



# Article Chronically Low Nutrient Concentrations in Tree Rings Are Linked to Greater Tree Vulnerability to Drought in Nothofagus dombeyi

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**Abstract:** Forest dieback and mortality episodes triggered by droughts are receiving increasing attention due to the projected increases in these extreme climate events. However, the role played by nutrient impairment in dieback is understudied, despite interactions among carbon-water balances and nutrition. Here, we followed a comparative analysis of long-term growth, intrinsic water-use efficiency (iWUE), oxygen isotopes ( $\delta^{18}$ O) and wood-nutrient composition patterns between living (L) and dead (D) trees of a *Nothofagus dombeyi* population, showing dieback in Argentina. The onset of the growth decline of D trees occurred ca. 40 years before death. These trees showed higher iWUE, pointing to higher drought stress. Their lower  $\delta^{18}$ O values, together with the uncoupling between  $\delta^{18}$ O and leaf-level processes, suggested a deeper source of water uptake for this vigor class. D trees showed a poorer nutritional status than L trees that likely amplified the dieback. This was supported by numerous positive associations of P- and K-concentrations in wood and related ratios with iWUE,  $\delta^{18}$ O and tree growth. Therefore, drought-related nutrient deterioration can significantly contribute to dieback and be an early warning signal of impending tree death.

**Keywords:** basal area increment; dieback; nutrient limitation; Patagonia; soil depth; water sources; water-use efficiency

# 1. Introduction

Episodes of forest dieback and tree mortality triggered by severe droughts have been reported worldwide [1], and they affected boreal [2], temperate [3], semi-arid [4] and tropical biomes [5]. The increase in temperature and vapor-pressure deficit will likely be accompanied by increased interannual variability in precipitation and/or reduced rainfall in some regions [6], leading to more frequent and severe dry spells [7,8]. These projected climatic trends have raised concerns about the fate of drought-sensitive forests under warmer and drier conditions [9].

Two main physiological processes have been proposed to explain drought-induced tree dieback: hydraulic failure (i.e., cessation of symplastic biochemical functioning and disruption of water transport due to xylem embolism), and carbon starvation (i.e., strong reduction of internal carbon pools that disrupts the maintenance of metabolism) [10,11]. Although both mechanisms are non-mutually exclusive, hydraulic failure has received stronger support across taxa, particularly in gymnosperms [12]. However, canopy dieback and tree mortality are complex phenomena in which water stress can interact with other factors, such as a deficient nutrient availability and biotic stressors [13,14].

Nutrient availability and uptake are largely controlled by soil moisture conditions. Drought decreases nutrient mineralization due to reduction in soil microbial activity and



**Citation:** González de Andrés, E.; Suárez, M.L.; Querejeta, J.I.; Camarero, J.J. Chronically Low Nutrient Concentrations in Tree Rings Are Linked to Greater Tree Vulnerability to Drought in *Nothofagus dombeyi. Forests* **2021**, *12*, 1180. https://doi.org/10.3390/ f12091180

Academic Editor: Martin Bader

Received: 29 June 2021 Accepted: 26 August 2021 Published: 31 August 2021

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**Copyright:** © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). ion mobility, so reducing nutrient availability for plants [15]. The acquisition of nutrients depends on nutrient diffusion and mass flow driven by the water potential gradient created by leaf transpiration that transports nutrients from soil to roots [16]. In addition, drought kills many fine roots which play a relevant role to uptake soil nutrients [10,14]. Consequently, prolonged droughts may cause nutrient deficiencies and stoichiometric imbalances that impair important physiological processes, such as photosynthesis, growth or hydraulic function [17,18], which can result in long-term increases of tree-drought sensitivity [19]. For instance, Salazar-Tortosa et al. [20] proposed the existence of a detrimental feedback loop between reduced transpiration and nutrient uptake in species with a tight stomatal regulation under drought stress. Besides the plastic behavior of many tree species capable of shifting water uptake to deeper soil layers, subsoil or weathered bedrock during drought [21] may exacerbate the contribution of nutrient impairment to drought-induced dieback, considering the lower abundance of nutrients as soil depth increases [22]. Subsequent nutrient imbalances would unleash physiological processes leading to forest dieback.

Tree rings are a powerful tool to evaluate long-term growth trends and patterns in tree-ring width, which have been used to identify early-warning signals of dieback and tree mortality [23,24]. Yet, additional information is required to disentangle the physiological mechanisms leading to growth decline. The analysis of carbon and oxygen isotopic ratios of wood have been commonly used for this purpose [25,26]. Carbon isotope signatures in tree rings are modulated by the regulation of gas exchange during ring formation, due to the linkages between carbon-isotopic discrimination during photosynthesis ( $\Delta^{13}$ C) and the ratio between intercellular CO<sub>2</sub> concentration (*Ci*) and atmospheric CO<sub>2</sub> concentration (*Ca*) [27,28]. Increasing *Ca* and water limitation result in stronger stomatal control, leading to higher intrinsic water-use efficiency (iWUE). However, those increases in iWUE have not usually translated into enhanced tree growth [29,30]. The analysis of oxygen isotopes ( $\delta^{18}$ O) has been proposed to help discern the effects of environmental changes on stomatal conductance, since  $\delta^{18}$ O is not substantially affected by photosynthesis, but depends on leaf-level evaporative effects and the transpirational dilution of evaporatively enriched leaf water [31].

Nevertheless, the interpretation of the so-called dual-isotope approach [26] presents with some caveats. Firstly, the oxygen-isotope signatures of tree rings may reflect not only evaporative processes at the leaf level, but also variations in the  $\delta^{18}$ O of source water and exchanges in oxygen between carbohydrate pools associated with transport and xylogenesis [32,33]. For instance, Treydte et al. [34] reported stronger signals of source water than leaf enrichment on a conifer species. In fact, variability in tree-ring  $\delta^{18}$ O has been associated with species-specific patterns in seasonal water uptake [35,36] and to water uptake from deeper soil layers with increasing drought [37,38]. Secondly, this isotope-based approach focuses on carbon and water relations in the tree, albeit other potential mechanisms may contribute to forest dieback, including nutrient availability and uptake [20] and related nutritional imbalances [39,40]. Therefore, the combination of different approaches that include growth patterns, wood isotopic signals and dendrochemistry data can provide insights into the growing knowledge that links nutrient-related processes to tree mortality triggered by droughts.

