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Measured estimates of semi-natural terrestrial NPP in Great Britain: comparison with modelled values, and dependence on atmospheric nitrogen deposition

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Abstract:	Plant growth in nitrogen (N)-limited, unfertilised terrestrial ecosystems should respond to additional N inputs from atmospheric deposition (Ndep). We investigated this for sites in Great Britain (GB) by compiling 796 estimates of net primary productivity (NPP) from measured biomass production over the period 1932-2014, although the great majority were for 1995 onwards. The sites were largely vegetated with shrubs, grass and bracken, and had a wide range of Ndep ($0.5 - 3.3$. gN m-2 a-1 in 2000). The measured NPP estimates were compared with calculated values from the biogeochemical ecosystem model N14CP, which predicts that NPP depends strongly upon Ndep. The measured and modelled average total NPP values (gC m-2 a-1) from all data were 387 (standard deviation, SD = 193) and 377 (SD = 72) respectively.							

	Measured and modelled averages for vegetation classes followed the sequence: broadleaved trees ~ needle-leaved trees > herbs (rough grassland + bracken) ~ shrubs. After averaging measured values for sites in individual model grid cells (5 km x 5 km) with 10 or more replicates, the measured and modelled NPP values were correlated (n=26, r2=0.22, p=0.011), with a slope close to unity. Significant linear relationships were found between measured In NPP and cumulative Ndep for both herbs (n=298, p=0.021) and shrubs (n=473, p=0.006), with slopes comparable to those predicted with the model. The results suggest that semi-natural NPP in GB depends positively upon Ndep, in a manner that agrees quantitatively with N14CP predictions. Calculations with the model, using modelled temporal variation in Ndep, indicate that fertilisation by Ndep caused average increases in semi-natural NPP over the period 1800 to 2010 of 30% for shrubs, 71% for herbs, and 91% for broadleaved trees. Combined with previous published results for forests, our findings suggest a general and widespread vegetation response to fertilisation by Ndep.
Response to Reviewers:	Please see attached file.

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To the Editor, Biogeochemistry

We wish to submit the manuscript entitled *Measured estimates of semi-natural terrestrial NPP in Britain: comparison with modelled values, and dependence on atmospheric nitrogen deposition* by *Edward Tipping, Jessica A.C. Davies, Peter A. Henrys, Susan G. Jarvis, Edwin C. Rowe, Simon M. Smart, Michael G. Le Duc, Robert H. Marrs, Robin J. Pakeman* for publication in Biogeochemistry.

The paper asks the question "Does the net primary productivity (NPP) of nutrient-limited ecosystems respond to atmospheric nitrogen deposition (Ndep)?" The first piece of evidence is that the measured estimates of NPP for herbs and shrubs are correlated with predicted values from the N14CP model, which assumes N limitation and an effect of Ndep. The second is that the measured estimates of NPP show a significant increase with modelled Ndep across Britain.

We think this work is significant because the sequestration of carbon by ecosystems depends upon nutrient availability, and N is probably the most widespread limiting nutrient. But the "use" of N deposition to remove CO_2 from the atmosphere may be reversible, if N pollution is mitigated. Also, N enrichment by N deposition has consequences for plant diversity. Therefore we need to understand and quantify the effects of N deposition.

The paper is concerned with the biogeochemistry of the fixation of atmospheric carbon in vegetation, as affected by atmospheric nitrogen deposition. It therefore falls within the area of interest of Biogeochemistry.

Sincerely yours

Ed Tipping, on behalf of all authors

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1	Submitted to Biogeochemistry, January 2019						
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3	Measured estimates of semi-natural terrestrial NPP in <mark>Great Britain</mark> :						
4	comparison with modelled values, and dependence on atmospheric nitrogen						
5	deposition						
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7	Smart · Michael G. Le Duc · Robert H. Marrs · Robin J. Pakeman						
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21 Abstract

22 Plant growth in nitrogen (N)-limited, unfertilised terrestrial ecosystems should respond to additional N inputs from atmospheric deposition (N_{dep}). We investigated this for sites in Great Britain (GB) by 23 24 compiling 796 estimates of net primary productivity (NPP) from measured biomass production over the period 1932-2014, although the great majority were for 1995 onwards. The sites were largely 25 26 vegetated with shrubs, grass and bracken, and had a wide range of N_{dep} (0.5 – 3.3. gN m⁻² a⁻¹ in 2000). 27 The measured NPP estimates were compared with calculated values from the biogeochemical ecosystem model N14CP, which predicts that NPP depends strongly upon N_{dep}. The measured and 28 modelled average total NPP values (gC m⁻² a⁻¹) from all data were 387 (standard deviation, SD = 193) 29 and 377 (SD = 72) respectively. Measured and modelled averages for vegetation classes followed the 30 31 sequence: broadleaved trees ~ needle-leaved trees > herbs (rough grassland + bracken) ~ shrubs. After averaging measured values for sites in individual model grid cells (5 km x 5 km) with 10 or more 32 33 replicates, the measured and modelled NPP values were correlated (n=26, r^2 =0.22, p=0.011), with a 34 slope close to unity. Significant linear relationships were found between measured In NPP and cumulative N_{dep} for both herbs (n=298, p=0.021) and shrubs (n=473, p=0.006), with slopes comparable 35 36 to those predicted with the model. The results suggest that semi-natural NPP in GB depends positively upon N_{dep}, in a manner that agrees quantitatively with N14CP predictions. Calculations with the model, 37 38 using modelled temporal variation in N_{dep}, indicate that fertilisation by N_{dep} caused average increases in semi-natural NPP over the period 1800 to 2010 of 30% for shrubs, 71% for herbs, and 91% for 39 40 broadleaved trees. Combined with previous published results for forests, our findings suggest a 41 general and widespread vegetation response to fertilisation by N_{dep}.

42

43 Key words: Net primary production · Nitrogen deposition · Modelling · Grass · Bracken · Shrubs ·

44 Long-term change

45 Introduction

46 The net primary productivity (NPP) of an ecosystem can be defined as the difference between the 47 annual carbon gain from the atmosphere in plant photosynthesis and the annual carbon (C) loss in 48 plant respiration (Chapin et al. 2006; Schlesinger & Bernhardt 2013). It is a key ecosystem flux (gC m⁻² 49 a^{-1}), equivalent to the net amount of carbon (C) fixed per year, and strongly related to the amount of 50 C, with its associated elements, transferred from living biomass to the soil each year. In agricultural 51 systems it is strongly related to yield. Another important aspect is the association between NPP and 52 plant species diversity; the well-known unimodal model of Grime (1973a,b) postulates maximum 53 species richness at intermediate NPP, although debate about this relationship continues (Adler et al. 2012; Fraser et al. 2015). The modelling and prediction of NPP is a crucial activity in trying to 54 55 understand natural and agricultural ecosystems, and to predict how they might respond to environmental change, and models need to be tested against data where possible. 56

