as there are no duplicate measurements on any

aggregate. The same method based on the AIC was used to assess the evidence for including each term in the model above the between-aggregate effect.

# 2.5. Data transformations

The REML estimator for random effects parameters makes an explicit assumption that the random variation in the model is normally distributed. This assumption is of particular importance in the model with nested random effects, because these must be modelled as independent additive components. This is plausible for normal random variables, but not in general otherwise. For this reason, it was necessary to transform the data so that the residuals from any fixed effects could be regarded as normal. To this end, a Box-Cox procedure from the MASS package in R (Venables and Ripley, 2002) was used to apply the Box-Cox transform. Under this transform the variable x is transformed to a normal variable y by finding a maximum likelihood estimate of the parameter  $\lambda$  such that

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$$y = \begin{cases} \frac{x^{\lambda - 1}}{\lambda}, & \lambda \neq 0 \\ \ln(x), & \lambda = 0 \end{cases}$$
 (6)

Note that the log-transformation is a special case of the Box-Cox, with  $\lambda$ =0. In this study, the log-normal transformation was used for cases where the 95% confidence interval of  $\lambda$  included 0. Where this was not the case, the Box-Cox transformation with the maximum likelihood estimate of  $\lambda$  was used.

A disadvantage of transformation is that the results are not on familiar scales, and that the relative and absolute magnitudes of the variances depend on the transformation and the mean. Section 2.6 describes how the variance components estimated on the transformed scales were used to calculate the width of confidence intervals for estimates of field means for forms of N estimated from different sensor arrays by a simple back-transformation of confidence limits on the transformed scale. However, it is not possible to perform a

comparable simple back-transformation of variance components for general interpretation of the variability at different scales. This would be true of any Box-Cox transformation.

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Two sets of results have been presented. The first are the variance components on the transformed scale, the scale of measurement on which each data set can most plausibly be modelled as an additive combination of random components at different scales (Panel A, Figs. 2 & 3; tables 3, 5 & 7). For the different scales, percentages of the total accumulated variances have been calculated, and these values are referred to in the results section to aid comparison between the different N forms and sampling events. Secondly, a non-parametric statistic was computed on the original scales of measurement for each nested analysis. Using the estimated variance components on the transformed scale, 10,000 data values in a nested configuration were simulated with, in the case of the spatially nested sample, pairs of points in contrasting strata, pairs of points at different locations within a stratum and pairs of points within strata separated by 2 m, 0.5 m, 0.1 m and 0.01m respectively. Each simulated value was then back-transformed to the original scale of measurement by inverting the Box-Cox transformation. All these pairs of observations were then examined on the original scale, computing a non-parametric and robust measure of the variability of the differences. This is the median absolute deviation from the median (MAD). If one considers all pair comparisons over 2 m, for example, the median difference is first computed and then the absolute difference between this median difference and each individual pair difference is computed, and the median of all these values is extracted. The MAD is a measure of variability on the original scale of measurement (like the standard deviation). These were extracted for all variables and plotted alongside cumulative plots of the variance components on the transformed scale to aid the interpretation of these latter plots and give an indication of the magnitude of variation that can be expected for measurements on the original scale (panel B, Figs 2 & 3). The same process was followed after the analysis of aggregate-scale variation to

compute the MAD of comparisons between two aggregates within a core, between two cores in a pair, between two sites within a stratum, and between pairs of strata, these are presented in Table 8. MAD values are also explicitly reported in the results section with units of  $\mu g \, N \, g^{-1}$ .

### 2.6. Optimising the design of an in-situ sensor network

The transformed variance components derived from the nested sampling and subsequent statistical analysis were used to examine the theoretical performance of different designs of in-situ soil NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> sensor networks. When considering the optimal design, two factors must be considered. Firstly, what is the required level of precision for the estimation of the field mean and how many sensors and data loggers are required to achieve this? Secondly, how can the design be optimised in-terms of achieving a desired level of precision at minimum cost? Alternatively, it may be useful to explore how to design the network to achieve the highest precision possible given a certain budget restriction.

To estimate the level of precision associated with a particular sensor network design, the between-sensor within-logger component of variance, where a cluster of  $n_e$  sensors are randomly located within a region of 2 m diameter around each of  $n_1$  data logging hubs, which are located by simple random sampling, can be approximated by

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$$\sigma_{\text{sensor}}^2 = \sigma_2^2 + \sigma_{0.5}^2 + \sigma_{0.1}^2 + \sigma_{0.01}^2,$$
 (6)

and the between-logger variance by

$$\sigma^2_{\text{logger}} = \sigma^2_{\text{s}} + \sigma^2_{\text{m}}.$$
 (7)

As such, the standard error of the field mean soil N concentration derived from the sensor network can be estimated as follows:

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$$\sigma_{\text{mean}} = \{ (\sigma_{\text{logger}}^2 / n_l) + (\sigma_{\text{sensor}}^2 / n_l n_e) \}^{1/2} . \tag{8}$$

This allows the 95% confidence interval of the field mean estimations to be calculated, given the variance components calculated from the nested sampling, for particular combinations

and numbers of data loggers  $(n_1)$  and sensors  $(n_e)$ . These calculations were performed on the transformed scale prior to back-transformation of the 95% confidence interval to the original scale of measurement.

In order to demonstrate how the design may be optimised on a cost basis it was necessary to decide on a unit cost for a data logger and a sensor. Given the potential of electrochemical sensors for in-situ monitoring, it was decided that the unit cost for the sensor would be £200, based on the cost of a commercially available NH<sub>4</sub><sup>+</sup> or NO<sub>3</sub><sup>-</sup> ISE (ELIT 8021, ELIT 003, Nico2000, Harrow, UK) and £2000 for the data logger, based on the cost of a commonly used data logger (DL2e DeltaT, Cambridge, UK). Whilst these costs are somewhat arbitrary, it does allow useful comparison between designs to be made. It would also be possible to change these unit costs to explore how using different sensors and loggers may affect the optimisation of the network.

The 95% confidence intervals were computed for sensor network designs that consisted of 1 to 10 data loggers with 2 to 15 sensors distributed equally among the loggers, 15 being the maximum number of sensor ports on the data logger (DL2e DeltaT, Cambridge, UK). This allowed construction of graphs (Fig. 4) which illustrate the total cost for each design plotted against the resulting 95% confidence interval of the estimated field mean.

