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2	A unifying concept for the dependence of whole crop N:P ratio on biomass:
3	theory and experiment
4	
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# 1 Abstract

2

Background and aims Numerous estimates have been made of the concentrations of N and P
 required for good growth of crop species but they have not been defined by any unifying
 model. Our aim was to develop such a model for the dependence of N:P ratio on crop mass,
 to test its validity and to use it to identify elements of similarity between different crop
 species and wild plants.

*Methods* A model was derived between plant N:P ratio (*R<sub>w</sub>*) and its dry biomass per unit area
 (*W*) during growth with near optimum nutrition by considering that plants consist of growth
 related tissue and storage related tissue with N:P ratios *R<sub>g</sub>* and *R<sub>s</sub>* respectively. Testing and
 calibration against experimental data on different crop species led to a simple equation
 between *R<sub>w</sub>* and *W* which was tested against independent experimental data.

*Key results* The validity of the model and equation was supported by 365 measurements of
 *R<sub>w</sub>* in 38 field experiments on crops. *R<sub>g</sub>* and *R<sub>s</sub>* remained approximately constant throughout
 growth, with average values of 11.8 and 5.8 by mass. The model also approximately
 predicted the relationships between leaf N and P concentrations in 124 advisory estimates on
 immature tissues and in 385 wild species from published global surveys.

*Conclusions* The N:P ratio of the biomass of very different crops, during growth with near optimum levels of nutrients, is defined entirely in terms of crop biomass, an average N:P ratio of the storage/structure related tissue of the crop and an average N:P ratio of the growth related tissue. The latter is similar to that found in leaves of many wild plant species and even microorganisms and terrestrial and fresh water autotrophs.

23

24 Key words: Biomass, growth rate, model, N:P ratio, plant allometry, plant mass, stoichiometry.

#### **INTRODUCTION**

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2

3 Elucidating changes in N:P ratios during plant growth and developing mechanistic models for 4 them could help solve major problems in crop, environmental and ecological sciences (Sterner 5 and Elser, 2002; Sadras, 2006). For example in the context of crop production, although 6 numerous measurements have been made on the N and P concentrations in leaves as an aid to 7 diagnosing deficiencies it is not clear how far the differences in estimated "optimum" 8 concentrations result from experimental error, differences in plant age, plant mass, leaf position 9 or differences between species. In addition uncertainty about the changes in the N:P ratio of 10 crop biomass during growth has proved to be a serious problem in simulating the interactive 11 effects of N and P on crop growth (Zhang et al., 2007). A model with a strong theoretical 12 background is needed to explain some of the uncertainties in the estimated optimum nutrient 13 concentrations and thus contribute to a more effective use of added nutrients. Such a model 14 might also be useful in detecting atmospheric N-deposition damage to natural vegetation 15 (Güsewell, 2005), and in establishing the effects of imbalance of nutrients on natural vegetation 16 (Güsewell, 2004).

17 The possibility that such a model could be devised is suggested by the discovery of 18 important principles and relationships that emphasise the similarities in the N and P dynamics in 19 different organisms. The specific growth rates (SGR) of heterotrophs are generally strongly and 20 positively related to their contents of RNA and thus to their P:N ratios (Sterner and Elser, 2002). 21 The Growth Rate Hypothesis was proposed to encompass this relationship (Elser et al., 1996, 22 Elser et al., 2000b, Sterner and Elser 2002) and it applies not only across different 23 microorganisms, but also at the level of the individual species (Karpinets *et al.*, 2006). Ågren 24 (2004) has further developed the theory for algae and tree seedlings. On the basis of biochemical considerations he predicts that their SGR may increase or decrease with increase in P:N ratio depending on whether SGR is high or low. Measurements on leaves of wild herbaceous species backed up by theory indicate that SGR increased with increase in P:N ratio (Niklas, 2006). Strong correlations were also found at the species level between shoot concentrations of P and N in 117 angiosperm species grown in hydroponic culture with a constant nutrient regime (Broadley *et al.*, 2004).

7 Lack of knowledge about the changes in N:P ratios during the growth of the entire 8 biomass of both crop and of natural vegetation is particularly conspicuous. Thus in one survey 9 of natural vegetation Kerkhoff and Enquist (2006) found that the N:P ratios were approximately 10 the same, about 11, over a wide range of plant weights whereas in another study it was found 11 that the N:P ratios were greater in the leaves than in the stems and roots of herbage crops which 12 implies that the N:P ratio decreased with increase in plant mass (Kerkhoff et al., 2006). 13 Agricultural studies are sparse but a recent review (Sadras, 2006) indicates that N:P ratios of 14 cereals varied between 1 and 20. A major cause of the variability lies in variations in the supply of nutrients to crops and, in particular, the tendency of crops to absorb far more P than is needed 15 16 to meet the immediate needs and to store it (Bollons and Barraclough, 1999). However this 17 cannot explain the variability in the N:P ratio when there is an optimal supply of nutrients. Our 18 work was therefore initially focussed on crop biomass N:P ratio during growth of very different 19 crop species receiving near optimum levels of nutrients.

