Nitrogen saturation and net ecosystem production


Magnani et al.1 found that net carbon (C) sequestration of temperate and boreal forests is clearly driven by nitrogen (N) deposition. From the positive relationship between average net ecosystem production (NEP) and wet N deposition, the authors further conclude that “no signs of N saturation were apparent” in the studied forests and that this is “casting doubts on the risk of widespread ecosystem nitrogen saturation”. Nitrogen additions can clearly alter net ecosystem production, but net ecosystem production cannot be used as an indicator of N saturation.

Nitrogen saturation implies a change in N cycling from a closed internal cycle to an open cycle1 where excess N is leached and/or emitted from the forest ecosystem. These changes in forest ecosystem functioning have been extensively documented3,4. Examples from literature lead us to suggest that some of the forest ecosystems discussed in the concerned article might be N saturated, irrespective of the increased net ecosystem production. Evidence for N saturation has been observed in forest ecosystems subject to N deposition levels similar to the relatively low deposition range reported in the concerned article1 (that is, less than 10 kg N ha\(^{-1}\) yr\(^{-1}\) wet deposition), including considerable nitrate loss (up to 10 kg N ha\(^{-1}\) yr\(^{-1}\)) through runoff or seepage water3,7 and elevated emissions of NO and N\(_2\)O.4 Because Magnani et al.1 did not measure any of these pathways of N loss, they cannot rule out N saturation in the studied forest ecosystems.

The demonstrated relationship between N deposition and C sequestration is an important finding, consistent with other literature4. In our opinion, however, Magnani et al.1 demonstrate an incorrect view on the phenomenon of N saturation and, in doing so, greatly ignore the effect of N deposition and saturation on soil acidification, groundwater and surface water quality, biodiversity, and ecosystem services other than C sequestration. Because the data presented by Magnani et al.1 do not allow an evaluation of the N saturation status of the studied forests, the expressed “doubts on the risk of widespread ecosystem nitrogen saturation” are not substantiated. At the moment, N saturation of forest ecosystems is probably not yet a widespread problem on a global scale, but it is surely a widespread problem in densely populated and more industrialized regions3,7. From the Europe-covering IFF (Indicators of Forest Ecosystem Functioning) and Level-II (UN-ECE/EC intensive monitoring plots) databases10,11, it can be deduced that more than 25% of the European forests included in these databases are N saturated (considering the nitrate seepage flux as indicator and 5 kg (357 mol) N ha\(^{-1}\) yr\(^{-1}\) as a threshold value of N saturation). Furthermore, from a large number of sites in the northeastern United States1, N saturation was indicated as a frequently occurring phenomenon. As substantial increases in global N emissions are predicted for the coming 50 yr\(^{-2}\), the potential risk of widespread N saturation of forest ecosystems in the long term cannot be denied.

An De Schrijver, Kris Verheyen, Jan Mertens, Jeroen Staelens, Karen Wuysts1 & Bart Muys2
1Laboratory of Forestry, Ghent University, Geraardsbergske Steenweg 267, 9090 Gontrode, Belgium.
2Division Forest, Nature and Landscape, Katholieke Universiteit Leuven, Celestijnenlaan 200E, 3001 Leuven, Belgium.

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Ecologically implausible carbon response?


Magnani et al.1 present a very strong correlation between mean lifetime net ecosystem production (NEP, defined as the net rate of carbon (C) accumulation in ecosystems3) and wet nitrogen (N) deposition. For their data in the range 4.9–9.8 kg N ha\(^{-1}\) yr\(^{-1}\), on which the correlation largely depends, the response is approximately 725 kg C per kg N in wet deposition. According to the authors, the maximum wet deposition level of 9.8 kg N ha\(^{-1}\) yr\(^{-1}\) is equivalent to a total deposition of 15 kg N ha\(^{-1}\) yr\(^{-1}\), implying a net sequestration near 470 kg C per kg N of total deposition. We question the ecological plausibility of the relationship and show, from a multi-factor analysis of European forest measurements, how interactions with site productivity and environment imply much smaller NEP response to N deposition.