#### 3. Results

3.1. June nested sampling - evaluating the spatial variation of soluble N in soil prior to application of N fertiliser

The mean concentrations of amino acid,  $NH_4^+$  and  $NO_3^-$  were found to be fairly similar with values of 1.44, 1.87 and 1.71 respectively (Table 2). All of the N-forms had a positively skewed distribution. This was especially the case for  $NH_4^+$ , which had a skewness value of 12.82 and a maximum value of 80.49  $\mu g$  N  $g^{-1}$  (histograms of distributions can be

viewed in the supplementary information, Fig. S1). The Box-Cox parameter,  $\lambda$ , for each variable had a 95% confidence interval which excluded zero, so the maximum likelihood estimate of  $\lambda$  (Table 2) was used to transform each variable. Plots of the profile likelihood for  $\lambda$ , with the 95% confidence interval can be viewed in the supplementary information, Fig. S2.

The different forms of N showed slightly different scale-dependencies, although in general, short-range variance dominated (Fig. 2 and Table 3). On the original units, the MAD for comparisons where all sources of variation contributed were largest for NO<sub>3</sub> (0.92 μg N  $g^{-1}$ ), followed by NH<sub>4</sub><sup>+</sup> (0.75 µg N  $g^{-1}$ ), and amino acids with (0.51 µg N  $g^{-1}$ ). For amino acids, the 1-cm scale had the largest variance component, constituting 59% of the total accumulated variance. The 10-cm and the between-mainstations within-strata term were also considered important spatial components (as judged by AIC; see Table 3 and Table S1). For NH<sub>4</sub><sup>+</sup> the 1-cm scale had the largest variance component, constituting 63% of the total accumulated variance. However, for spatial scales greater than 1 cm, only the betweenmainstations within-strata term was considered important. For NO<sub>3</sub>, the relative importance of variation at scales coarser than 1cm was larger than for other forms of N (Fig 2) with the 10-cm scale having the largest variance component, constituting 28% of the total accumulated variance on the transformed scale. A comparable pattern was seen with the MAD values. Furthermore, all the spatial scales, with the exception of the 2-m scale, exhibited variance that was considered important. Short-range scale variation still dominated though, with 70% of the variance occurring at spatial scales up to 50 cm. It should be noted that the 1-cm scale component will also include any measurement error.

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3.2. July nested sampling - evaluating the spatial variation of soluble N in soil after application of N fertiliser

The mean concentrations of amino acid,  $NH_4^+$  and  $NO_3^-$  were found to be fairly similar with values of 1.25, 1.96 and 1.36 respectively (Table 2). All of the N-forms had slight positively skewed distributions.  $NH_4^+$  displayed the largest positive skew, with a skewness value of 3.28 and a maximum value of 9.88  $\mu g$  N  $g^{-1}$  (histograms of distributions can be viewed in the supplementary information, Fig. S3). The Box-Cox parameter,  $\lambda$ , for each variable had 95% confidence interval which excluded zero, so the maximum likelihood estimate of  $\lambda$  (Table 4) was used to transform each variable. Plots of the profile likelihood for  $\lambda$ , with the 95% confidence interval can be viewed in the supplementary information, Fig. S4.

The different forms of N showed slightly different scale-dependencies, although in general short-range variance dominated (Fig. 3 and Table 5). On the original units, the MAD for comparisons where all sources of variation contributed were largest for NH<sub>4</sub><sup>+</sup> (0.88 µg N  $g^{-1}$ ), followed by NO<sub>3</sub><sup>-1</sup> (0.64 µg N  $g^{-1}$ ) and amino acids with (0.41 µg N  $g^{-1}$ ). For amino acids, the between mainstations within-strata had the largest variance component, constituting 36% of the total accumulated variance, although 58% of the total accumulated variance occurred at scales up to 10 cm. The 1-cm, 10-cm and the between-mainstations within-strata term were considered important spatial components (as judged by AIC; see Table 5 and Table S2). For NH<sub>4</sub><sup>+</sup>, the 1-cm scale had the largest variance component, constituting 55% of the total accumulated variance. Spatial scales greater than 10 cm accounted for only 13% of the total accumulated variance. Only the 1-cm and the between-mainstations within-strata terms were considered important spatial components. For NO<sub>3</sub>, the between-mainstations within strata scale was the largest variance component, constituting 39% of the total accumulated variance. The 1-cm, 10-cm and the between-mainstations within-strata term were considered important spatial components. Short-range scale variation still dominated though, with 61% of the variance occurring at spatial scales up to 50 cm.

Duplicate measurements on 4 samples from each mainstation allowed the 1-cm spatial variance component to be resolved from the subsampling and measurement error. As this residual term formed the ultimate term in the model, it allowed an assessment of the importance of the 1-cm spatial component. For all of the N forms, the 1-cm scale was considered an important spatial component, and was larger than the residual variance. However, the residual variance, which was similar for all N forms, constitutes a substantial component of the accumulated variance and was, for all N forms, larger than the variance at 50 cm and 2 m.

### 3.3. Aggregate-scale variability of soluble N in soil

In the case of the aggregate-scale data, the 95% confidence interval for the Box-Cox parameter,  $\lambda$ , for each variable included zero (see supplementary material Fig. S6), so the log-transformation was applied.

In all cases, the largest variance component was found to be the between-aggregate within-core scale (table 7 & 8). For NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup>, 91% and 80% respectively of the total accumulated variance occurred at this scale, which was an order of magnitude higher than the variance at the between core scale. The variance at the aggregate scale for amino acid-N was lower at 66%. It should be noted that any analytical error that occurred will also appear in this variance component. The between-core component, which represents the 1-cm spatial scale, was considered important (as judged by AIC; see Table 7 and Table S3) for amino acids and NO<sub>3</sub><sup>-</sup>, but not NH<sub>4</sub><sup>+</sup>. Neither the between-pair component, which is similar to the between-mainstations scale, nor the between-strata component, were considered to be important spatial components. However, the stratum and mainstation scale in this analysis were based on limited replication. The focus of this particular sampling exercise was on the aggregate and core scale, so general conclusions from these results about the importance of coarser-

scale variation are not made.

