## Forest fine-root production and nitrogen use under elevated CO<sub>2</sub>: contrasting responses in evergreen and deciduous trees explained by a common principle

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## Abstract

Despite the importance of nitrogen (N) limitation of forest carbon (C) sequestration at rising atmospheric CO<sub>2</sub> concentration, the mechanisms responsible are not well understood. To elucidate the interactive effects of elevated CO<sub>2</sub> (eCO<sub>2</sub>) and soil N availability on forest productivity and C allocation, we hypothesized that (1) trees maximize fitness by allocating N and C to maximize their net growth and (2) that N uptake is controlled by soil N availability and root exploration for soil N. We tested this model using data collected in Free-Air CO<sub>2</sub> Enrichment sites dominated by evergreen (Pinus taeda; Duke Forest) and deciduous [Liquidambar styraciflua; Oak Ridge National Laboratory (ORNL)] trees. The model explained 80–95% of variation in productivity and N-uptake data among eCO<sub>2</sub>, N fertilization and control treatments over 6 years. The model explains why fine-root production increased, and why N uptake increased despite reduced soil N availability under eCO<sub>2</sub> at ORNL and Duke. In agreement with observations at other sites, the model predicts that soil N availability reduced below a critical level diminishes all eCO<sub>2</sub> responses. At Duke, a negative feedback between reduced soil N availability and N uptake prevented progressive reduction in soil N availability at eCO<sub>2</sub>. At ORNL, soil N availability progressively decreased because it did not trigger reductions in N uptake; N uptake was maintained at ORNL through a large increase in the production of fast turnover fine roots. This implies that species with fast root turnover could be more prone to progressive N limitation of carbon sequestration in woody biomass than species with slow root turnover, such as evergreens. However, longer term data are necessary for a thorough evaluation of this hypothesis. The success of the model suggests that the principle of maximization of net growth to control growth and allocation could serve as a basis for simplification and generalization of larger scale forest and ecosystem models, for example by removing the need to specify parameters for relative foliage/stem/root allocation.

*Keywords:* allocation, elevated carbon dioxide, FACE experiments, fine-root longevity, forest growth model, optimization, plant theory, soil N availability, soil N uptake

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#### Introduction

The long-running forest Free-Air  $CO_2$  Enrichment (FACE) experiments have provided substantial

evidence of ecosystem-level responses to elevated CO<sub>2</sub> (eCO<sub>2</sub>), induced by the primary effects of CO<sub>2</sub> on leaf photosynthesis (Gifford, 2004). In the longer term, the response of forests to eCO<sub>2</sub> is a product of direct CO<sub>2</sub> effects and interactions with other resources that influence forest growth and carbon (C) flux. Soil nitrogen (N) availability is of particular importance for longer term responses because it limits forest production and C sequestration (Vitousek & Howarth, 1991), as well as their CO<sub>2</sub> responses (Oren et al., 2001; Reich et al., 2006a) in many temperate ecosystems. Soil N availability may also be subject to negative feedbacks associated with increased soil and plant N immobilization at eCO<sub>2</sub>, leading to progressive N limitation (Comins & McMurtrie, 1993; Luo et al., 2004). Soil N availability also modulates the effect of eCO<sub>2</sub> on forest growth through changes in C allocation, (i.e. shifting proportions of C invested in fine root, leaf and wood production). Increased C allocation to wood at eCO2 could increase the potential carbon sink in forest biomass due to the long mean residence time of wood compared with other tissues, whereas C allocation to root systems may enhance C transfer to soil organic matter pools. Clearly, understanding the interactive effects of eCO2 and soil N availability is essential for accurate projections of forest responses to rising atmospheric CO2. Furthermore, to enable such projections, the understanding needs to go beyond qualitative results towards mechanistic formulations that can be used in quantitative models.

In two mature forest FACE experiments located at Duke University and Oak Ridge National Laboratory (ORNL), net primary production (NPP), wood production, fine-root production (RP) and N uptake (Nup) all increased in response to eCO<sub>2</sub> (Hamilton et al., 2002; Norby et al., 2004; Norby & Iversen, 2006). The relative increase in NPP at eCO<sub>2</sub> was similar at the two sites. However, RP responded more and wood production less at ORNL than at Duke (DeLucia et al., 2005). At ORNL, annual RP was 91% higher at [CO<sub>2</sub>] of  $550 \,\mu\text{mol}\,\text{mol}^{-1}$  than at  $375 \,\mu\text{mol}\,\text{mol}^{-1}$ , whereas at Duke, the mean difference was only 19% (Norby et al., 2004; Finzi et al., 2006). It has been hypothesized that increased RP at eCO2 is a response to increasing N limitation (Norby et al., 2004). However, measured N uptake (Nup) increased at eCO<sub>2</sub> at both FACE sites, which is not consistent with the N limitation hypothesis and contrasts to predictions of reduced N uptake at eCO<sub>2</sub> by earlier biogeochemical cycling models (e.g. Rastetter et al., 1997; Medlyn et al., 2000). This has led to the suggestion that models must be reformulated to allow increased soil N uptake via increased C allocation to fine roots and their means of N acquisition, directly or via microbial activities (Schimel & Bennett, 2004; Finzi et al., 2007). However, although N uptake can be increased via root C allocation in different ways at the microscopic level, the ecosystem level mechanisms controlling the total root C allocation and associated N uptake at  $eCO_2$  are not yet well understood.

Here, we analyse the interaction between eCO<sub>2</sub> and soil N availability using a forest C-N model previously described (Franklin, 2007), extended by including N uptake. Whereas, the model by Franklin (2007) included soil effects only indirectly through measured fine-root/ leaf ratios; here, the plant is dynamically and directly connected to the soil through fine-root C allocation that responds to soil N availability and plant N demand. This new development of the model is essential for understanding the soil-plant feedback and its consequences for plant growth and soil N availability. In addition to standard modelling of production and respiration, our model uses a controlling principle of plant allocation. Based on evolutionary principles, we assume that maximization of net growth controls tree growth and allocation. This hypothesis successfully predicted responses of NPP and leaf area index (LAI) to eCO<sub>2</sub> at four forest FACE experiments including ORNL and Duke (Franklin, 2007). Here, our objective is to mechanistically explain how the interaction of eCO<sub>2</sub> and soil N availability controls N uptake and C allocation in forests. We test the model by explaining the differences in root allocation and generation of N limitation between the FACE experiments in an evergreen (Pinus taeda) forest at Duke and a deciduous forest (Liquidambar styraciflua) at ORNL, both with fully developed canopies and occurring in similar climates and latitudes.