57 The main factors usually considered to determine, or limit, NPP are vegetation type, light, 58 temperature, and the availability of water and nutrients (Chapin et al. 2011; Schlesinger & Bernhardt, 59 2013). In temperate natural and semi-natural (i.e. managed but not fertilised) ecosystems, N is the 60 nutrient element that is most often limiting, according to evidence from experimental nutrient additions at individual locations (Elser et al. 2007; LeBauer & Treseder 2008; Lee et al. 2010; Field et 61 62 al. 2017); see also McGuire et al. (1992). It therefore follows that additions of N, via atmospheric deposition (N_{dep}, gN m⁻² a⁻¹), over recent decades and centuries could have brought about increases in 63 NPP, and consequent changes in ecosystem functioning and composition. Observations of temporal 64 changes in the above-ground biomass of woodlands receiving different levels of N_{dep} support this 65 66 contention (De Vries et al. 2006, 2009, 2015; Magnani et al. 2007; Kahle et al. 2008; Thomas et al. 67 2010). To demonstrate effects of N_{dep} on non-woodland ecosystems is more difficult, because the 68 biomass does not accumulate, and therefore the most straightforward temporal evidence would 69 comprise repeated measurements of annual production over a sufficiently long time period, at 70 locations with different N_{dep}. Such measurements have not been made, but it may be possible to 71 compare data from different locations with the same vegetation, sampled at different times. Another 72 approach is to test whether spatial variations in annual production are related to spatial variations in N_{dep} . This can be supported by comparison with model expectations, which give an idea of the 73 74 magnitudes of effects over both space and time.

The ecosystem model, N14CP (Davies *et al.* 2016a,b) was developed to link soil and plant processes and explore and predict how changes in nutrient status may affect pools and fluxes of C, N and phosphorus (P), within 5 km × 5 km grid cells, and over a temporal scale focusing on changes over the last several hundred years. The model simulates the stoichiometrically-interlinked cycles of C, N and P in temperate and boreal ecosystems over space and time, driven by climate (mean annual
temperature and precipitation, MAT and MAP), N_{dep} and vegetation class. According to N14CP,
fertilisation of GB semi-natural ecosystems by N_{dep} over the past two centuries led to increases in NPP,
thence to increased rates of input of organic matter into soils, and an increase in soil organic carbon
(SOC) pools and concentrations. The modelling results agree, in terms of both SOC changes with time
and the dependence of the changes on N_{dep}, with soil survey-resurvey results for samples collected
over the period 1959 to 2010 (Tipping *et al.* 2017).

86 We tested the model's simulations of NPP in GB, focusing on its prediction that N_{dep} has 87 increased semi-natural NPP. GB is well-suited for this test, in view of the substantial variations in N_{dep} 88 across the island (NEGTAP 2001; Smith et al. 2000), and plentiful measurements of plant production, 89 which can be used to estimate NPP. We analysed data from a large number (771) of measurements of 90 shrub, rough grass and bracken production made at different sites over recent decades, the majority of the data coming from Pearsall & Gorham (1956), Marrs et al. (1998), Le Duc et al. (2000), Milne et 91 92 al. (2002), Rowe et al. (2016) and Smart et al. (2017). Additional, although relatively few (27), individual site data for woodlands were also analysed, and tree growth data published by the Forestry 93 94 Commission (2002) for the United Kingdom were also used to estimate average NPP values.

- 95 Methods
- 96
- 97 *Production data*
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99 The 771 non-woodland sites comprised grassland, shrubland and mixed moorland, most of which are 100 grazed by sheep, managed for grouse shooting, or unmanaged. Except for six sites with more detailed 101 measurements, production estimates were confined to above-ground biomass. The largest single set 102 of results came from Milne et al. (2002), who measured above-ground production at 66 sites for 103 shrubs and 17 for herbs in six regions of England and Wales (NE England, NW England, North Pennines, 104 South Pennines, SW England and Wales), giving a total of 597 individual plots. The Milne *et al.* (2002) data form a coherent set, with replication at different sites, eight different dominant plant species or 105 106 plant development stages, and data for three years (1995-1997). At each site, replicate measurements 107 were made in two or three different years (1995, 1996, 1997) and at between two and nine plots, 108 three in most cases. Other major (results for > 10 sites) non-woodland data sets were from Pearsall & 109 Gorham (1956), Paterson et al. (1997), Marrs et al. (1998), LeDuc et al. (2000), Rowe et al. (2016), and 110 Smart et al. (2017). The remainder of the data were from studies at fewer than 10 locations each. The 111 sites in these studies refer to areas of representative vegetation, from which quadrats with areas 112 between 0.0625 and 1 m² were sampled to quantify above-ground production (g dry mass m⁻²).

We found NPP data for 27 individual GB woodland sites, obtained by various methods. In three cases, total NPP was estimated from biometric or CO₂ exchange (Reichle, 1981; Morison *et al.* 2012; Fenn *et al.* 2015). In the studies of Ovington & Pearsall (1956), Smart *et al.* (2017) and others, aboveground NPP was determined. We also used data summarising tree wood yields at the national scale (Forestry Commission 2002).

The production measurements were for an initial total of 811 sites, although some were discounted (see Results) so the final total was 796 (see Results). Of the 811 values, 90 were for the period 1932-1989, 646 for 1990-2000, and 75 for 2001-2014. The full collated data set, including information on sampling methods, is given in Table S1, and the geographical distribution of the data is shown in Fig. 1.

123

124 Estimation of NPP

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The results are expressed and analysed in terms of total net primary production, referred to as NPP, including above and below ground production (ANPP and BNPP), with units of g C m⁻² a⁻¹, which is what the N14CP model predicts. The **literature** data included only a few instances in which all components of NPP had been estimated. In the great majority of cases, conversions and assumptions
 were applied as follows (see Table S1 for the conversion procedure for each sampling site).

(a) For all data reported as dry mass, it was assumed that 50% was C (cf. Schlesinger &Bernhardt 2013).

133 (b) For herbs, reported production data were mainly measurements of peak above-ground 134 biomass, at sites without grazing or with exclosures erected to prevent grazing (e.g. Milne et al. 2002). 135 The above-ground peak biomass was equated with the year's growth, and therefore taken to 136 represent ANPP. The results for Agrostis-Festuca grassland reported by Milne et al. (2002) were 137 obtained from three clippings per growing season, to simulate grazing. According to Scurlock et al. (2002), for grasslands, peak biomass is a "reasonable benchmark indicator of the magnitude of 138 139 productivity for study sites within a particular sub-biome" (see also Lauenroth et al. 1986). The 140 measured above-ground values of dry mass were converted to ANPP (g C m⁻² a⁻¹), and these were 141 converted to NPP by multiplying by 2.0, this factor being based on measurements (Scurlock et al. 2002; 142 Olsen et al. 2013; Sims & Singh, 1971) and modelling (Gill et al. 2002; Hall et al. 1995; del Grosso et al. 143 2008).

(c) For shrubs, sampled at sites without grazing or with exclosures erected to prevent grazing,
the current season's new biomass had been identified and isolated after sampling the entire plant by
clipping with secateurs, and then quantified. We made the same assumptions as for herbs, i.e. that
peak biomass provides a reasonable measure of annual production, and that ANPP can be converted
to NPP by multiplying by 2.0.