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# 3.5. Optimisation of a within-field sensor network for monitoring soluble N in soil

The optimisation of the design of an in-situ network of NO<sub>3</sub> and NH<sub>4</sub> sensors can be explored using the graphs in Fig. 4. The graphs show how increasing both the number of sensors per data logger, and increasing the number of data loggers, reduces the width of the 95% confidence interval of the estimated field mean derived from the sensor network. There are differences in the results between NO<sub>3</sub> and NH<sub>4</sub> and between sampling events. For example, to achieve a 95% confidence interval width of no more than 1 µg N g<sup>-1</sup> for a NO<sub>3</sub><sup>-1</sup> sensor network, given the spatial variation observed in the June sampling event, would require 3 data loggers each with 11 sensors at a cost of £8200. For the July sampling, 2 data loggers each with 7 sensors would be sufficient, at a lower cost of £5400. For a NH<sub>4</sub><sup>+</sup> sensor network, given the spatial variation observed in the June sampling event, 2 data loggers each with 6 sensors, at a cost of £5200 would be required to achieve a 95% confidence interval width of no more than 1 µg N g<sup>-1</sup>. For the July sampling, 2 data loggers each with 8 sensors, at a slightly higher cost of £5600, would be required. Reducing the width of the 95% confidence interval substantially below 1 µg N g<sup>-1</sup> dry soil would result in a large cost increase, with small marginal improvement on increasing the size of the network. For a NO<sub>3</sub> sensor network, given the spatial variation observed in the June sampling event, reducing the width of the confidence interval to less than  $0.5~\mu g$  N  $g^{-1}$  would require 10~loggers each with 12 sensors, at a cost of £22400.

An alternative approach is to optimise the sensor network design within the constraints of a fixed budget. A budget of £5000 for a NO<sub>3</sub><sup>-</sup> sensor network could provide a single data logger with 15 sensors or 2 data loggers each with 5 sensors. This could be used to provide a single logger with 15 sensors on each, or two loggers with 5 sensors on each.

The width of the confidence interval for these two options is 2.12 and 1.70  $\mu g$  N  $g^{-1}$  dry soil respectively, so the second option is the rational choice.

### 4. Discussion

# 4.1. Spatial variation of soluble N at within-field scales

The dominance of short range variation (i.e. less than 2 m) for all the N forms may be attributed to the relatively random and uneven deposition of N from sheep excreta within the context of a broadly homogenous field. The proportion of the total accumulated variance attributed to the 1-cm scale was much larger for amino acid-N and NH<sub>4</sub><sup>+</sup>-N than NO<sub>3</sub>-N which may be related to their relative diffusion coefficients, interactions with the solid phase (Owen and Jones, 2001) and the rapid rate of amino acid turnover and mineralisation in this soil (Jones et al., 2004; Wilkinson et al., 2014).. Similar small-scale variation of NO<sub>3</sub><sup>-</sup> in grazed pastures has been identified in previous studies, with semi-variograms exhibiting the range of spatial dependency of less than 5 m (White et al., 1987; Broeke et al., 1996; Wade et al., 1996; Bogaert et al., 2000), and a nugget variance of 60% (Bogaert et al., 2000). These results contrast with similar studies performed on arable soils, which were characterised by ranges of spatial dependencies for NO<sub>3</sub><sup>-</sup> of greater than 39 m (Van Meirvenne et al., 2003; Haberle et al., 2004). The observed variation at larger spatial-scales, especially NO<sub>3</sub><sup>-</sup>, could be due to the habit of sheep to frequent certain areas of the pasture such as paths, a drinking trough and areas of shade (Bogaert et al., 2000).

It is unlikely that the observed variation at the "aggregate" scale is driven by the deposition of sheep excreta. Previous studies of spatial variation in soil N, in the context of within-field scales, have not investigated variation over such small scales. This small-scale variation is likely due to the inherent micro-heterogeneity of soil properties, for example, the abundance of plant roots and mycorrhizal hyphae (Stoyan et al., 2000), availability of labile

organic matter (Parkin, 1987; Wachinger et al., 2000), earthworm channels and the composition and abundance of the microbial community (Grundmann and Debouzie, 2000; Nunan et al., 2002), which in turn will affect biogeochemical processes controlling soil N concentrations.

There was also some suggestion of a spatio-temporal interaction as evidenced by small differences in the spatial dependencies of the N forms between the June and July nested sampling events. In the case of NO<sub>3</sub><sup>-</sup>, the total accumulated variance was lower, with more of the observed variance attributed to scales greater than 2 m for the July sampling. This change may be attributed to the removal of sheep and the associated local inputs of N, combined with N fertilisation of the field (60 kg N ha<sup>-1</sup>) that occurred 3 weeks prior to the second nested sampling event.

# 4.2. Optimisation of designing a within-field soil N sensor network

This study clearly demonstrates how nested sampling combined with geostatistical analysis can be used to optimise the design of an in situ sensor network. Furthermore, given knowledge of logger and sensor costs it is possible to rationalise planning decisions on a cost-precision basis. Essentially, the shape of the curves within the optimisation graphs (Fig. 5) reflects the observed spatial variation. The largest observed accumulated variance was for NO<sub>3</sub><sup>-</sup> from the June nested sampling. Consequently, a NO<sub>3</sub><sup>-</sup> sensor network based on this spatial variation would require a greater number of data loggers and sensors (with a resultant cost increase), to achieve a desired level of precision when compared to a NO<sub>3</sub><sup>-</sup> sensor network based on the July nested sampling results and a NH<sub>4</sub><sup>+</sup> sensor network based on either the June or July nested sampling results. The distribution of variance across the spatial scales also affects the most efficient use of a specific budget, in terms of the choice of number of data loggers and sensors used and the resulting width of the 95% confidence interval. For

example, a budget of £7000 could be used to purchase either 2 data loggers with 15 sensors on each or 3 data loggers with 5 sensors on each. Given the observed variation from the June nested sampling, the former of the above choices gives the most efficient design for  $NH_4^+$  and the later for  $NO_3^-$ . This reflects the fact that, compared to  $NO_3^-$ , a greater proportion of the total accumulated variance for  $NH_4^+$ , is attributed to scales less than 2 m.

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It is important to note that the data used for these calculations were derived from the nested sampling which used a soil corer of 1 cm diameter. As such, these calculations are based on the assumption that any given sensor used for the in-situ network would have a similar sized zone of influence. Results derived from the aggregate-scale sampling exhibited large variation at the sub 1-cm scale, which for NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> was an order of magnitude larger than the 1-cm scale. This will have significant implications when using sensors, for which the measurement represents an integration over a sensor-soil contact area of diameter less than 1 cm, as they will be affected by the observed "aggregate" scale variation. If this variation is not considered when designing the sensor network, it is likely that the precision of the estimated field mean would be overestimated. To compensate for this, more local replication at the sub 1-cm scale and hence an increase in the size of the sensor network would be required for an acceptable level of precision to be achieved, resulting in increased costs. To explore optimisation of a network of such sensors, further sampling at the sub 1-cm scale would be required. Ideally this would involve a similar level of replication, across all scales, to that which was used in the July nested sampling campaign. This evidence may also be quite instructive for optimising sensor design, as sensors with larger sampling areas will encompass more of this small-scale variation.