Models considering plants to consist of at least two types of tissues have been developed for the decline of N concentration of the whole plant with increasing plant mass (Caloin and Yu, 1982; Caloin and Yu, 1984; Hardwick, 1987; Lemaire and Gastal, 1997). Caloin and Yu (1984) developed a model that predicted a linear increase of plant N concentration with rate SGR. The prediction was confirmed by a controlled environment experiment and later by field experiments of others (Greenwood *et al.*, 1991). These studies suggest that an approach based on the relative
 rates of growth of different plant tissues could lead to a unifying mechanistic insight into
 relationships between N:P concentrations and plant biomass.

4 The objective of our work was to develop a mechanistic model along these lines for the 5 changes in N:P ratio of whole crop biomass during growth with near optimum nutrition and to 6 test its validity against a wide range of experimental data. We established that processes 7 governing N:P ratios are defined by the same equations for different species. On this basis we 8 have developed a model for the changes in N:P ratio with increase in crop mass. We have tested 9 its validity and calibrated it with experimental data obtained with near optimum nutrients. This 10 led to a simple equation which we tested against data that was entirely independent of that used 11 for developing the model. It enables predictions to be made about the N:P ratios of plant 12 biomass and of leaves. We believe that the proposed model and the equation and associated 13 principles could be useful in advancing the crop, environmental and ecological sciences.

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## MATERIALS AND METHODS

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18 Model development

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Definitions of the symbols and the units are given in the Appendix Table A1. The model is concerned with changes in the N:P ratio of whole crop dry weights, excluding fibrous roots, that occur during growth with optimum levels of nutrients before the onset of senescence. We consider that most of the changes arise because of a decline in the tissues related to growth relative to those that are related to storage and structure. More specifically we consider that

1 growth related tissues predominate in leaves and the storage and structure related tissues, which 2 will be referred as the storage related tissue, in the remaining parts of the plant. According to 3 this classification, Kerkhoff et al., (2006) found that in natural herbage the N:P ratio of the 4 storage related tissue is lower than in the growth related tissue. Thus, as plants get larger the 5 proportion of storage related tissue to growth related tissue would be expected to increase and 6 the N:P ratio to decline. We propose that at the beginning of growth, plants are mainly 7 comprised of the proliferating cells organized in metabolically active, mainly growth plant 8 tissues, which will be further referred as growth related tissue ( $W_g$ ). The N:P ratio of the plant 9 organism at this stage is considered to be similar to those of exponentially growing unicellular organisms. As the plant gets bigger the proportion of  $W_{\rm g}$  in the plant biomass decreases with a 10 11 concomitant decrease in SGR; at the same time there is an increase in the proportion of storage 12 related tissue  $(W_s)$ . On this basis we derive a simple relationship between N:P ratio of the whole 13  $\operatorname{crop}(R_w)$  and  $\operatorname{crop} dry$  biomass per unit area (W) with two constants that represent an average N:P ratio in the growth related tissue  $(R_g)$  and an average N:P ratio in the storage related tissue 14 15  $(R_s)$ . The relationship is based on the following three assumptions. 16 17 A. The growth rate of the crop is always proportional to the weight of growth related tissue  $(W_g)$ 18  $dW/dt = \alpha \cdot W_{g}$ 19 (1) 20 where W is in Mg ha<sup>-1</sup> and t is in days and  $\alpha$  is a proportionality constant. 21 22 **B**. Growth rate with the optimum levels of nutrients is defined by 23 24

$$dW/dt = K_2 \cdot W/(K_1 + W) \tag{2}$$

1

where  $K_2$  (Mg ha<sup>-1</sup> d<sup>-1</sup>) is the growth rate coefficient and  $K_1$  (Mg ha<sup>-1</sup>) is a constant. The last 3 coefficient represents the value of W when growth rate is half the maximum. The equation was 4 5 derived by considering that interception of radiation by the crop and thus dW/dt increases 6 asymptotically with W until the onset of senescence (defined as when growth rate starts to 7 decline) and it has been validated by experiments on field vegetables during the main UK 8 growing season (Greenwood et al., 1977). 9 10 By considering the previous assumption (assumption A) dW/dt can be eliminated from eqns (1) 11 and (2), which gives 12  $\alpha \cdot W_{g} = K_{2} \cdot W / (K_{1} + W)$ 13 (3) 14 C. At the beginning of growth the crop biomass (W) is comprised only of growth related tissues, 15 i.e. as  $W \rightarrow 0$ ,  $W = W_g$ 16 17 18 This assumption about the initial growth condition is important as it permits the estimation of the coefficient " $\alpha$ " in the eqn (3). Indeed, if crop biomass is small, i.e.  $W \rightarrow 0$ ,  $K_2 \cdot W/(K_1 + W)$  can 19 20 be replaced with  $W K_2/K_1$ . But according to the assumption C, crop biomass at the beginning of growth is comprised only of growth related tissue or  $W=W_g$ . Therefore  $\alpha \cdot W_g = W_g \cdot K_2/K_1$  or  $\alpha =$ 21 22  $K_2/K_1$ . As  $\alpha$  remains constant throughout growth (assumption A),  $K_2/K_1 = \alpha$  may be substituted 23 in eqn (3) to give 24