## Materials and methods

## Model

The model described here simulates processes of radiation interception, canopy photosynthesis, autotrophic respiration, C allocation to leaves, fine roots and wood, litterfall and N uptake by roots. In this model, we integrate our previous plant model [i.e. plant production and N demand; Franklin (2007)] with newly developed models of soil N availability, soil N uptake, and the interaction between N uptake and demand. Equations for each process are kept simple so that we can analytically ascertain the plant's integrated response to  $eCO_2$  and soil N availability.

## Plant production and N demand

Canopy photosynthesis is calculated from the nonrectangular hyperbolic light response of leaf photosynthesis (Cannell & Thornley, 1998). Light-saturated photosynthetic rate ( $A_{max}$ ) is linearly related to leaf nitrogen per unit area ( $N_A$ ),  $A_{max} = a(N_A - N_{min})$ , where a is the slope of the relationship and  $N_{min}$  is its x-intercept. The initial slope of the photosynthetic light response is the quantum efficiency ( $\phi = 2.73 \,\mu g \, C \, J^{-1}$ , Wong *et al.*, 1979). Effects of [CO<sub>2</sub>] on photosynthesis are introduced as an increase in the leaf photosynthetic capacity per unit N (a) and an increase in  $\phi$  (Cannell & Thornley, 1998). Leaf photosynthesis is integrated over the canopy to evaluate gross primary production (GPP), assuming optimal  $N_A$  distribution and optimal LAI (L) as described in Franklin (2007). where  $I_a$  is absorbed photosynthetically active radiation (PAR), h increases the C costs per  $N_c$  and leads to a lower optimal  $N_c$  ( $N_c^*$ ) and lower GPP, as illustrated in Fig. 1a. Whereas, canopy C costs are tied linearly to  $N_c$ , root C costs and production (*RP*), determined by the optimality condition for *G*, has a maximum with respect to  $N_c$  as illustrated in Fig. 1a.

The choice of net plant growth G as optimization target rests on the assumption that maximizing G (size increase and reproduction) is a plausible strategy for maximizing fitness in the face of competition over the lifetime of a tree (Franklin, 2007). It is assumed that canopies that have reached steady state (i.e. peak LAI), such as at Duke and ORNL, thereafter always have

$$GPP = h \frac{\phi I_{a} + a(N_{c} - N_{\min}L) - \sqrt{[\phi I_{a} + a(N_{c} - N_{\min}L)]^{2} - 4\phi I_{a}a(N_{c} - N_{\min}L)\theta}}{2\theta}$$
(1)

is day length,  $N_c$  is canopy N content and  $\theta$  is the curvature of photosynthetic light response. Because GPP is co-limited by  $N_c$  (through photosynthetic capacity) and incoming PAR, GPP is a saturating function of  $N_c$  at constant PAR (mathematical derivations are given in Appendix).

NPP is calculated from GPP by subtracting maintenance respiration  $R_{\rm m}$  and growth respiration: NPP =  $y(\text{GPP}-R_m)$ , where growth respiration is a fixed fraction (1/y-1) of NPP and y = 0.72 (Choudhury, 2001).  $R_{\rm m}$  is expressed as a linear function of  $N_{\rm c}$  and the amount of N in other respiring tissues (Reich et al., 2006b). Total maintenance respiration is  $R_{\rm m} = rN_{\rm c}(1 + f_{\rm r}q_{\rm r} + f_{\rm s})$ , where r is respiration rate per unit N,  $f_r$  is the root to leaf N ratio  $(f_r = N_r/N_c)$ ,  $f_s$  is the sapwood to leaf N ratio  $(f_s = N_s/N_c)$ and  $q_r$  is a factor that accounts for the higher respiration rate per unit N in fine roots relative to foliage (Ryan et al., 1996). Litter production (T) of foliage (FP) and fine roots (RP) is determined from mean residence times and N:C ratios of leaves  $(t_{cr}, n_c)$  and fine roots  $(t_{rr}, n_r)$ , and is expressed as a linear function of tissue N contents,  $T = FP + RP = N_c[(1/n_ct_c) + f_r/(n_rt_r)]$ . As GPP is a saturating function of  $N_{c'}$  whereas both  $R_{m}$  and T are proportional to  $N_{c}$ , net plant growth G, defined as:

$$G = NPP - T = y(GPP - R_m) - T = yGPP - wN_c \quad (2)$$

has a maximum with respect to  $N_c$ , where  $N_c$  is optimal  $(N_c^*)$  [Appendix Eqn (A5)]. *G* includes woody tissue (stem, branches and coarse roots) increment and reproductive production.

The parameter *w* in Eqn (2) represents the carbon costs per  $N_{cr}$  and is a function of  $f_r$ :  $w = yr(1 + f_rq_r + f_s) + [1/(n_ct_c) + f_r/(n_rt_r)]$ . An increase in  $f_r$  increases *w* through an increased in fine-root N requirement per  $N_{cr}$  which optimal  $N_c$ . The optimal value of canopy N ( $N_c^*$ ) can be combined with equations above for GPP [Eqn (1)],  $R_m$ and T to determine values of GPP, NPP, G and LAI for optimized canopies (Franklin, 2007). Plant productivity is then controlled by changes in (1) photosynthetic parameters a and  $\phi$ , which affect GPP; (2) allocation parameters ( $f_r$  and  $f_s$ ), leaf and fine-root N : C ratios ( $n_c$ and  $n_r$ ) and residence times ( $t_c$  and  $t_r$ ), which affect  $R_m$ and T and (3) environmental parameters such as incident PAR ( $I_0$ ). Because T is assumed to be in steady state at fixed parameter values, productivity should be evaluated for a time period longer than  $t_c$  and  $t_r$  and not for shorter term fluctuations.

The N demands (*Nd*, Fig. 1b and d) associated with the carbon fluxes in Fig. 1a are determined by the N:C ratios and turnover times of the plant parts:

$$Nd = Gn_{\rm G} + \frac{N_{\rm c}(1 - q_{\rm rf})}{t_{\rm c}} + \frac{N_{\rm c}f_{\rm r}}{t_{\rm r}},\tag{3}$$

where  $n_G$  is the mean N:C ratio of tissues other than leaves and fine roots, (i.e. mainly wood), and  $q_{rf}$  is the fraction of N resorbed before leaf senescence. N demand of *G* and foliage [first and second term of Eqn (3)] are monotonically decreasing with  $f_r$  (through decreasing  $N_c^*$ ), whereas root N demand [last term of Eqn (3)] has a maximum, due to the maximum of root C allocation (Fig. 1a).