Within this optimisation, no consideration has been made to the observed depth effects. The resulting estimate of the field mean derived from the sensor network would therefore, only be applicable to the 5-10 cm depth. Rooting depth, and therefore, nutrient

uptake, in fields adjacent to the study site has previously been observed to a depth of 30 cm (Jones et al., 2004) and a decrease in plant-available with depth has been observed in the study field (see supplementary information, Fig. S4). As such, any quantification of plant-available N derived from the sensor network should be adjusted for observed depth effects. In the case of cereals, which may root to depths in excess of 1.5 m, both topsoil and subsoil sensors will probably be required to avoid bias and gain a representative pattern of soluble N within the field. Logistically, however, the deployment of sensors in subsoils represents a significant challenge.

## 4.3. Potential for use of in-situ sensor networks within precision agriculture

Adoption of in-situ networks not only relies on the development of suitable sensors but also on evidence of the cost-benefit. Given the results of the sensor network optimisation, it is probable that uptake of this approach will incur significant costs; especially if a high level of precision is required. As such, it is likely that this approach will be limited initially to high value horticultural crops. As the cost of sensor and data logging technology continues to fall, adoption by arable and pastoral agriculture may increase. One key factor that will affect the cost of a sensor network is the required precision of the resulting estimated mean and this in turn may depend on how the generated data is used to improve fertiliser management. The creation of a decision support tool that will bring about improved NUE, and hence reduce input costs and/or increase profits, requires significant future research and is not an insignificant challenge.

It is important to consider how the approach used in this study could be applied to field exhibiting significant random and non-random (i.e. a gradient) large-scale variation. It is possible that the field could be split into management zones each with their own sensor network. These management zones could be delimitated on the basis of *a priori* knowledge of

variables that may affect or indicate soil N status such as topography (Kravchenko and Bullock, 2000), soil type (Moral et al., 2011), yield variability (Diker et al., 2004) and farmers knowledge (Fleming et al., 2000). Alternatively, proximal or remote sensing, such as electromagnetic induction, may allow rapid and cost effective identification of large scale heterogeneity of soil physical properties (Hedley et al., 2004; King et al., 2005). However, the extent to which these variables correlate to soil N concentration is likely to be site specific and so may require some ground truthing. A further broad question which needs to be addressed with respect to management zones, is at what point does the magnitude and the spatial-scale of soil N variation become sufficiently large enough to justify site-specific agriculture?

The success of the approach used here to optimise a sensor network requires temporal stability of spatial variation (Sylvester-Bradley et al., 1999; Shi et al., 2002). Given significant spatio-temporal interaction, the results from any sensor network could no longer be considered accurate or precise. In this study there was evidence of a slight spatio-temporal interaction which was related to the removal of sheep from the field and the application of N fertiliser. An alternative approach to that advocated here, would be the implementation of a grid network, with sensor arrays at each node to account for small-scale soil variation. This would enable temporal, large-scale spatial variation and their interaction to be monitored. Kriging techniques could then be used to produce dynamic maps of soil N concentrations which could be used to inform variable-rate fertiliser management. However, this approach is likely to require significantly more sensing units and data loggers with a resulting cost increase.

### 5. Conclusions

This study demonstrates how a network of in-situ soil N sensors could be efficiently designed and optimised on the basis of cost and precision. To achieve this, the spatial variation of plant available N – amino acids,  $NH_4^+$  and  $NO_3^-$  – within the soil of a grazed grassland field was investigated using a nested sampling approach and geo-statistical analysis. Variation of all N forms at small scales (less than 2 m) was shown to be dominant, with further large variance evident at scales less than 1 cm. The observed variation was considered in line with previous work and was attributed to the random input of N to the soil via sheep excreta and the inherent heterogeneity of soil at the aggregate scale. Based on the observed spatial variance observed in the June nested sampling, and given a budget of £5000, the  $NO_3^-$  field mean could be estimated with a 95% confidence interval width of 1.70  $\mu$ g N g<sup>-1</sup> using 2 randomly positioned data loggers each with 5 sensors. Achieving a 95% confidence interval width substantially lower than 1.70  $\mu$ g N g<sup>-1</sup> would require significant extra cost. Adoption of in-situ sensor networks will depend upon the development of suitable low-cost sensors, demonstration of the cost-benefit and the construction of a decision support system that utilises the generated data to improve the NUE of fertiliser N management.

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## Figure legends

**Figure 1.** The optimised sampling design of a mainstation used to perform spatial nested sampling in a 1.9 ha grassland field. Distances between sampling points were fixed but angles were randomized, with the exception of the 2 m vectors.

Figure 2. Accumulated variance components of the Box-Cox transformed data (Panel A) from the finest to coarsest spatial scale, derived from the June nested sampling results (before fertiliser addition). Panel B shows median absolute deviation from the median (MAD) of differences over the nested spatial intervals on the original scale of measurement. Source is the spatial-component in meters, with M and S representing the between-mainstation and between-strata components respectively.

Figure 3. Accumulated variance components of the Box-Cox transformed data (Panel A) from the finest to coarsest scale, derived from the July nested sampling results (after fertiliser addition). Panel B shows median absolute deviation from the median (MAD) of differences over the nested spatial intervals on the original scale of measurement. Source is the spatial-component in meters, with M and S representing the between-mainstation and between-strata components respectively.

Figure 4. Width of the 95% confidence interval for alternative sensor network designs of different cost computed to facilitate monitoring of soil N in a 1.9 ha grassland field. Values are computed from variance components from nested sampling of nitrate in (a) June (before fertiliser addition) and (b) July (after fertiliser addition) and of ammonium in (c) June (before fertiliser addition) and (d) July (after fertiliser addition) and on the basis of unit costs for a sensor and a data logger of £200 and £2000 respectively. Note that the arrays comprise 1-10 loggers and a maximum of 15 sensors per logger. To allow a common range of values on the ordinates of these graphs, and to facilitate interpretation, arrays with fewer than five sensors in total have been excluded from

Figure 4(a) and arrays with fewer than three sensors have been excluded from Figures 4(b-d).