1
$$W_g = K_1 \cdot W/(K_1 + W)$$
(4)2and3and455 $W_s = W \cdot K_1 \cdot W/(K_1 + W)$ (5)677Let  $P_g$  and  $P_s$  be the total weights of P and  $N_g$  and  $N_s$  the total weights of N in the growth and the  
storage related tissues respectively, then the ratio of N:P in the whole plant  $(R_w)$  is given by9 $R_w = (N_g + N_g)/(P_g + P_g)$ (6)10 $R_w = (N_g + N_g)/(P_g + P_g)$ (6)11Multiplying throughout by  $(P_g + P_s)$  and then dividing by  $P_g$  gives13 $R_w + R_w P_g / P_g = N_g / P_g$  and  $R_s = N_g / P_s$  gives14which on rearranging and substituting the N:P ratios of the growth and storage related tissues15respectively, namely  $R_g = N_g / P_g$  and  $R_s = N_g / P_s$  gives161117 $R_w = [R_g + R_s (P_s / P_g)]/(1 + P_s / P_g)$ (7)181219Let  $C_{P_g}$  and  $C_{P_g}$  be the concentrations of P in the growth and storage related tissues then  
 $P_s = W_s \cdot C_{P_s}$  and  $P_g = W_g \cdot C_{P_g}$  which when substituted in eqn (7) gives21 $R_w = [R_g + R_s (\frac{W_s \cdot C_{P_g}}{W_g \cdot C_{P_g}}]/[1 + (\frac{W_s \cdot C_{P_g}}{W_g \cdot C_{P_g}}]]$ 

1 Measured differences in P concentration between different plant organs usually differ by a factor 2 of less than 2 (e.g. Ingestad and Lund, 1979; Greenwood et al., 1980b; de Groot et al., 2003; 3 Kerkhoff et al., 2006). The effect of varying  $C_{Ps}/C_{Pg}$  on  $R_W$  was calculated with W varying from 0.1 to 20 Mg ha<sup>-1</sup>, with  $W_s$  and  $W_g$  calculated from W by eqns (4) and (5) and with  $R_g$  and  $R_s$ 4 5 having the average values found for the different species of field vegetables. This sensitivity analysis given in [Table S1 in Supplementary Information] indicated that  $R_w$  was almost 6 7 unaffected by  $C_{Pg}/C_{Ps}$  when W was either small or large. And even over the entire range, a 8 change in  $C_{Pg}/C_{Ps}$  by ±50% on average only resulted in a less than 10% change in  $R_w$ . Thus 9 although  $C_{Pg}/C_{Ps}$  deviates considerably from unity, setting  $C_{Pg}/C_{Ps} = 1$  will result in little error. 10 We therefore consider that  $W_g/W_s$  can be taken as equal to  $P_g/P_s$  which when substituted in eqn 11 (7) gives 12  $R_{\rm w} = [R_{\rm g} + R_{\rm s}(W_{\rm s}/W_{\rm g})] / [(W_{\rm g} + W_{\rm s})/W_{\rm g}]$ 13 (8) 14 and as  $W = W_g + W_s$ , eqn (8) can rewritten as 15 16  $R_{\rm w} = [R_{\rm g} \cdot W_{\rm g} + R_{\rm s} \cdot W_{\rm s}]/W$ 17 (9) 18 19 Substitution in this equation of  $W_g$  and  $W_s$  by eqns (4) and (5) gives 20  $R_{\rm w} = [(K_1(R_{\rm g}-R_{\rm s})/(K_1+W)] + R_{\rm s}]$ 21 (10)22 23 The last equation is a linear relationship between the crop N:P ratio and the dry biomass related 24 function  $K_1/(K_1+W)$  with two constants  $(R_g-R_s)$  and  $R_s$ .

### 2 Datasets

3

#### 4 Details of all three datasets are given in [Table S2 in Supplementary Information].

5 Dataset A. Most measurements in the dataset are from 24 experiments carried out at

6 Wellesbourne, UK. In these experiments 14 different species of field vegetables were grown in 7 the same field with near optimum levels of N, P and K fertilizers (Greenwood et al., 1977; 8 Greenwood et al., 1980a). In each experiment there were between 5 and 21 harvests with 9 measurements of W, and of N and P contents, made at intervals during the growing period until 10 the commercial harvest date which is before the onset of senescence. The near optimum levels 11 of fertilizer had, depending on the crop, N:P ratios of from 0.7 to 5.4 and gave average yields 12 that were 92% of the maximum achieved with any combination of levels of fertilizer-nutrients 13 in an adjacent multilevel NPK fertilizer experiment on the same crop. The remaining 14 experiment in this dataset was at Lerida, NE Spain (Bosch Serra, 1999) on a Xerofluvent aquic, fine silty, mixed (calcareous) messic soil (Soil Survey Staff, 1975). Onions were drilled in 15 March, and at 50% emergence were thinned to 80 plants m<sup>-2</sup>. They were drip irrigated to 16 maintain the water potential higher than -18kPa and were fertigated at 334 kg N ha<sup>-1</sup>, 62 kg P 17 ha<sup>-1</sup>, and 403 kg K ha<sup>-1</sup>. Pest disease and weed control were effective. 18

Altogether there were 287 measurements on 15 different species (Table 1). A special feature of these experiments is that there is a sufficient proportion of measurements (43%) with small values of W (W< 2 t h<sup>-1</sup>) to provide a sensitive test of the model.

