

# Towards a common methodology for developing logistic tree mortality models based on ring-width data

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**Abstract.** Tree mortality is a key process shaping forest dynamics. Thus, there is a growing need for indicators of the likelihood of tree death. During the last decades, an increasing number of tree-ring based studies have aimed to derive growth–mortality functions, mostly using logistic models. The results of these studies, however, are difficult to compare and synthesize due to the diversity of approaches used for the sampling strategy (number and characteristics of alive and death observations), the type of explanatory growth variables included (level, trend, etc.), and the length of the time window (number of years preceding the alive/death observation) that maximized the discrimination ability of each growth variable. We assess the implications of key methodological decisions when developing tree-ring based growth–mortality relationships using logistic mixed-effects regression models. As examples, we use published tree-ring datasets from *Abies alba* (13 different sites), *Nothofagus dombeyi* (one site), and *Quercus petraea* (one site). Our approach is based on a constant sampling size and aims at (1) assessing the dependency of growth–mortality relationships on the statistical sampling scheme used, (2) determining the type of explanatory growth variables that should be considered, and (3) identifying the best length of the time window used to calculate them. The performance of tree-ring-based mortality models was reasonably high for all three species (area under the receiving operator characteristics curve, AUC > 0.7). Growth level variables were the most important predictors of mortality probability for two species (*A. alba*, *N. dombeyi*), while growth-trend variables need to be considered for *Q. petraea*. In addition, the length of the time window used to calculate each growth variable was highly uncertain and depended on the sampling scheme, as some growth–mortality relationships varied with tree age. The present study accounts for the main sampling-related biases to determine reliable species-specific growth–mortality relationships. Our results highlight the importance of using a sampling strategy that is consistent with the research question. Moving towards a common methodology for developing reliable growth–mortality relationships is an important step towards improving our understanding of tree mortality across species and its representation in dynamic vegetation models.

**Key words:** growth–mortality relationship; logistic model; sampling; survival; tree mortality; tree ring; *Abies alba*; *Nothofagus dombeyi*; *Quercus petraea*.

## INTRODUCTION

Accelerating rates of background tree mortality and related die-off events have recently been reported for

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many forest ecosystems and have been associated with the progressive increase in temperatures, drought, and the duration and frequency of extreme climatic events (e.g., van Mantgem et al. 2009, Allen et al. 2010). As this mortality trend is expected to continue due to global change (Collins et al. 2013), there is a growing demand for early warning signals that allow anticipating where

and when forest die-off is likely to occur (cf. Scheffer et al. 2012, Allen et al. 2015). The interest for indicators of tree mortality probability is large in both basic and applied science. First, they can improve our understanding of the causes and mechanisms that lead to tree mortality. Second, predicting which trees or populations are prone to die would help forest managers to design and implement plans to harvest them before their expected death or to preserve them, for example, by thinning to reduce the intensity of competition for resources such as light or water. Finally, this type of information is key for the development of more robust and reliable dynamic vegetation models whose projections depend critically on how tree mortality is modeled (e.g., Wyckoff and Clark 2002, Friend et al. 2014, Bircher et al. 2015).

Three approaches are commonly used to assess and simulate tree mortality probability. First, the spatio-temporal analysis of mortality patterns from ground surveys or remote sensing can be used to derive empirical relationships between mortality probabilities and indices that reflect environmental conditions such as drought or competition intensity (e.g., Bravo-Oviedo et al. 2006, Clifford et al. 2013, Das et al. 2013, Vilà-Cabrera et al. 2013, Dorman et al. 2015). This method has the advantage of providing large sample sizes and covering large environmental gradients. However, the temporal extent is limited (<50 yr), and this phenomenological approach provides few insights into the mechanisms underlying the mortality process itself. Second, physiological indicators such as changes in the plant hydraulic system (reduced water transport capacity due to xylem embolism, potentially causing hydraulic failure) or the carbon budget (decrease of non-structural carbohydrate concentrations potentially leading to carbon starvation) can be used to reveal a decrease in tree vitality and hence an increase in the likelihood of mortality (McDowell et al. 2013, Tague et al. 2013). Although this mechanistic approach is highly promising, the development of such indicators requires intensive, long-term tree monitoring involving measurements of the pools and fluxes of carbon and water at high temporal and structural resolution (i.e., in different organs), and even in this case the predictive ability of current mortality models is limited (McDowell et al. 2013). Moreover, this type of monitoring is very costly and can be implemented for a few trees at highly instrumented research sites only, thus making results for a broad suite of species rather unlikely. Third, focusing on tree radial growth is a promising alternative as it can be investigated easily for many individuals, has the advantage of covering long time periods, thus including the potential of considering lagged mortality (e.g., Bigler et al. 2007, Camarero et al. 2015), and is highly appropriate to reflect the spatio-temporal variability of environmental conditions and tree vitality (Dobbertin 2005). Indeed, cell growth and formation are among the first processes impacted by environmental stress (low temperature or decline in turgor due to drought; Palacio

et al. 2014, Lempereur et al. 2015), are highly sensitive to crown defoliation (Puri et al. 2015), and may be associated with non-structural carbohydrates reserves (Heres et al. 2014, but see Palacio et al. 2014). Radial growth data also show high potential to predict individual mortality from tree rings (e.g., Bigler and Bugmann 2004) or forest inventory data (e.g., Holzwarth et al. 2013). Forest inventories usually cover a wide range of tree species and large areas, but they almost always lack an annual resolution, whereas tree-ring data are available from a very large number of studies, each of which typically covers one to a few species growing at one to a few sites (e.g., Gillner et al. 2013, but see Kane and Kolb 2014).

During the last 15 years, an increasing number of tree-ring-based studies aimed at deriving growth–mortality functions (Appendix S1: Table S1.1). Most of them applied logistic regression models; however, they did not pay attention to several methodological issues that affect the structure and performance of such models. (1) The number of trees and the ratio between alive and death observations used to calibrate the model are usually chosen arbitrarily despite having a strong impact on model fit (Hartmann et al. 2007). (2) A range of statistical schemes may be employed to sample alive and death observations within a stand or even within a single tree, from the inclusion of all available observations (i.e., longitudinal data) to a specific focus on the years during which mortality occurred (i.e., cross-sectional data). (3) The type and combination of explanatory growth variables used may matter as well. They can be calculated using ring width (RW) or basal area increment (BAI) data, the latter depending less on tree diameter and being more closely related to biomass increment (Bowman et al. 2013). Growth level variables, i.e., averages of RW or BAI over certain time windows, are commonly used, as prolonged periods of low radial growth tend to be associated with an increased tree mortality probability (Pedersen 1998, Wyckoff and Clark 2000, Bigler and Bugmann 2004). However, as some slow-growing trees may survive for very long periods (e.g., old *Pinus longaeva*; Schulman 1958) and some fast-growing trees have been observed to die after a rapid growth decrease, models may also include growth trend variables that reflect the rate of change in mean growth within a given time period (see Bigler and Bugmann 2004). Similarly, as mortality probability is highly related with tree size (Das et al. 2007, Holzwarth et al. 2013), relative growth variables may be used to consider that small trees are relatively more productive than large ones given a similar diameter increment, or tree diameter (size) may be used directly. Other growth variables have been used as well, such as the interannual variability of growth, which was positively related with mortality probability in several studies (e.g., Suarez et al. 2004, Macalady and Bugmann 2014) or the temporal autocorrelation in the ring-width series (Camarero et al. 2015).

All these factors make the comparison of results among mortality studies and subsequent biological

interpretations difficult and hinder the feasibility of robust meta-analyses that would be highly useful to detect differences in tree growth patterns prior to mortality among sites, among species, and between the different factors triggering tree death.