The N demand curves in Fig. 1b and d represent the rate of N uptake required to support annual growth predicted by the growth model. It includes  $CO_2$  effects on photosynthesis and canopy N, and consequent changes in allocation to foliage, wood and fine roots. However, it does not describe how N uptake (*Nup*) is related to soil N availability.

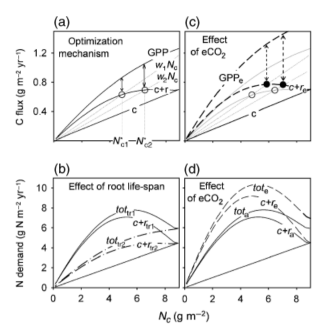


Fig. 1 Mechanisms of canopy optimization and allocation of C and N in response to fine-root: canopy N ratio  $(f_r)$ . (a) Canopy C gain [gross primary production (GPP)] and canopy C costs (c), i.e. respiration and litter production, are fixed functions of canopy N ( $N_c$ ). Root C costs are added to c, which gives the total C costs (c + r), shown here for two fixed values of  $f_r$  ( $w_1N_{cr}$ )  $w_2N_c$ ;  $w = \text{total C costs per } N_c$ ). Optimal  $N_c$  ( $N_c^*$ ) occurs where net growth (G, arrows) is maximized, i.e. where  $wN_c$  and GPP are parallel (circles). Varying  $f_r$  through a range of values shifts the slope of  $wN_c$  and, subject to the optimality condition, depicts the total C costs for canopy and roots for the *optimized* canopy (c + r). For clarity, effects on tissue N:C ratios (used in the modelling) are not included in this illustration. Values represent Oak Ridge National Laboratory (ORNL). (b) Solid lines show N demands corresponding to the C fluxes in (a) for the canopy (c), root + canopy (c + r) and total N demand (tot). The subscripts tr1 and tr2represent fine-root lifespan  $(t_r)$  roughly corresponding to ORNL (solid lines) and Duke (dadot lines) values, respectively.  $t_r$  for Duke is here three times longer, while root respiration per N is higher than at ORNL, keeping total root C costs per  $N_c$  the same for both sites. Other parameters are kept the same as in (a), i.e. representing ORNL. (c) Curves as in (a) for aCO<sub>2</sub> (thin curves, open circles) and curves and symbols for eCO2 (dashed curves, closed circles). The primary effect of eCO<sub>2</sub> is to raise GPP to GPP<sub>e</sub>. According to the mechanism described in (a), for the same values of  $w_1$  and  $w_2$  as in (a), the raised GPP raises both  $N_c^*$ (closed circles), G (dotted arrows), and root C costs ( $c + r_e$ ). (d) N demand for aCO<sub>2</sub> (subscript a, solid lines) and for eCO<sub>2</sub> (subscript e, dashed lines) corresponding to the eCO2 effect on the C fluxes shown in (c).

## Soil N uptake

We define soil N availability (*Nav*) as the maximum rate of N uptake per root carbon ( $C_r$ ) when  $C_r$  is small;  $Nav = Nup/C_r$  when  $C_r \rightarrow 0$ . Although other, soil-

centric perspectives on *Nav* are more commonly used, our plant-centric definition of *Nav* is more relevant for our plant growth modelling. For simplicity, N uptake is represented by an hyperbolic function of  $C_r$ , where the mechanism for that relationship is not specified, though it may relate to exploration of the soil volume by roots (McClain *et al.*, 2003) including increased rooting depth, solute transport to roots as a function of inter-root distance (Yanai, 1994), increased competitiveness of tree roots for soil N (Schimel & Bennett, 2004) or N uptake via C allocation to mycorrhizal fungi or exudates, all which may scale with  $C_r$  (Finzi *et al.*, 2007):

$$Nup = Nav \frac{C_{\rm r}}{(C_{\rm r}/\lambda) + 1} \tag{4}$$

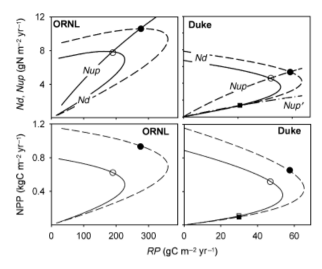
 $\lambda$  represents  $C_r$  required to achieve half-maximum N uptake (cf. McMurtrie, 1985). Saturation of N uptake may be related to limited soil volume, or decreased root-uptake efficiency as the density of roots increases, or spatial variation in N availability, where N rich parts of the soil volume are explored first. *Nup* has a theoretical maximum = *Nav* $\lambda$  (when  $C_r \rightarrow \infty$ ), whose value is of no significance as our model is parameterized and used within a range of much lower *Nup* values.

#### Balancing N demand and uptake

Rates of N uptake (*Nup*) and N demand (*Nd*) are shown in Fig. 2 as functions of root production (*RP*). The operating point of N use is the intersection Nup = Nd, which is dynamically stable with respect to changes in soil N availability (*Nav*) or *Nd* (see Appendix).

Elevated CO<sub>2</sub> steepens the relationship between GPP and  $N_{cr}$  which increases the optimal value of  $N_{c}$  for a given  $f_{r}$  as well as the N demand, as illustrated in Fig. 1c and 1d. The effect is to raise the Nd curve illustrated in Fig. 2, and to shift it to the right. This means that for a given Nav (which controls the Nup curve), RP, Nd and NPP all increase at eCO<sub>2</sub>, although the effect is small at extremely low Nav (Nup', Fig. 2). The largest increase in RP at eCO<sub>2</sub> occurs at intermediate Nav, where N use (intersection of *Nup* and *Nd* curves) is near the peak value of RP. With decreasing Nav, plant responses to eCO<sub>2</sub> decline dramatically after the value of RP passes its peak on the Nd curve (Fig. 2). This change in  $CO_2$ response can be explained through the effect of CO<sub>2</sub> on root C allocation shown in Fig. 1c (vertical distance between c + r and c curves). To obtain the same root C allocation for eCO<sub>2</sub> as for aCO<sub>2</sub> at high  $N_c^*$  (high Nav, low  $f_r$ ) requires little change in  $f_r$ , yielding a higher  $N_c^*$  at  $eCO_2$  than at  $aCO_2$ , which enhances the effect of  $eCO_2$ on production. To obtain the same root C allocation at low  $N_{c'}^*$  fr must be larger for eCO<sub>2</sub> than for aCO<sub>2</sub>, yielding a lower  $N_c^*$  at eCO<sub>2</sub> than at aCO<sub>2</sub>. This lowering