22 Datasets A was also used to test the validity of the integrated form of eqn (2) namely

23

24  $K_2 t = K_1 \cdot \ln W + W + C$  (11)

2 where C is the constant of integration and where W is crop dry weight (excluding fibrous roots in Mg ha<sup>-1</sup> at time t in days and  $K_1$  is equal to 1 Mg ha<sup>-1</sup> for field vegetables as this had been 3 4 found to give good fits to past field experimental data (Greenwood et al., 1977; Greenwood et 5 al., 1985).  $K_2$  is a growth rate coefficient which is generally constant throughout growth until senescence; in effect it corrects growth for differences in plant mass.  $K_1 lnW + W$  was regressed 6 against t for each experiment to provide estimates of  $K_2$ ; on average the regressions had an r<sup>2</sup> of 7 8 0.96 (Table 1).

9

10 Dataset B. This dataset was obtained by searching the literature for measurements at intervals, 11 during growth with near optimal nutrition, of W, %N, and %P that were not included in dataset 12 A. It consists of 78 sets of measurements from 13 experiments in 5 countries; there were 6 13 different crops of which 5 were not in dataset A. The data was also different from that in dataset A, because it contained few measurements where W was less that 2 Mg ha<sup>-1</sup> and, thus, did not 14 allow satisfactory estimates of  $R_g$ . It was used to test the predicted relationship between N:P 15 ratio and W for W between 0.1 and 27 Mg ha<sup>-1</sup>. It was also used to confirm the finding from 16 17 dataset A that the time course dry crop weight increase followed eqn (11) [Table S2 in 18 Supplementary Information].

#### 19 The measured average N:P ratio is 7.9 in dataset A and 7.2 in dataset B compared with 20 an average of 7.2 in the particulate matter in the sea which is the same as the average ratio in sea 21 water (Sterner and Elser, 2002, p29).

22 Dataset C. Plant analyses are often used to guide fertilizer practice. Many experiments have been carried out to find the "optimum or adequate concentrations" at different stages of growth 23 24 of many crop species. Tcierling (1978) deduced such concentrations from cereal experiments in Russia and Bergman (1992) deduced them for numerous crop species from experiments in a wide range of countries. Dataset C includes measurements taken directly from the Tcierling's review (Tcierling 1978). Bergman (1992) gives the upper and lower values of the adequate range of %N and %P; the mean values of each of these two values are also given in dataset C and these were compared with the predictions as described below. Tcierling's data consisted of values of %N and %P ( for the above ground parts of the plant) at different growth stages of cereals.

8 Neither the Tcierling nor the Bergman data included values of W, which is an essential input
9 into eqn (10) after calibration to form eqn (15), as will be described in the Results section.
10 Apparent values of W in the dataset were calculated from inputs of %N by

11

12 *Optimum* 
$$\%N = 1.35(1 + \beta e^{-0.26W})$$
 (12)

13

14 (Greeenwood and Draycott, 1989) with  $\beta = 3$  which had been found to be satisfactory for many field vegetables and has been used widely (Zhang et al., 2007). For the Tcierling data eqn (12) 15 16 was used to produce a set of corresponding values of %N, and W.  $R_w$  was calculated by inserting 17 these calibrated values of W into eqn (15). The Bergman (1992) data covered a considerable 18 range of crop species and of %P and %N. These mean values for each species were compared 19 with a predicted relationship between %P and %N. To generate the prediction we first used eqn. (12) to calculate %N for  $W = 2 \text{ t ha}^{-1}$  to 20 t ha<sup>-1</sup> in steps of 1 t ha<sup>-1</sup>. The same values of W were 20 21 then used to calculate  $R_w$  by inserting into eqn (15) and to predict %P multiplying  $1/R_w$  by %N. 22 These values of %P were regressed against those of %N to give a predicted linear relationship 23 that was compared with Bergman's data in Fig. 5. The same procedure but with a different 24 range of values of W was used to calculate a predicted relationship between %P and %N for

1	comparison with experimental data on grasses reported by (Salette, 1990) given in Fig.6. Thus,
2	these predicted linear relationships were independent of the data with which they were compared.
3	
4	Regression analysis. Whenever both variables were subject to error regressions were by the
5	Reduced Major Axis (RMA) procedure (Niklas, 2006) with software by Bohonak (2004). If only
6	one variable was subject to error, regression was by the standard procedure.
7	
8	RESULTS
9	
10	1. The N:P ratio of the whole plant declines with increase in plant mass during growth
11	
12	The validity of eqn (10) was tested by regression of plant N:P ratio ( <i>Rw</i> ) against $1/(1+W)$ , as $K_1$
13	= 1 Mg ha <sup>-1</sup> , for the measurements in dataset A. Fig. 1 gives regressions for those experiments
14	with most measurements and illustrates that the points were generally randomly scattered about
15	the best fitting line; Table 1 gives values of $r^2$ averaged over the experiments for each crop.
16	They show that measurements of $W$ and N:P ratio during growth fit eqn (10) closely. Thus this
17	theoretical equation gives good predictions for all crops of the linear increase in the ratio as a
18	function of $1/(1+W)$ . Therefore the ratio decreases with increase in $W$ .
19	
20	2. The N:P ratio of the whole plant declines with decrease in the plant specific growth rate
21	during growth
22	
23	Strong evidence that, for a given experiment, eqn (2) with the same value of $K_2$ held throughout
24	growth is provided by the excellent fits (average $r^2$ for each experiment = 0.96) to eqn (11), the