(d) *Pteridium aquilinum* possesses an extensive rhizome system (Marrs & Watt, 2006), which
supplies approximately half of the carbon for the growth of (above-ground) fronds, the rest being
obtained by photosynthesis (54 % from rhizome reserves from the data of Williams & Foley 1976). We
assumed that the rhizome biomass is essentially the same at the start and end of the year in question
(Pakeman *et al.* 1994), which means that the net annual production (dry mass) can be equated with
the peak mass of the above-ground standing crop. Thus, NPP was estimated by multiplying the peak
dry mass by 0.5 to convert to g C m⁻² a⁻¹.

(e) In four studies of woodland, total NPP was reported by the original authors, although even
in these cases the derivation of NPP involved some assumptions and estimations. In all other cases,
we estimated woodland NPP from three partial measures of production, i.e. ANPP, wood increment,
and litterfall, using relationships (Fig. S1) derived from data for boreal and temperate woodlands
published by Scarascia-Mugnozza *et al.* (2000), Gower *et al.* (2001) and Fenn *et al.* (2015). The
calculations yielded an average ANPP/NPP ratio of 0.69 for all the tree plots, which means that 31%
of the production is below-ground. This approach ignores the successional state of the woodland, and

the tendency for NPP to rise then decline as trees mature (Ovington, 1962; Peterken & Newbould,
1966; Gower *et al.* 2001). Clark *et al.* (2001) discuss in detail the problems inherent in estimating
woodland NPP.

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167 *Modelling with N14CP*

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169 A full description of the N14CP model is available in Davies et al. (2016a). A brief summary, with a 170 focus on NPP is provided here. N14CP describes plant-soil pools and fluxes of C, N and P, driven by 171 climate, N_{dep}, sulphur deposition (S_{dep}), base cation deposition (BC_{dep}), weathering of base cations (BC) 172 and P, and known or assumed vegetation history. The model simulates on a quarterly time step the growth and decay of plant biomass, N fixation, the production of litter and its incorporation into soil 173 174 organic matter (SOM), the immobilisation of nutrients (N, P) by SOM, SOM turnover, leaching losses, 175 and denitrification. N fixation is positively related to temperature and P availability, but does not 176 depend on vegetation type. It is down-regulated by N_{dep} such that no additional N from N_{dep} becomes

177 available until the N_{dep} rate outstrips the potential fixation rate.

Four vegetation classes are represented: broadleaved trees, needle-leaved trees, herbs and 178 179 shrubs, with varying stoichiometric demands and litter characteristics based on literature values 180 (Tipping et al. 2012). Bracken was modelled as a herb. Plant biomass is divided into coarse and fine 181 tissues; the C:N:P stoichiometry of the former (wood and coarse roots) is constant, while the stoichiometry of the latter varies with N and P availability. Biomass is not divided into above and below 182 ground fractions, and only total NPP is simulated. Following von Liebig's Law of the Minimum, NPP is 183 184 determined by one of four factors, MAT, MAP, N availability, or P availability. First, the maximum NPP 185 is calculated dependent upon MAT or MAP, maximum NPP being estimated by quantile regression, 186 taking the 90% quantile of NPP data collated by Chapin *et al.* (2011), as functions of MAP and MAT 187 (Davies et al. 2016a). Then, if nutrients are insufficient to meet the stoichiometric requirements of the maximum plant growth, whichever nutrient is in the shorter supply is taken to be limiting. Plant 188 189 growth occurs in the second and third quarters of each year, and therefore there is no variation in 190 growing season length. 191 Davies et al. (2016a) performed a non-site-specific parameterisation of the model by fitting 192 plot scale soil and soil water C, N and P data (pools and fluxes) for 44 sites vegetated with broadleaved 193 trees, needle-leaved trees, herbs and shrubs in northern Europe, then tested the model against data 194 from a separate set of 44 northern European sites. The model yielded reasonable average values of

195 element pools and fluxes, but inter-site variations could only be reproduced by allowing the

196 weatherable P pool to vary on a site-by-site basis, suggesting that the availability of site-scale P

197 weathering data may improve prediction. However, in the absence of such data, a crude distinction 198 on the basis of current soil type is used. Separate weatherable P pools, available at the start of soil 199 formation, were assigned to podzols and rankers on the one hand (smaller initial P pool), and all other 200 soil types on the other (larger initial P pool). The parameterisation approach did not use measured 201 NPP data as a constraint; instead, the fitting aimed to achieve an overall mean NPP of 75% of the 202 maximum values, which was roughly in line with available data (Tipping et al. 2012; Davies et al. 203 2016a). Whilst this was the applied constraint, the non-site-specific parameterisation produced an 204 average value for the 88 sites that was 83% the maximum NPP. 205 In the present work, modelling was performed as follows. 206 (a) The parameterisation by Davies et al. (2016a) was used to predict NPP on a 5 km x 5 km 207 grid across GB. Simulations started at the beginning of the Holocene (10,000 BCE), at which point soil 208 C, N and P began to accumulate. 209 (b) The UK Land Cover Map 2007 (LCM2007, Morton et al. 2007) was processed to provide 210 fractional covers on this grid scale for the broad habitat types examined here. We modelled both grass 211 and bracken as herbs, and shrubs were modelled as a separate vegetation class. Using the LCM2007 212 as a contemporary starting point, we defined the history of each land use fraction in each grid cell 213 using a range of sources and assumptions. Spatially resolved land cover data for 1931 and 1600 was 214 available from Stamp (1931) and Thirsk (1989) allowing us to estimate land use change over these 215 periods for each grid cell fraction. It is assumed in the model land use history that all contemporary 216 needle-leaved trees in GB are plantation, as native pine-leaved forests make up ~1% of the existing 217 stand. The planting date for needle-leaved trees was assumed to be either 1920 or 1955 (determined 218 by changing land fractions between the Dudley Stamp and LCM2007), as these were two periods of 219 marked acceleration in planting across GB (P. Crow, Forest Research, personal communication). 220 Fractions with shrubs in 1600 were assumed to have had the same vegetation prior to this date. Land 221 use fractions classed as rough grassland in 1600 were assumed to have been converted to grassland 222 from broadleaf forest at some point in the past, with a clearance date based on data from Roberts 223 (2013). Fractions classed as broadleaf forest in 1600 were assumed to have been ancient woodland at 224 this time and to have developed naturally, having succeeded from herbaceous plant cover in ~6000 225 BCE. This succession date was also applied to fractions where forest was cleared prior to 1600. The 226 sensitivity of outputs to these forest clearance and succession assumptions has been explored 227 previously (Tipping et al. 2017), where the model outputs were shown to be much more sensitive to 228 variation in contemporary N_{dep} than the assumptions about land use in the distant past. 229 (c) Climate inputs were defined using gridded Meteorological Office data between 1910 and

230 2011. For earlier dates, MAP at each site was assumed constant, while MAT variation was estimated