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**Fig. 2** Balancing N uptake (*Nup*) and demand (*Nd*). Upper panels: *Nup* and *Nd* vs. root production (*RP*) for aCO<sub>2</sub> (solid lines, open symbols) and eCO<sub>2</sub> (dashed lines, closed symbols). The symbols indicate the operating point where *Nup* = *Nd*. The slope of *Nup* is controlled by N availability (*Nav*). For Duke, *Nup'* (dadot line, squares) represents a hypothetical extremely low *Nav*. Lower panels: net primary production (NPP) corresponding to the *Nd* of the upper panels. Each curve, except *Nup*, depicts an increase in root N/canopy N ratio (*f*<sub>r</sub>) and associated reduction in tissue N:C ratios, starting from *f*<sub>r</sub> ≈ 0 at the upper left endpoints.

of  $N_c^*$  suppresses the effect of eCO<sub>2</sub> on production and, therefore, on *Nd*. This closes the gap between the *Nd* curves of aCO<sub>2</sub> and eCO<sub>2</sub> at low *Nav* (i.e. for N use on the lower arm of the *Nd* curve in Fig. 2). Hereafter, values of *Nav* that result in an N use on the upper and lower arm of the *Nd* curve relative to that at peak *RP* are referred to as higher and low *Nav*, respectively.

#### Experiments and measurements

The FACE experiments (Table 1) and datasets are described in detail in Finzi et al. (2007). For ORNL FACE, annual data for the years 1998-2003 were used directly, while for Duke, annual data were aggregated to represent averages for the years 1998-1999 and 2002-2004. The time periods selected were based on the availability of fine-root data. The aggregation of annual data at Duke was done because of the relatively long lifespan of roots and leaves in Duke, to better match data with the model assumption that foliage and fine-root biomass are in equilibrium with the optimal state of the plant (see 'Model'). The data include both overstory and understory trees, but only at Duke does the understory contribute significantly to forest production (13% of total aboveground NPP). The ORNL fertilization experiment was performed on a previous FACE plot

Table 1 Characteristics of the two FACE experiment	nts
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Name	Duke	ORNL
Location	Durham, NC, USA	Oak Ridge, TN, USA
Latitude, longitude	35°58'N, 79°05'W	05A 35°54'N, 84°20'W
Annual precipitation (mm)	1140	1390
Annual temperature (°C)	15.5	14.2
Growing season* (days)	200	190
Soil texture	Clay loam	Silty clay loam
Total soil N ( $g kg^{-1}$ )	0.79	1.12
Overstory vegetation	Pinus taeda L.	Liquidambar styraciflua L.
Peak leaf area index <sup>†</sup> $(m^2 m^{-2})$	3.4	5.5
Day length ( $h$ ; s day <sup>-1</sup> )	50 400‡§	43 200‡¶
Incident PAR ( $I_0$ ; J s <sup>-1</sup> m <sup>-2</sup> )	184‡§	211‡¶

Data taken from Norby et al. (2005).

\*For deciduous stands, the growing season is the duration that trees have leaves; for the evergreen system, it is the period of active stem growth.

<sup>†</sup>Values of leaf area index are expressed as projected leaf area per ground area.

‡Growing season average values.

SNorby et al. (2003).

¶Delucia et al. (2002).

FACE, Free-Air CO<sub>2</sub> Enrichment; ORNL, Oak Ridge National Laboratory; PAR, photosynthetically active radiation.

without  $CO_2$  treatment, where 200 kg ha<sup>-1</sup> of N as urea was added and data collected in 2004 and 2005 (Iversen & Norby, 2008). Measurement of net N mineralization at Duke is described in Finzi *et al.* (2006). Photosynthesis measurements are described for Duke in Crous & Ellsworth (2004) and Crous *et al.* (2008) and for ORNL in Sholtis *et al.* (2004).

## Model parameterization and input data

The model was parameterized for six FACE plots at Duke and five FACE plots and two fertilization treatments (six plots pooled per treatment) at ORNL. Constant site mean values were determined for leaf photosynthetic capacity per unit (a = 73,Ν  $26 \,\mu g \,C \,g^{-1} \,N \,s^{-1}$ ), respiration rate per N (r = 0.187,  $0.147 \,\mathrm{g}\,\mathrm{C}\,\mathrm{g}^{-1}\mathrm{N}\,\mathrm{d}^{-1}$ ), sapwood *N*/canopy *N* ( $f_{\rm s} = 1.5$ , 0.5), fine-root lifespan ( $t_r = 0.53$ , 3 years) and fine-root respiration/foliage respiration per N ( $q_r = 1, 3.57$ ), where the values represent ORNL and Duke, respectively. These parameters were estimated by fitting all the modelled and measured values of RP, G, N<sub>c</sub> and Nd simultaneously (inverse modelling). This model parameterization approach focuses the subsequent model evaluation on prediction of the variation among plots and treatments, in particular the  $CO_2$  effects, whereas, biases due to errors in parameter mean values are minimized (Franklin, 2007).

Soil N availability. Soil N availability (Nav) was determined for each plot and year using Eqn (4) and measured values of total N uptake and root mass carbon  $(C_r)$ . In estimating the effects of plot and year on soil N availability (Nav), we assumed a constant halfsaturation  $C_r$  ( $\lambda$ ). This assumption is based on the findings that soil volume and root physiology did not vary significantly among CO<sub>2</sub> treatments, plots and years (Norby et al., 2004; Pritchard et al., 2001), whereas soil N and N mineralization varied. Furthermore,  $\lambda$  was constrained by the assumption that RP must never be higher than the RP that maximizes net N uptake (=Nup-N in root litter production), which sets a lower limit for  $\lambda$ . A further increase in RP would reduce net N uptake and therefore be suboptimal. Because a first fit of  $\lambda$  and *Nav* to the complete dataset yielded a  $\lambda$  that was lower than the limit described above,  $\lambda$  equal to the limit (460 and  $59.2 \,\mathrm{g}\,\mathrm{C}\,\mathrm{m}^{-2}$  for ORNL and Duke, respectively) was chosen.  $\lambda$  values higher than this would also be possible but would increase the variation in Nav among plots and years. Time and treatment effects on Nav were compared using ANCOVA.

