# Spatial pattern of trees influences species productivity in a mature oak-pine mixed forest 

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## Title:

"Spatial pattern of trees influences species productivity in a mature oak-pine mixed forest"

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#### Abstract

: Spatial pattern has a key role in the interactions between species in plant communities. These interactions influence ecological processes involved in the species dynamics: growth, regeneration and mortality. In this study, we investigated the effect of spatial pattern on productivity in mature mixed forests of sessile oak and Scots pine. We simulated tree locations with point process models and tree growth with spatially explicit individual growth models. The point process models and growth models were fitted with field data from the same stands. We compared species productivity obtained in two types of mixture: a patchy mixture and an intimate mixture. Our results show that the productivity of both species is higher in an intimate mixture than in a patchy mixture. Productivity difference between the two types of mixture was $11.3 \%$ for pine and $14.7 \%$ for oak. Both species were favored in the intimate mixture because, for both, intraspecific competition was more severe than interspecific competition. Our results clearly support favoring intimate mixtures in mature oak-pine stands to optimize tree species productivity; oak is the species that benefits the most from this type of management. Our work also shows that models and simulations can provide interesting results for complex forests with mixtures, results that would be difficult to obtain through experimentation.


Keywords: Point process model; Spatially explicit growth model; Intimate mixture; Patchy mixture; Quercus petraea; Pinus sylvestris

## 1. Introduction

Since the beginning of the 1990s when the worldwide fight against biodiversity loss gained recognition (Earth summit, Rio de Janeiro, 1992), interest in mixed forests has been growing. Species composition has become a key criterion of sustainable forest management, as defined at the 2003 Vienna conference on forest protection in Europe (MCPFE et al. 2011). Moreover, several scientific studies have shown the advantage of setting up mixed stands compared to pure stands. For example, a mixture of tree species can reduce damage by phytophagous insects (Jactel and Brockerhoff 2007). Mixing species can also lead to an increase in stand productivity (Pretzsch and Schutze 2009; Vallet and Perot 2011) thanks to better resource exploitation and facilitation mechanisms between species (Kelty 2006). More recently, the question of how ecosystems will adapt to climate change has strengthened the interest in mixed forests (Lenoir et al. 2008). According to the insurance principle (McNaughton 1977), mixing tree species could mitigate the consequences of future climatic changes on forest ecosystem functioning by distributing the risks over the different species. In Europe, mixed-stand management is also a very important economic issue because the surface area these stands cover is considerable (MCPFE et al. 2011).

How to optimize the productivity of mixed forests, while at the same time preserving them, is therefore an important question for forest research. To reach this goal, managers need better knowledge and a more precise description of the factors that influence trees and species growth in mixtures. Spatial pattern is known to have a significant impact on species interactions which in turn impact ecological processes in plant communities (Mokany et al. 2008; Begon et al. 2006; Dieckmann et al. 2000). Spatial pattern refers to the organization of individuals in space and therefore reflects the local environment around each individual. This local environment modifies the expression of dynamic natural processes such as growth, mortality and regeneration (Barot et al. 1999; Courbaud et al. 2001). Thus, spatial pattern can
modify species productivity. For herbaceous species, Lamosova et al. (2010) showed that the type of spatial organization affected species productivity in mixtures, and depended on complicated interplay between interspecific and intraspecific competition: generally, in a random pattern the dominant species (superior competitors) increased their productivity, while the aggregated pattern was more favorable for the subordinate species (inferior competitors). However, few studies have dealt with the relationship between spatial pattern and productivity in forest stands, much less in mixed forest stands, partly because experimental approaches which take tree spatial patterns into account is difficult to set up for mixed forests (Vanclay 2006). Some authors used model simulations to overcome this difficulty. For example, Pukkala (1989) studied the effect of spatial pattern type on productivity in monospecific forest stands. To differentiate the effects of intra- and interspecific competitions in mixed stands, spatially explicit models have been developed (e.g. Vettenranta 1999). These growth models use competition indices that require to know the spatial position of trees in the stand. Spatial point processes, which are stochastic models that governs the location of points in space (Cressie 1993), were used to model the spatial structure of mixed forests (e.g. Pretzsch 1997). An approach using simulations with these kinds of realistic models is therefore an interesting way to investigate the impacts of spatial structure on mixed forests productivity (Pretzsch 1997).