231	using a historical anomaly based on the temperature record for northern Europe derived by Davis <i>e</i>
232	al. (2003) from pollen records. Inputs of N_{dep} , S_{dep} and BC_{dep} resolved spatially (5 km x 5 km gric
233	squares) over the period 1800-2010, and taking vegetation type into account, were estimated as ir
234	previous modelling reported in Tipping <i>et al</i> . (2017).
235	(d) The initial pool of weatherable P was determined by the soil classification (podzols and
236	rankers vs. other soils) as described above; the fraction of the two soil classes in each grid cell, as
237	calculated from soil survey data (National Soil Resources Institute 2013; Soil Survey of Scotland Staff,
238	1981), yielded a weighted average weatherable P pool. The weathering of BC in a cell was estimated
239	from the local rock mineralogy.
240	
241	Statistics
242	
243	Summary statistics and conventional linear regression analyses were computed with Microsoft Excel
244	Generalised Linear Models (GLMs, McCullagh & Nelder, 1989) were used to model NPP against
245	cumulative N_{dep} for herbs and shrubs. In view of the skewed nature of the data, even after taking
246	logarithms, a gamma distribution of the data was assumed, rather than the Gaussian distribution. A
247	log link function was employed, which dampens large skew whilst also respecting the bounded nature
248	of the observed data, i.e. that NPP values cannot fall below zero. Regressions for each vegetation type
249	were conducted independently and any evidence of spatial autocorrelation, resulting from close
250	proximity of some sites, was assessed in each case. For both herbs and shrubs, the variation betweer
251	sites separated by large distances was no different to the variation in sites separated by smal
252	distances. All GLMs were fitted using the R statistical programming language (R Core Team 2017). This
253	approach does not yield conventional r ² values because the parameters are estimated via maximum
254	likelihood rather than by minimising variance through ordinary least squares. Rather, pseudo r ² values
255	can be obtained that are comparable to conventional r^2 in the sense that they quantify the
256	improvement of the fitted model over a null model and work on a 0 to 1 scale. Here we used
257	McFadden's pseudo r^2 , which is based on a ratio of model deviance to null deviance.

258 Results

In the following text, "measured NPP" means total NPP, i.e. ANPP+BNPP, estimated from
measurements, e.g. from peak above-ground biomass, as detailed in Methods. Modelled NPP also
refers to total NPP.

262 The measured NPP values were put into perspective by plotting all values against MAP and 263 MAT (Fig. 2), to allow comparison with the theoretical climate-dependent maximum values, from 264 global observations (see Methods) derived by Davies et al. (2016a) for the N14CP model. These 265 maxima apply when nutrients are not limiting. Considering all the data (811 points), there were few 266 exceedances; for MAP only 2% of the values exceeded the theoretical maximum, while for MAT 7% 267 showed exceedance. The fact that in the great majority of cases the NPP values fell below the theoretical climate-based maxima is consistent with NPP in GB semi-natural ecosystems being limited 268 269 by factors other than climate. This agrees with results obtained with the N14CP model, which 270 predicted that N availability was the factor limiting NPP at all but two of the 811 sites, which were P-271 limited.

Before further analysis of the results, nine outlier points (two for herbs, seven for shrubs) with high NPP (> 1000 gC m⁻² a⁻¹) were removed. Furthermore, in 6 cases there was not a match with modelled values, because the land cover map (Morton *et al.* 2011) used in the simulations (see below) did not include the relevant vegetation type in the grid cell in question. These sites were also omitted from the analysis. Therefore, the final data set for analysis comprised 796 points. The removed data are indicated in Table S1.

We compared average measured values of NPP with the corresponding averages of the model 278 279 predictions (Table 1). The measured estimates for herbs and trees tended to exceed the predictions, 280 whereas the opposite was true for shrubs. The overall measured:modelled ratio was 1.03. Only for the 281 small number of needle-leaved sites was the ratio appreciably different from unity, the measurements 282 exceeding the predictions by a factor greater than 2.0. However when the average measured and predicted values for needle-leaved trees in the national forest inventory were compared, the 283 284 agreement was much closer (Table 1). For each vegetation class the variability in NPP was high 285 (average relative standard deviation, RSD = 0.43), as expected from previous work by Milne et al. 286 (2002). The modelled values were appreciably less variable (average RSD = 0.19), which reflects the simple assumptions of the model, i.e. that NPP depends mainly upon nutrient availability and 287 vegetation class. 288

The average measured and modelled values for different vegetation classes show the expected relationship (Fig. 3), which arises mainly because trees have higher NPP than non-trees. For herbs and shrubs, there were sufficient data to obtain average NPP values for the 5 km x 5 km grid cells used by the N14CP model (see Methods). If cells with 10 or more measurements were used for averaging, data for a total of 26 cells were obtained, covering a reasonably wide geographical range (Fig. S2). As shown in Fig. 4, the average measured values were significantly correlated with the modelled values ($r^2 = 0.22$, p = 0.011), and the regression slope is close to unity. A similar relationship was obtained if cells with five or more measurements were used (n = 40, $r^2 = 0.16$, p = 0.010), but with 15 measurements as the lower limit, the relationship, although positive, was insignificant (n = 20, p =0.13).

299 The main reason for modelled variation in NPP is variation in N_{dep} and therefore the results of 300 Fig. 4 are consistent with N fertilisation causing increases in NPP. To test this further we regressed 301 measured and modelled NPP against modelled cumulative N_{dep} (to the date of sampling) which is a 302 quantitative indicator of the N-enrichment of a site, taking into account both intensity (N_{dep}) and time 303 (cf. Duprè et al. 2010). Since herbs and shrubs were modelled separately, and since they have different 304 average measured:modelled ratios (Table 1), separate testing was performed. Both vegetation classes gave significant positive NPP relationships to N_{dep} , with slopes for the measured data between 50-60% 305 306 of those from modelled values (Table 2, Fig. 5). The pseudo-r² value for the herbs plot was 0.018, that 307 for the shrubs plot was 0.023.

We could not establish any temporal trends in the measured values of NPP, principally because there were no instances in which the same vegetation type was monitored at the same site over a sufficiently long period of time. The longest sequences, which apply to bracken, are for less than 25 years, and little change in NPP is calculated with N14CP for the monitoring periods (Fig. S3). Short term variation in NPP is appreciable; for the 67 sites in the database where NPP had been measured at different times, the average RSD was 0.28, which is already greater than the modelled RSD, and more than half the total RSD of 0.45 (from data in Table 1).

315 Modelled time series (Fig. 6) suggest substantial changes in NPP over the period 1800-2010, 316 almost entirely driven by the increases in N_{dep} . Very modest NPP increases are calculated in the 317 absence of N_{dep}, resulting from slightly faster nutrient cycling due to increasing temperatures (Tipping 318 et al. 2017). The model calculations suggest that fertilisation by N_{dep} caused average increases in semi-319 natural NPP over the period 1800 to 2010 of 30% for shrubs, 71% for herbs, and 91% for broadleaved 320 trees. Results for needle-leaved trees are not shown, since many sites are in conifer plantations, established in the 20th century, which means that modelling long-term variations in NPP is 321 322 inappropriate.