**Table 1** Background properties of the agricultural grassland Eutric Cambisol used in the study. Values represent means  $\pm$  SEM (n=4). All soil values are expressed on a dry weight soil basis.

Site property	Mean $\pm$ SEM
рН	$6.57 \pm 0.05$
EC (µS cm <sup>-1</sup> )	$26.5\pm1.0$
Basal soil respiration (mg CO <sub>2</sub> kg <sup>-1</sup> h <sup>-1</sup> )	$12.61 \pm 1.04$
Total soil C (g C kg <sup>-1</sup> )	$25.35 \pm 1.47$
Total soil N (g N kg <sup>-1</sup> )	$2.95 \pm 0.06$
Soil C:N	$8.62 \pm 0.64$
DOC (mg C kg <sup>-1</sup> )	$70.08 \pm 2.57$
DON (mg N kg <sup>-1</sup> )	$10.48 \pm 1.07$
Mineralisable N (mg N kg <sup>-1</sup> d <sup>-1</sup> )	$3.92 \pm 0.54$
Microbial C (g C kg <sup>-1</sup> )	$1.03 \pm 0.10$
Microbial N (g N kg <sup>-1</sup> )	$0.16 \pm 0.01$
Exchangeable Ca (mg Ca kg <sup>-1</sup> )	$501 \pm 122$
Exchangeable K (mg K kg <sup>-1</sup> )	$46.05 \pm 12.61$
Exchangeable Na (mg Na kg <sup>-1</sup> )	$25.43 \pm 5.13$
Available P (mg P kg <sup>-1</sup> )	$7.38 \pm 2.02$
Above ground biomass (t DM ha <sup>-1</sup> )	$1.56 \pm 0.14$

Table 2 Summary statistics describing the spatial variability of soluble N ( $\mu g$  N  $g^{-1}$ ) derived from the nested sampling of a grassland soil prior to the application of N fertiliser. Alongside the raw data, an estimate of the Box-Cox transformation parameter ( $\lambda$ ) is also provided.

Variable	Mean	Median	Minimum	Maximum	Skewness	λ
Nitrate	1.71	1.10	0.29	22.51	5.41	-0.426
Ammonium	1.87	1.27	0.29	80.49	12.82	-0.541
Amino acid	1.44	1.39	0.65	5.20	3.37	-0.492

Table 3

Variance components for the (Box-Cox transformed) variables and associated Wald tests describing the spatial variability of soluble N derived from the nested sampling of a grassland soil prior to the application of N fertiliser. The Wald statistic and associated p-value describe differences between the two sampling days. Those variance components marked with an asterisk are ones which caused an increase in AIC if they were dropped from the model (finest scale cannot be dropped).

X7 ' 11		Variance component						
Variable	$\sigma_{\rm s}^2$	$\sigma^2_{m}$	$\sigma^2_2$	$\sigma^2_{0.5}$	$\sigma^2_{0.1}$	$\sigma^2_{0.01}$	statistic	<i>p</i> -value
Nitrate	$0.0629^*$	$0.0362^{*}$	0.0	$0.0795^{*}$	$0.0937^{*}$	0.0628	0.001	0.974
Ammonium	0.0087	$0.0121^{*}$	0.0078	0.00008	0.0153	0.0751	6.8	0.009
Amino acid	0.0058	$0.0035^{*}$	0.0	0.0	$0.0124^{*}$	0.0307	1.89	0.17

Table 4 Summary statistics describing the spatial variability of soluble N ( $\mu g$  N  $g^{-1}$ ) derived from the nested sampling of a grassland soil after the application of N fertiliser. Alongside the raw data, an estimate of the Box-Cox transformation parameter ( $\lambda$ ) is also provided.

Variable	Mean	Median	Minimum	Maximum	Skewness	λ
Nitrate	1.36	1.25	0.26	3.45	0.89	0.302
Ammonium	1.96	1.71	0.26	9.88	3.28	-0.424
Amino acid	1.25	1.18	0.56	4.40	2.58	-0.481

Table 5

Variance components for the (Box-Cox transformed) variables and associated Wald tests describing the spatial variability of soluble N derived from the nested sampling of a grassland soil after the application of N fertiliser. The Wald statistic and associated *p*-value describe differences between the two sampling days. Those variance components marked with an asterisk are ones which caused an increase in AIC if they were dropped from the model (finest scale cannot be dropped).

X7 1- 1 -		Variance component							1
Variable	$\sigma_{s}^{2}$	$\sigma^2_{m}$	$\sigma^2_2$	$\sigma^{2}_{0.5}$	$\sigma^2_{0.1}$	$\sigma^{2}_{0.01}$	$\sigma^2_{\epsilon}$	statistic	<i>p</i> -value
Nitrate	0.0	0.0638*	0.0	0.0052	0.049*	0.031*	0.0131	7.89	0.005
Ammonium	0.0039	$0.0069^{*}$	0.0	0.0	0.015	$0.045^{*}$	0.0109	15.43	$8.60 \times 10^{-15}$
Amino acid	0.002	$0.0241^{*}$	0.0025	0.0	$0.0086^*$	$0.0199^{*}$	0.0103	0.708	0.4

Table 6 Summary statistics describing the aggregate-scale variability of soluble N ( $\mu g$  N  $g^{\text{-1}}$ ) within a grassland soil.

Variable	Mean	Median	Minimum	Maximum	Skewness
Nitrate	1.20	1.04	0.19	3.13	0.80
Ammonium	2.00	1.78	0.30	5.85	1.24
Amino acid	1.56	1.50	0.77	2.69	0.49

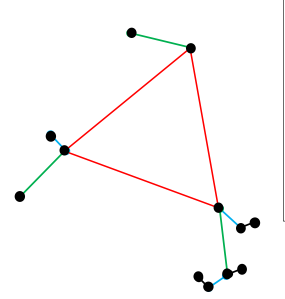
Table 7

Variance components for the (log-transformed) variables describing the aggregate-scale spatial variability of soluble N in a grassland soil. Those variance components marked with an asterisk are ones which caused an increase in AIC if they were dropped from the model (finest scale cannot be dropped).