1 integral of eqn (2) for dataset A (Table 1). Similar results were obtained for dataset B [Table S2 2 in Supplementary Information]. From eqn (2) 3  $SGR = \frac{dW}{dt} \frac{1}{W} = K_2 / (K_1 + W)$ 4 (13)5 Substitution of  $K_2/SGR$  for  $(K_1+W)$  of this eqn in eqn (10) gives 6 7 8  $R_{w} = [K_{1}(R_{g}-R_{s}). SGR/K_{2}] + R_{s}$ (14)9 Assuming all of the parameters except  $R_w$  and SGR are constant as is indicated by Table 1 it 10 11 follows that during a given experiment  $R_w$  decreases with decrease in SGR. 12 13 3. Average N:P ratios are similar among crops at the level of growth related tissue or of storage related tissue and do not change significantly during plant growth 14 15 16 According to eqn (10) the calculated regression coefficients (intercept and slope) allow 17 estimates to be made of the N:P ratio in the growth related tissue  $(R_g)$  and in the storage related 18 tissue  $(R_g)$  for each crop (Table 1). Because there were two independent experiments on most 19 crops, an analysis of variance of the values of  $R_{\rm g}$  and of  $R_{\rm s}$  was carried out. The standard errors for the differences between crop means,  $R_g$  and  $R_s$  are given at the bottom of Table 1. Both  $R_g$ 20 21 and  $R_s$  did not differ significantly between most crops. Given this similarity the average values of  $R_g$  and  $R_s$ , which are 11.83, and 5.57 respectively, were used as constants in the theoretically 22 derived eqn (10), with  $K_1 = 1$  t ha<sup>-1</sup> to give 23  $R_{\rm w} = 6.26/(1+W) + 5.57$ 24 (15)14

2 This calibrated equation was further verified by considering the N:P ratio in different crops at
3 different stages of growth using experimental data in datasets B and C.

4

# 5 4. The validity of the model and equation (15) is supported by agricultural and ecological 6 experiments

7

Agricultural experiments (dataset B) that are entirely independent of those used for developing the model give the relationship between N:P ratio and W that is similar to calculated from eqn (15) (Fig. 2). The dataset B includes measurements made throughout the growth period on the C4 crop maize and on five C3 crops. The values of W range from 0.1 to 27 Mg ha<sup>-1</sup> and yet are in reasonably good agreement with eqn (15).

As N:P ratios are particularly sensitive to changes in W, when W is small, we tested the validity of the model against measurements made in the early stages of crop growth. In dataset A, the average N:P ratio of crop seedlings, for values of W < 0.2 Mg ha<sup>-1</sup>, is 9.8 compared with 11.16 obtained by substituting W=0.1 Mg ha<sup>-1</sup>in eqn (15). The average ratio of seedlings in dataset B with W < 2 Mg ha<sup>-1</sup> was 9.3, which is close to the predicted value of 8.65 obtained by substituting W=1 Mg ha<sup>-1</sup>in eqn (15). [Tables S3 and S4 in Supplementary Information].

According to the model, the average N:P ratio of the growth related tissue,  $R_g$ , is an important plant characteristics that does not vary much during growth (Fig.1, Table 1). As a result changes in N:P ratio during the growth is mainly driven by changes in the biomass of the growth related tissue, but not by its N:P ratio. Because leaves consist mainly of growth related tissue the average values of N:P ratio in leaves should be close to the  $R_g$  found in the study. We have studied ecological measurements on leaves to check this proposition, namely, to confirm

1 that leaf %P is proportional to leaf %N with a proportionality constant of 1/11.83 = 0.0843. Fig. 2 3a gives leaf %P plotted against leaf %N on logarithmic axes for the 206 sets of data on the 3 leaves of herbage species (Kerkhoff et al., 2006) and Fig. 3b gives a similar plot for 177 sets of 4 data on the leaves of evergreen trees of deciduous shrubs (Wright et al., 2004). The lines are 5 calculated by assuming the N:P ratio is always equal to that of the growth related tissue namely, 6 0.0843 and appear to give a good representation of the average values of the data. Leaf 7 photosynthesis usually ceases when the leaf nitrogen concentrations are low (Nátr, 1975) so 8 such leaves could not be regarded as growth related tissue. For this reason we have confined 9 formal statistical analyses on leaf measurements to leaves with a %N greater than one. The 10 details of these analyses are given in Table S5 in supplementary information. RMA 11 regressions of log<sub>10</sub>%P against log<sub>10</sub>%N were carried out separately on the Kerkhoff and the 12 Wright data. The means sums of squares of the deviations between measured and calculated 13 values (from the regressions) of  $\log_{10}$ %P and vice versa were determined. They were also 14 determined when the values were calculated from P/N = 0.0843. Although the gradients of the 15 RMA fits were greater than one indicating a curvature in the relationships, the degree of 16 agreement between measured and calculated values was not significantly different whether 17 calculations were from the RMA fitted equations or from P/N = 0.0843. The ratio of the mean 18 sum of squares of deviations for calculations with the latter equation to that with the former equations was 1.14 which is not significant at p = 0.05. Thus P/N = 0.0843 gives a good 19 20 description of the data. This is further supported by the finding that the average P:N ratios of the 21 Kerkhoff and of the Wright leaves were not significantly different from 0.0843. Therefore the 22 P:N ratio of wild leaves having a %N greater than one is similar to that inferred from our model 23 (eqn 10) and experiments on field vegetables.