We use data from published mortality studies using tree rings and systematically compare different methodologies for developing and assessing growth–mortality relationships. We aim to provide specific recommendations on good practices to predict tree mortality from radial growth data, depending on the objectives of the study and data availability. Specifically, we address the following questions, all of which affect the predictive capacity of the resulting statistical models.

- (1) What scheme should be used to sample alive and death observations?
- (2) What type of explanatory growth variables should be considered, and which specific metrics should be used?
- (3) How can the best length of the time window be determined to calculate the growth variables?
- (4) How should the best multi-variable logistic model be built, considering the collinearity between growth variables, and how can the impact of each of them on predicted mortality probability be best quantified?

## MATERIALS AND METHODS

### Tree-ring datasets

We re-analysed tree-ring datasets covering three species and several sites. In all cases, living and dead trees

were sampled at the same sites. Each species dataset differed in terms of the number of sites considered, number of trees sampled, ratio between living and dead trees, and distribution of the death events over time (Table 1). These datasets cover ecologically contrasting species including *Abies alba* Mill., *Nothofagus dombeyi* (Mirb.) Oerst., and *Quercus petraea* (Matt.) Liebl., which were selected to reflect the diversity of sampling strategies observed in the literature (Appendix S2).

*Abies alba* is a gymnosperm from European temperate mountain forests. Tree-ring data were taken from the studies of Bigler et al. (2004), Cailleret et al. (2014), Linares and Camarero (2012), and Lombardi et al. (2008). This dataset includes 195 dead and 321 living trees from 13 different sites with a wide range of DBH (diameter at breast height) and cambial age. Trees died between 1955 and 2008.

*Nothofagus dombeyi* is an evergreen angiosperm from South America with diffuse-porous wood. The trees were cored and ring widths measured by Suarez et al. (2004). The sampling size is relatively low, as only 43 dead and living trees growing at one site were analyzed, but the variability in tree size and age was high; trees died in a single event in 1998.

Finally, *Quercus petraea* is a deciduous angiosperm with ring-porous wood, dominant in dry European temperate forests. Tree-ring data were available from one site (I. Mészáros, unpublished data). In this case, the number of dead trees sampled is low compared to the living ones (25 dead and 194 living trees), and their DBH was not measured. Mortality occurred over a relatively long period, extending from 1953 to 2000.

At every site, tree-ring widths had been measured to the nearest 0.01 mm on increment cores taken at breast

TABLE 1. Main characteristics of the datasets used to assess growth–mortality relationships. The ranges in DBH and cambial age (defined at coring height; 1.3 m) are 95% confidence intervals.

Species	Sites	Article	Formation year of last ring	<i>N</i> dead trees	<i>N</i> living trees	DBH range (cm)	Cambial age range	Period of mortality
<i>Abies alba</i>	Bistra	Bigler et al. (2004)	2001	2	17	38–59	100–167	1988–2001
	Ravnik			16	16	27–57	73–189	
	Issole2	Cailleret et al. (2014)	2009	8	19	15–38	66–166	1998–2008
	Ventoux_Dvx1			8	12	18–40	53–95	
	Ventoux_Dvx2			11	19	19–45	51–87	
	Ventoux_Dvx3			10	15	16–31	39–113	
	Vesubie3			9	17	19–56	81–112	
	Vesubie6			1	12	31–76	135–184	
	Ventoux_TC	Linares and Camarero (2012)	2000	69	158	15–50	46–168	1996–2000
	Paco_Ezpela_High			3	7	28–47	73–116	
Paco_Ezpela_Low	5			7	38–51	92–154		
Lopet	2			7	26–51	79–128		
Canalicchio	Lombardi et al. (2008)	2005	51	15	10–71	46–163	1955–2002	
<i>Nothofagus dombeyi</i>	Cerro_Otto	Suarez et al. (2004)	1997	43	43	11–57	21–124	1998
<i>Quercus petraea</i>	Sikfokut	Meszaros, unpublished manuscript	2012	25	194	22–41†	55–104	1953–2009

†The DBH of dead *Quercus petraea* trees was not measured.

height (1.3 m). When a core did not reach the pith, the total missing width and the number of missing rings were estimated by interpolating the distance to the pith using the curvature of the innermost rings of the sample, which allowed us to estimate tree cambial age (at breast height). DBH data were available for two species only (*A. alba* and *N. dombeyi*). Annual DBH values were reconstructed for these species by subtracting DBH measured at the time of coring by the ring width produced earlier (from outside to inside; annual DBH values were rescaled in case of negative DBH calculated at age = 1).

*Modeling approach*

Generalized linear models with binomial error distribution (i.e., logistic regression) were used to predict the survival probability of a tree *i* at time *t*, hereafter labeled  $\Pr(Y_{i,t} = 1)$ , and to analyze the corresponding growth–mortality relationships (Bigler and Bugmann 2004). Logistic regression models with fixed effects were used for *N. dombeyi*, as trees were growing at one site and died on the same date

$$\log \left[ \frac{\Pr(Y_{i,t} = 1)}{(1 - \Pr(Y_{i,t} = 1))} \right] = \alpha_0 + \beta_0 * G_{i,t,p} \tag{1}$$

where  $\Pr(Y_{i,t})$  follows a binomial distribution with  $Y_{i,t} = 0$  indicating that tree *i* is dead at time *t*, while  $Y_{i,t} = 1$  indicates that the tree is alive;  $\alpha_0$  is the intercept, and  $\beta_0$  is the coefficient for the growth variable calculated over a time window of length *p* ( $G_{i,t,p}$ ; *p* for period).

When the tree-ring dataset included multiple sites and/or the mortality events occurred during several years (i.e., *A. alba* and *Q. petraea*), logistic regression models with mixed effects were used. The species-specific growth–mortality relationship was provided by the fixed effects of the model. To consider that growth–mortality relationships may change among sites due to the different relative contributions of environmental stress and biotic attacks, random effects were estimated for the growth variable (i.e., the slope) with study site as the grouping variable. In order to remove the potential impact of annual disturbances that can modify tree survival at a given site irrespective of growth (e.g., windthrow), random effects were also estimated for the intercept in groups of years nested in each study site

$$\log \left[ \frac{\Pr(Y_{i,t} = 1)}{(1 - \Pr(Y_{i,t} = 1))} \right] = \alpha_0 + \alpha_{st} + (\beta_0 + \beta_s) * G_{i,t,p} \tag{2}$$

where  $\alpha_{st}$  and  $\beta_s$  are vectors of random effects normally distributed (N), with  $\alpha_{st} \sim N(0, \sigma^2_{\alpha_{st}})$  being the random effect for the intercept grouped by year (*t*) nested in each study site (*s*), and  $\beta_s \sim N(0, \sigma^2_{\beta_s})$ , the random effect for the growth variable grouped by study site (random effect of the slope).

Both fixed-effects (Eq. 1) and mixed-effects (Eq. 2) models were designed for one single growth variable as

detailed previously, but they were also used with a combination of up to *X* growth variables. Thus, the mixed-effects model was structured as follows:

$$\log \left[ \frac{\Pr(Y_{i,t} = 1)}{(1 - \Pr(Y_{i,t} = 1))} \right] = \alpha_0 + \alpha_{st} + \sum_{x=1}^{x=X} (\beta_{0x} + \beta_{sx}) * G_{i,t,p,x} \tag{3}$$

where  $\beta_{0x}$  are the coefficients for each growth variable calculated over a time window *px* ( $G_{i,t,p,x}$ ), and  $\beta_{sx}$  are the vectors of random effects for each growth variable grouped by study site, with  $\beta_{sx} \sim N(0, \sigma^2_{\beta_{sx}})$ .

*Statistical sampling schemes used for model calibration*

We tested different approaches for sampling the alive observations, depending on the research questions being addressed. Note that the ratio between death and alive observations that were derived from these samplings is not representative of stand-scale mortality rates observed in the field. Evidently, this precludes the use of these logistic models to predict mortality at the population level, which was not intended here.