Here, we focused on *Nothofagus dombeyi* (Mirb.) Blume, an evergreen broadleaf species inhabiting sites showing a wide range of climatic and edaphic conditions from mesic-tocool temperate rainforests in Patagonia, South America. The patchy canopy dieback and mortality episodes of this species, at the eastern side of the Andes range (Argentina), have attracted the attention of scientists for decades [41–43]. High growth variability has been identified as a predisposing factor to tree mortality [42]. However, the ecophysiological basis behind such conspicuous patterns remains poorly understood (but see [44]). So, we aimed at analyzing patterns of radial growth, carbon and oxygen isotope discrimination and nutrient composition in tree rings of conspecific living and dead individuals to attain a long-term perspective of drought-induced dieback in *N. dombeyi* and to infer its underlying mechanisms. Our specific objectives were to assess the potential differences between treevigor classes regarding (1) carbon and water relations, and (2) nutrient status, and (3) to determine mechanistic linkages among them and with radial growth patterns.

# 2. Materials and Methods

# 2.1. Study Area and Species

The study area is located by the Lake Gutiérrez (41°10′35″ S, 71°24′05″ W, 850 m a.s.l.) in the Nahuel Huapi National Park, northern Patagonia, Argentina. It encompasses the dry eastern edge of the species' distribution, where *N. dombeyi* co-exists with the conifer *Austrocedrus chilensis* in even-aged stands, established after devastating wildfires about 150–170 years ago [45]. Their diameters at breast height (DBH), range between 30 and 135 cm, with a mean value of  $63.2 \pm 23.0$  cm. The stand density is 375 trees ha<sup>-1</sup>. The soils are Andosols, derived from layers of volcanic ash covering the glacial topography.

The climate in the study area is strongly determined by the rain–shadow effect exerted by the Andes, which leads to a precipitation gradient with decreasing rainfall eastwards [46]. Climatic data were retrieved from the nearest weather station (Bariloche Airport,  $41^{\circ}09'00''$  S,  $71^{\circ}10'12''$  W, 840 m a.s.l.) that covers the period 1931–2018. The mean annual temperature is 8.2 °C. January and February are the months with higher temperatures (average temperature of 14.4 °C), while July is the coldest month (average temperature of 2.4 °C). The mean annual precipitation is 860 mm, and it is seasonally distributed, with the period between May and August providing more than 60% of the annual precipitation. Identification of drought events was assessed using a multi-scalar drought index, the Standardized Precipitation Evaporation Index (SPEI) [47]. The SPEI was calculated on a six-month time scale, including the growing season (previous October-March [43]). We calculated SPEI using Bariloche climate data and the *SPEI* package [48] in R software [49].

Monthly  $\delta^{18}$ O of precipitation was retrieved from the Global Network of Isotopes in Precipitation (GNIP, [50]). We selected the closest station with long-term  $\delta^{18}$ O data (1967–2017), i.e., Puerto Montt, Chile (41°28′12″ S, 72°55′48″ W, 45 m a.s.l.), located 130 km away from the study area.

#### 2.2. Field Sampling and Dendrochronological Data

We sampled 17 couples of dominant healthy living (L) and dead (D) trees (34 trees in total). Only dead trees showing evidence of being killed by climatic events and exhibiting bark or recently fallen bark were selected, in order to avoid the inclusion of old (rotten) dead trees [51]. Two cores were extracted at 1.3 m height from each tree using increment borers. The wood samples were air-dried, glued onto wooden mounts, and polished until the xylem cellular structure was visible [52]. We visually cross-dated all samples, and tree-ring width was measured to a precision of 0.001 mm, using a Velmex Measuring System (Velmex Inc. Bloomfield, NY, USA). The quality of the cross-dating was examined using the software COFECHA, which calculates moving correlations among individual tree series [53]. We followed the convention for the Southern Hemisphere, which assigns to each tree ring the date of the year in which growth started. The mortality dates of dead trees (i.e., the date of the outermost tree ring) were defined using the master chronology developed for the site [54]. DBH of each sampled tree was measured at field at the time of sampling using tapes.

Tree-ring width series were transformed to basal area increment (BAI) series since it is a two-dimensional measure that better represents the growth trend of the whole tree than one-dimensional tree-ring width [55]. We assumed concentric rings and used the following equation:

$$BAI = \pi (R_t^2 - R_{t-1}^2), \tag{1}$$

where  $R_{t}^{2}$  and  $R_{t-1}^{2}$  are the radii corresponding to the years *t* and *t* - 1, respectively.

#### 2.3. Tree-Ring Isotopic Composition

Carbon and oxygen isotope analysis was performed in groups of five contiguous rings for the period 1915–2018 (e.g., 1915–1919, 1920–1924, etc.) pooled by tree-vigor class level (n = 21 samples per vigor class). For that purpose, the wood radii of each tree were carefully separated using a scalpel in 5-year segments, milled and homogenized to a fine powder using a ball mill (Retsch ZM1, Haan, Germany). Stable isotope composition was measured at the Stable Isotope Facility (University of California, Davis, CA, USA). The isotopic composition of carbon ( $\delta^{13}$ C) and oxygen ( $\delta^{18}$ O) were expressed relative to Vienna Pee Dee Belemnite and Vienna Standard Mean Ocean Water standards, respectively [56]. The accuracy of the analyses (SD of working standards) was 0.05–0.21‰.