323 Discussion

324 Measured estimates of NPP largely agree with the modelled values, based on overall average values for different vegetation classes (Table 1, Fig. 3), and a significant regression with a slope near to unity 325 326 for averaged data within model cells (Fig. 4). Significant regressions of measured NPP against 327 cumulative N_{dep} (Fig. 5) are consistent with the major role of fertilisation by atmospherically deposited 328 N being the main reason for NPP variation. Although the slopes from measured data in Fig. 5 are 329 smaller than the model-based values, the standard errors (Table 2) mean that the ranges overlap, and 330 so we cannot conclude that there is significant disagreement. The extra supply of N has increased the 331 flux of N that can cycle through the soil-plant system, permitting more C to be fixed by photosynthesis. 332 Whereas previous field-based work has shown effects of N_{dep} on trees and, to a lesser extent, shrubs, 333 based on temporal increases in standing biomass (see Introduction), this is the first study to do so for 334 herb ecosystems, and to be based on direct measurements of annual production. Taken together with 335 the previous results, the case for a widespread vegetation response to fertilisation by N_{dep} is strong.

336 The quantitative agreements between modelled and measured average values, both overall 337 and for different vegetation classes (Table 1, Fig. 3), arise partly because in parameterising the N14CP 338 model Davies et al. (2016a) set as a fitting target an average NPP value, at 75% of the mean maximum 339 NPP estimated for the northern European sites for which observations were available (see Methods), 340 and this would tend to generate a modelled average NPP in the right range. Nonetheless, the sites 341 considered in the present work are different from those used for parameterising the model, and so 342 the results provide a partially independent test. The vegetation class variation of Fig. 3 arises because 343 the model assumes the different vegetation classes to have different stoichiometric (C:N:P) 344 compositions.

345 The NPP-N_{dep} trends of Fig. 5 are obscured by the high degree of scatter in the observations, 346 which leads to low pseudo-r² values, and so little of the variance is explained the statistical model. Some of this scatter likely arises because different plant species within a vegetation class have 347 348 different NPP, as noted by Milne et al. (2002), extending to different phases of Calluna vulgaris. Milne 349 et al. (2002) also noted various dependences among their data on measured physical and soil 350 properties, although not applicable to all species, and not generalisable, since trends that applied to one dominant species did not apply to others. A further contribution to the data scatter is short-term 351 352 temporal variation, with an RSD of 28%. Scatter could be caused by numerous additional factors 353 including mineral fertilisation (either deliberate or by proximity to other land receiving fertiliser), 354 nutrient deficiencies (P, molybdenum, potassium etc.), herbivory, animal excreta, pests, disease, light 355 availability, ozone levels, soil moisture, soil fauna, and small-scale climatic variation. Scatter could also 356 arise from the approximate nature of the conversion of production values to NPP.

357 Notwithstanding the high scatter, the trends of Fig. 5 are statistically significant, and provide evidence that N_{dep} affects NPP. This is reinforced by the agreement between measured and modelled 358 359 averaged NPP (Fig. 4), which arises because the averaging reduces variations among sites and years. 360 Undoubtedly, the best way to investigate variability in NPP would involve a sampling programme that 361 allows appropriate spatial averaging of production data, while covering wide ranges of climatic and biogeochemical factors, and N_{dep}. Although the data assembled here are imperfect in these respects, 362 363 their large number, and the wide range of N_{dep} covered, justify our meta-analysis. 364 Milne et al. (2002) suggested that the spatial variation of NPP in their dataset reflected the 365 length of the growing season, which would also explain why Calluna vulgaris NPP in Scotland, based on data reported by Miller and Watson (1978), Miller (1979) and Grant et al. (1982), was low compared 366

367 to other parts of GB. However, Milne et al. (2002) reported low Calluna vulgaris production in SW 368 England, which has the longest growing season in GB. We checked the dependence of measured NPP 369 on latitude (approximately inversely proportional to growing season length) for the averaged data of 370 Fig. 4, and found no relationship. Variations in NPP are better explained by the predictions of N14CP 371 (Fig. 4), which attributes them to variations in N_{dep} and does not take variation in growing season length into account. The model could potentially have predicted that average temperature in the 372 373 second and third annual quarters was the factor limiting production, but instead it predicted N 374 availability, dependent upon N_{dep} , to be the limiting factor in these unfertilised semi-natural

375 ecosystems.

376 There have been other studies in GB relevant to the present work. Rowe et al. (2012) showed 377 that in semi-natural habitats, readily-mineralisable N (a measure of plant N availability) increased with 378 N_{dep} , which agrees with the assumptions of the N14CP model, and is clearly relevant to plant growth. 379 However, Rowe et al. (2014) found that both bicarbonate-extractable P stock and mineralisable N 380 were predictors of Ellenberg N score, taken to be an independent metric of productivity, but that the 381 P variable was superior. In a more spatially-limited study of bracken productivity, Rowe et al. (2016) 382 found no dependence on either total soil N or soil organic P. Field et al. (2017) found that N additions 383 increased *Calluna vulgaris* growth in an experimental field study. Thus, there is some confirmatory 384 evidence for the limitation of productivity by N, but it is not conclusive. The reason that the N14CP 385 model rarely predicts P limitation is that semi-natural soils in GB are young and calculated still to have 386 significant weatherable apatite.

Although the measured NPP values analysed in this work cover the period 1932 to 2014, i.e. 82 years, 80% referred to the period 1990-2000, and so the agreement achieved with modelled results depends on spatial NPP variation, which according to the model is due to variation in N_{dep}. Therefore, the temporal predictions of Fig. 6 must be justified by time-for-space substitution. However, there is 391 a temporal link through the study of Tipping et al. (2017) who used N14CP to simulate increases in the 392 SOC concentration of GB soils, owing to increased litter inputs associated with increases in NPP caused 393 by N_{dep} fertilisation. Simulated SOC increases over the period 1959 to 2010 agreed with statistically-394 significant measured changes from sample-resample observations at nearly 2000 field sites (different 395 to those of the present work). Moreover, increases in broad-leaved woodland SOC were highly 396 significantly related to N_{dep} (increases for non-woodland SOC were positively related, but not 397 statistically significant). These results for SOC provide support for the increases in NPP modelled here. 398 Furthermore, the measured and modelled increases in SOC beneath broadleaved trees were found to 399 be greater than those under non-tree vegetation, which is consistent with the larger modelled NPP 400 response of broadleaved trees. Taken together, the present results and the previous SOC modelling 401 provide a coherent quantitative description of GB semi-natural ecosystems, consistent with two large 402 field data sets. As already pointed out by Tipping et al. (2017), the N14CP modelling shows that the additional 403 404 storage of SOC, resulting from increased NPP caused by N_{dep}, cannot be regarded as permanent burial, 405 because of SOC turnover. Maintenance of the elevated litter inputs would be needed to prevent the extra SOC returning to the atmosphere over decades to centuries. This implies a need to maintain N_{dep} 406 407 at present levels, which is undesirable with respect to plant diversity, since that is known to have been 408 reduced by N_{dep} in GB (Stevens et al 2004; Maskell et al. 2010). Therefore, there is a conflict between 409 the goals of carbon sequestration and the conservation of plant species diversity in semi-natural 410 ecosystems.