*All years are considered.*—We first used a sampling strategy that considers all the years of each tree’s life as recorded in the tree-ring data (sampling *T* for total number of years; Appendix S3: Fig. S3.1). Here, the random effect for the intercept grouped by year was not considered in the logistic model (vector  $\alpha_{st}$  of Eqs. 2 and 3). This sampling is preferable if one wants to better understand the growth patterns prior to the mortality of a specific tree in a specific year, derive growth–mortality relationships that are reliable for all types of mortality and all tree developmental stages, or use these relationships to simulate individual-tree mortality in dynamic vegetation models (Bircher et al. 2015).

*Focus on the years for which mortality occurs.*—To predict which tree will die during a specific mortality event (e.g., drought-induced mortality in a certain period) and to compare the growth patterns of the living trees and now-dead trees prior to this event, we sampled only the death/alive observations and the associated growth variables corresponding to the years when mortality occurred. We thus generated unequal samplings (*U*) in which all the alive observations of a given year were used (Appendix S3: Fig. S3.1), which usually leads to a lower ratio of alive/death observations than sampling *T*.

To build models that are able to predict well both survival and mortality, rather than only survival, the prevalence of the alive observations in the data has to be reduced (see Lawson et al. 2014). Thus, as a third approach a paired sampling (*P*) was generated, i.e., for each dead tree that died at time *t*, we sampled only one alive observation from the same site at the same *t*



(Appendix S3; Fig. S3.1). The pair was removed from the dataset if one of the two trees did not contain a given explanatory variable. The alive observation was sampled from both living and now-dead trees and chosen either randomly (*P-R*; with replacement) or from a tree that was as similar as possible to the dead tree in terms of cambial age (*P-A*; note that an alive observation could be used repeatedly if the number of death events was higher than the number of survival events). The *P-A* approach is often employed in studies to help control for the temporal changes in micro-environmental conditions.

We also assessed if the structure and performance of the mortality models was impacted by the status of the trees used to sample the alive observations (e.g., living trees only vs. dead trees only; Appendix S3), sample size (Appendix S4), and (3) the random selection of the alive observations (Appendix S5). For *Nothofagus dombeyii*, all the trees died at the same date (last ring formed in 1997). As a consequence, alive observations arise from living trees only, and the determination of the pairs (randomly or per age) was not relevant. Moreover, as the number of dead trees was equal to the number of living ones, unequal sampling could not be drawn, which restricted us to study the sampling schemes *P-A* and *T*.

#### Assessment of model performance

Model-predicted survival probabilities ( $\text{Pr}_{i,t}$ ) were compared with the individual binary alive/death information from the observed data set to assess model performance.

Although calibration metrics such as Akaike's information criterion (AIC) are commonly used (Appendix S1), AIC is strongly positively correlated with the number of samples used to calibrate the model and depends on the equitability of the sampling (i.e., prevalence of the living observations; see Lawson et al. 2014; Appendix S6). This is highly problematic when comparing the performance of models fitted to different sample sizes and properties. Using a minimum AIC approach would lead to a preferential selection of models with low sample size, i.e., models for which the growth variables are calculated using a large time window (Appendix S6).

Discrimination metrics are more suitable for our purpose as they reflect the ability to separate survival ( $Y_{i,t} = 1$ ) and death ( $Y_{i,t} = 0$ ) observations based on  $\text{Pr}_{i,t}$  (Lawson et al. 2014). Binary metrics such as sensitivity (fraction of true positive cases, i.e., the tree survived and the model predicts a survival event) or specificity (fraction of true negative cases, i.e., the tree died and the model predicts a death event) can be used, but they require determining a threshold that would classify a tree as dead or alive, whose value strongly depends on the rather arbitrary method used for its calculation (cf. Bircher et al. 2015).

As an alternative, continuous discrimination metrics such as the area under the receiver operating characteristic (ROC) curve (AUC) are preferable, especially due to their independence of the equitability of the sampling.

An ROC curve is constructed by plotting the ratio of true positives to false positives ( $1 - \text{specificity}$ ) for all possible threshold values of the estimated survival probability (Fielding and Bell 1997). AUC values range from 0 to 1, with 0.5 indicating a random model. A model providing an excellent prediction has  $\text{AUC} \geq 0.9$ , a fair model  $0.7 \leq \text{AUC} < 0.9$  (Swets 1988), and  $\text{AUC} < 0.7$  indicates a poor model.

All mixed-effects models were fitted using the package lme4 of the software R (R Core Team 2015; Bates et al. 2015), and their AUC was calculated using the package ROCR (Sing et al. 2013). In the case of very large eigenvalues and a perfect model ( $\text{AUC} = 1$ ), the significance of the effect of each growth variable was assessed using Markov chain Monte Carlo techniques (package MCMCglmm; Hadfield 2010), and Firth's penalized-likelihood logistic regression (package logistf; Heinze et al. 2013), respectively.

#### Growth variables

Trees can die during the growing season when ring formation is not finished, which induces an incomplete outermost ring. As the precise (intra-annual) timing of tree death was not available, we did not consider the last ring of the dead trees to avoid a bias in performance of the models that use recent growth to discriminate dead from living trees. Thus, the year of death considered here was a proxy and was defined as the year prior to the formation of the outermost ring. Basal area increments (BAI) were calculated from ring-width (RW) data for all trees whose DBH information was available (i.e., all trees except for dead *Q. petraea*), and both variables were used to reflect different growth characteristics of tree *i* at time *t* (Fig. 1).

*Absolute and relative growth level.*—Growth level variables were defined as the average growth of the last (outermost) *p* rings (years)

$$\text{RW}_{i,t,p} = \text{mean} \left( \sum_{y=t-p+1}^{y=t} \text{RW}_{i,y} \right), \quad (4)$$

$\text{RW}_{i,t,p}$  corresponds to the width of the last ring when  $p = 1$ . The same equation was used to calculate  $\text{BAI}_{i,t,p}$ , and the relative growth variable  $\text{relBAI}_{i,t,p}$  was calculated as the ratio of  $\text{BAI}_{i,t,p}$  and  $\text{BAI}_{i,t}$ . We also tested the impact of the log-transformation of these growth level variables on model performance, since logarithms are frequently used to reduce the effect of high growth values on  $\text{Pr}(Y_{i,t} = 1)$  (see Appendix S1).

*Growth trend.*—The slopes of local linear regressions fitted over the last *p* years were used to characterize growth trends with RW ( $\text{sloRW}_{i,t,p}$ ) and BAI ( $\text{sloBAI}_{i,t,p}$ ) and were used as independent variables in the growth-mortality models (Fig. 1).

*Growth variance and autocorrelation of annual ring-widths.*—The mean sensitivity (MS) of the raw data or of the detrended chronology is often used to assess