5. The calibrated theoretical relationship in eqn (15) predicts optimal N and P
 concentrations in the dry biomass of different crops

3

4 Relatively few experiments (given in Fig.2) have been published where sequential 5 measurements were made of W, %N and %P. on crops grown with near optimum levels of 6 nutrients. However, a considerable amount of other data exists on the optimal %N and %P that 7 does not include W for different crops and conditions. We therefore sought to find how far they 8 could be predicted with eqn (15). The methods were as previously described under dataset C. 9 The data includes N:P ratios (dataset C) at each of four growth stages of four different C3 10 cereals, grown with 'adequate nutrition', that were derived from Tcierling's (1978) review of 11 nationwide Russian experiments. Comparisons between measured N:P ratios and those 12 calculated with eqn (15) are given in Fig.4. An upper limit for the s.e of the measurements is 0.7 13 (see legend to Fig.4) indicating that the least significant difference is about 1.4. As in 12 out of 14 the 15 comparisons the difference between measurement and calculated N:P ratio is less than 1.4, 15 it follows that agreement between measurement and calculation is reasonably good. Bergman 16 (1992) in his review of world-wide literature gives the ranges of %N and %P of immature 17 tissues of numerous species required for good growth. Altogether the data set includes 124 18 records (dataset C), the average of the range of %P of each record was plotted against the 19 average %N. They are linearly related and are plotted in Fig. 5 which also gives the linear 20 relationship predicted with eqn (15). The slopes of the two relationships are almost identical but 21 the intercept of the predicted relationship is about 0.1% higher that that for the measurements. 22 Measurements of %P and %N of the above ground biomass made at intervals during the growth 23 of tall fescue and rye grass grown with near optimum conditions (Salette, 1990) are also in a 24 good agreement with that obtained with eqn (15) (Fig. 6).

#### DISCUSSION

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4 Our study indicates that the N:P ratios in the growth related tissues of many crops are about 5 11.83, and are similar to those of the leaves of many naturally grown species (Fig.3). This value 6 is in the range of that found for micro-organisms (Sterner and Elser, 2002) and is close to the 7 averages of 12.65 and 13.64 for terrestrial and fresh water autotrophs (Elser et al., 2000a). 8 These similarities imply that the same cellular processes, such as changes in the transcribed 9 rRNAs (Elser et al., 2000b) and in RNA:protein ratios (Karpinets et al., 2006), may underlie the 10 relationships between the N:P ratios of growing cells and their growth rate whether in plants or 11 micro-organisms. The nutrient requirements of this conserved molecular machinery could 12 dominate the N and P requirements for cellular growth and thus explain their close balance and 13 synergy in natural ecosystems (Elser et al., 2007).

14 An alternative to the proposed mechanistic modelling of the crop N:P ratio is to fit an 15 empirical model to the experimental data. One widely used model in crop nutrition (Lemaire, 16 1997) considers the relative changes in %N and %P during growth of the whole crop in terms of simple scaling relationships with W; these are  $\% N = aW^b$ ;  $\% P = cW^d$  and N:P =  $(a/c)W^{(b-d)}$  where 17 18 a, b, c and d are coefficients that are fitted separately for each experiment. The model gave good 19 fits to each of 5 sets of our test data [Table S6 in supplementary information]; an example of 20 which is given in Fig.7. As both the coefficients b and d were always negative and b was less 21 than d the relative decline in %N was always greater than that of %P. The model also indicates 22 that N: P will never remain constant unless b = d. Our mechanistic model also gave a good 23 description of the dependence of N:P on W. In addition, the calibrating coefficients have a clear 24 biological sense representing the N:P ratios in each of the main plant tissues. One of these

parameters appears to be constant, as mentioned above, not only across plants, but also across
 microbial organisms which demonstrates that proposed mechanistic model quantifies global
 regularity in biological plant species.