The C response to N deposition is restricted by the C:N stoichiometry of the forest ecosystem compartments. The implied NEP response of 470 kg C per kg N would require that the fate of the deposited nitrogen was exclusively in stem wood, which is the only carbon sink with a C:N ratio of this magnitude. This is unreasonable because N-limited forest stands, as suggested1, invest primarily in roots3, with C:N ratios near 50–100. The impossibility of near-total

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storage of deposited N in stem wood follows also from the expected N leaching rates, varying between 10–50% of the N input in a range of 10–25 kg N ha$^{-1}$ yr$^{-1}$ (refs 4, 5), which is the likely range for total N deposition at the plots of Magnani et al.$^1$ where the high C:N response was found (that is, their European sites where wet N deposition was estimated at 4.9–9.8 kg N ha$^{-1}$ yr$^{-1}$). This total N deposition range follows from an application of the EMEP (European Monitoring and Evaluation Programme of the Long-range Transmission of Air Pollutants) model, used in the NITROEUROPE project, for the year 2000 to the sites of Magnani et al.$^1$. Using this model, the estimates of which are in close agreement with measured atmospheric N deposition at forest sites$^6$, we found that total N deposition at the sites of Magnani et al.$^1$ is 2–7 times greater than wet deposition. This analysis implies a relationship of approximately 175 kg C per kg N in the mentioned total N deposition range.

However, even this lower response is unlikely.$^{15}$N-labelled tracer experiments in temperate forests indicate that N retention hardly occurs in stem wood but mainly in the soil.$^7$ Considering the fate of N and the ranges in C:N ratios in forest ecosystem compartments, this implies a carbon response near 50 kg C per kg N in forest ecosystems.$^7$ Even though the above-ground C sequestration may be underestimated by Nadelhoffer et al.$^7$, owing to neglecting the effect of direct foliar uptake$^{7,10}$, this effect is likely to be small, as above-ground foliar uptake is generally less than 5 kg N ha$^{-1}$ yr$^{-1}$ (ref. 11), whereas below-ground uptake is generally more than 50 kg N ha$^{-1}$ yr$^{-1}$. Furthermore, similar results are found in long-term (15–30 yr) nitrogen-fertilizer trials at rates of nitrogen addition below 30 kg N ha$^{-1}$ yr$^{-1}$ (refs 12, 13) and in process-based model simulations$^{14}$. The reason for the extremely high influence of N deposition on NEP suggested by Magnani et al.$^1$ is probably due to the contribution of other factors—which co-vary with wet N deposition—to the derived relationship. The authors filtered out the effects of age and investigated the separate effect of temperature but they aggregated all tree species and site characteristics, such as site fertility and stand density, into one relationship.

We carried out a multi-factor analysis of measured forest growth data at nearly 400 intensively monitored forest plots in Europe, including Norway spruce, Scots pine, common beech and oak. The influence of nitrogen and acid deposition was considered by using values during the growth period (1993–2000), whereas the impacts of temperature, precipitation and drought were addressed by taking the deviation of these climatic parameters in the growth period (1993–2000) from the 30-yr mean. We simultaneously accounted for site factors influencing measured tree growth, including site productivity, stand age and stand density. We also applied a multi-factor analysis with measured basal-area increment of each individual tree as responding factor. The influencing factors used in this study and the results of the multivariate analyses at stand and individual tree level are shown in Table 1. The approach at tree level indicated a 1.2–1.5% increase in basal-area increment, depending on tree species, in response to 1 kg of N ha$^{-1}$ yr$^{-1}$. The approach at stand level indicated only a significant response of Norway spruce and Scots pine to N with roughly a 1–2% increase in volume growth in response to 1 kg of N ha$^{-1}$ yr$^{-1}$, depending on the C:N ratio of the plots. We recalculated these responses in terms of C sequestration by multiplying the mean measured volume growth at each stand with the estimated growth increase and the mean wood density of each tree species, assuming a C content of 50%. The results of our analyses at both tree and stand level indicate a response of trees between approximately 20–40 kg C per kg N. Additionally, results of long-term nitrogen addition experiments indicate soil responses of 10–30 kg C per kg N$^{12,13,15}$. Thus, the total NEP response would be around 30–70 kg C per kg N, which is much smaller than that estimated by Magnani et al.$^1$.

### METHODS

The multivariate regression at stand level was carried out by a backward stepwise method, where the model was reduced step-by-step by removing nonsignificant effects. The results of the hypothesis testing of the effects are based on partial $F$-tests.