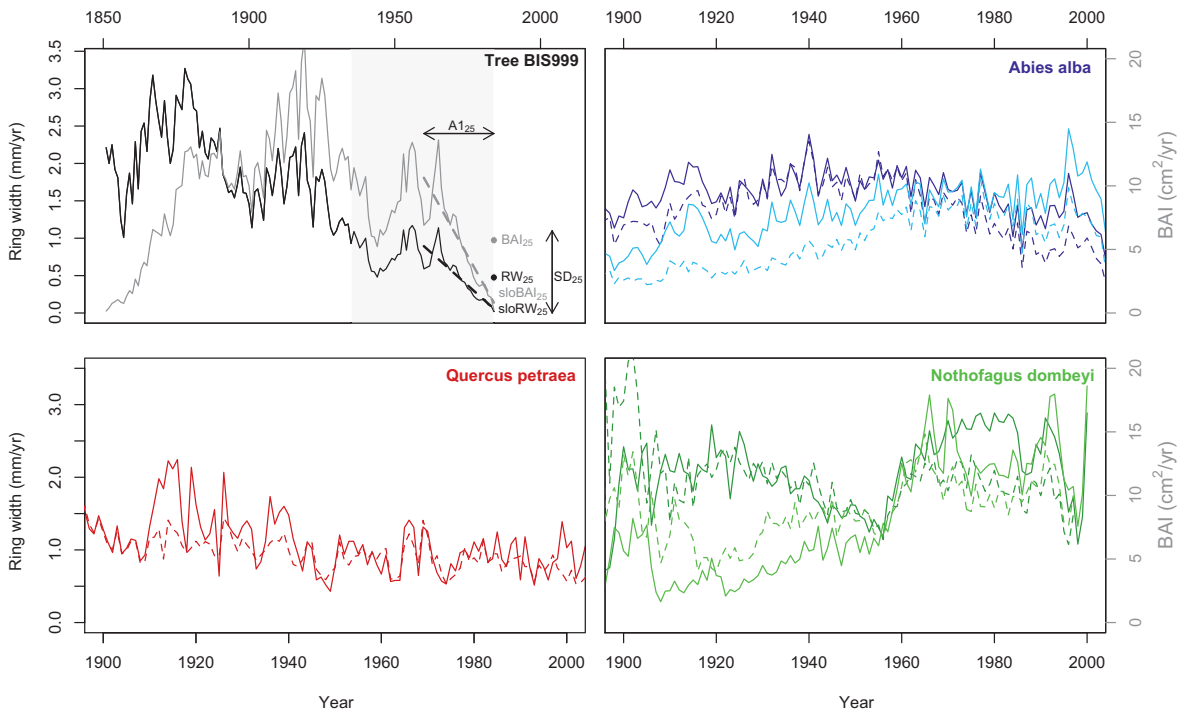


FIG. 1. Top left: Ring-width (black) and basal area increment (gray) chronologies of tree BIS999 (*Abies alba*; site Bistra) and the growth variables used as predictors of tree survival probability. Variables that reflect growth level (mean RW, mean BAI), growth trend (sloRW, sloBAI; slope of the linear regression represented by the dotted line), and the variance (SD) and lag-1 autocorrelation (A1) in the ring-width series are shown for a time window ( $p$ ) of 25 yr (dark gray polygon). The gray polygon reveals the different time windows tested, from 1 to 50 yr. Top right and bottom: Change over time of the median RW (dark colors) and median BAI (light colors) of living (solid line) and dead trees (dashed line) for *A. alba* (blue), *Q. petraea* (red), and *N. dombeyi* (green).

the interannual variation in the ring-width chronology. However, MS is strongly correlated with the lag-1 autocorrelation of the rings (A1; see Bunn et al. 2013) and is not necessarily a good proxy of growth variance. Thus, after detrending each series using a power transformation (see Cook and Peters 1997), we estimated both variance and autocorrelation in tree-ring growth using the standard deviation and the AR1 coefficient of an autoregressive model fitted to each detrended chronology (Bunn et al. 2013; Appendix S7).

#### Determination of the best-fitting time window for calculating growth variables

The next step for deriving reliable logistic mortality models was to define the number of rings ( $p$ ) that should be considered to calculate the growth variables. All growth variables were calculated using  $p$  rings that ranged between  $p_{\min}$  and  $p_{\max}$ , with  $p_{\min} = 1$  for growth level variables,  $p_{\min} = 2$  for growth trend variables, and  $p_{\min} = 10$  for the growth variance and autocorrelation variables.  $p_{\max}$  was set to 50 rings so as to consider potential long-term negative effects of intense stress and disturbances on tree status.

*Logistic models with a single growth variable.*—In the classical approach, all trees were used to calibrate the

logistic models, using every combination of  $p$  and growth variables (Eqs. 1 and 2). However, with the increase of  $p$ , young trees (or trees for which partial tree-ring data were available) were increasingly excluded from the calibration dataset, inducing a decrease of sample size (Fig. 2). The change of AUC with changing  $p$  may thus be biased due to differences in the set of trees that were selected in the sample (number and characteristics of the trees). Therefore, we assessed the change of model AUC with  $p$  for identical samplings using an iterative process, hereafter called constant sampling approach. We firstly used the trees with at least  $p_{\max}$  rings and calculated model AUC obtained for every  $p$  with this dataset. Then, we used the same approach with the trees with at least  $p_{\max} - 1$  rings (same or higher number of trees than previously), etc., down to  $p_{\min}$ . We ended up with  $(p_{\max} - p_{\min})$  different patterns of changes in model AUC over  $p$  (Fig. 2), where AUC at each  $p$  was calculated from the same sample of trees. For each of these curves, the  $p$  that maximized model AUC was recorded. Finally, from the set of all these curves, we determined the proportion of samples whose AUC was maximized at each  $p$ . The  $p$  with the highest proportion of samples was then considered as the best-fitting time window (hereafter called best-fitting  $p$ ). As only one sample was available at  $p_{\max}$ , we defined that the best-fitting  $p$  was equal to  $p_{\max}$  only when model AUC increased monotonically with  $p$ .

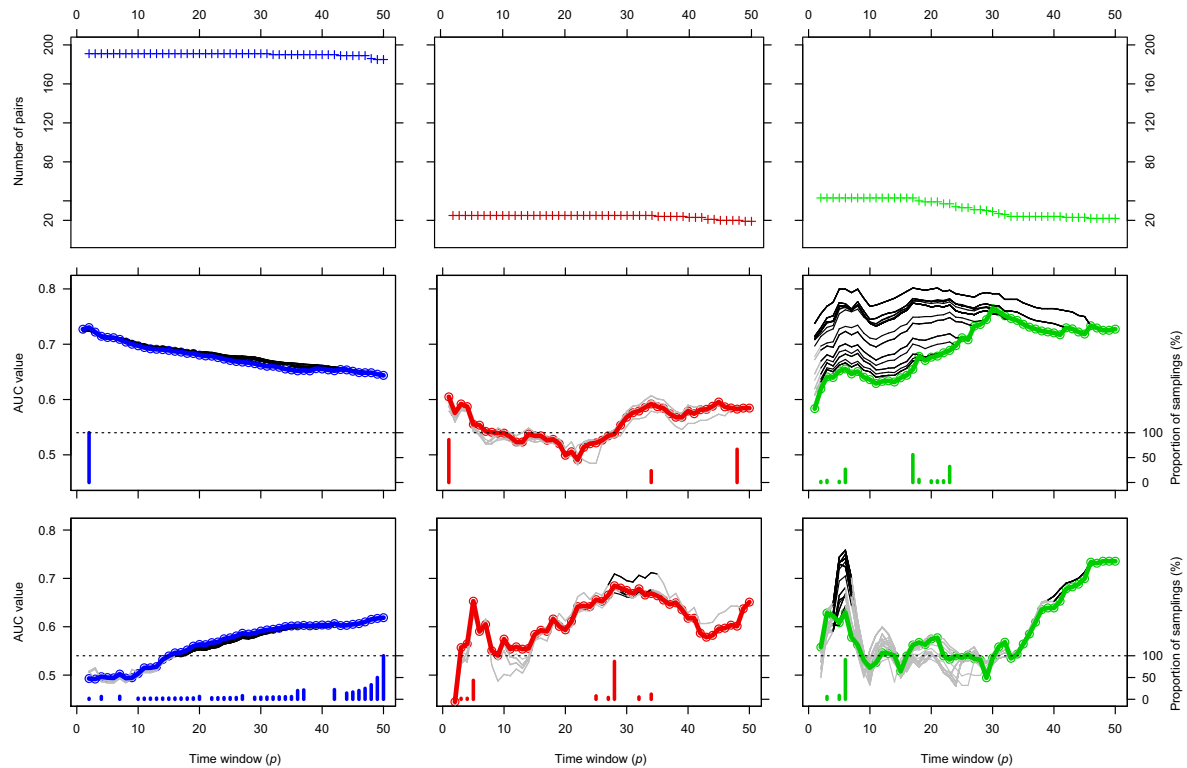


FIG. 2. Top: Change in the number of pairs used (crosses) to calibrate the logistic model when increasing the number of rings  $p$  used to calculate  $RW_{i,t,p}$  for *A. alba* (left, blue, sampling *P-A*), *Q. petraea* (center, red, sampling *P-A*) and *N. dombeyi* (right, green, sampling *P-A*). Change in the AUC value of logistic models that use  $RW_{i,t,p}$  (middle row) and  $sloRW_{i,t,p}$  (bottom row) as single predictor of tree mortality according to the length of the time window ( $p$ ) used to calculate this growth variable. The colored curves reflect the change in AUC using the classical approach. The different black and gray curves reflect the change in AUC with different samples whose size is constant over  $p$  (gray lines indicate that the growth–mortality relationship was not significant at this  $p$ ). At each  $p$ , we calculated the proportion of these samples for which AUC is maximized at this specific  $p$  and reported this value in the histograms shown in the bottommost panels.