Intrinsic water-use efficiency (iWUE) was calculated from  $\delta^{13}$ C as the ratio between the photosynthetic rate (*A*) and the stomatal conductance rate (*g*<sub>*S*</sub>), according to the next equation [27]:

$$iWUE = Ca \times [1 - (Ci/Ca)] \times 0.625,$$
 (2)

where *Ca* and *Ci* are  $CO_2$  concentrations in the atmosphere and the intercellular space, respectively, and 0.625 is the relation among conductance of H<sub>2</sub>O and CO<sub>2</sub>. To determine *Ci*, we used the following equation:

$$Ci = Ca \left[ (\delta^{13}C_{\text{tree}} - \delta^{13}C_{\text{atm}} + 1) / (b - a) \right],$$
(3)

where  $\delta^{13}C_{tree}$  and  $\delta^{13}C_{atm}$  are the tree and atmospheric C isotope compositions, respectively, *a* is the diffusion fractionation across the boundary layer and the stomata (+4.4‰) and *b* is the Rubisco enzymatic biologic fractionation (+27.0‰). Estimated values of  $\delta^{13}C_{atm}$  were obtained from [57].

#### 2.4. Nutrient Analyses

Xylem nutrient concentrations were measured in the same milled wood segments than isotope composition, that is, 5-year pools grouped by tree-vigor classes for the period 1915–2018. Phosphorus (P), potassium (K), calcium (Ca), manganese (Mn), magnesium (Mg), aluminum (Al) and zinc (Zn) concentrations were measured by inductively coupled plasma optical emission spectrometry (ICP-OES; Thermo Elemental Iris Intrepid II XDL, Franklin, MA, USA) after a microwave-assisted digestion with HNO<sub>2</sub>:H<sub>2</sub>O<sub>2</sub> (4:1, v/v). Wood nitrogen (N) mass-based concentration (%) was measured with an elemental analyzer (Elementar VarioMAX N/CM, Hanau, Germany). In addition, seven meaningful molar ratios were calculated (N:P, N:K, P:Mn, K:Ca, Ca:Mn, Ca:Al, and Mg:Mn), as they have been proposed to better reflect the environmental changes than simple elemental concentrations [39,58].

## 2.5. Data Analyses

Trends in climatic variables, isotopes and nutrient composition were assessed using the Kendall  $\tau$  statistic. Breakpoint analysis was applied to detect structural changes in BAI series corresponding to the onset of growth decline in each tree-vigor class. Then, trends were calculated for the periods before and after break point date. Differences between L and D trees regarding DBH, isotopic composition, elemental concentrations and molar ratios were assessed for the whole series using the Mann–Whitney *U* test. We used the Wilcoxon rank-sum test to check if the changes through time of BAI differed between tree-vigor classes.

In order to avoid spurious relationships that may arise from strong temporal dependence, we detrended BAI and isotope composition series by means of the Ensemble Empirical Mode Decomposition (EEMD) technique. EEMD is an empirical but highly efficient and adaptive method for processing non-linear and non-stationary signals [59]. This methodology has been successfully applied to ecological time series including dendrochronological [60,61], phenological [62], and nutrient content and stoichiometric ratios [63] data. The EEMD decomposes a time series into a small number of oscillatory components, from high to low frequencies, and a residual (trend) component using a splinebased iterative shifting process. Detailed description of EEMD can be found in [59,64,65]. Detrended BAI and isotopic signal series were constructed by summing the three and two first oscillatory components, respectively. Finally, we obtained mean detrended BAI chronologies of L and D trees using a bi-weight robust mean.

Climate-growth relationships were evaluated through bootstrapped correlations between monthly mean temperature and precipitation and mean detrended BAI chronologies for the period 1931–2018, which corresponds to the period with available instrumental records. We also computed correlations between detrended iWUE and  $\delta^{18}$ O and climate variables averaged in 5-year periods. The window of analysis spanned from previous April to April of the year of tree-ring formation.

We performed a non-metric multidimensional scaling (NMDS, [66]) with Euclidean dissimilarity for nutrient concentrations (N, P, K, Ca, Mn, Mg, Al and Zn) and stoichiometric ratios (N:P, N:K, P:Mn, K:Ca, Ca:Mn, Ca:Al and Mg:Mn) to obtain a multidimensional overview of the long-term nutritional status and nutrient imbalances, respectively, of the two vigor classes. Significant differences between L and D trees were tested with permutational multivariate analysis of variance (PERMANOVA, [67]). The NMDS axes were correlated with detrended iWUE and  $\delta^{18}$ O and projected into the ordination space. Additionally, relationships between detrended isotopic signals and individual nutrient concentrations and stoichiometric ratios were assessed using pairwise Spearman rank correlations ( $\rho$ ).

To integrate the effect of nutrient status and isotopic signals on tree-level BAI, we fitted linear mixed-effects models (LMMs, [68]). A different LMM was fitted for each vigor class. Fixed effects were DBH, iWUE,  $\delta^{18}$ O and scores of the NMDS1 and NMDS2 of ordinations of nutrient concentrations and stoichiometric ratios. A random intercept associated with tree and a first-order autocorrelation structure were included in the models [69]. BAI was log-transformed (log (x + 1)) prior to analyses in order to achieve normality assumptions. Fixed effects were standardized prior to model fitting to enable direct comparison among coefficients of predictors. The goodness-of-fit of the models was evaluated with the coefficient of determination for GLMMs ( $R_GLMM^2$ ) proposed by Nakagawa et al. [70]. Marginal R<sup>2</sup> ( $R_GLMM(m)^2$ ) accounts for the proportion of variance explained by the fixed effects, and conditional R<sup>2</sup> ( $R_GLMM(c)^2$ ) is interpreted as the variance explained by the entire model, that is, fixed plus random effects.

We performed all statistical analyses in R (version 3.6.3, Vienna, Austria) [49]. Processing of dendrochronological data was conducted with the R package *dplR* [71]. Trends in temporal series and break points analysis were assessed with the R packages *Kendall* [72] and *strucchange* [73], respectively. EMMD detrending was conducted using the R-package *Rlibeemd* [74]. Climate-growth associations were evaluated using the *treeclim* package [75]. Multivariate analysis was performed with the R package *vegan* [76]. The R packages *nlme* [77] and *MuMIn* [78] were used to fit LMMs and calculate goodness of fit, respectively.