4 In the Introduction we emphasised that in accord with the Growth Rate Hypothesis the 5 specific growth rate of heterotrophic organisms increases with decrease in N:P ratio (Elser et al., 6 1996; Elser et al., 2000b; Sterner and Elser, 2002). It was also pointed out that a similar 7 relationship had been found for leaves (Niklas, 2006). In fact we found limited support for the 8 relationship as the N:P ratio of the growth related tissue, which is thought to consist mainly of 9 leaves, for the experiments in data set A was negatively, although weakly, correlated with the growth rate  $K_2$ . Nevertheless in our studies the N:P ratio of the whole plant declines as plants 10 grew and their specific growth rate fell, which is in accord with Ågren's predictions (Ågren, 11 2004) for whole plants with SGR's (<0.1 day<sup>-1</sup>) that are similar to the SGR's of our data. 12 13 According to our model this difference occurs because whole plants, unlike unicellular 14 organisms and individual parts of plants, consist of different tissues, each with different functions and thus different demands for N and P. During growth the proportions of the tissues 15 16 change leading to changes in the total requirement of the whole plant for N relative to that of P. 17 Our model is a simplification of the various processes but its validity is supported by much 18 experimental information covering widely different species grown in different ways. It 19 illustrates how strongly changes in the relative proportions of the growth and storage related 20 tissues dominate the N:P ratio of the whole crop. For example, according to eqn (4), the weight 21 of growth related tissue as a fraction of total plant weight, W, is equal to 1/(1+W) which 22 indicates that the fraction declines sharply with small increases in W; in fact the fraction drops from 1 to 0.5 with an increase in W from 0 to 1 Mg ha<sup>-1</sup>. As the average N:P ratio of this tissue 23 24 11.83 and that of storage related tissue is 5.58 it follows that there is a very sharp fall in the N:P

ratio of the whole plant with small increases in *W* as found experimentally (Fig. 2). It thus
 supports the view that N:P ratio of crop biomass grown with optimum concentrations of nutrient
 is greatly affected by the partition of assimilates to different tissues during growth.

4 The low N:P ratio of field vegetables  $(R_w)$  compared with that of much standing mass of 5 natural communities deserves comment. The low N:P ratio is dominated by the low N:P ratio of 6 the storage related tissues  $(R_s)$  which is subsequently used for reproduction and thus requires a 7 high %P. As the average weight of the standing biomass of many natural communities is much 8 greater than that of crops most of it could have consisted of wood which contains only very 9 small amounts of nutrients (Altman and Dittmer, 1964). In consequence the N:P ratio could be 10 largely determined by that of the growth related tissue and be approximately 11.83 irrespective 11 of the total weight of biomass. Indeed, a survey of such communities indicated that they had an 12 N:P ratio of approximately 13 irrespective of the total weight of biomass (Kerkhoff et al., 2005).

13 Ecological studies emphasise the similarities in the relationships between %N and %P in 14 the dry matter of leaves grown on soils of differing fertility (Niklas et al., 2005), whereas crop studies emphasise the dependence of crop %N and of %P on soil nutrient levels (Lemaire, 1997; 15 16 Bollons and Barraclough, 1999). In this paper we show that the average N:P ratio of naturally 17 grown leaves in a large dataset is on average similar to the N:P ratio of the growth related tissue 18 of crops grown with near optimum levels of nutrients. As growth related tissue is thought to 19 consist mainly of leaves, the average N:P ratio of natural leaves appears to be that expected with 20 optimal plant nutrition. Presumably the leaves of natural vegetation have adapted over long 21 periods to the stable but low soil nutrient levels, by various feed back mechanisms in such a way 22 that the different nutrients limit growth simultaneously (Bloom et al., 1985; Gleeson and Tilman, 23 1992; Knecht and Göransson, 2004) and that when this occurs the N:P ratios are similar to those 24 obtained for plants grown with the optimum fertilizer levels. Arable crops are in the ground for

1 much shorter periods, have much less time to adapt to the nutrient regimes, so homeostasis is 2 less marked and plant nutrient concentrations are more dependent on soil nutrient levels. They have also been bred for rapid growth and high yields, possibly at the expense of their root 3 4 systems and their ability to withstand soil nutrient stresses. Thus inter-site variation in growth 5 of crops could be much more dependent on soil nutrient levels than that of natural vegetation. 6 Also it is possible that the N:P ratio of the growth tissue is approximately the same over a wide 7 range of natural and crop species and the soil nutrient affects on plants are entirely 8 accommodated by differences in the N:P ratios of the storage related tissues.

9 Our model could improve crop nutrition by providing general diagnostic criteria for 10 nutrient imbalance and by its incorporation into dynamic models for crop response to fertilizers, 11 as described in the Introduction. Nevertheless some characteristics of the experimental data used for developing eqn (15) impose limits on its application. It may not apply when W is 12 greater than 20 Mg ha<sup>-1</sup> as most values of W in the experiments were less than this. It may not 13 14 apply after the onset of appreciable leaf senescence as the field vegetables were harvested before 15 the onset of senescence. An obvious major uncertainty in these studies is the extent to which 16 deviations from optimum applications of N and P fertilizers have affected the measured N:P ratios. Less obvious is the error that may have resulted from setting  $K_I = 1 \text{ Mg ha}^{-1}$  for all crops. 17 It is the value of W when growth rate is half the maximum. Such a low value seems to be 18 19 appropriate for most of the field vegetables grown in the field experiments as these were grown 20 at high plant densities, and complete crop cover and maximum growth rates were attained whilst W was small. Setting  $K_1 = 1$  Mg ha<sup>-1</sup> for all crops was based on the excellent fits that were 21 obtained with this value of  $K_1$  in fitting eqn (11) (Table 1). However,  $K_1$  and  $K_2$  are highly 22 23 correlated in such fits and equally good fits can be obtained with different combinations of values of each of these coefficients and it may be that  $K_1$  should be greater than 1 for widely 24

1 spaced crops. The same uncertainty exists for eqn (15) governing the dependence of N:P ratio 2 on W. Increasing  $K_1$  in eqn (10) and also in eqn (15) results in the N:P ratio declining less rapidly 3 with increase in W. This could explain why  $R_w$  of Brussels sprouts declined less with increase in W as it was grown at a far wider spacing than the other crops. Wide plant spacing depressed the 4 5 decline of %N of lucerne with increase in plant mass compared with close spacing (Lemaire 6 and Gastal, 1997) which could be explained in terms of the interception of light (Hirose and 7 Werger, 1987; Lynch and Gonzalez, 1993). Spacing might therefore be expected to affect the 8 dependence of N:P ratio on W.