** ' 11		Variance component					
Variable	$\sigma_{s}^{2}$	$\sigma^2_{p}$	$\sigma^2_{c}$	$\sigma_{a}^{2}$			
Nitrate	0.0	0.0	0.072*	0.295			
Ammonium	0.0	0.0293	0.003	0.3074			
Amino acid	0.0	0.0031	0.0132*	0.0321			

**Table 8** Median absolute deviations from the median (MAD) describing the aggregate-scale spatial variability of soluble N ( $\mu g$  N  $g^{-1}$ ) in a grassland soil. Comparisons are nested, so the stratum-scale MAD includes the pair, core and aggregate-scale.

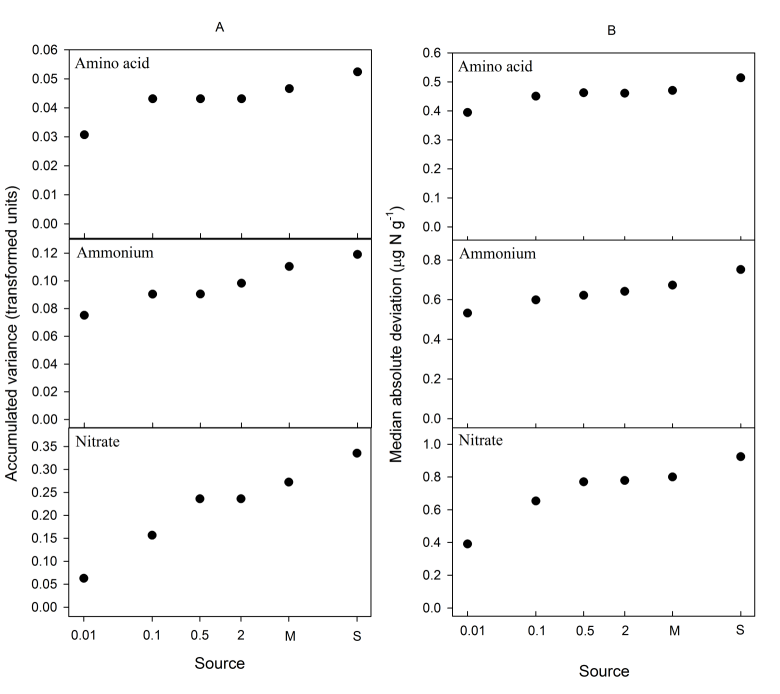
		MAD (με	g N g <sup>-1</sup> )		
Variable	Aggregate	Core	Pair	Stratum	
Nitrate	0.75	0.85	0.85	0.85	
Ammonium	1.27	1.30	1.34	1.35	
Amino acid	0.39	0.45	0.46	0.46	

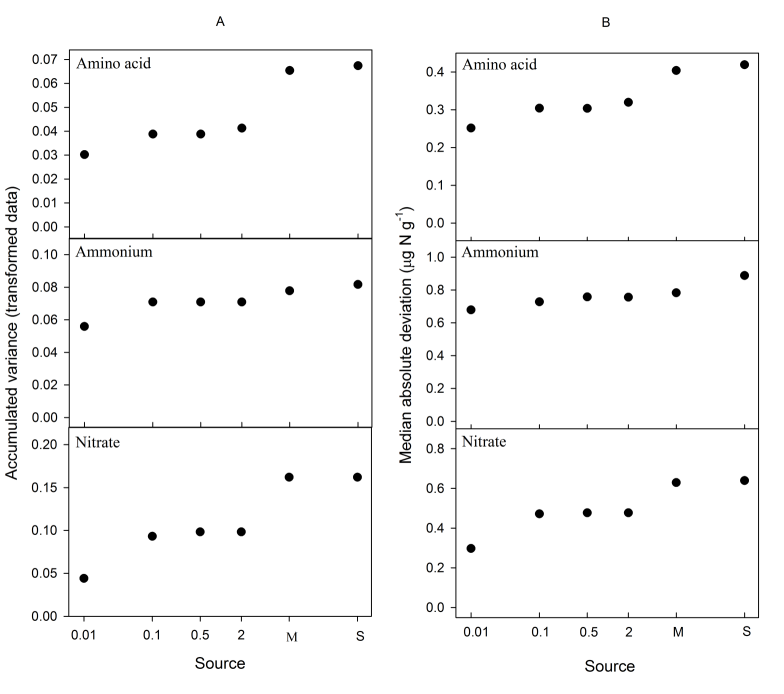


## Key

Length of a line joining two sample points

Sample •





## Nitrate (a) (b) Number of loggers Width of confidence interval ( $\mu g \ N \ g^{-1} dry \ soil)$ Width of confidence interval ( $\mu g \ N \ g^{-1} dry \ soil)$ 2 3 က 9 $\sim$ 10 0 0 5000 10000 15000 20000 5000 10000 15000 20000 Cost (£) Cost (£) Ammonium (c) (d) Width of confidence interval ( $\mu g \; N \; g^{-1} dry \; soil)$ Width of confidence interval ( $\mu g \; N \; g^{-1} dry \; soil)$ က 7 $^{\circ}$ 0 0 5000 10000 15000 20000 5000 10000 15000 20000 Cost (£) Cost (£)

Figure S1. Histograms of the absolute ( $\mu g \ N \ g^{-1}$ ) and Box-Cox transformed units of soil nitrate, ammonium and amino acid concentrations from the June nested sampling prior to fertiliser addition sampling (n = 192 for each N form).