#### 3. Results

# 3.1. Patterns and Climate-Drivers of Growth and Tree-Ring Isotope Signature

We found significant temporal trends of increasing growing season temperature ( $\tau = 0.275$ , p < 0.001) and decreasing precipitation ( $\tau = -0.162$ , p = 0.026) along the 1931–2018 period (Figure A1). Extreme drought events were identified during growing seasons of 1954, 1979, 1999, 2008 and 2015–2016 (dates are referred as the end year of the growing season), based on SPEI values of the growing season below the -1.5 threshold.

We did not find significant differences in DBH between L trees ( $58.6 \pm 9.82$  cm) and D trees ( $55.8 \pm 15.36$  cm) (U = 118, p = 0.533). Mortality dates of D trees ranged from 1998 to 2016, being 1999 the most repeated year of death (n = 8 trees). Radial growth of the two tree-vigor classes ran closely until the early 1960s, when L trees displayed an increase in BAI not shown by D trees (Figure 1). Nevertheless, we found a common trend break point in both vigor classes corresponding to 1962–1963. No significant trends in BAI were

found neither in L trees ( $\tau = 0.087$ , p = 0.311) nor in D trees ( $\tau = -0.069$ , p = 0.421) before 1963, but afterwards trends became significantly negative in both vigor classes (L trees:  $\tau = -0.546$ , p < 0.001; D trees:  $\tau = -0.692$ , p < 0.001). A second break point was detected in mean BAI series of L trees in 1998–1999, when the prior negative trend turned positive ( $\tau = 0.463$ , p = 0.005). BAI significantly differed between vigor classes during the 1992–1995 period and from 2006 onwards.



**Figure 1.** Basal area increment (BAI) of living (L; green) and dead (D; red) *Nothofagus dombeyi* trees. Solid lines represent the mean BAI for each tree-vigor class and shaded areas around them the standard error of the mean. Dash lines show significant (p < 0.05) trends in mean BAI before and after detected break point dates. The grey filled areas indicate the periods when BAI of L and D trees significantly (p < 0.05) differed according to Wilcoxon rank-sum tests.

Considering the whole period together, D trees showed significant higher iWUE (U = 317, p = 0.016) and lower  $\delta^{18}$ O (U = 120, p = 0.048) than L trees (Figure 2). Both vigor classes presented a positive temporal trend of iWUE that was slightly more pronounced in D trees.  $\delta^{18}$ O decreased along the study period in both vigor classes. iWUE and  $\delta^{18}$ O were not correlated in L trees and negatively correlated in D trees (Figure 3a), which suggests that tree-ring oxygen isotopes could reflect variability of water sources  $\delta^{18}$ O. Indeed, tree-ring  $\delta^{18}$ O of D trees was positively correlated with oxygen isotope signature of June precipitation (Figure 3b).



**Figure 2.** (a) Intrinsic water-use efficiency (iWUE) and (b) oxygen isotopic ratios ( $\delta^{18}$ O) in tree rings of living trees (L; green symbols and lines) and dead trees (D; red symbols and lines) of *Nothofagus dombeyi* along the 1910–2018 period. Symbols represent values of five consecutive rings pooled at vigor class level and lines indicate significant trends based on the Kendall  $\tau$  statistic. Kendall  $\tau$  statistic and associated *p*-value (\* *p* < 0.05; \*\* *p* < 0.01; \*\*\* *p* < 0.001) is indicated in each graph for both vigor classes.

The correlation between EEMD detrended BAI chronologies and monthly climatic variables for 1931–2018 identified a significant negative effect of high temperatures during

February (summer) on growth of both tree-vigor classes. The growth of both vigor classes positively responded to previous April and October precipitation, which corresponded to previous autumn and current spring, respectively. In addition, D trees showed a positive correlation with previous August (winter) precipitation (Figure 4a). The effects of climatic conditions on iWUE differed between vigor classes. It was positive for summer temperature in D trees and negative for spring precipitation in L trees (Figure 4b). Finally,  $\delta^{18}$ O of L trees showed significant positive correlations with previous winter temperature. Meanwhile, October precipitation had a positive effect on  $\delta^{18}$ O of D trees (Figure 4c).



**Figure 3.** Relationships between tree ring oxygen isotopic ratio ( $\delta^{18}$ O) and (**a**) intrinsic water-use efficiency (iWUE), and (**b**) rainfall  $\delta^{18}$ O in June for the period 1964–2017 in living trees (L; green symbols) and dead trees (D; red lines and symbols) of *Nothofagus dombeyi*. Values correspond to groups of five consecutive rings pooled at vigor-class level. Spearman  $\rho$  statistic and associated *p*-value (\* *p* < 0.05) are indicated in each graph for correlations at vigor class level. Significant correlations are highlighted with lines.



**Figure 4.** Bootstrapped correlations between monthly mean temperature (left) and precipitation (right) and basal area increment (BAI) (**a**), intrinsic water-use efficiency (iWUE) (**b**) and oxygen isotope composition ( $\delta^{18}$ O) (**c**) of tree rings of living (green symbols) and dead (red symbols) *Nothofagus dombeyi* trees. BAI, iWUE and  $\delta^{18}$ O were detrended using ensemble empirical mode decomposition (EEMD) technique prior analysis. Error bars are 95%-confidence intervals and significant and non-significant effects are represented by solid and dash bars, respectively. Grey shaded areas indicate growing season of the year of ring formation.

# 3.2. Long-Term Reconstruction of Tree Nutrient Status

Temporal evolution of nutrient concentrations in tree rings followed specific patterns according to each nutrient (Figure 5). K was the only element showing a concentration increasing along the period 1915–2018 for both tree-vigor classes. Concentrations of P, Ca, Al, Mg, and Mn showed a positive temporal trend only in D trees, whereas N increased in L trees (Table 1). In the case of stoichiometric ratios, N:K and K:Ca showed negative and positive trends, respectively, in both vigor classes. D trees showed increasing P:Mn, Ca:Mn and Mg:Mn ratios over time (Table 1).



**Figure 5.** Temporal evolution of nutrient concentration in tree-rings of living (green symbols and lines) and dead (red symbols and lines) *Nothofagus dombeyi* trees including: (**a**) nitrogen (N), (**b**) phosphorus (P), (**c**) potassium (K), (**d**) calcium (Ca), (**e**) aluminum (Al), (**f**) magnesium (Mg), (**g**) manganese (Mn), and (**h**) zinc (Zn). Symbols represent values of five consecutive rings pooled at vigor class level and lines indicate significant trends based on the Kendall  $\tau$  statistic (see Table 1).