*Photosynthetic effects of eCO\_2*. The effects of  $CO_2$  on plant growth were introduced through the observed relative changes in the leaf photosynthetic capacity per unit N (effect on a = a/site mean of a), and a proportional change in  $\phi$  (effect on  $\phi$  = effect on a/3). Effects on afor the different rings and years were estimated using measured  $A_{max}$  and  $N_A$  data using the relationship  $A_{\text{max}} = a(N_{\text{A}} - N_{\text{min}})$ , where  $N_{\text{min}}$  is constant across all treatments. The estimated effects on a for ORNL were 1.29, 1.29, 0.79, 0.79, 1.00, 0.84, 0.84 for the plots 1-7, respectively, where plots 1 and 2 are eCO<sub>2</sub> in the FACE, plots 3–5 are aCO<sub>2</sub> in the FACE experiment, and plots 6 and 7 represent means for six replicate plots each in a fertilization trial (at aCO<sub>2</sub>). For Duke, there was no significant difference among plots of the same CO<sub>2</sub> treatment or among years, so only two values for the effect on *a* were used, 0.85 and 1.15 for  $aCO_2$  and  $eCO_2$ , respectively.

Other treatment effects. Changes in tissue N:C ratios (*n*) for estimates in Figs 3–6 were taken from measurements. For the illustration of theoretical response curves (Fig. 2), changes in *n* in response to *Nav* were modelled using an empirically determined relation between  $f_r$  and *n* that passes through the observed values.

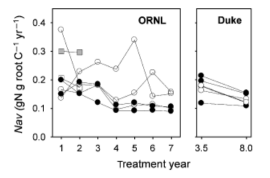
 $n_x = n_{x-obs}(1 - (f_r/p_1)^{p_2})/(1 - (f_{r-obs}/p_1)^{p_2})$ , where *x* stands for leaf, fine roots or wood and *obs* are observed values.  $p_1$ ,  $p_2$  are parameters, with values 7, 0.35 and 7, 0.7 for ORNL and Duke, respectively. For ORNL, *n* was related to  $f_r$  for all the tissues (i.e. leaves, fine roots and wood). For Duke, *n* varied only for foliage, while *n* of fine roots and wood were constant. For the fertilized stands at ORNL,  $f_s$  = sapwood N/canopy N was estimated to have decreased by 40% compared with control stands due to increased LAI and lagging increase in sapwood area, due to the short duration of the fertilization exposure. For longer time responses, sapwood area/LAI is not changed by fertilization (Hubbard *et al.*, 2004; Samuelson & Stokes, 2006).

## Results

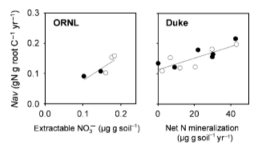
## Model validity

The model was evaluated using input data on soil N availability (*Nav*) derived for each plot and year, tissue N:C ratios (*n*) taken from measurements, and photosynthetic effects of  $eCO_2$  on *a* and  $\phi$  derived from independent measurements – see 'Materials and methods'. Of these parameters, the model was most sensitive to *a* and *Nav*, and was relatively insensitive to tissue N:C ratios.

We did not use direct measures of soil N availability for Nav because of the paucity of data and because, our definition of Nav is not equivalent to any available soil measure. Nav does not correspond to common views of N availability as a pool or flux of free N in the soil, or as N taken up by the plants. Instead, Nav represents the maximum potential N flux that can be extracted from the soil per root mass by a single root in the soil [Eqn (4)], which presents a rhizo-centric view of plant-N uptake feedback (Phillips, 2007). The exact mechanisms, which may include absorption of mineralized N, stimulation of N mineralization via C exudation and competition with microbes for N (Schimel & Bennett, 2004), are of subordinate importance for this ecosystem level analysis (see also 'Discussion'). However, modelled trends in Nav (Fig. 3) correlate with observed N mineralization over time at Duke (Fig. 4), relative plot differences in extractable NO<sub>3</sub><sup>-</sup> at ORNL (Fig. 4) and a reduction in gross N mineralization at eCO<sub>2</sub> observed for 2 years at ORNL (Zak et al., 2003). These relationships indicate that Nav is not only a useful representation of a virtually un-measurable entity but also is linked to independently measured soil properties. Nav declined over time and was significantly (P = 0.0114)lower at eCO<sub>2</sub> than at aCO<sub>2</sub> and at ORNL but not at Duke (Fig. 3). Nav declined faster at eCO<sub>2</sub> than at aCO<sub>2</sub> at ORNL, although the difference was not significant



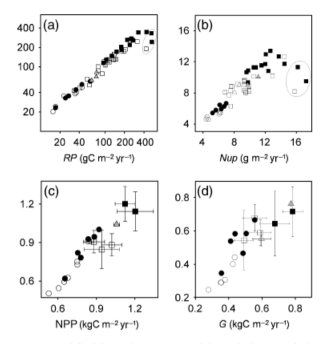
**Fig. 3** Estimated soil nitrogen availability (*Nav*). Symbols: Open –  $aCO_2$ , closed –  $eCO_2$ , grey – fertilized, circles – Free-air  $CO_2$  enrichment (FACE) plots, squares – fertilization experiment at Oak Ridge National Laboratory (ORNL). For ORNL, points show annual values for the period 1998–2004 and 2004–2005 for the FACE and fertilization plots, respectively. For Duke, the points represent mean values for the periods 1998–1999 and 2002–2004. Values calculated from measured N uptake and fineroot C using Eqn (4).



**Fig. 4** Modelled soil N availability (*Nav*) vs. measured extractable  $NO_3^-$  in Oak Ridge National Laboratory (ORNL) and Net N mineralization in Duke. Symbols as in Fig. 3.

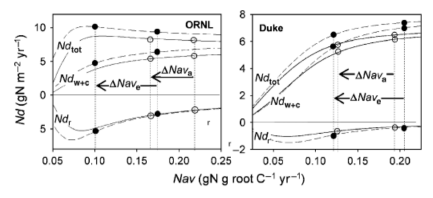
(Figs 3 and 4). To test the model's validity, modelled results were compared with measured data. Modelled and measured root production (*RP*), N uptake, NPP and net growth (*G*) are shown in Fig. 5. Results are shown for each year at ORNL and for periods of 2–3 years at Duke.