In our work, we focused on the case of a mixed forest of sessile oak (Quercus petraea L.) and Scots pine (Pinus Sylvestris L.) in central France. In a previous study, the spatial pattern of these stands had already been accurately described (Ngo Bieng et al. 2006). The authors identified different spatial patterns of canopy trees: the two species showed an intraspecific spatial pattern characterized by a gradient from random to strong aggregation while the interpecific spatial pattern was characterized by a gradient from independence to interspecific repulsion. Moreover, Ngo Bieng et al. (2011) built point process models in order to simulate
the different spatial patterns identified in these stands. In another previous work in the same forest, Perot et al. (2010) developed individual growth models based on local competition indices and showed that within these stands, intraspecific competition had a more negative effect on growth than interspecific competition for both species. According to these results, species productivity may be enhanced in a mixture where intraspecific competition is minimized.

The aim of the present study was to clarify and quantify the impact of tree spatial pattern on species productivity in a mature mixed forest. To do this, we used point process models to simulate two contrasting types of existing spatial pattern that had been identified by Ngo Bieng et al. (2006). We then simulated tree growth with a spatially explicit individual based model using the point process realizations as the initial state, then we compared the productivity obtained in each type of spatial pattern. Finally, we assessed the contribution of spatial pattern to productivity variability of each species.

## 2. Methods

### 2.1 Study site and types of spatial pattern for the simulated oak-pine mixed stands

Our work focused on oak-pine mixed stands in the Orléans forest located in central France $\left(47^{\circ} 51^{\prime} \mathrm{N}, 2^{\circ} 25^{\prime} \mathrm{E}\right.$ ). With 35,000 hectares, the Orléans forest is France's largest public woodland. The forest is dominated by oaks (mainly Quercus petraea L.) and Scots pine (Pinus sylvestris L.). Between 2004 and 2007, 30 plots in the Orléans forest of between 0.5 and 1.25 ha were fully inventoried and mapped so as to run an in-depth study on the horizontal spatial pattern in these stands (Ngo Bieng et al. 2006). These plots were characterized by a mixed canopy composed of oak and pine, and by an understory dominated by oak. Between 2006 and 2007, nine of the 30 plots were selected to study growth in mixed oak-pine stands (Perot et al. 2010). In these plots (Table 1), the mean oak age as determined
by cores taken at breast height ranged from 52 to 78 years, and that of pines from 50 to 112 years. In any given plot, all the trees of the same species were approximately the same age, thus indicating a single cohort for pines and a single cohort for oaks. Pines were restricted to the canopy of the stands while oaks occupied both the canopy and the understory. In order to quantify the effect of tree spatial pattern on species productivity, in this study we focused on two contrasted types of canopy spatial pattern identified by Ngo Bieng et al. (2006). The first type of mixture is characterized by monospecific clusters (clusters of oaks and clusters of pines) with interspecific spatial repulsion (Fig. 1a). For this first type, repulsion occurs between clusters of individuals. This "patchy mixture" is henceforth called Type 1. In the second type of mixture, individual oaks and pines are randomly scattered (or only slight aggregated) (Fig. 1b). Here, the interspecific structure is characterized by repulsion between individuals at short distances and results in an intimate mixture at the plot scale. This "intimate mixture" is referred to as Type 2 in the following sections.

We also took the understory trees into account since they participate in stand productivity and are involved in local competition. In the studied stands, the understory is mainly composed of oak. Several types of spatial pattern have been identified for the understory in these stands (Ngo Bieng 2007). However, in eight of the nine plots where we measured tree growth, the spatial pattern of the understory was the same. Consequently, we chose only one type of spatial pattern for the understory and applied it to both types of mixture (Type 1 and Type 2 ).