Wood-nutrient concentrations for the whole series (1915–2018) differed between treevigor classes for P, Al and Zn, which showed higher values in L trees. Regarding stoichiometric ratios, we also found differences between vigor classes for N:P, Ca:Mn, Ca:Al, Mg:Mn (L trees < D trees), and P:Mn (L trees > D trees) (Table 1).

Ordination analyses helped us to characterize the nutritional status and nutrient imbalances of the target trees (Figure 6). Regarding nutrient concentrations, the first axis (NMDS1) was positively related to K, P and Al, and negatively to Ca, Mg and N. The second axis (NMDS2) encompassed most of the variability of Zn. Considering nutrient concentrations, both vigor classes significantly differed (PERMANOVA test F = 6.485, p = 0.001). In agreement with individual comparisons, L trees were associated with higher concentrations of K, P, Zn and Al, whereas D trees showed higher Ca and Mg (Figure 6a). The NMDS1 derived from the ordination of stoichiometric ratios was associated with P-and K-related ratios (K:Ca with positive loadings, N:K and N:P with negative loadings).

Meanwhile, the NMDS2 mainly explained the variability of Ca-related ratios (K:Ca with positive loadings, Ca:Mn with negative loadings). We found significant differences in the ordination space of nutrient ratios between vigor classes (PERMANOVA test F = 4.628, p = 0.005). L trees scores were located towards higher values of K:Ca, and D trees showed greater N:P and Ca:Mn (Figure 6b).

**Table 1.** Mean ( $\pm$  standard errors) of nutrient concentrations and molar stoichiometric ratios. Different letters indicate significant differences (p < 0.05) between living (L) and dead trees (D) according to Mann–Whitney U tests. Trends are based on the Kendall  $\tau$  statistic and significance in shown with asterisks (\* p < 0.05; \*\* p < 0.01; \*\*\* p < 0.001).

		Living Trees		Dead Trees	
		$\mathbf{Mean} \pm \mathbf{SE}$	Trend	$\mathbf{Mean} \pm \mathbf{SE}$	Trend
Nutrient	$N (mg g^{-1})$	$1.6\pm0.01~\mathrm{a}$	0.438 **	$1.6\pm0.01~\mathrm{a}$	0.076
concentrations	$P(\mu g g^{-1})$	$474.4\pm22.19$ a	0.124	$376.9\pm31.9\mathrm{b}$	0.638 ***
	K ( $\mu g g^{-1}$ )	$243.8 \pm 75.73$ a	0.638 ***	$114.8\pm30.09~\mathrm{a}$	0.829 ***
	Ca ( $\mu g g^{-1}$ )	$546.1 \pm 26.04$ a	-0.076	$627.6 \pm 94.05$ a	0.600 ***
	Al $(\mu g g^{-1})$	$15.4\pm2.51~\mathrm{a}$	0.200	$7.2\pm0.94\mathrm{b}$	0.543 ***
	Mg ( $\mu g g^{-1}$ )	$331.5 \pm 16.59$ a	0.029	$346.1 \pm 23.56$ a	0.667 ***
	$Mn (\mu g g^{-1})$	$9.9\pm0.73~\mathrm{a}$	0.067	$8.7\pm0.75$ a	0.600 ***
	$Zn (\mu g g^{-1})$	$18.3\pm2.42$ a	-0.200	$6.5\pm0.33$ b	0.133
Stoichiometric	N:P	$7.7\pm0.43$ a	0.143	$11.0\pm1.17~\mathrm{b}$	-0.328
ratios	N:K	$198.2\pm77.08~\mathrm{a}$	-0.619 ***	$318.3\pm94.62\mathrm{b}$	-0.790 ***
	P:Mn	$89.3 \pm 3.96 \text{ a}$	0.114	$77.6\pm4.38\mathrm{b}$	0.448 **
	K:Ca	$0.4\pm0.12~\mathrm{a}$	0.762 ***	$0.2\pm0.03~\mathrm{a}$	0.733 ***
	Ca:Mn	$78.6 \pm 2.73$ a	-0.133	$94.5\pm4.44~\mathrm{b}$	0.390 *
	Ca:Al	$34.9\pm4.06~\mathrm{a}$	-0.286	$62.1\pm4.06~\mathrm{b}$	-0.067
	Mg:Mn	$79.9\pm3.98~\mathrm{a}$	0.095	$91.9\pm3.51~\text{b}$	0.371 *



**Figure 6.** Non-metric multidimensional scaling (NMDS) biplots of nutrient concentrations (**a**) and stoichiometric molar ratios (**b**) of living (L; green symbols and areas) and dead (D; red symbols and areas) trees. Points represent the values of each group of five consecutive rings for the period 1915–2018. Shaded areas represent the centroid of the values for each tree-vigor class. Black arrows indicate loadings of nutrient concentrations (**a**) and molar ratios (**b**) in NMDS axes; those very close to the point (0,0) are not shown for the sake of clarity. The correlation between intrinsic water-use efficiency (iWUE) and oxygen isotope composition ( $\delta^{18}$ O) and NMDS axes was projected in the ordination diagram and represented with blue arrows. iWUE and  $\delta^{18}$ O were detrended using ensemble empirical mode decomposition (EEMD) technique prior analysis.

# 3.3. Integration of Growth, Isotopic Signals and Nutrient Status

Pairwise Spearman rank correlations between detrended isotope signatures and nutrients in tree rings revealed a lack of significant associations between iWUE and nutrient concentrations or stoichiometric ratios in L trees (Table 2). In contrast, iWUE of D trees showed a significant positive correlation with P, K, Al, Mn, and Zn concentrations and K:Ca ratio, that became negative in the case of N:P and N:K ratios. Similarly,  $\delta^{18}$ O showed a stronger response to nutrient variables in D trees than in L trees. Concentration of P, K and Mn was positively correlated with  $\delta^{18}$ O in D trees, while the correlation was negative with Ca. Both vigor classes showed significant responses to N:P ratio, although it was negative for D trees and positive (marginally significant) for L trees. Again, marginally significant correlations were found between tree-ring  $\delta^{18}$ O and N:K (negative) and K:Ca (positive) in D trees (Table 2). Therefore, our results suggest a tighter relationship between isotopic signals and nutrients in D trees since we found more frequent significant correlations in this vigor class.