The multivariate regression analysis at tree level was carried out by using tree size and tree competition variables on tree level and site factors and environmental factors on plot level, including plot as a random effect and applying the restricted maximum likelihood (REML) method for parameter estimation. Parameters in the model, which exhibited significant ($P < 0.05$) coefficients and behaved according to their known impacts, are included in Table 1b. We accounted for correlations, such as those between climatic parameters and N deposition (for example, we found a quadratic relation between N deposition and temperature; $R^2 = 0.62$).

**Wim de Vries**, Svein Solberg*, Matthias Dobbertin*, Hubert Sterba*, Daniel Laubhahn**, Gert Jan Reinders, Gert-Jan Nabuurs, Per Gunderson* & Mark A. Sutton*

1. Alterra, Wageningen University and Research Centre, PO Box 47, 6700 AA Wageningen, The Netherlands.
2. e-mail: wim.devries@wur.nl

### Table 1 | Multivariate regression results at stand level and individual tree level

<table>
<thead>
<tr>
<th>Tree species</th>
<th>Site productivity$^*$</th>
<th>Age†</th>
<th>SDI‡</th>
<th>N deposition§</th>
<th>Drought¶</th>
<th>Temperature change∥</th>
</tr>
</thead>
<tbody>
<tr>
<td>All plots</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Norway spruce</td>
<td>0.054</td>
<td>−0.005</td>
<td></td>
<td>0.020#</td>
<td>−0.0032</td>
<td>0.524</td>
</tr>
<tr>
<td>Scots pine</td>
<td>−</td>
<td>−0.017</td>
<td></td>
<td>0.010</td>
<td>−0.0032</td>
<td>−</td>
</tr>
</tbody>
</table>

Sensitive plots

<table>
<thead>
<tr>
<th>Tree species</th>
<th>Site productivity$^*$</th>
<th>Age†</th>
<th>SDI‡</th>
<th>N deposition§</th>
<th>Drought¶</th>
<th>Temperature change∥</th>
</tr>
</thead>
<tbody>
<tr>
<td>Norway spruce</td>
<td>0.039</td>
<td>−0.004</td>
<td>0.001</td>
<td>0.022</td>
<td>−0.002</td>
<td>0.32</td>
</tr>
<tr>
<td>Scots pine</td>
<td>−</td>
<td>−0.017</td>
<td></td>
<td>0.013</td>
<td>−0.002</td>
<td>−</td>
</tr>
</tbody>
</table>

Multivariate regression results indicating the relative change in stem volume growth per unit change in influencing factor (for example, a value of 0.013 for N deposition implies an increase in stem growth of 1.3% for each additional 1 kg ha$^{-1}$ yr$^{-1}$ of N deposition). Note that a dash (−) implies that the effect was insignificant ($P > 0.05$). Sensitive plots are plots with a C:N ratio above 25.

* Site productivity is a variable (m$^3$ ha$^{-1}$ yr$^{-1}$) derived from selected European site index curves, with input variables being age and top height.

† Stand age (yr).

‡ SDI, stand density index (number of trees per ha).

§ N deposition is total N deposition (unit change: kg ha$^{-1}$).

¶ Drought is a variable describing drought given as a relative value (unit change: %) to the normal (30-yr mean) drought stress at each site.

∥ The temperature difference during the growing period compared with the 30-yr average temperature (unit change: °C).