*Logistic models with multiple growth variables.*—Under the constant sampling approach, there are  $(p_{\max} - p_{\min})^X$  combinations of models that include  $X$  growth variables that can be calculated using  $p_{\min}$  to  $p_{\max}$  different time windows (Eq. 3). This results in  $>6000000$  ( $=50^4$ ) models that have to be fitted to assess the best-fitting time window of four variables with  $p_{\min} = 1$  and  $p_{\max} = 50$ . In order to reduce simulation time, we used differential-evolution optimization algorithms (R package DEoptim; Mullen et al. 2011) to detect the combination of best-fitting  $p$  that optimizes model AUC while escaping local minima (Storn and Price 1997), developed one logistic model including all RW-related growth variables (RW, sloRW, SD, A1), and used three possible  $p_{\max}$  (20, 35, 50) that would allow us to disentangle the effects of changing calibration dataset and changing the maximum length of the time windows. To assess the relative importance of each growth variable on predicted  $pr_{i,t}$ , we calculated the proportion of the total variance in  $pr_{i,t}$  explained by the variation in each explanatory variable (i.e., effect size of the fixed effects) using variance analysis techniques.

## RESULTS

### *Differences in best-fitting p among approaches*

Two main patterns were obtained when comparing the best-fitting  $p$  determined using the classical and the constant sampling approaches. On the one hand, in some cases the best-fitting  $p$  was consistent in both approaches, e.g., when using growth-level variables for *A. alba* and on the paired sampling schemes for *Q. petraea* (Figs. 2 and 3; Appendix S8). For instance, on samplings *P-A* for *A. alba* and *Q. petraea*, the maximum AUC of models using  $RW_{i,t,p}$  was obtained at  $p = 2$  and  $p = 1$ , respectively, irrespective of the approach. By contrast, with  $sloRW_{i,t,50}$  for *A. alba* (*P-A*), all samples for which the logistic model could be calibrated showed the highest performance at  $p = 50$  (Fig. 2). Admittedly, this was based on a single sample, but it was coherent with the increase in model performance over  $p$  observed for all other samples and also when using the classical approach.

On the other hand, the determination of the best-fitting  $p$  on the sampling scheme *P-A* of *N. dombeyi* led to different results (Figs. 2 and 3; Appendix S8). For  $RW_{i,t,p}$ ,

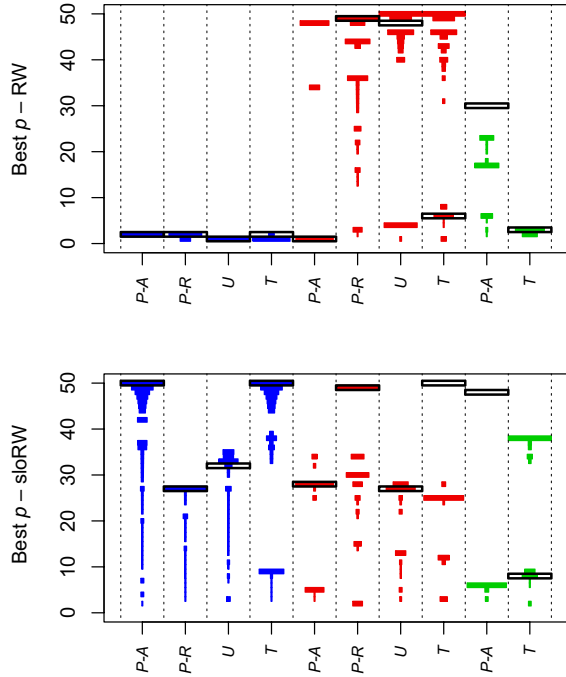


FIG. 3. Change in the frequency of  $n$  that maximizes AUC of models that used  $RW_{i,t,p}$  (top) and  $sloRW_{i,t,p}$  (bottom) to predict survival probability of *A. alba* (blue), *Q. petraea* (red), and *N. dombeyi* (green) according to the statistical sampling scheme used. The larger is the colored rectangle, the higher is the number of samples for which maximum AUC is obtained at this  $p$ . Open rectangles with black borders reveal the best-fitting  $p$  obtained with the classical approach.

the classical approach revealed a lower AUC of models that used  $p < 30$  rings, with an optimal AUC at  $p = 31$ . This pattern was not found when analyzing the change in AUC over  $p$  for samples with constant size, in which maximum AUC occurred at  $p = 6$  and  $p = 17$  for 27% and 56% of these samples, respectively (Fig. 2). Samples that included only trees  $>30$ -yr old showed much higher AUC at  $p < 30$  than samples with both old and young trees. For instance, at  $p = 17$ , AUC ranged between 0.65 and 0.80 depending on the sampling that was applied (Table 2). The reduction in AUC towards lower  $p$  that was observed with the classical approach for *N. dombeyi* (*P-A*) was caused by a strong increase in the number of young trees in the sample for which the relationship between  $RW_{i,t,p}$  and tree status was not significant while it was significant for older trees (divergence in the late 1960s; Figs. 2 and 4). This was not observed for *A. alba* and *Q. petraea* for which the relationships between  $RW_{i,t,p}$  and tree status were similar among tree size classes and the reduction in the number of pairs was not as pronounced (i.e., 95% of *A. alba* trees were older than 50 yr; Fig. 2).

Differences among statistical sampling schemes

Within a species and for a given growth variable, the determination of best-fitting  $p$  depended also strongly on

TABLE 2. Range of AUC of the single-variable models obtained at best-fitting  $p$  (in brackets) with the constant sampling approach for each species, statistical sampling scheme, and growth variable studied.

	<i>A. alba</i>				<i>N. dombeyi</i>				<i>Q. petraea</i>			
	<i>P-A</i>	<i>P-R</i>	<i>U</i>	<i>T</i>	<i>P-A</i>	<i>T</i>	<i>P-A</i>	<i>T</i>	<i>P-R</i>	<i>U</i>	<i>T</i>	
RW (+)	[0.72–0.73] (2)†	[0.72–0.74] (2)	[0.83–0.84] (1)††	[0.86–0.90] (1)†	[0.65–0.80] (17)	[0.67–0.82] (3)	[0.58–0.60] (1)	[0.67–0.67] (49)	[0.67–0.67] (49)	[0.58–0.59] (4)	[0.62–0.65] (46)	
BAI (+)	[0.70–0.71] (1)†	[0.72–0.73] (2)	[0.83–0.83] (1)††	[0.69–0.79] (1)†	[0.63–0.75] (5)	[0.52–0.77] (2)						
relBAI (+)	[0.70–0.71] (1)†	[0.68–0.71] (1)	[0.81–0.82] (1)††	[0.82–0.91] (1)†	[0.61–0.81] (5)	[0.65–0.81] (3)						
sloRW (+)	[0.62] (50)††	[0.64–0.65] (27)†	[0.76–0.77] (33)††	[0.78] (50)†	[0.63–0.76] (6)	[0.73–0.75] (38)	[0.67–0.71] (28)	[0.68–0.70] (30)	[0.67–0.70] (27)	[0.65–0.72] (25)		
sloBAI (+)	[0.67] (50)†	[0.68–0.72] (32)†	[0.78–0.78] (48)††	[0.80–0.80] (48)†	[0.81–0.83] (43)	[0.77–0.79] (37)						
SD (+)	[0.58–0.59] (13)	[0.49–0.56] (17)†	[0.74–0.75] (10)††	[0.69–0.69] (44)†§	[0.61–0.70] (34)	[0.52–0.67] (16)	[0.52–0.54] (15)	[0.52–0.58] (10)	[0.51–0.53] (10)	[0.52–0.54] (10)		
AI (–)	[0.60] (50)†	[0.58–0.62] (10)†	[0.77] (50)	[0.63–0.65] (11)	[0.53–0.63] (24)	[0.56–0.61] (19)	[0.66–0.71] (25)	[0.67] (50)	[0.56–0.63] (25)	[0.61–0.67] (23)		