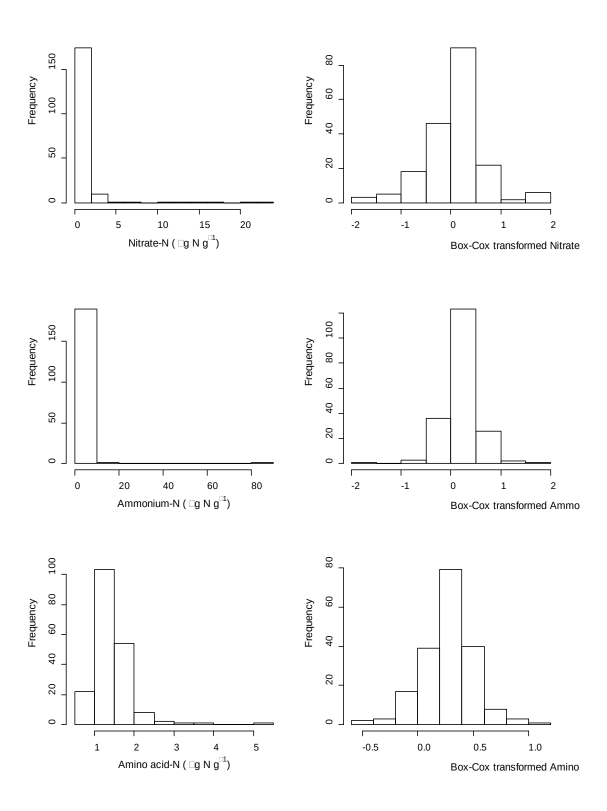
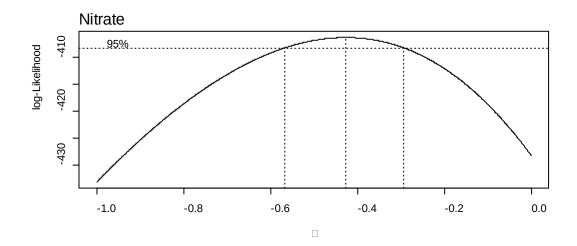
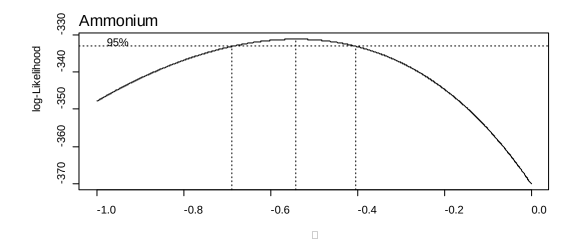


Figure S2. Profile likelihood plot for the  $\lambda$  parameter of the Box-Cox transformation for soil nitrate, ammonium and amino acid concentrations from the June nested sampling prior to fertiliser addition.





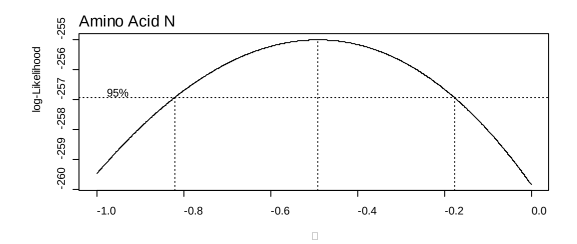


Figure S3. Histograms of the absolute ( $\mu g \ N \ g^{-1}$ ) and Box-Cox transformed units of soil nitrate, ammonium and amino acid concentrations from the July nested sampling following fertiliser addition sampling (n = 192 for each N form).

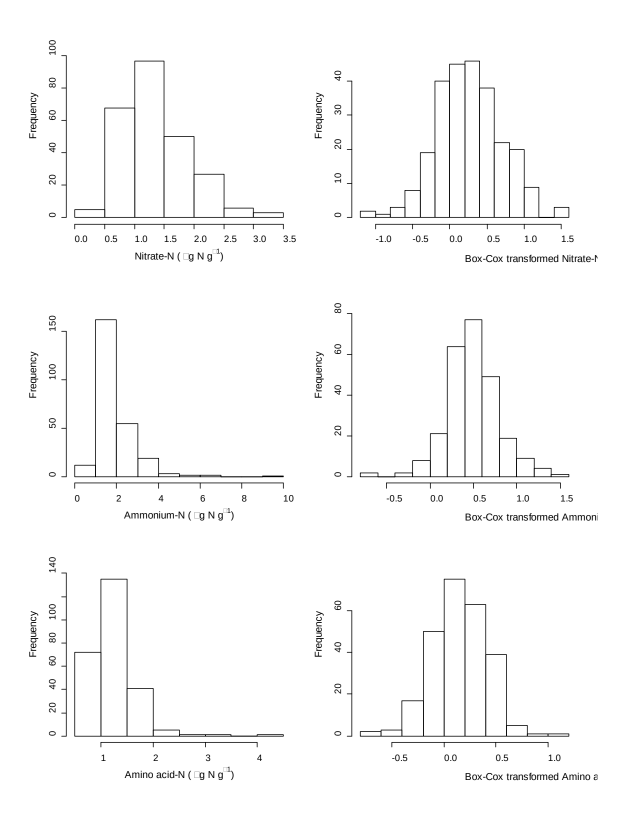
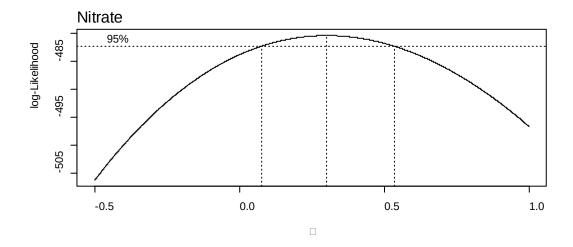
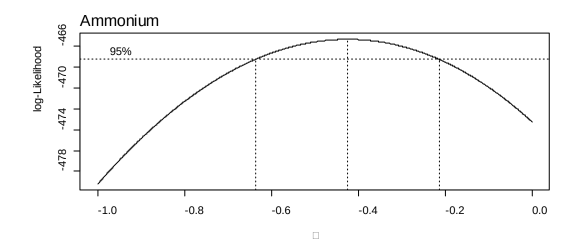


Figure S4. Profile likelihood plot for the  $\lambda$  parameter of the Box-Cox transformation for soil nitrate, ammonium and amino acid concentrations from the July nested sampling prior to fertiliser addition.





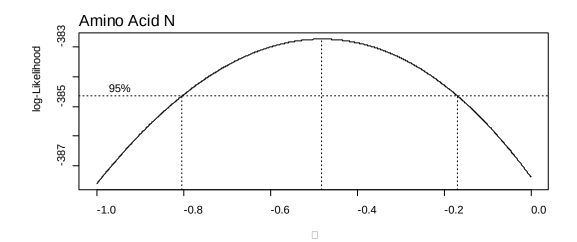


Figure S5. Histograms of the absolute ( $\mu g \ N \ g^{-1}$ ) and Box-Cox transformed units of soil nitrate, ammonium and amino acid concentrations from the aggregate-scale sampling (n = 192 for each N form).