### 2.2 Point process models of oak-pine mixed stands.

### 2.2.1 Point process models

The point process model we used in our study was a combination of classic point processes. In forestry applications, as in this study, the spatial pattern of the trees in a stand is assumed to result from a given point process. We therefore used known point processes to reproduce the
spatial features observed in the studied stands. In order to generate clustered or aggregated spatial point patterns, we used the Neyman-Scott (NS) point process (Tomppo 1986; Ngo Bieng et al. 2011). In order to generate the repulsion between individuals or groups of individuals, we used the "soft core" (SC) point process, which is a pairwise interaction process where pairs of points should not be closer than a threshold distance or "soft core" distance (Illian et al. 2008; Ngo Bieng et al. 2011). With the combination of these two point processes, Ngo Bieng et al. (2011) developed point process models fitted on field data to reproduce the spatial patterns of oak-pine mixed stands. These models took into account the spatial pattern of the two species when reproducing the observed spatial features, thus describing the spatial interactions between qualitative marks associated to the simulated spatial point process. For our work, we used the point process models developed by Ngo Bieng et al. (2011) to simulate oak-pine mixed stands. These models are described in the following subsections.

### 2.2.2 Point process model for spatial pattern of Type 1: patchy mixture

This point process model is a combination of Neyman-Scott processes (NS) and soft core processes (SC). Oak locations were simulated by an NS process. Pines locations were simulated by a NS process with an additional regularity constraint obtained through a SC process. The regularity constraint takes into account regularity at short distances, which is typical of the spatial pattern of pines (Ngo Bieng 2006). The regularity constraint is a threshold distance of regularity (dreg) which corresponds to the minimum distance allowed between two pines. To generate a more realistic regularity, if the distance between two pines is below the threshold distance, tree locations can be retained with a probability depending on the distance between the two trees (principle of the SC process). This probability varies linearly from 0 at a null distance to 1 at the threshold distance dreg. Interspecific repulsion was also simulated with a SC process and a repulsion distance drep. The Type 1 model has
six parameters (Table 2): the number of oak aggregates ( $n c l_{\text {oak }}$ ), the radius of the oak aggregates $\left(r c l_{o a k}\right)$, the number of pine aggregates ( $n c l_{\text {pine }}$ ), the radius of pine aggregates $\left(r c l_{\text {pine }}\right)$, the minimal intraspecific distance between pines or regularity distance (dreg), the minimal repulsion distance between oaks and pines or repulsion distance (drep).

### 2.2.3 Point process model for spatial pattern of Type 2: intimate mixture

This model is a combination of a NS process and a SC process. Pine locations were simulated with a NS process with a regularity constraint obtained with a SC process as explained for the previous model. Oak individuals were then randomly simulated with a repulsion distance also ensured with a SC process. Contrary to the previous model, the probability of accepting an oak closer to a pine than the threshold repulsion distance is constant and does not vary with the distance. This model has five parameters (Table 2): the number of pine aggregates ( $n c l_{\text {pine }}$ ), the radius of pine aggregates ( rcl $l_{\text {pine }}$ ), the intraspecific minimal distance between pines or distance of regularity (dreg), the minimal repulsion distance between oaks and pines (drep) and $p$ the constant probability to accept an oak tree at a distance lower than drep from a pine.

### 2.2.4 Point process model for oak understory

As mentioned previously, the understory was mainly composed of oak, and its spatial pattern did not vary much among the studied plots. We therefore chose to simulated only one type of spatial pattern for understory oaks: the most frequent type in the plots where growth was measured. For Type 1 and Type 2 mixtures, the simulated spatial pattern of understory oaks was therefore identical. As we did for the canopy trees, we used a point process model fitted on field data to simulate the locations of understory oaks (Ngo Bieng et al. 2011). This point process model simulates an attraction with the oaks in the canopy and a repulsion with the pines in the canopy. The point process model for the understory oaks was a combination of