Notes: Boldface indicates that the effect of growth<sub>*i,t,p*</sub> on  $Y_{i,t}$  was significant ( $P$  value  $< 0.05$ ) for the model calibrated at best-fitting  $p$  with all the possible samples. (+) and (–), respectively, reveal a positive and negative relationship between each growth variable with survival probability. †The effect of the growth variable differed among sites (best model includes the random effect of growth grouped by site). ‡The effect of the intercept grouped by site nested in year. §The sign of the relationship is different from the general pattern.



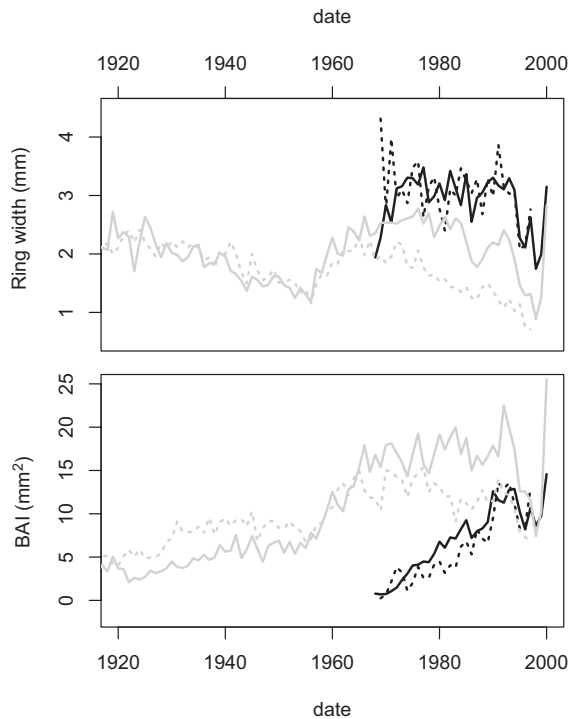


FIG. 4. Change over time of the median RW (top) and median BAI (bottom) of living (solid line) and dead (dashed line) *N. dombeyi* for both young ( $\leq 30$  yr old, black line, 13 pairs) and old trees (gray, 30 pairs) included in the sampling *P-A*.

the statistical sampling schemes used to calibrate the models. For *A. alba*, the best-fitting  $p$  for calculating  $\text{sloRW}_{i,t,p}$  ranged between 27 and 33 yr for the samplings *P-R* and *U*, but was 50 yr for the samplings *P-A* and *T* (Table 2). This instability in the determination of the best-fitting  $p$  was also observed depending on whether alive observations were selected from living trees or dead trees only (Appendix S3) and within statistical sampling schemes, especially when sample size was limited (e.g.,  $< 40$  pairs; Appendices S3 and S4). In addition, model performance also varied among statistical sampling schemes (Table 2; Appendix S3). Model AUC obtained from unpaired samplings was higher than AUC obtained from paired samplings. For instance, models that related  $\text{BAI}_{i,t,1}$  with survival probability of *A. alba* showed an AUC of 0.83 and 0.71 for the *U* and *P-A* samplings, respectively (Table 2). This result simply reflects that alive observations are more easily predicted than death observations, and thus should not be interpreted as unpaired samplings being superior for predicting mortality.

#### Differences among growth variables in univariate models

For the growth level variables, there was no evidence of higher prediction accuracy of survival probability by BAI than by RW. For both variables the best fits were typically obtained using the last year or the last two years for *A. alba* (Fig. 3 and Table 2). Using the *P-A* sampling scheme for

*N. dombeyi*, the best-fitting  $p$  changed depending whether the model used  $\text{RW}_{i,t,p}$  or  $\text{BAI}_{i,t,p}$  as explanatory variable (17 and 5 yr, respectively; Table 2). However, this difference was not robust because of the high uncertainty in best-fitting  $p$ , and as AUC values obtained at  $p = 5$  for  $\text{RW}_{i,t,p}$  were almost identical to those at  $p = 17$  (Fig. 2 and Table 2). Similarly, the log transformation of the growth level variables did not significantly improve model AUC (Appendix S9). The use of relative growth level variables instead of absolute ones slightly decreased model AUC on average ( $-0.05$  to  $+0.06$  between models that used BAI and  $\text{relBAI}$ ; Table 2), except for the samplings *T*, which strongly benefitted from this change ( $+0.04$  to  $+0.12$ ).

The temporal trend in RW series was not a suitable predictor of survival probability (AUC  $< 0.7$ ), except for the samplings *U* and *T* for *A. alba* and *N. dombeyi*. For the other samplings, BAI should be preferred for fitting linear regressions (i.e.,  $\text{sloBAI}$ ). Between 25 and 30 yr should be considered to calculate growth trend variables of *Q. petraea*. No unique best-fitting  $p$  emerged for *N. dombeyi*, whereas for *A. alba* it was  $> 25$  yr (cf., Fig. 3).

For the sampling schemes *P* and *T*, logistic models that used SD or A1 did not predict survival probability correctly. However, although the averaged relationships between these growth variables and survival probability were not significant for unpaired samples, the models' AUC were  $> 0.7$  revealing that the sign and magnitude of these relationships changed among sites (Table 2). Despite the large uncertainty in best-fitting  $p$ , usually short time windows ( $< 20$  yr) were obtained.

#### Best multivariate models: sign of the growth–mortality relationships

We derived multivariate models to determine to what extent multivariate models perform better than univariate ones. Moreover, as most growth variables were correlated (Appendix S10), they were helpful to assess and rank the true effect of each of them, i.e., to assess the sign of the growth–mortality relationships and the contribution of each growth variable to the model-predicted survival probability.

Similarly to models with one single growth variable, the structure and performance of the best multivariate models changed as the maximum length of the time window ( $p_{\max}$ ) considered to calculate the growth variable increased and with the statistical sampling scheme (Table 3). For *A. alba*, AUC was lower for paired than for unequal samplings, while this was not clear for *Q. petraea*, probably because of the low sample size. For paired samplings, mortality models that included four growth variables showed higher AUC than the best univariate models ( $+0.06$ ,  $+0.09$ , and  $+0.11$ , for *A. alba*, *Q. petraea*, and *N. dombeyi*, respectively). On average, this increase in AUC was lower in the case of samplings *U* and *T* ( $+0.02$ ,  $+0.03$ , and  $+0.05$ ).

As expected, survival probability was positively correlated with growth-level and growth-trend variables, the former being predominant for predicting survival

TABLE 3. Time windows ( $p$ , number of years preceding the dead or live observation) used to calculate the growth variables included in the best multivariate mortality model, including: mean ring width (RW), the temporal trend of RW (sloRW), and the standard deviation (SD) and lag-1 autocorrelation (A1) of the RW time series.