There was strong agreement between model and measurement data at both sites. Between 80% and 95% of variation in measurements is described by the model (Fig. 5), indicating that the model was well suited to describe linkages between photosynthetic processes and soil N availability, which were parameterized from independent datasets (see 'Methods'). The agreement for *RP* may seem predetermined as root C was used in the estimation of *Nav*. However, the fitting of *Nav* constrains only one of the curves determining *RP*, [i.e. N uptake (*Nup*) but not the demand (*Nd*, Fig. 2)], which depends on the other independent parameters of which *a* is the most influential. For ORNL,



**Fig. 5** Modelled (*y* axis) vs. measured (*x* axis) data. Symbols: circles – Duke Free-air CO<sub>2</sub> enrichment (FACE), Squares – Oak Ridge National Laboratory (ORNL) FACE, triangles – ORNL fertilization trial, Open symbols – aCO<sub>2</sub>, black – eCO<sub>2</sub>, grey – N fertilized. Encircled points are outliers. (a) Annual fine-root production (*RP*) (logarithmic scale),  $r^2 = 0.95$ . (b) Nitrogen uptake (*Nup*),  $r^2 = 0.85$ . (c) net primary production (NPP). Results for three FACE aCO<sub>2</sub> and two eCO<sub>2</sub> plots, and two fertilized and two unfertilized treatments at ORNL are represented by means and bars. Bars represent standard deviation among years,  $r^2 = 0.92$ . (d) Net growth (G). Symbols and bars as in (c),  $r^2 = 0.80$ . Each point represents, for Duke – one plot and one period (mean over 2–3-year period), for ORNL FACE *RP* and *Nup* – 1 year and one plot, for ORNL FACE NPP and *G* – mean values over 7 years, and for ORNL fertilization 1 year and a 6 plot average.

there are three outliers (Fig. 5a and b) for which RP and Nup are underestimated because measured RP values are much higher than the maximum possible model RP (cf. Fig. 2). The outliers correspond to observations of unusually deep roots and may be influenced by rapid dynamic changes in root mass, not compatible with the equilibrium assumption in the model. However, given the small number of outliers and the difficulty of measuring RP, measurement error can be a contributing factor, which may also have contributed to the slight divergence between modelled and measured NPP at low NPP (Fig. 5c), where RP is a large fraction of NPP. In order to focus on the interaction of eCO<sub>2</sub> effects and soil N availability, the model does not include effects of annual climate variation on productivity, especially the effects of droughts on NPP and its allocation (McCarthy



**Fig. 6** Changes in N use (*Nd*) and soil N availability (*Nav*) over time modelled for an aCO<sub>2</sub> (open circles, solid lines) plot and an eCO<sub>2</sub> plot (black symbols, dashed lines) at each site.  $\Delta Nav$  (arrows) show the reduction in *Nav* between treatment years 3.5–8 and 1–5 for Duke and Oak Ridge National Laboratory (ORNL), respectively (cf. 1 Fig. 3).  $Nd_{tot}$ ,  $Nd_{w+c}$  and  $Nd_r$  are total, wood + foliage and fine-root N use, respectively.

*et al.*, 2006). It is, therefore, not surprising that the model does not capture the inter-annual variation in NPP and *G* at ORNL, and that modelled and measured *G* diverge for some periods under  $eCO_2$  at Duke (Fig. 5d). In summary, despite the highly aggregated and simplified representation of processes in the model, it appears to provide a working mechanistic explanation of the interaction of forest production and N uptake in the studied forests.

## Discussion

#### Potential uncertainties

The simplicity of the model makes it clear that the qualitative conclusions are sensitive to only a few key model assumptions. These assumptions are (1) photosynthesis and maintenance respiration are linked to tissue N, (2) C and N allocation is regulated to maximize *G* (or some property closely related to *G*) and (3) *Nup* increases with root C and soil N availability (*Nav*). The first assumption, although not valid for all leaves (Maier *et al.*, 2008), has strong support (e.g. Reich & Ellsworth, 1998; Reich *et al.*, 2006b; Crous *et al.*, 2008). The second assumption has been evaluated in Franklin (2007) and the last assumption corresponds to the observations used here and reported previously (Norby *et al.*, 2004).

The N uptake model based on root C and *Nav* is by necessity a simplified aggregation of many processes, such as allocation to mycorrhiza and exudates that are involved in plant N uptake. For example, from the relationship between *Nav* and N mineralization at Duke (Fig. 4), it is clear that N mineralization alone does not explain total soil N availability to the plant (*Nav*). Although separating the processes involved in N uptake would be desirable, such a model would not be

testable based on currently available data. This means that we are implicitly assuming that the total effect of all processes on N uptake is proportional to fine-root C. Globally, this assumption is supported by observations that arbuscular mycorrhiza is correlated to fine-root biomass rather than soil organic matter (Treseder & Cross, 2006). Our model is consistent with observations that mycorrhizal root colonization (Parrent & Vilgalys, 2007; Garcia et al., 2008) and soil microbial activity (Sinsabaugh et al., 2003) are not strongly affected by the CO<sub>2</sub> treatments in our sites, in contrast to increases observed in fine roots and N uptake. Another indication that fine roots exert strong influence on soil activity, which may be linked to N uptake, is that the spatial distribution of soil respiration reflected the fine root distribution at Duke (Andrews et al., 1999).

In a C budget perspective, enhanced allocation to mycorrhiza and root exudation per fine-root C at  $eCO_2$  (Norby *et al.*, 1987) would constitute a part of NPP not accounted for in the observations (Schäfer *et al.*, 2003). In this model, an underestimation of measured below ground C export would primarily result in a model parameterization overestimating the root respiration per N ( $q_{rr}$  see 'Model'), whereas other parameters are more constrained by aboveground data. This potential error may have consequences for the soil C budget, but it has no impact on plant growth and N uptake within our model framework.

A potentially more important uncertainty for this study pertains to the hypothesis that fine-root mean residence time ( $t_r$ ) decreases with soil N availability (Nadelhoffer, 2000). Although shifts in  $t_r$  with eCO<sub>2</sub> have not been found in the experiments evaluated here (Pritchard *et al.*, 2001; Norby *et al.*, 2004), the consequence would be slight shifts of the N demand (*Nd*) curves due to increased *RP* at high *Nd* and decreased *RP* at low *Nd* (Fig. 2). Because of the relative flatness of

the upper arm of the *Nd* curves, the effect would be largest at low *Nd*. But more importantly, even a doubling of  $t_r$  at very low *Nd* would not qualitatively change the conclusions for the effects of CO<sub>2</sub> and *Nav*.