411 Conclusions

- 412 Semi-natural NPP in GB is generally lower than maximum possible values as determined by • 413 climate, and modelling with N14CP suggests that this is due to N limitation. 414 • Mean measured estimates of NPP are in reasonable quantitative agreement with mean modelled values, and the modelled order of NPP by vegetation class (herbs and shrubs < trees) 415 416 is seen in the data. 417 • Measured NPP, although displaying high variability, is significantly positively correlated with N_{dep} for both herbs and shrubs, approximately as predicted with N14CP. 418 If space-for-time substitution is accepted, there have been substantial (30-100%) increases in 419 •
- If space-for-time substitution is accepted, there have been substantial (30-100%) increases in
 NPP in British semi-natural habitats over the past several hundred years, due to N_{dep}.

421 Acknowledgements

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- 429
- 430 Supplementary material
- 431 Table S1. The data base
- 432 Fig. S1. Regression relationships in data for temperate and boreal broadleaf and conifer woodlands,
- taken from Scarascia-Mugnozza et al. (2000), Gower et al. (2001) and Fenn et al. (2015).
- Fig. S2. Locations of model grid cells containing >= 10 measurements; (see Fig. 4). Left, herbs; right,
 shrubs.
- 436 Fig. S3. Bracken time series; points are measured values, lines are model predictions. Data at Cannock

437 Chase are from two separate areas.

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- 582 Table 1. Averaged values of NPP (g C m⁻² a⁻¹), estimated from observations and modelled with N14CP.
- 583 Numbers of sites are given by *n*. National averages for woodland are based on Forestry Commission
- 584 (2002) data for England, Wales and Scotland.

		<u>measured</u>		<u>modelled</u>		<u>measured</u>
dates	n	mean	SD	mean	SD	<u>/modelled</u>
1932-2014	19	<mark>585</mark>	<mark>176</mark>	444	59	<mark>1.32</mark>
1932-2014	6	688	280	305	125	2.25
1952-2014	298	378	199	306	48	1.24
1958-2014	473	382	184	419	42	0.91
1932-2014	796	<mark>388</mark>	<mark>195</mark>	377	72	<mark>1.03</mark>
1995-1999	-	574	-	497	84	1.15
1995-1999	-	538	-	463	141	1.16
	1932-2014 1932-2014 1952-2014 1958-2014 1932-2014 1995-1999	1932-2014191932-201461952-20142981958-20144731932-20147961995-1999-	dates n mean 1932-2014 19 585 1932-2014 6 688 1952-2014 298 378 1958-2014 473 382 1932-2014 796 388 1995-1999 - 574	dates n mean SD 1932-2014 19 585 176 1932-2014 6 688 280 1952-2014 298 378 199 1958-2014 473 382 184 1932-2014 796 388 195 1995-1999 - 574 -	dates n mean SD mean 1932-2014 19 585 176 444 1932-2014 6 688 280 305 1952-2014 298 378 199 306 1958-2014 473 382 184 419 1932-2014 796 388 195 377 1995-1999 - 574 - 497	dates n mean SD mean SD 1932-2014 19 585 176 444 59 1932-2014 6 688 280 305 125 1952-2014 298 378 199 306 48 1958-2014 473 382 184 419 42 1932-2014 796 388 195 377 72 1995-1999 - 574 - 497 84

586

Table 2. Summary of regression analysis results, ln NPP (gC m⁻² a⁻¹) vs cumulative N_{dep} (gN m⁻²). SE =

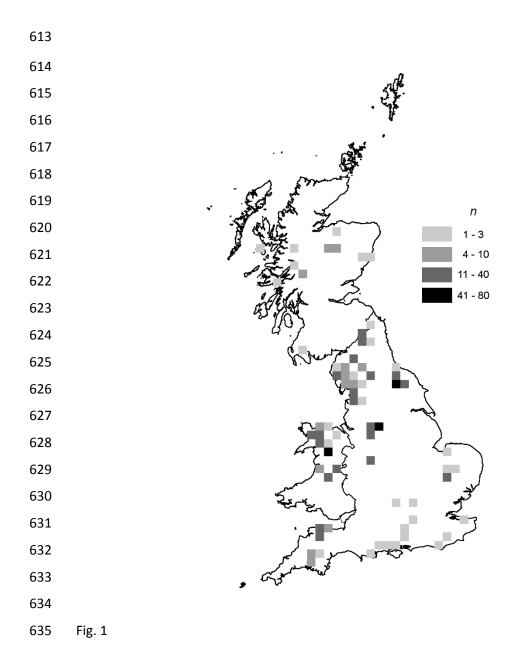
588 standard error.

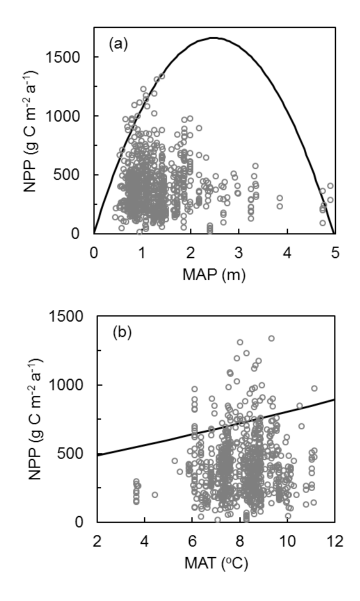
Vegetation class	measured			modelled			
		value	SE	р	value	SE	р
Herbs	intercept	5.73	0.09	0.000	5.35	0.016	0.000
	slope	0.00101	0.00043	0.021	0.00188	0.00008	0.000
Shrubs	intercept	5.72	0.084	0.000	5.67	0.008	0.000
	slope	0.00111	0.00040	0.006	0.00179	0.00004	0.000

589

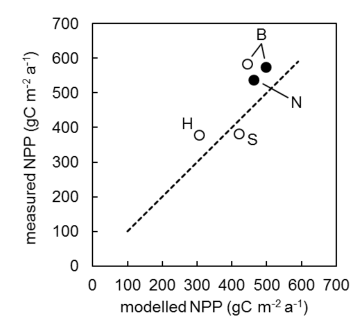
590

- 591 Figure captions
- Fig. 1. Map showing densities of sampling sites within 20 km x 20 km squares (16 model grid cells). See
 Table S1 for details.
- 594 Fig. 2. Measured NPP compared with (a) MAP limits and (b) MAT limits used in the N14CP model. The
- limits, shown by solid lines, are fits of the 90% quantiles of reported NPP values collated by Chapin *et al.* (2011). Estimated NPP values at individual sites are shown by open circles.
- Fig. 3. Measured vs modelled mean NPP for different vegetation classes, using data from Table 1. Open
 circles are averages from individual sites (needle-leaved trees omitted), filled ones are Forestry
 Commission (2002) data. Key: B = broadleaved trees, N = needle-leaved trees, H = herbs, S = shrubs.
- 600 The 1:1 line is shown.
- Fig. 4. Average measured vs modelled NPP for 5km x 5km grid cells with 10 or more observations. The time periods covered for each point were up to 17 years, although most were for 3 years, between 1978 and 1997. All but one of the 26 points arise from the Milne *et al.* (2002) data set, the other is bracken data from 1978-1996 (Marrs *et al.* 1998). Cell locations are shown in Fig. S2. The open circles show results for herbs, the closed ones for shrubs. Standard errors are shown. The regression line is 1.01x - 17.7, $r^2 = 0.22$, p = 0.011, n = 25.
- Fig. 5. Regressions of ln NPP vs cumulative N_{dep} for (a) herbs and (b) shrubs. The solid lines indicate
 regressions of the measured values, the dashed lines are modelled trends. See Table 2 for a summary
 of the regression statistics. All plotted points are open circles.
- Fig. 6. Modelled changes of N_{dep} and NPP over time for herbs (a,d), shrubs (b,e) and broadleaved trees
 (c,f) in semi-natural land areas of Great Britain. The central line is the mean, dotted lines show 5 and
- 612 95 percentiles.