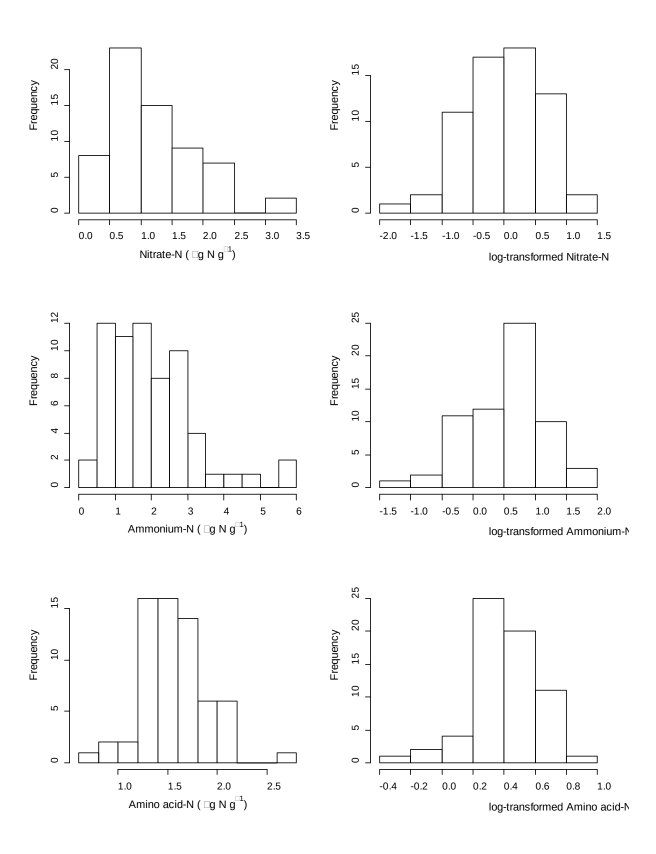
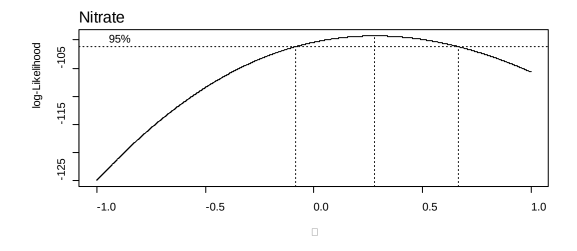
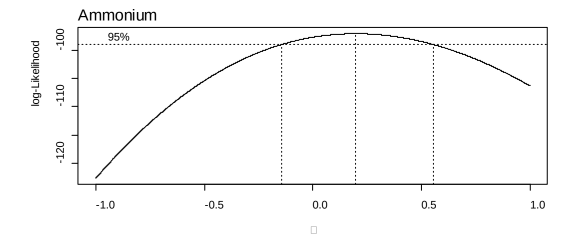


Figure S6. Profile likelihood plot for the  $\lambda$  parameter of the Box-Cox transformation for soil nitrate, ammonium and amino acid concentrations from the aggregate-scale sampling.





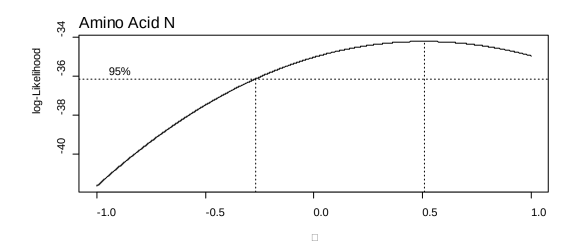


Figure S7. Variability of soil nitrate, ammonium and amino acid with soil depth. Data points represent means  $\pm$  SEM (n = 12) of soil N concentrations ( $\mu g N g^{-1}$ ) for each 5 cm depth increment.

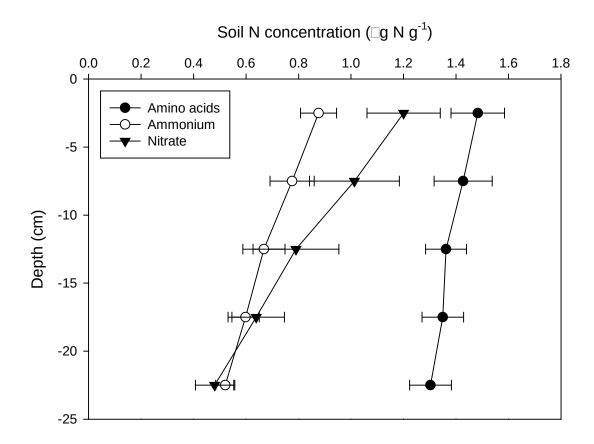


Table S1. Akaike's information criterion (AIC) values for the full model used to describe the spatial variation of soil N forms in a grassland soil derived from the June nested sampling event. The resulting AIC values when each variance component is dropped in turn from the full model are also shown. Where the variance for variance components is zero, no AIC value is reported. Variance components refer to distances in meters, with m and s representing the between-mainstation and between-strata components respectively.

*** * 1.1	AIC value for full	AIC value if variance component is dropped from model						
Variable	model	$\sigma_{s}^{2}$	$\sigma^2_{\ m}$	$\sigma^2_2$	$\sigma^2_{0.5}$	$\sigma^2_{0.1}$		
Nitrate	-88.35	-86.78	-87.19		-85.85	-72.06		
Ammonium	-217.59	-218.44	-216.81	-219.12	-219.59	-218.91		
Amino Acid	-369.1	-370.01	-368.81			-368.21		

Table S2. Akaike's information criterion (AIC) values for the full model used to describe the spatial variation of soil N forms in a grassland soil derived from the July nested sampling event. The resulting AIC values when each variance component is dropped in turn from the full model are also shown. Where the variance for variance components is zero, no AIC value is reported. Variance components refer to distances in meters, with m and s representing the between-mainstation and between-strata components respectively.

*** * 11	AIC value	AIC value if spatial component is dropped from model					
Variable	for full model	$\sigma_{s}^{2}$	$\sigma^2_{\ m}$	$\sigma^2_{2}$	$\sigma^2_{0.5}$	$\sigma^2_{0.1}$	$\sigma^2_{0.01}$
Nitrate	-412.87	_	-395.20		-414.69	-400.52	-376.95
Ammonium	-461.87	-463.08	-460.95			-462.02	-408.89
Amino Acid	-560.68	-562.60	-547.14	-562.11	_	-560.48	-533.60

Table S3. Akaike's information criterion (AIC) values for the full model used to describe the aggregate-scale spatial variation of soil N forms in a grassland soil. The resulting AIC values when each variance component is dropped in turn from the full model are also shown. Where the variance for variance components is zero, no AIC value is reported. The variance components s, p and c representing the between-strata, the between-pair and between core components respectively.

Variable	F. 11 1.1	Terr	n dropped from m	odel
	Full model —	$\sigma_{s}^{2}$	$\sigma^2_{p}$	$\sigma^2_{c}$
Nitrate	10.71		_	11.8
Ammonium	7.27	_	5.82	5.28
Amino Acid	-123.0		-124.9	-121.0