## Soil N availability effects on the responses to eCO<sub>2</sub>

The model implies that increased fine-root production (RP) at eCO<sub>2</sub> is an unavoidable consequence of the combination of increased N demand (Nd) and constant N uptake (Nup) as functions of RP (Fig. 2). The increase in *RP* at eCO<sub>2</sub> is greatest at a value of Nav ( $Nav_0$ ) that, in Fig. 2, makes Nup cross Nd near the maximum RP. If Nav exceeds Nav<sub>0</sub>, then RP is a decreasing function of *Nav*, whereas if *Nav* is less than  $Nav_0$  (lower than any observation at Duke or ORNL), RP is an increasing function of Nav. Although none of our data falls on the lower arm of the Nd curve ( $Nav < Nav_0$ ), the general shape of the curve is plausible, because if Nd approaches zero, so must RP and NPP. The different responses at higher vs. low Nav may be a reason for the variable, both positive and negative, root production responses to soil N availability observed (Nadelhoffer, 2000). In the case of progressive reduction in Nav, our model predicts that aboveground eCO<sub>2</sub> responses will decrease, whereas RP will increase until Nav drops below  $Nav_{0}$ , where all CO<sub>2</sub> responses will decline dramatically. This decline is caused by reduced allocation of N to canopy photosynthesis (reduced  $N_c^*$ ), which reduces the impact of the leaf photosynthetic stimulation at eCO<sub>2</sub> (see 'Model'). This type of declining CO<sub>2</sub> response has been observed in a nutrient-poor woodland (Day et al., 2006; Hungate et al., 2006) and provides an explanation for the observations that CO<sub>2</sub> responses of both roots and aboveground parts are declining at low N availability at some sites (Pregitzer et al., 2000; de Graaff et al., 2006).

The model implies that, unless Nav is very low, N use efficiency (NUE = NPP/N use) increases with  $CO_2$  due to increased production per canopy N (Fig. 1c) and with *Nav* due to increased relative allocation to G (mainly wood), because wood has a lower N:C ratio than litter (Fig. 2). In accordance with this, the strongest NUE increase at eCO2 among forest FACE sites was observed at the site where N was least limiting (PopFACE; Calfapietra et al., 2007; Finzi et al., 2007), whereas the absence of an increase in NUE at eCO2 at ORNL is explained by the concurrent decrease in Nav (Fig. 4). Our prediction of increasing NUE at higher soil N availability is in line with recent findings regarding resource use efficiency (Binkley et al., 2004; Stape et al., 2004; Franklin, 2007). However, our results contrast to the earlier methods of NUE estimation based on aboveground litter production only (Pastor & Bridgham, 1999), which, therefore, do not capture the allocation shifts between fine roots and wood that strongly contribute to the relationships predicted here. Generalizing our model implies that increased resource use efficiency follows when increased availability of a limiting resource (here N), given sufficient time for plant acclimation, reduces the acquisition and maintenance C costs (here root C allocation, cf. Fig. 1) per unit resource.

# Differences in the CO<sub>2</sub> response between an evergreen forest (Duke) and a deciduous forest (ORNL)

On average, the  $CO_2$  response of fine-root production (*RP*) is smaller and the response of wood production is larger at Duke compared with ORNL (DeLucia *et al.*, 2005). This allocation difference is of significance as wood production results in longer term biomass carbon sequestration than fine-root production. In our framework, the allocation difference between the sites is mainly related to two factors: the different effects of  $CO_2$  on *Nav* in the two sites and the difference in fine-root lifespan ( $t_r$ ).

The model predicts that reduced Nav at eCO<sub>2</sub> contributed to increased RP at ORNL, while for Duke, there is no consistent effect of eCO<sub>2</sub> on Nav. At both sites and under both aCO<sub>2</sub> and eCO<sub>2</sub>, Nav as well as observed N mineralization in Duke and extractable  $NO_3^-$  in ORNL declined over time (Figs 3 and 4). However, a reduction in Nav gives rise to different feedbacks on N use (Nd) in the two sites, caused by the strongly differing  $t_r$  between the species [0.53 year at ORNL and 3 years at Duke, roughly matching measured values (Norby et al., 2004; Strand *et al.*, 2008)]. The large difference in  $t_{\rm rr}$ which also has been confirmed using an isotope tracer (Matamala & Schlesinger, 2000), means that most of the fine-root C (75%) at ORNL is allocated to litter production with associated N use, whereas at Duke, most of the fine-root C allocation (82%) is ultimately respired with no associated N use (Fig. 1b). As illustrated in Fig. 6a (and explained by the shape of the upper arm of *Nd* in Fig. 2a), despite decreasing Nav, the short  $t_r$  and associated high production and N use of fine roots maintain a high total Nd at ORNL. This high Nd at declining Nav should contribute to continuing reduction in Nav, unless root litter N is efficiently remineralized. However, observations of increased total soil N at eCO2 (Johnson et al., 2004), despite increased N uptake and unchanged leaf litterfall, indicate that additional inputs of root litter N to the soil are not quickly recycled. In accordance with predicted consequences of reduced Nav, extractable  $NO_3^-$  and observed wood/root production ratio (G/RP) declined over time at ORNL, where the decline was stronger at eCO<sub>2</sub> than at aCO<sub>2</sub>. At Duke, due to low root N use, reductions in Nav generate reductions in Nd (Figs 6b and 2b), providing a negative feedback on further reduction in Nav. This feedback may have prevented or delayed a similar decline in the relative allocation to wood at eCO<sub>2</sub> as at ORNL. Generalizing this result implies that species with slow root turnover and low fine-root N use, such as evergreens, are less likely to cause progressive reduction in Nav. These species may, therefore, be less prone to progressive N limitation of wood production and associated biomass carbon sequestration at eCO<sub>2</sub> than species with fast root turnover, such as many deciduous trees (Withington et al., 2006). However, for a thorough evaluation of this hypothesis, longer term studies are recommended, as well as studies that quantify parallel below ground C inputs and their effect on N uptake.

## Conclusions

We modelled effects of eCO<sub>2</sub> and soil N availability based on essentially only two independent input variables, mean photosynthetic capacity per leaf N (a) and soil N availability (Nav). Despite this simplicity, model predictions were consistent with measurements of N uptake, production and allocation at ambient and eCO<sub>2</sub> in two FACE experiments and a fertilization experiment. In addition, the model provides an explanation for declining CO<sub>2</sub> responses observed at other more strongly nutrient limited sites. We attribute the models' ability to integrate responses to eCO<sub>2</sub> and soil N availability over a range of conditions to the applied optimization perspective. In addition to commonly used statistical analysis of what happened and process modelling of *how* things happen, here we hypothesized *why* trees behave as they do. The hypothesis, that trees maximize their net growth and reproduction (G), integrates all individual plant processes and responses to eCO<sub>2</sub> and ensures that their joint behaviour is optimal. In this integrated framework, empirical results may be placed into a bigger picture, not limited by the range of experimental growing conditions. For example, the negative effect of aboveground productivity on root C allocation in the FACE sites under current conditions (Palmroth et al., 2006) is predicted to switch to a positive effect at lower soil N availabilities (Nav) (Fig. 2). Furthermore, our model suggests that dominance of LAI effects on aboveground eCO2 responses (McCarthy et al., 2006) is mainly limited to expanding canopies, low Nav and low LAI [see also Franklin (2007)], while for higher Nav, nitrogen use efficiency responses are larger.