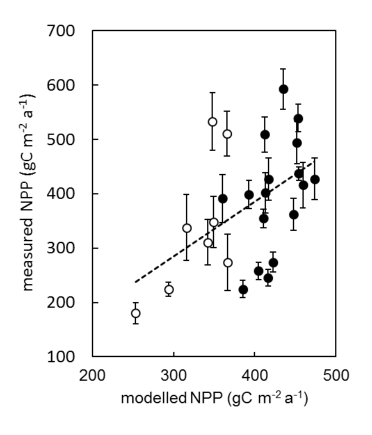




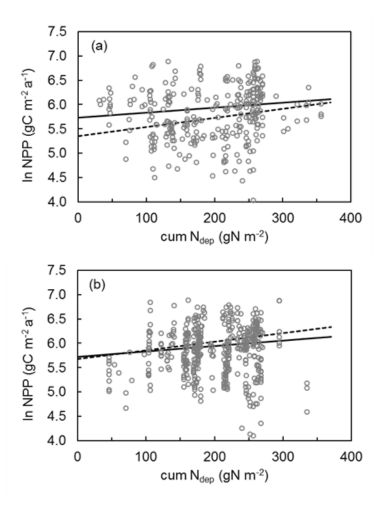
637 Fig. 2



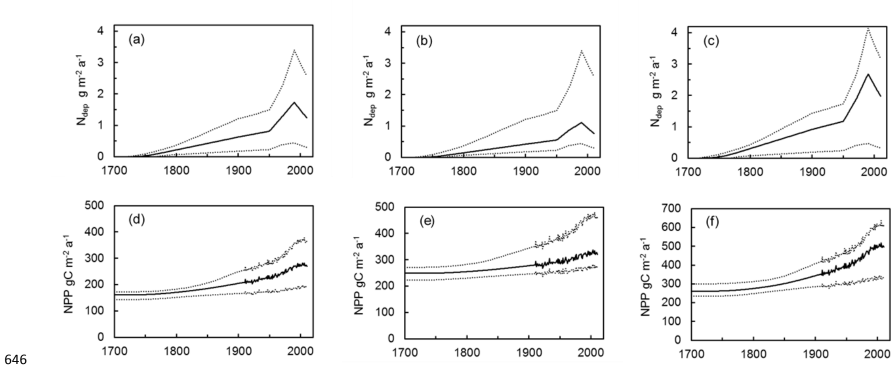




641 Fig. 4









RESPONSES TO EDITOR'S AND REFEREES' COMMENTS

Line numbers in the following refer to the Revision. Original comments are in plain text, responses in italics.

In the highlighted Revision, changes in response to comments are highlighted in yellow. Our own additional corrections and improvements are highlighted in blue.

We realised that the references were not in the correct style for BG, and corrected them. We also change "Figure" to "Fig." throughout, in accordance with BG practice.

In answering the Editor's question about tree AG and BG NPP, we noticed an error in 3 of the broadleaved tree estimates of TNPP, in Table S1. These have been corrected, and also the consequent errors in Table 1 and Fig. 3. The text in lines 279-280 has been modified (see response to one of the Editor's points below).

Editor comments:

I find this to be an interesting study, and I applaud the large dataset to compare your model with. In addition to the two reviewers' comments, I have a few others that you should address in your revision.

Lines 142-146. I would argue that using a constant factor of 2.0 to convert ANPP to NPP would estimate a systematic bias. For example, if you look at the cited Chapin et al. (2011) text, the belowground portion of NPP is strongly dependent on ecosystem type, being much lower for forests than for grasslands.

You are correct about the BNPP being lower for forests. The ratio of TNPP to ANPP in our data (Table S1) for broadleaved trees is 1.45, which means that the BNPP is only 31% of the total. For the needleleaved trees the TNPP/ANPP ratio is 1.44, so BNPP is also 31% of the total. We have now explained this in the section on trees (was d, now e) in Methods, lines 160-162. The factor of 2.0 for TNPP/ANPP was only used for herbs and shrubs, and so is only mentioned in sections (b and c).

Make sure that all abbreviations used in the text are defined on first usage.

We edited the text on the following lines 23, 77, 127, 171, 174, 209. We did not explain CO₂ (line 114), because it is very well known. Nor did we explain directional abbreviations (S, NW etc).

Lines 266-267. If I read the text and table correctly, then the description for trees is reversed, i.e., the measured values for trees exceed the predictions. Also, some description for the poor fit with needle-leaved trees is called for, although in the end, this PFT is a small part of the paper.

You are correct, the measured values for trees do exceed the predictions, apologies. We have modified the text, also to comment on the needle-leaved trees (lines 279-280).

Similar to Reviewer 3, I think that you should give the r2 for Fig. 5 and Table 2. It looks quite poor. With this many points, regressions can be statistically significant and explain very little of the actual variance. You discuss the poor fit in the discussion but you never actually quantify it. I must admit that based on the fit in Fig. 5, it gives me much less confidence in the predictions of Ndep effects on NPP in Fig. 6. Perhaps this could bear greater justification in the discussion.

Technically, the statistical approach does not produce a conventional r2, which why we did not report values in the original submission. But we can report pseudo-r2 values, and these are included in the revised version (lines 306-307). We have also explained what pseudo-r2 means in the Methods section (lines 252-257).

Please note that the use of the model to run long-term simulations is also justified by the results in Fig. 4, not Fig. 5 alone. We are not sure that our data and results can bear "greater justification" – they are what they are, and they show something that is statistically significant. But we have tried to explain this more cogently in rewritten text in the Discussion (Lines 345-363).

In Fig. 5., what do the open and closed symbols represent?

There are no closed symbols, but it may appear so because there are many overlapping points. The same applies to Fig. 2. We added text to the caption to draw readers' attention to this (lines 596, 609).

Reviewer #2

In this manuscript the authors have compiled biomass data from (semi)natural mainly low vegetation ecosystems in UK and converted the biomasses into estimates of NPP. These NPP data are then compared to results from a previously developed model. The estimated NPPs show an increase and based on the model simulations the authors conclude that this is 'a fertilizer' response to the increased N deposition.