NS and SC processes. First, understory oaks were simulated with a NS process. During this simulation, repulsion with the pines in the canopy was ensured with a SC process containing an additional constraint of attraction with canopy oaks. This attraction constraint between understory and canopy oaks was simulated by checking that each understory oak was at a distance below or equal to a given attraction distance. This model had four parameters (Table 2): the number of oak aggregates in the understory $\left(n c l_{u n d}\right)$, the radius of oak aggregates in the understory $\left(r c l_{\text {und }}\right)$, the distance of intraspecific attraction between understory oaks and canopy oaks (dattr), the distance of interspecific repulsion between understory oaks and canopy pines (drep).

Fig. 1 presents simulated stands for the patchy (Type1) and the intimate (Type2) mixtures.

### 2.3 Spatially explicit individual growth models

As mentioned above, we developed our growth model from data collected from nine plots in the Orleans forest. The nine plots cover the two types of mixture simulated in this work (Table 1). In each plot, we selected 30 oaks and 30 pines based on a stratified sampling method. The stratification variables were tree size and local environment (see Perot et al. 2010 for details). Sampled trees were cored to the pith at a height of 1.3 m . The cores were scanned and analyzed using the WinDENDRO software, version 2005a (Regent 2005), and ring width was measured to the nearest 0.01 mm . The COFECHA software (Grissino-Mayer 2002) was used to cross-date the individual ring-width series. The ring width analyses were performed on a final total of 230 oaks and 269 pines. Detailed information on past disturbances was not available for our plots (location and size of suppressed trees) so we chose the 6 years period from 2000 to 2005 to study tree growth because there had been no thinnings or storms during that time.

The growth model we developed is a spatially explicit individual based model based on local competition indices (Uriarte et al. 2004b). This model is similar to that presented by Perot et
al. (2010) but for the present study we added a plot random effect to account for factors influencing tree growth at the plot level (soil quality, stand age, stand density). The final model for each species was a linear mixed effect model. For both species, the competition indices were the basal areas of the oaks and pines belonging to the neighborhood of the target tree ( $C I_{o a k}$ et $C I_{p i n e}$ ). In a previous work on the same plots (Perot et al. 2010), several radii (5, 10 and 15 m ) were tested for the neighborhood so as to cover the range of radii reported in other studies (Canham et al., 2004; Stadt et al., 2007; Uriarte et al., 2004a) and to minimize the influence of edge effects when computing the competition indices. Based on model comparisons, the authors concluded that indices computed with a 10 m radius gave the best results. Based on this work, we defined the neighborhood as a 10 m radius circle around the target tree. These competition indices account for both intra- and interspecific competitions. For each species the final model was written as follows:

$$
\begin{equation*}
\Delta r_{i, k}=\left(\alpha_{0}+\alpha_{k}\right)+\left(\beta_{0}+\beta_{k}\right) \text { girth }_{i, k}+\lambda_{\text {oakk }} C I_{i, \text { aak }}+\lambda_{\text {pine }} C I_{i, \text { pine }}+\varepsilon_{i, k} \tag{1}
\end{equation*}
$$

where $\Delta r_{i, k}$ is the radial increment of tree $i$ for plot $k$ over a growth period of 6 years, girth $_{i, k}$ is the girth of tree $i$ at $1.3 \mathrm{~m}, C I_{i, o a k}$ and $C I_{i, p i n e}$ are the competition indices for oak competitors and pine competitors respectively, $\left\{\alpha_{0}, \beta_{0}, \lambda_{\text {oak }}, \lambda_{\text {pine }}\right\}$ are the parameters estimated for the fixed effects of the model, $\left\{\alpha_{k}, \beta_{k}\right\}$ are the parameters corresponding to the random part of the model (plot effect) and $\varepsilon_{i, k}$ is the residual part of the model.