### Notes

1. Results from a linear regression; in the multivariate analysis the coefficient was just not significant ($P < 0.05$).

2. BAL is basal area of larger trees, which affects tree competition (m$^2$ ha$^{-1}$).

3. C:N$_{ initializing}$ is the C:N ratio of the mineral topsoil (0–30 cm).

4. Temperature is average yearly temperature during the investigation period 1993–2000 (°C).

5. BAL is basal area of larger trees, which affects tree competition (m$^2$ ha$^{-1}$).

6. C:N$_{ initializing}$ is the C:N ratio of the mineral topsoil (0–30 cm).

7. Temperature is average yearly temperature during the investigation period 1993–2000 (°C).

8. C:N$_{ initializing}$ is the C:N ratio of the mineral topsoil (0–30 cm).


10. C:N$_{ initializing}$ is the C:N ratio of the mineral topsoil (0–30 cm).

11. Temperature is average yearly temperature during the investigation period 1993–2000 (°C).

12. C:N$_{ initializing}$ is the C:N ratio of the mineral topsoil (0–30 cm).

13. Temperature is average yearly temperature during the investigation period 1993–2000 (°C).

14. C:N$_{ initializing}$ is the C:N ratio of the mineral topsoil (0–30 cm).

15. Temperature is average yearly temperature during the investigation period 1993–2000 (°C).
Magnani et al. reply


Nitrogen (N) deposition alters ecosystem function in several ways, with important effects on N leaching and water quality, as well as on interspecific competition and biodiversity. These changes have been attributed to ecosystem N saturation, defined as the alleviation of N limitations on rates of biological function1. After an initial fertilization effect, N saturation has also been suggested to reduce plant function and growth2, eventually leading to forest dieback. Although our observation of a substantial positive effect of N deposition on forest carbon (C) sequestration3 does not imply the absence of nitrate losses or other negative effects, as rightly stressed by De Schrijver et al.4, the sustained response observed demonstrates that the fear of a generalized forest decline in response to N fertilization could be overstated, at least within the rather broad N deposition range explored in our analysis. The nature of the observed response of forest C sequestration to N deposition, however, has been questioned outright by de Vries et al.5, who suggested that it could be an artefact resulting from the covariation between N deposition and other environmental variables. The arguments proposed against an overwhelming N effect, however, do not seem to stand up to close scrutiny.

We agree that ecosystem gross primary production (GPP) and plant growth are, to a large extent, controlled by local climate, drought and fertility (that is, N mineralization associated with soil organic matter decomposition), although fertility could be itself influenced by current and past N deposition6. However, the same environmental factors would modulate in parallel ecosystem respiration, and as a result do not seem to affect net ecosystem production (NEP), which is the difference between GPP and ecosystem respiration and is the subject of our analysis6. Both components of NEP seem to be also affected by N deposition, but in opposite directions: apart from the positive effects on plant growth considered by de Vries et al.5, respiration is known to be significantly reduced by N fertilization, as demonstrated by manipulation experiments7,8 as well as regional transect studies8. The combined effect at the ecosystem level is largely missed when focusing on tree growth alone.

The question remains of the magnitude of the observed response to N deposition. Assuming a linear relationship between NEP and N deposition, a slope of 445 ± 38 kg C per kg N of wet N deposition can be inferred from our entire data set (n = 20, rather than the subsample of 8 data points in the analysis by de Vries et al.5). If we assume, rather conservatively, that wet deposition constitutes 40–50% of total N deposition10, this would imply a NEP sensitivity to total N deposition of approximately 175–225 kg C per kg N, which is consistent with the stoichiometry of plant tissues and soil organic matter. Although it is true that fine roots account for a significant fraction of forest growth, it should be noted that one of the main effects of increased N availability is an increased allocation to woody tissues (with a high C:N ratio of up to 500:1) away from fine roots11. This mechanism could indeed represent an important component of the observed response to N deposition.

Far from implausible, a 200:1 sensitivity is nevertheless higher than suggested by long-term forest fertilization experiments12. Potential problems with N manipulation studies have already been discussed13. In particular, they overlook the role of canopy N uptake, which enables plants to absorb a relevant fraction of incoming N without any competition from soil microbes. Canopy N uptake amounts to up to 70% of N deposition, providing as much as one-third of tree N requirements14,15. The critical comparison of results from ecosystem manipulation and observational studies could be providing a rare, unforeseen insight into the key factors controlling C–N relations in forest ecosystems.

Federico Magnani1, Maurizio Mencuccini2, Marco Borghetti3, Frank Berninger4, Sylvain Delzon5, Achim Grelle6, Pertti Hari7, Paul G. Jarvis8, Pasi Kolar1, Andrew S. Kowalski9, Harry Lankreijer10, Paul E. Law11, Anders Lindroth12, Denis Loustau13, Giovanni Manca14, John B. Moncrieff15, Marco Tedeschi16, Riccardo Valentini17 & John Grace18

1Department of Fruit Tree and Woody Plant Science, University of Bologna, Bologna I-40127, Italy. e-mail: federico.magnani@unibo.it
2School of GeoSciences, University of Edinburgh, Edinburgh EH93JU, UK.
3Department of Crop Systems, Forestry and Environmental Sciences, University of Basilicata, Potenza I-85100, Italy.

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