**Table 2.** Pairwise correlations between detrended isotopic signals (intrinsic water-use efficiency (iWUE) and oxygen isotope composition ( $\delta^{18}$ O)) and nutrient concentrations and stochiometric ratios. The Spearman  $\rho$  statistic and associated *p*-value (<sup>+</sup> *p* < 0.1; \* *p* < 0.05; \*\* *p* < 0.01; \*\*\* *p* < 0.001) is shown for pair of variables for living (L) and dead (D) trees.

		iWUE		δ1	<sup>8</sup> O
		L Trees	D Trees	L Trees	D Trees
Nutrient concentrations	N (mg g <sup>-1</sup> ) P ( $\mu$ g g <sup>-1</sup> ) K ( $\mu$ g g <sup>-1</sup> ) Ca ( $\mu$ g g <sup>-1</sup> )	$0.131 \\ -0.126 \\ -0.139 \\ -0.100$	-0.232 0.490 * 0.548 * 0.317	$0.162 \\ -0.265 \\ -0.190 \\ 0.347$	0.068 0.473 * 0.458 * 0.432 *
	Al ( $\mu g g^{-1}$ )	-0.074	0.713 ***	0.293	0.311
	Mg (μg g <sup>-1</sup> ) Mn (μg g <sup>-1</sup> ) Zn (μg g <sup>-1</sup> )	-0.016 0.017 0.164	-0.016 0.457 * 0.576 **	-0.321 0.009 0.288	0.579 * 0.473 * 0.144
Stoichiometric ratios	N:P N:K P:Mn K:Ca Ca:Mn Ca:Al Mg:Mn	$\begin{array}{c} 0.138\\ 0.095\\ -0.121\\ -0.166\\ -0.306\\ 0.134\\ -0.029\end{array}$	$\begin{array}{r} -0.473 \\ -0.541 \\ * \\ 0.221 \\ 0.468 \\ * \\ 0.329 \\ -0.343 \\ -0.029 \end{array}$	$\begin{array}{r} 0.362 \\ + \\ 0.213 \\ - 0.321 \\ 0.243 \\ - 0.140 \\ 0.282 \\ - 0.345 \end{array}$	$\begin{array}{r} -0.423 * \\ -0.394 + \\ 0.165 \\ 0.393 + \\ -0.351 \\ 0.034 \\ 0.364 \end{array}$

We only found significant correlations between detrended isotopic signals and NMDS axes in the nutrient concentration ordination. Both iWUE (r = 0.183; p = 0.039) and  $\delta^{18}$ O (r = 0.163; p = 0.045) were positively correlated with NMDS1, so higher iWUE and  $\delta^{18}$ O values were associated to greater K, P and Al concentrations and lower Ca concentrations (Figure 6a).

The higher the DBH, the higher the radial growth of both vigor classes (Table 3). BAI of D trees responded negatively to increasing iWUE (Figure 7a), whereas  $\delta^{18}$ O did not show significant effects on BAI of any vigor class. Our results show that stoichiometric ratios showed stronger effect on growth of L trees, and that nutrient concentrations were more important for modulating growth of D trees. Higher concentrations of K, P and Al and lower concentrations of Ca (i.e., higher NMDS1 scores of nutrient concentrations ordination) resulted in reductions of growth of D trees (Figure 7b). Meanwhile, growth of L trees positively responded to increasing K:Ca ratio, whereas decreasing N:P, N:K, and Ca:Mn had a negative effect on BAI of L trees (Figure 7c).

**Table 3.** Linear mixed-effects models (LMMs) characterizing basal area increment (BAI) at tree level. The table shows coefficient estimates  $\pm$  standard errors (Coef.  $\pm$  SE) and the *F* statistic with its associated *p*-value (\* *p* < 0.05; \*\* *p* < 0.01; \*\*\* *p* < 0.001). Fixed effects included diameter at breast height (DBH), tree age, intrinsic water-use efficiency (iWUE), oxygen isotope ratio ( $\delta^{18}$ O) and scores of NMDS1 and NMDS2 of ordination analysis of nutrient concentration and stoichiometric ratios. The explanatory power of each model is indicated by *R\_GLMM(m)*<sup>2</sup> (proportion of variance explained by the fixed factors) and *R\_GLMM(c)*<sup>2</sup> (proportion of variance explained by the entire model) [76].

Variables	Living Trees		Dead Trees		
	Coef. $\pm$ SE	F-Value	$\textbf{Coef.} \pm \textbf{SE}$	F-Value	
DBH	$0.350\pm0.038$	72.868 ***	$0.499 \pm 0.050$	78.751 ***	
iWUE	$-0.035 \pm 0.046$	0.763	$-0.117 \pm 0.060$	3.950 *	
$\delta^{18}O$	$0.051\pm0.038$	2.569	$0.004 \pm 0.028$	0.025	
NMDS1 <sub>Concentration</sub>	$-0.280 \pm 0.161$	0.328	$-0.265 \pm 0.183$	29.072 ***	
NMDS2 <sub>Concentration</sub>	$-0.094 \pm 0.059$	0.208	$0.101\pm0.161$	0.256	
NMDS1 <sub>Ratio</sub>	$0.397\pm0.075$	16.830 ***	$0.059\pm0.045$	2.991	
NMDS2 <sub>Ratio</sub>	$0.143\pm0.039$	7.986 **	$0.016\pm0.036$	0.505	
$R_GLMM(m)^2$	0.319		0.482		
$R_GLMM(c)^2$	0.387		0.523		



**Figure 7.** Relationships between basal area increment (BAI) and (**a**) intrinsic water-use efficiency (iWUE), (**b**) NMDS1 of nutrient concentration ordination and (**c**) NMDS1 of stoichiometric ratios ordination in living (green symbols and lines) and dead (red symbols and lines) *Nothofagus dombeyi* trees. Solid lines represent predicted relationships by linear mixed-effects models (see Table 3) and dash lines are 95% confidence intervals.