The model suggests that the sensitivity of biomass C sequestration to soil N availability (*Nav*) differs among species differing in root lifespan ( $t_r$ ). Because a short  $t_r$  leads to high N uptake despite declining *Nav*, the model

suggests that species with short  $t_r$  (e.g. at ORNL) are more prone to progressive reduction in *Nav* and therefore N limitation of carbon sequestration in woody biomass, than long  $t_r$  species, such as evergreen trees (e.g. at Duke). In order to evaluate the importance of this result for long-term forest C balance, the principles for integrating responses to different factors presented here could be incorporated into forests ecosystem models that include explicit modelling of the soil. Ultimately, our model could provide a basis for the improvement of large-scale forest and vegetation dynamics models by replacing fixed but uncertain parameters (such as fine root/leaf production) with dynamic optimization.

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## Appendix

## Derivation of canopy photosynthesis, GPP

Canopy photosynthesis (GPP) is calculated by integration of leaf photosynthesis (GPP<sub>leaf</sub>) over canopy depth (*z*).  $A_{\text{max}}$  is a function of N per leaf area ( $N_A$ ),  $A_{\text{max}} = a(N_A - N_{\text{min}})$ , where  $N_A$  is derived as a function of *z* assuming an optimal nitrogen distribution [Eqn (A2)] as described in (Franklin & Ågren, 2002). PAR absorbed by a leaf is related to canopy depth according to  $I(z) = I_0 k e^{-kz}$ , where *k* is the light extinction coefficient and  $I_0$  is incident PAR above the canopy. The integral in Eqn (A1) is easily calculated by first separating out the *z* dependence through the factor  $ke^{-kz}$ , which occurs in all terms of the integrand.

$$GPP = \int_{0}^{L} GPP_{\text{leaf}} dz$$
$$= \int_{0}^{L} \frac{\phi I(z) + A_{\max}(z) - \sqrt{\left[\phi I(z) + A_{\max}(z)\right]^{2} - 4\phi I(z)A_{\max}(z)\theta}}{2\theta} dz$$
(A1)

$$N_{\rm A}(z) = \frac{(N_{\rm c} - N_{\rm min}L)k {\rm e}^{-kz}}{1 - {\rm e}^{-kL}} + N_{\rm min}. \tag{A2}$$

#### Derivation of optimal $N_{\rm c}$

Optimal  $N_c$  is derived through maximization of *G* with respect to  $N_c$ . To simplify calculations, the substitution  $N_P = N_c - N_{\min}L$ is made, so that optimal  $N_c$  is given by

$$\frac{\mathrm{d}G}{\mathrm{d}N_{\mathrm{c}}} = \frac{\mathrm{d}G}{\mathrm{d}N_{\mathrm{P}}} \frac{\mathrm{d}N_{\mathrm{P}}}{\mathrm{d}N_{\mathrm{c}}} = \frac{\mathrm{d}G}{\mathrm{d}N_{\mathrm{P}}} = 0. \tag{A3}$$

Using the last expression in Eqn (A3) and G = yGPP– $w N_c$ 

$$\frac{\mathrm{d}}{\mathrm{d}N_{\mathrm{P}}}G = y \frac{\mathrm{d}}{\mathrm{d}N_{\mathrm{P}}}\mathrm{GPP} - w$$
$$= \frac{hya}{2\theta} \left( 1 - \frac{\phi I_{\mathrm{a}} + aN_{\mathrm{P}} - 2\phi I_{\mathrm{a}}\theta}{\sqrt{(\phi I_{\mathrm{a}} + aN_{\mathrm{P}})^2 - 4\phi I_{\mathrm{a}}aN_{\mathrm{P}}\theta}} \right) - w. \quad (A4)$$

Solving Eqn (A4) = 0 for  $N_{\rm P}$  gives two solutions where one is negative and the other is

$$N_{\rm P} = \frac{I_{\rm a}}{a} \phi \left[ \sqrt{\frac{1-\theta}{\frac{dhy}{w} - \theta}} \left( \frac{dhy}{w} - 2\theta \right) + 2\theta - 1 \right]. \tag{A5}$$

Substituting again  $N_{\rm p} = N_{\rm c} - N_{\rm min}L$  gives optimal  $N_{\rm c}~(N_{\rm c}^*)$ 

$$\begin{split} N_{\rm c}^* &= \frac{I_{\rm a}}{a} \phi \left[ \sqrt{\frac{1-\theta}{ahy}}_{\frac{dw}{w}-\theta} \left( \frac{ahy}{w} - 2\theta \right) + 2\theta - 1 \right] \\ &+ N_{\rm min}L = \frac{I_{\rm a}\varepsilon_{\rm max}}{a} + N_{\rm min}L \quad \text{where} \\ \varepsilon_{\rm max} &= \phi \left[ \sqrt{\frac{1-\theta}{ahy}}_{\frac{dhy}{w}-\theta} \left( \frac{ahy}{w} - 2\theta \right) + 2\theta - 1 \right] \end{split}$$

## Stability of N uptake and demand

In Fig. 2, there is a single value of *RP* where *Nd* is equal to *Nup*. For other values of *RP*, *Nd* is either less than or greater than *Nup*.

If *Nup* is less than *Nd*, the plant will experience an N deficit, in response to which the plant will increase the ratio of root N/ canopy N ( $f_r$ ) leading to a decline in  $N_c^*$ . As  $f_r$  increases, the value of *Nd* will move along the N-demand curve in Fig. 2 towards the intersection where Nd = Nup. Conversely, if *Nup* exceeds *Nd*, then  $f_r$  will decline over time and the value of *Nd* will move in the opposite direction along the N-demand curve towards the intersection shown in Fig. 2. Thus, in both cases, C allocation will change over time so that N demand approaches N uptake. In that sense, the intersections in Fig. 2, where Nd = Nup, represent stable operating points in terms of root production and N use.