	<i>Abies alba</i>				<i>Nothofagus dombeyi</i>		<i>Quercus petraea</i>			
	<i>P-A</i>	<i>P-R</i>	<i>U</i>	<i>T</i>	<i>P-A</i>	<i>T</i>	<i>P-A</i>	<i>P-R</i>	<i>U</i>	<i>T</i>
$p_{\max} = 20$										
RW	2 (+)***	2 (+)***	1 (+)***	1 (+)***	18 (+)**	3 (+)***	20	1	1	1 (+)*
sloRW	4	20	5	16	3 (-)*	7 (+)***	5 (+)*	20	13	6
SD	10	18	17	18 (-)**	17	20	12	16	15	10
A1	19	11 (-)*	11	10 (-)***	13	11 (-)*	10	19	13	20
AUC	0.742†‡	0.799†	0.850†‡	0.916†	0.751	0.781	0.731	0.688	0.673	0.675
$p_{\max} = 35$										
RW	3 (+)***	2 (+)***	2 (+)***	3 (+)***	17 (+)**	3 (+)***	13	13	1	4
sloRW	4 (+)*	24	31 (+)***	3 (+)***	12	35 (+)*	5 (+)*	34 (+)*	27 (+)**	28 (+)*
SD	24	25 (-)*	35	17 (-)***	29	12	22	10	11	10
A1	22	10 (-)*	35	12 (-)***	35	11 (-)**	24 (-)*	29	35	23
AUC	0.737†‡	0.808†‡	0.854†‡	0.911†	0.849	0.856	0.754	0.762	0.701	0.700
$p_{\max} = 50$										
RW	6 (+)***	42 (+)***	2 (+)***	3 (+)***	10 (+)**	12 (+)***	7	50	5	50
sloRW	4 (+)**	25 (+)***	31 (+)***	4 (+)***	41 (+)*	7 (+)**	34 (+)*	9	27 (+)**	28 (+)*
SD	24	23	43 (-)**	23 (-)*	33	31	41	26	17	28
A1	41	10 (-)*	43	12 (+)***	36 (+)*	13	11	50	11	23
AUC	0.743†‡	0.794†‡	0.853†‡	0.903†	0.926	0.868	0.798	0.798	0.719	0.750

Notes: Models were calibrated using different  $p_{\max}$  and statistical samplings of dead and live observations. We used the total sample, i.e., all the rings ( $T$ ), or focused only on those years when mortality occurred. In the latter case, we considered all live observations ( $U$ ) or generated dead/live pairs (trees of same age,  $P-A$ ; or randomly,  $P-R$ ). (+) and (-), respectively, signify positive and negative effects of each growth variable on tree survival probability. \* $P < 0.05$ ; \*\* $P < 0.01$ ; \*\*\* $P < 0.001$ . † and ‡ indicate significant random effects of the growth variables grouped by site and of the intercept grouped by site nested in year on survival probability, respectively.

probability of *A. alba* while the pattern was reversed for *Q. petraea* (Appendix S11). When the relationship was significant, growth-level variables of *A. alba* were mainly calculated with  $p$  ranging from 1 to 3 yr for  $RW_{i,t,n}$  (Tables 3 and B3). This best-fitting  $p$  was much more variable for growth trend variables (from 3 to 31 yr), although it was mostly between 25 and 34 yr for *Q. petraea* (Table 3).

The relationships with SD and A1 were slightly significant and were only reported for specific combinations of species,  $p_{\max}$ , and sampling scheme. For instance, SD was never related to survival probability of *Q. petraea* and *N. dombeyi*, and only five of the 12 samples of *A. alba* showed a negative relationship, with the time window used to assess SD being rather uncertain. Results were even less clear for A1. For *A. alba* and *N. dombeyi*, the sign of the relationship between A1 and survival changed with  $p_{\max}$  and among sampling schemes (Table 3) and was mainly negative when alive observations could arise from dead trees (Appendix S3).

DISCUSSION

Potential and limitations of tree-ring widths as predictors of tree death

As revealed by the high AUC values of the multivariate models, there is a high potential of using tree rings as a

robust indicator of tree mortality probability (Bigler and Bugmann 2004). In the present study, AUC was calculated using the same data as for model calibration and optimized using appropriate algorithms. As our aim was to assess growth–mortality relationships per se and not to predict mortality probability by applying the models, they were not validated on an independent dataset (such as in Bigler and Bugmann 2004), and thus their performance may be exaggerated. Nevertheless, AUC can be quite low, e.g., for *Q. petraea*, because growth may not reflect all environmental influences that drive mortality probability (Das et al. 2008, Kane and Kolb 2010, Ferrenberg et al. 2014) and because the date of tree death estimated using ring observations on cores may be biased due to partial cambial dieback (Bigler and Rigling 2013).

If prediction is the goal, the performance of tree-ring-based mortality models may be improved by standardizing the growth estimates for each site, e.g., using the regional curve standardization method (Esper et al. 2003). As the species-specific effect of growth on mortality may depend on the local conditions of the study site, as revealed by some significant random effects for *A. alba*, these relative growth values are probably better indicators of the vitality of a single tree within a specific site (see Das and Stephenson 2015) and are of higher interest to forest managers (e.g., the mortality probability is higher than 50% if the last ring width was below a given percentage of mean growth). However, model calibration

and conclusions become more vulnerable to the sampling strategy. When the data are not representative, i.e., not all trees are sampled, or the sampled plots do not fully portend actual forest dynamics, it may be difficult to use growth variables that are expressed relative to those of other trees.

Another way to improve the statistical determination of growth patterns prior to mortality would be to use a different statistical modelling approach (e.g., Wyckoff and Clark 2000). In the present study, we assumed that the response variable, (i.e., tree status) follows a binomial distribution, but a Weibull distribution could also be used (Wyckoff and Clark 2002). Other methods may also be powerful, such as the quickest detection method (Carpenter et al. 2014), autologistic models, neural networks (Hasenauer et al. 2001), multivariate adaptive regression splines, classification and regression trees, or general additive models (Austin 2007).

#### *Data requirements: sample size and statistical sampling schemes*

Tree-ring-based analyses are strongly influenced by sampling schemes.

First, the number of sampled trees affects growth–mortality relationships analogously to the quantification of climate–growth associations (Mérián et al. 2013). A large instability in the best-fitting  $p$  of the different growth variables was observed among and within the different statistical schemes for *Q. petraea* (Appendix S5) and within samplings of *A. alba* with 40 pairs (*P-R*; Appendix S4), revealing that a small sampling size may bias the structure and performance of the best mortality models.

Second, growth–mortality relationships depend on the characteristics of the sampled trees, including their age, size, and social status (cf. Martín-Benito et al. 2008 for climate–growth associations), independently of the total number of samples. The sampling design (e.g., random or stratified selection of the trees) can bias the determination of the best mortality models, similar to tree-ring-based metrics of stand-scale forest productivity (Nehrbass-Ahles et al. 2014).

Third, as revealed by the significant random effect of the site on the growth variables of single- and multi-variable logistic models of *A. alba*, growth–mortality relationships are most likely not only species-specific, but depend on the combination site  $\times$  species  $\times$  mortality driver (Wunder et al. 2008, Macalady and Bugmann 2014, Camarero et al. 2015).

In consequence, in order to assess reliable growth–mortality relationships for a given species, we recommend to study multiple sites in contrasting environmental conditions, either to calibrate the model and/or to validate it (e.g., Bigler and Bugmann 2004) and to choose the field and statistical sampling design based on the research questions (see *Materials and Methods*; Hartmann et al. 2007).