# 4. Discussion

We found that D trees showed a declining growth trend after the early-1960s (Figure 1). This agrees with [42], who identified this date as the onset of dieback after analyzing these and other populations of *Nothofagus dombeyi*. The long-lasting growth decline prior to death has been observed as a typical response during drought-induced dieback in *Nothofagus* species [79], in which the cumulative effect of successive drought events reduces resilience and increases the risk of mortality [24,80]. In contrast, L trees showed a sharp increase in growth rate in the early-1960s and again in 1999. These growth releases were likely explained by reductions in competition pressure resulting from extensive mortality events of conspecific individuals triggered by severe droughts after 1962–1963 and 1998–1999 severe drought events [42].

Climate-growth relationships supported the evidence for drought sensitivity of *N. dombeyi* trees [45]. Growth was enhanced by wet spring and cool summer conditions, and also by wet conditions during prior autumn (Figure 4a). Contrary to other species showing distinct climate sensitivities between declining and non-declining trees due to contrasting water-use strategies (e.g., [81]), we did not find substantial differences in growth response to climatic conditions between L and D trees. In contrast, D trees showed higher iWUE (Figure 2a) and higher responsiveness to summer temperatures (Figure 4b) and steeper increases over the last decades, suggesting more severe drought stress in this tree class [82]. These results are consistent with previous studies of drought-induced dieback on broadleaf [83,84] and conifer species [85]. However, the opposite pattern has been also found [86,87], evidencing the insufficient information provided by tree-ring carbon isotopes alone to elucidate the mechanisms prompting dieback.

The relative contribution of *A* and  $g_5$  to carbon isotope signatures of tree-rings has been commonly assessed using  $\delta^{18}$ O data [26]. However absent (L trees) or even negative (D trees) iWUE- $\delta^{18}$ O relationships revealed the uncoupling between leaf-level processes and oxygen isotope ratios in tree-rings (Figure 3a). In addition, tree-ring  $\delta^{18}$ O of D trees was positively correlated with the  $\delta^{18}$ O of winter precipitation (Figure 3b). Caution is needed when interpreting this result, owing to the large distance and distinct climatic conditions of Puerto Montt, located in a very wet region in the Chilean Pacific. In any case, altogether this study contributes to the growing evidence supporting a predominant effect of the variability in source-water  $\delta^{18}$ O on tree-ring and leaf isotopic signals in many ecosystems [21,34,36,81,88].

The approximate water uptake depth of trees can be inferred from the  $\delta^{18}$ O signals imprinted on tree ring cellulose by the isotopic composition of their xylem water (not measured in the present study), which in turn reflects that of the soil/bedrock water sources. This interpretation is based on well-established knowledge of the steep vertical gradients in evaporative isotopic enrichment of soil water that develop in drying soils during prolonged rainless periods (with strong isotopic enrichment near the surface that exponentially decreases with depth; [89]). The lower tree-ring  $\delta^{18}$ O signature in D trees suggested their utilization of deeper water sources for water uptake (Figure 2b) since deep water pools usually show more depleted  $\delta^{18}$ O values than topsoil water due to their lower exposure to evaporative isotopic enrichment in deep layers [35]. Climate- $\delta^{18}$ O correlations further support this hypothesis (Figure 4c). The heavier reliance of L trees on shallower soil layers is confirmed, at least in part, by the correlation of tree-ring  $\delta^{18}$ O with temperature because topsoil water is more exposed to evapotranspiration [90]. On the contrary, positive associations in D trees between tree-ring  $\delta^{18}$ O and early spring (October) rainfall amount and  $\delta^{18}O$  variations of June precipitation were consistent with extraction of water from deeper subsoil water pools, which are usually replenished in winter [91]. As the strong climate-warming trend in the study area causes enrichment in precipitation  $\delta^{18}$ O values over time [92], declining trends in tree-ring  $\delta^{18}$ O of both vigor classes may be related to increasing depth of water uptake due to reductions in topsoil-water availability over time, in line with those reported in seasonally dry Mediterranean ecosystems [37].

The exploitation of water reserves stored in deeper soil/bedrock layers by D trees may imply disadvantages in terms of nutrient availability and uptake, considering that most nutrients are more abundant in shallow soil layers [22]. This is evidenced by the frequent positive correlations in D trees between tree-ring  $\delta^{18}$ O and nutrient concentrations (Table 2). Therefore, our results fit well within the conceptual model recently put forward by Querejeta et al. [93], who postulated that heavier utilization of subsoil/bedrock water induced by topsoil desiccation leads to deterioration of vegetation mineral nutrition due to the vertical decoupling of water and nutrient availability in soil/bedrock profiles. Differences between vigor classes may arise from the location of D trees on exposed microsites and/or sites with shallow soils with low water-holding capacity, as has been proposed before for *N. dombeyi* [42] and Mediterranean *Quercus* species [86,94] undergoing dieback. Alternatively, higher drought-induced defoliation due to the total loss of leaf conductance and dehydration [44], and tighter stomatal control (higher iWUE) of D trees may reduce transpiration-driven water flux, thus impairing nutrient uptake [20]. Such divergences in foliar safety margins and stomatal response between vigor classes can lie in a genetic background and resulting physiological adjustments, in view of the dendrogenomic differences between *N. dombeyi* individuals inhabiting similar microenvironmental conditions but showing different degrees of crown defoliation [95].