9 We conclude that the same physiologically derived equation with two constants gives a 10 good prediction of the changes in N:P ratio in the biomass of a wide range of crops grown, in 11 different ways, with near optimal nutrition. The model also gave good estimates of the relation 12 between %P and %N in the leaves of natural vegetation provided the leaf %N was greater than 13 one. The N:P ratio of crop biomass declines sharply with increase in plant mass per unit area 14 because of an increase in the proportion of storage related tissue which has a low N:P ratio 15 relative to growth related tissue which has a high N:P ratio.

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11	APPENDIX
12	Table A1
13	
14	
15	Supplementary Information
16	
17	<b>Table S1</b> Effect of variation in the ratio $C_{Ps}/C_{Pg}$ on $R_w$
18	
19	Table S2 Details of datasets A, B and C with references to the sources of data
20	
21	<b>Table S3</b> N:P ratios for values of $W < 0.2$ Mg ha <sup>-1</sup> from dataset A
22	
23	<b>Table S4</b> N:P ratios of $W < 2$ Mg ha <sup>-1</sup> from dataset B with references to the sources of data
24	

1	Table S5 Statistical analyses of predictions of P:N ratios of wild leaves
2	
3	Table S6 Empirical model fits measurements of %N and %P during crop growth
4	
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17	Legends to figures
18	
19	Fig. 1. Relationships between $N/P$ and $1/(1+W)$ with W in Mg ha <sup>-1</sup> for each experiment from
20	database A; the lines fitted by the RMA procedure. The data are for single experiments, namely
21	French beans 73, pea 72, onions 92, swede 72, and turnip 72 and the corresponding values of $r^2$
22	are 0.85, 0.90, 0.80, 0.91, 0.73 respectively. Each of the three different symbols for the onion
23	data refers to a different variety.
24	

Fig. 2. Literature measurements (database B) of the ratio, N/P and W in Mg ha<sup>-1</sup> during 1 2 growth of crops grown with near optimum levels of nutrients and predicted dependence of the 3 ratio on W by eqn (15). Details and sources of data are in [Table S2 in Supplementary 4 Information]. 5 6 Fig. 3. Comparison between predicted relationships (lines) calculated for the growth related 7 tissue (%P = %N/11.83) and measurements (symbols) of the leaves of wild plants obtained in 8 surveys.). A gives all 206 measurements for a survey of the leaves of herbage wild species 9 referred to in Kerkhoff et al. (2006) and kindly provided by Professor Kerkhoff. B. gives the 10 147 leaf measurements with %N > 0.5 % on evergreen trees (open circles) and 30 11 measurements on deciduous shrubs (closed circles) reported in Wright et al. (2004). 12 13 Fig. 4. Estimated (closed columns) and predicted values of %N/%P (hashed columns) at tillering 14 ear (Stage of growth 1), ear emergence (Stage of growth 2), booting (Stage of growth 3) and flowering (Stage of growth 4) of C3 cereals grown with optimum nutrients in Russia. An upper 15 16 limit for the average s.e. for each of the closed columns is 0.702 (d.f. =10). It was calculated 17 from the high order interactions in an analysis of variance for this data when combined with 18 associated data for the same crops grown with excess nutrients (Tcierling, 1978). 19 Fig. 5. The predicted relationship (continuous line) between optimum %P and %N (%P =  $0.1152 \times \%$ N + 0.0788 r<sup>2</sup> = 0.99) as described in Dataset C of the Dataset section and 124 20

21 advisory estimates (symbols) from the literature for leaves and whole plants of many species

22 (Bergman, 1992). RMA regression (dashed line) for these advisory relationships gives %P = 0.

23  $1208 \times \%$ N -0.03436, r<sup>2</sup> = 0.61; the s.e. of the gradient is 0.0068.

1 Fig. 6. Predicted relationship (continuous line) between optimum %P and %N ( %P =

0.0981 %N + 0.1151, r<sup>2</sup> = 0.97and measured values (calculated from authors regressions which
had r<sup>2</sup> >0.89) for grasses grown with near optimum levels of nutrients in France (Salette, 1990)

**Fig.7.** The calculated relationship between N:P and W (continuous curve) and the measured values of N:P for the turnip 72 experiment. Calculations were made with N:P = $(a/c)W^{(b-d)}$  in which the coefficients a and b were obtained by fitting %N =  $aW^b$  and c and d by fitting %P =  $CW^d$ . The fitted values of a and b were 4.375 and -0.157 (r2=0.742) and of c and d were 0.614

9 and -0.0488 (r2=0.644).