The coring of trees, measuring ring widths and cross-dating each individual growth series is quite time

consuming, and therefore the number of samples is always limited. The present study highlights the necessity and utility of assessing the uncertainty in the structure and performance of the best mortality models using re-sampling techniques (e.g., for *Q. petraea*; see Appendix S5) or employing different statistical sampling designs. The latter option may also reveal effects that were not accounted for in the initial model structure. For instance, comparing the performance of models calibrated with *P-A* and *P-R* samplings may indicate any effect of tree size/age and/or a temporal shift in growth mortality relationships (e.g., for *N. dombeyi*), while sampling the alive observations from dead trees only or from living ones can reveal specific properties of the tree-ring series of dead trees (Appendix S3). Site-specific effects of growth variables can be accommodated using hierarchical models with random site effects of the intercept and the slopes, as illustrated in our models for *A. alba* (cf. Eq. 2).

#### *Determination of the best-fitting time window*

Instead of arbitrarily selecting and comparing different time windows ( $p$ ), we tested all possible  $p$  between 1 ( $p_{\min}$ ) and 50 ( $p_{\max}$ ) years using an objective statistical approach that optimized the discriminatory power of each explanatory variable. This procedure allowed for considering long-term changes in growth dynamics and detecting local maxima in model performance. However, with the increase in  $p_{\max}$ , young trees (or trees whose innermost rings are lacking) were progressively excluded from the calibration dataset. This may induce a change in the structure and performance of the best model if trees of different ages have different growth–mortality relationships and when sample size is low (e.g., for *N. dombeyi*). To account for both the effect of increasing  $p_{\max}$  and the change of the calibration dataset, we suggest that one should follow the constant sampling approach, which may highlight potential developmental shifts in species-specific growth–mortality relationships. Such a shift can occur gradually as the main drivers of tree mortality usually change as trees grow and their micro-environment changes (e.g., Das et al. 2007). Trees mostly suffer from excessive shading during the suppressed juvenile stage, but with increasing size, they become less shade tolerant and less able to survive at low growth rates (Valladares and Niinemets 2008). Tall trees may also be more prone to drought-induced mortality than small ones (Bennett et al. 2015) due to lower whole-tree hydraulic conductance (hydraulic limitation hypothesis; Ryan et al. 2006). The change in growth–mortality relationship may be sudden as a result of an extreme climatic event or due to the abrupt emergence of pathogens or insect outbreaks in the study area. In the case of insect-related mortality events, we expect partial decoupling between growth and mortality processes, as mortality probability would be more related to defense characteristics (e.g., density of resin ducts against bark beetle attacks) than to radial growth per se (Ferrenberg et al. 2014).

### Growth variables

We deliberately did not test all the growth variables used in earlier studies but selected the most common ones or those that had shown a strong relationship with mortality probability in previous studies (Bigler and Bugmann 2004, Kane and Kolb 2014, Macalady and Bugmann 2014). In the case of low performance of univariate models (AUC < 0.7; e.g., based on paired samplings for *Q. petraea* and *N. dombeyi*), mortality models that included four growth variables performed better than the best univariate ones (increase in AUC > 0.1) due to the additional information provided by each variable. Variance decomposition analyses performed on multivariate models highlighted that growth level variables were a better indicator of tree survival probability in two of the three species (*A. alba* and *N. dombeyi*), while growth-trend variables should be considered in the case of *Q. petraea*. No specific recommendation can be made regarding the time windows that should be used to calculate growth-level and growth-trend variables as they varied among the three species and among the sampling schemes.

BAI is often considered to be a biologically more meaningful growth-level variable than RW, as it depends less on tree size and approximates biomass increase better (Bowman et al. 2013). This may be an important issue when the trees have different DBH or age, such as in *P-R* and *U* sampling schemes. However, and similar to Macalady and Bugmann (2014), in our case BAI did not perform better than RW in segregating living and dead trees. The reverse pattern was even found for the sampling *T*, in which all available rings were used. As BAI is very low during the juvenile phase compared to RW (e.g., first 20 yr; Figs. 1 and 4), models using BAI tend to overestimate the death probability of young trees. In this case, relative growth level variables such as relBAI should be preferred (Das et al. 2007, Holzwarth et al. 2013).

Using RW or BAI data also mattered for calculating growth trend variables. When linear regressions were fitted to RW series, the differences in the growth trend prior to the alive vs. death observations could be missed (or underestimated). Living trees usually have higher growth rates than dying trees, and therefore their radial increment in mm would decrease faster for similar volume growth (see Fig. 4; Bowman et al. 2013). This effect explains why the increase in model AUC between models that used sloRW and sloBAI was reported only for samples in which the alive observations arose from living trees.

Finally, *A. alba* trees exhibited higher interannual growth variability prior to death than surviving trees, and the magnitude of this relationship varied among sites, while the survival probability of *Q. petraea* and *N. dombeyi* was not correlated with growth variance. Interestingly, Suarez et al. (2004) found that dead *N. dombeyi* showed higher interannual growth variability than living ones. This contrasting result was due to the different metrics used to reveal growth variance, as they

used MS while the SD of an AR1 model fitted to the detrended ring-width series was preferred here (see also Macalady and Bugmann 2014). This highlights the importance of carefully selecting the metrics, especially when estimating growth variance in rather short time series ( $p < 50$  yr). Other metrics may be used to characterize growth variance, such as the Gini index (Biondi and Qeadan 2008), the skewness in the ring-width distribution, or the number of pointer years (e.g., Das et al. 2007; see Appendix S1). We encourage methodological studies devoted to this question (e.g., Bunn et al. 2013), but also the use of early warning detection methods (Dakos et al. 2012) that would reveal temporal trends in variance and autocorrelation in ring-width series, as both seem to increase before tree death (Appendix S3; Camarero et al. 2015).

### Towards a general methodology

Currently, it is not possible to carry out a meta-analysis of the growth–mortality relationships already published, as most of the authors followed different methodologies to derive them (Appendix S1). The use of different field and statistical sampling schemes, growth variables, and model performance metrics strongly influences model selection, and especially the time window that should be used to calculate each growth variable. Our study provides some recommendations based on general good modelling practice in order to obtain reliable results that can be compared among studies.

First of all, large sampling designs are important, i.e., hundreds of trees of various sizes growing at different sites should be sought (see Appendix S4). Our study further highlights the necessity of selecting a field and statistical sampling scheme that is consistent with the research question and quantifying sampling-related biases. We suggest following a constant sampling approach to determine the time window that best discriminates growth patterns of living vs. dead trees, as changing the calibration dataset while increasing the maximum length of the time window tested may bias this selection. Applying this method is all the more important since it can detect intra-specific shifts in growth–mortality relationships.

Due to the exponential increase of computational power and of the robustness of statistical methods over the past years, novel approaches can now be used to derive growth–mortality relationships and optimize their performance. First, mixed-effect logistic models are strongly encouraged as they allow for quantifying the variability in growth–mortality relationships among sites and can remove the potential impact of local disturbances on mortality that are independent of tree growth. Second, multivariate models should include different types of growth variables (growth level, trend, variance, temporal autocorrelation, etc.). Third, the time windows used to calculate the growth variables should be selected using appropriate optimization procedures.



Finally, we strongly recommend using AUC as the model performance metric for univariate models, as it depends less on sample size and on the ratio of alive/death observations than alternative metrics (e.g., AIC). However, in case of models with multiple intercorrelated variables, penalizing for extra parameters is important. In these cases, we suggest evaluating models by considering multiple criteria, e.g., combining AUC with AIC to handle multi-collinearity. Lastly, if the main goal of a study is to implement the mortality model in a dynamic vegetation model (e.g., calibration with the sampling scheme  $T$  and considering observed stand-scale mortality rates), binary metrics should be used to prevent any systematic over- or underestimation of mortality rates (see Bircher et al. 2015).

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## SUPPORTING INFORMATION

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## DATA AVAILABILITY

Ring-width data associated with this paper have been deposited in a Dryad digital repository <http://dx.doi.org/10.5061/dryad.1bv6n>