I am not a modeler so my evaluation is mostly focused on the data part. I find the description of the data compilation and all the manipulations needed to get to NPP estimates a bit sloppy, so that I lose confidence in the whole exercise. This although that I am very positive to the approach and applaud the work done by the authors to try to estimate the fertilization effect from N deposition in non-forest system.

We have tried to improve the description of data compilations to make it less "sloppy", as explained in response to the Referee's following specific comments.

Attention to the following points may improve the manuscript (mentioned by line number):

99-121: Most of the data are on standing aboveground biomass but that is not at all always clear here although it appears from some of the explanation in the next sections. The descriptions of the original data need to be clear, so that the estimations and assumptions made to come to NPP also become clear.

This paragraph is an overview of data sets, and so does not include details of the original data, estimations and assumptions – these come next in the Method section. But please note that Table S1 gives the measurement method for each site and this is stated on line 121. We altered the text here (lines 100-101) to make it clear that the non-tree site data were almost entirely above-ground biomass.

114-15: 'In studies of' give a few words on what these studies are about

These studies were investigations of primary production, in one case with a view to correlating it with plant diversity. With respect, we do not think there is any point in going into any detail about the studies' aims in a Methods section..

141: I understand the argument that peak biomass can be equal to NPP in grassland (line 139-141), but I think assuming the same for shrubs is very rough and doubtful. The characteristics of shrubs are that they have woody stems and branches that grow and persist for many years. So this is not a valid assumption. Also no argument is given.

We are puzzled by this comment, since the text (lines 137-139 of the original) describes how shrub biomass was quantified. But we suppose that this could be clarified by separating grass and shrubs in the description – thus we now have sub-sections (a) to (e) in our list. The revised text is in lines 133-148.

145-6: delete or reformulate; I do not understand how an assumption can be 'free of systematic error' when you have very limited documentation for the assumed value....

We agree that this should be deleted.

223, 307: here PFT is used for what elsewhere is 'vegetation class'. Please be consistent and do not use undefined abbreviations .

Thanks for pointing this out, we used PFT in an earlier draft and it's persisted. We edited the text at lines 220 and 325 of the Revision.

342, 344: 'relative low' or 'low' values are mentioned; compared to what, it is not completely clear.

We have rewritten this paragraph, we agree that it was poorly expressed in the original (lines 364-375).

347-8: The model does not account for variations in growing season length. But increasing growing season length due to climate change could alone also explain the observed increase in NPP. This point need some lines of discussion and arguments why Ndep is the most likely reason for the observed (if it is).

We added a line in the model description to explain that the model does not include variation in growing season see lines 188-190. We rewrote the text in the Discussion to explain more clearly that there is no evidence for a dependence of NPP on growing season, and that N_{dep} is a better predictor (lines 364-375).

376: I think this should read..... 'be maintained otherwise the extra SOC is returned to the atmosphere'

Yes, this was not well expressed. Please see our response to the last comment of Referee 3, which entailed rewriting much of this text.

569: 20 x 20 km is 16 model grid cells, not 4!!!! (4x4)

You are right, apologies. This has been corrected (line 592).

571-2: The legend is only partly explaining the figure. What is a) and b) and what does the lines show. I can figure it out but better if you guide the reader.

We agree that the caption was poorly-written. We have improved it, lines 594-596.

Reviewer #3

The authors explore the influence of Ndep on NPP using a data-model comparison approach. Broadscale NPP data are difficult to synthesize and the authors do a good job at describing the datasets and their associated caveats. I also appreciate that the data are provided in the supplemental materials. The modeled and measured equations for Ndep influence on NPP will no doubt be useful for future modeling efforts.

I have some minor comments:

Lines 163-232: it would also be effective if you provided a model structure schematic in addition to the description. Having some subheadings within the model methods section would also help the reader

We accept that the description of the model and its application was quite dense, and we have substantially edited this sub-section, in particular aiming to break up the text (lines 167-239). We do not feel that showing a model schematic would help, because it mostly shows soil processes, which although vital to the simulation of NPP are not strongly relevant to the subject of the present paper.

Line 205: Could you explain a bit more on how N fixation is modeled (is it only a function of Ndep, is it parameterised differently for your vegetation types, etc.)?

How N fixation is modelled is now described. It does not depend upon vegetation type (lines 175-177).

Line 238: Could you elaborate on the gamma distribution/log link use for a general audience?

We expanded the Statistics section in Methods (lines 241-257)

Line 242: Typo or missing word

Should be was not as, have amended (line 251).

Line 289 (re: Table 2/Fig 5): Why not also report R2?

Please see our response to the Editor's comment above. We have now reported pseudo- r^2 values (lines 306-307), as well as explaining what pseudo- r^2 is (line 252-257).

Lines 364-367: I am not sure I understand this entirely. Are you saying that because Tipping et al. 2017 found a temporal trend in SOC, that supports that your present results could be more temporal than what can be inferred from space-for-time? See also last comment below that is somewhat related.

This was poorly expressed, we have rewritten the last two paragraphs, also in response the Referee's last comment – see below.

Two broad comments:

1) Does the model provide any information on the relationship between below:aboveground allocation and Ndep? Based on your model description, I think that it does. A valuable contribution would be to test the hypothesis that increased Ndep would decrease below to aboveground fine-tissue ratios. This would provide an interesting discussion and additional insight into Ndep influences on NPP.

Our model is not so sophisticated, we treat above and below ground biomass as a combined entity, and therefore only calculate total NPP. We have clarified the model description to say this (lines 182-3).

2) I found that the last two paragraphs and the focus on SOC takes away from the study. I don't know how useful it is to talk about the link between NPP and long-term SOC when the authors do not have any information on decomposition. I understand that SOC is the big-picture implication, but I think the last paragraph should focus more on Ndep-NPP links than on the ancillary/inconclusive SOC topic. Authors can still keep the SOC discussion but just end with a last paragraph on the most

interesting aspect of their investigated Ndep-NPP links. Or, if authors really want to end on SOC, could they provide a graph from the model on the relationship between Ndep and SOC?

We are disappointed by this comment because to us the connection between NPP and SOC via N_{dep} is very important, especially in terms of the biogeochemistry of these ecosystems. When the Referee says "the authors do not have any information on decomposition" (s)he is not correct, because the N14CP model is built and parameterised on information about SOC turnover, including the use of radiocarbon for quantification. We do not think the SOC topic is ancillary or inconclusive – there is a strong link between the NPP-N_{dep} results reported here and the previous work on SOC.

We suppose that the last two paragraphs were not written convincingly enough, at least for this referee, and therefore we have rewritten them to bring out better the links between the present study and the previous SOC one (Tipping et al 2017). In particular we have tried to make it clear that the N14CP model deals with both NPP and SOC turnover. Turnover is now mentioned explicitly in the model description (line 174).

We do not think the introduction of a plot relating SOC to N_{dep} would be appropriate for the present paper, but we have now mentioned (lines 395-397) that we found such a direct relationship for SOC under broad-leaved trees.

Electronic Supplementary Material

Click here to access/download Electronic Supplementary Material Tipping et al_NPP_SM_REVISED.docx **Electronic Supplementary Material**

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