Whatever the mechanism, lower long-term concentrations and imbalances of essential nutrients such as P, K and Zn (Table 1; Figures 5 and 6) reflect poorer long-term nutritional status and the nutrient limitations of D trees. Such nutritional impairment of D trees likely exacerbated and amplified the negative effects of drought on physiological functioning and productivity and increased the risk of tree mortality [14]. Positive correlations between iWUE and P, K, and associated stoichiometric ratios in D trees (Table 2; Figure 6) indicate a strong nutrient limitation of A and/or  $g_s$ , consistent with the relevant role of these nutrients on photosynthesis, transpiration and hydraulic conductance [17,18]. Tree-water transport has been found to be regulated by changes in xylem ion concentrations [96]. Taking into account the prominent role of K in the hydraulic function [97], K deficiencies probably contribute to hydraulic failure during droughts. The acquisition of nutrients would also be impaired by the mortality of fine roots and mycorrhizal fungi caused by topsoil desiccation under prolonged drought [98]. In addition, droughts have a stronger impact on the availability and uptake of nutrients with limited mobility and diffusion rates in soil (e.g., P, K and Zn), compared to the highly mobile N [20]. The reduced photosynthetic capacity of highly defoliated trees showing poorly functional leaves due to low nutrient content could foster the growth decline patterns found in D trees.

The unexpected increases of tree-ring P and K concentrations in D trees over the last decades (Figure 5) may have several explanations. The increase of K concentrations in the sapwood of L trees was much steeper than in D trees (Figure 5c). Firstly, nutrient remobilization can increase K concentrations from inner to outer rings to compensate for losses in hydraulic conductivity [97] with ongoing drought and the deterioration of nutrition. Since remobilization of nutrients has been proposed to be source-driven [99], it would reflect a greater need for nutrient remobilization from older to younger rings in the more nutrient-limited D trees. Secondly, the decreasing growth rate of D trees progressively reduced nutrient dilution effects over the last decades [100]. Thirdly, greater exudation of organic acids and carboxylates by D tree's roots for improving mineralization and mobilization of inorganic and organic P may have also contributed to this pattern [101]. Organic acids and carboxylates exudation by roots also enhances the availability of other micronutrients such as Mn, thereby increasing its concentration in plant tissues [102] together with an accumulation of Ca [103]. This interpretation is further supported by the sharp recent increase of Ca and Mn concentrations and Ca:Mn ratio (Figure 5) and negative correlation between Ca concentration and  $\delta^{18}$ O observed in D trees (Figure 6). The elevated carbon cost of releasing organic acids and carboxylates to rhizosphere soil likely contributed to carbon depletion and loss of vitality in D trees. Nutrient imbalances regarding tree-ring Ca and Mn have been consistently related to dieback and growth decline [39,40], thus supporting the usefulness of Ca:Mn ratio as early warning of forest dieback in response to drought. However, different results between this and previous studies highlight the need for further research to fully understand the role of long-term nutrient imbalances on tree mortality.

Finally, tree growth results from the integrative effect of multiple physiological processes and environmental conditions. Although we did not find differences in DBH between vigor classes, and the BAI of both L and D trees positively responded to increasing size (Table 3). This size effect agrees with previous studies reporting the smaller size (DBH and height) of declining over non-declining trees in Quercus species experiencing drought-dieback [82,94,104]. The negative effect of iWUE on growth was observed only in D trees (Figure 7a), which supports the the greater drought-stress experienced by this vigor class. The more water-use conservative strategy of D trees could be the result of stronger reduction in soil moisture, related to lower-quality microsites that cause stomata closure to avoid hydraulic function impairment at the expense of reduced carbon uptake [105]. Meanwhile, the lack of significant effects of tree-ring  $\delta^{18}$ O on BAI can be related to the fact that it indicates variations on source-water  $\delta^{18}$ O instead of stomatal regulation, thus affecting nutrient availability and acquisition. Indeed, NMDS1 of nutrient concentrations (D trees; Figure 7b) and stoichiometric ratios (L trees; Figure 7c) showed large positive effects on tree growth. The counterintuitive negative effect of increasing K, P and Al concentrations on BAI of D trees is explained by the opposite trends that these variables experienced over the last decades, i.e., increases in P and K as a response to nutrient limitation (see explanation above) and ongoing growth decline.

# 5. Conclusions

This study provides insights into the underlying mechanisms that drove tree dieback and mortality of some individuals in a *N. dombeyi* population located at a dry edge of the species' distribution. The growth decline of dead trees started almost 40 years before death. The higher water-use efficiency and lower oxygen isotope composition of dead trees, and the uncoupling of the latter from leaf-level processes, pointed to their increased drought stress and water uptake from deeper soil layers. Considering the vertical decoupling between water and nutrient availability in soil/bedrock layers [93], dead trees experienced a long-term deterioration of nutritional status aggravating hydraulic and growth impairment triggered by successive droughts that could have played a critical role in dieback and mortality processes. In view of the outstanding patterns in Ca and Mn concentrations and related stoichiometric ratios, they may be useful early-warning signals of drought-induced dieback and impending tree death.

Author Contributions: Conceptualization, E.G.d.A., J.I.Q. and J.J.C.; methodology, E.G.d.A., M.L.S. and J.I.Q.; software, E.G.d.A.; validation, E.G.d.A.; formal analysis, E.G.d.A.; data curation, M.L.S., J.I.Q. and J.J.C.; writing—original draft preparation, E.G.d.A.; writing—review and editing, M.L.S., J.I.Q. and J.J.C.; funding acquisition, M.L.S. and J.J.C. All authors have read and agreed to the published version of the manuscript.

**Funding:** This research was funded by Ministerio de Ciencia y Tecnología (Spain), grant number RTI2018-096884-B-C31, and by Agencia Nacional de Promoción Científica y Tecnológica (Argentina), projects PICT 2017-3843 and CONICET fund PIP 2017-0484.

**Data Availability Statement:** Data will be made available upon reasonable request to the corresponding author.

**Acknowledgments:** The authors are thankful to Yamila Sasal for field work assistance. We thank Michele Colangelo for his help in the laboratory.

Conflicts of Interest: The authors declare no conflict of interest.

# Appendix A. Climatic Trends



**Figure A1.** Mean temperature (**a**), total precipitation (**b**) and Standardized Precipitation Evaporation Index (SPEI) (**c**) of the growing season (previous October–March) calculated with data obtained from Bariloche airport weather station. In (**a**) and (**b**) straight thick lines represent significant temporal trends. In (**c**) black dots indicate severe drought events under the -1.5 threshold.

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