Preliminary results showed that the variance of the residuals increased with the adjusted values. To correct for this heteroscedasticity, we modeled the variance of the residuals with the fitted values and a power function (Eq. 2), as suggested by Pinheiro and Bates (Pinheiro and Bates 2000):

$$
\begin{equation*}
\operatorname{Var}\left(\varepsilon_{i, k}\right)=\sigma^{2} \mid\left.\left(\text { fitted value }_{i, k}\right)\right|^{2 \delta} \tag{2}
\end{equation*}
$$

Where $\delta$ is the parameter of the variance model. The model was fitted using the R software version 2.14.0 (R Development Core Team 2011) with the lme function of the nlme package (Pinheiro et al. 2011).

### 2.4 Simulation experiment design

Initial stands for the two types of mixture were simulated with the point process models presented in section 2.2. Since stand density and tree size influence individual growth (see Eq. 1), in order to have exactly the same number of trees of each species and exactly the same dendrometric characteristics for the two types of mixture, we used the same tree list to simulate the initial stands for both mixture types. With this method, we ensured that the only parameter that changed between Type 1 and Type 2 mixtures was the spatial pattern of the trees. We carried out our simulations on a 1-ha plot (Table 4).

Both the spatial pattern within a mixture and growth show some variability. This variability was estimated from field data and was included in the point process models as well as in the individual growth model. To account for the different sources of variability, it was necessary to carry out several simulations with each model. We proceeded as follows: a) to account for variability in the spatial pattern within a mixture type, each type was simulated 200 times, b) to account for growth variability at the plot level, for each initial stand the parameters $\alpha_{k}$ et $\beta_{k}$ (Equation 1) were simulated 50 times, c) to account for variability in individual growth (residual variability), for each initial stand and each pair of values $\left\{\alpha_{k}, \beta_{k}\right\}$, individual tree growth was simulated 10 times following Equation 1. In all, we performed 200,000 simulations $(2 * 200 * 50 * 10)$. For each simulation, we calculated the basal area productivity for oak and for pine. All the simulations were performed in the Capsis platform with the oakpine1 module (Dufour-Kowalski et al. 2012).

### 2.5 Decomposition of the basal area productivity variability

Thanks to our simulation design, we were able to estimate the effects of several factors on the productivity of both species: 1) an effect related to the type of mixture, 2) an effect related to the variability in the spatial pattern within the type of mixture, 3 ) an effect related to the growth variability between plots (plots are nested in the type of mixture) and 4) an effect related to tree growth variability within the plot:

$$
\begin{equation*}
y_{i j k l}=\mu+\text { type }_{i}+p p_{i j}+\text { plot }_{i j k}+\varepsilon_{i j k l} \tag{3}
\end{equation*}
$$

Where $y_{i j k l}$ is the basal area productivity of one species, $\mu$ is the general mean, type $i_{i}$ is the type of mixture effect, $p p_{i j}$ is the spatial pattern random effect in the type, plot $_{i j k}$ is the plot random effect of the growth model in each point process realization, $\varepsilon_{i j k l}$ is the residual and corresponds to tree level variability in the growth model. The structure of our simulation design (balanced nested design) made it possible to decompose the variability of species productivity into different components and to estimate the contribution of each component to variability as follows (for simplicity, the variance $\sigma^{2}$ and the estimate of the variance are denoted identically):
$\sigma_{\text {total }}^{2}=\sigma_{\text {type }}^{2}+\sigma_{p p}^{2}+\sigma_{\text {plot }}^{2}+\sigma_{\text {res }}^{2}$ with $\left\{\begin{array}{l}\sigma_{\text {res }}^{2}=M S D_{\text {res }} \\ \sigma_{\text {plot }}^{2}=\left(M S D_{\text {plot }}-\sigma_{\text {res }}^{2}\right) / n_{\text {res }} \\ \sigma_{p p}^{2}=\left(M S D_{p p}-\sigma_{\text {res }}^{2}-n_{\text {res }} \sigma_{p l o t}^{2}\right) /\left(n_{\text {res }} n_{p l o t}\right) \\ \sigma_{t y p e}^{2}=\left(M S D_{\text {type }}-\sigma_{\text {res }}^{2}-n_{\text {res }} \sigma_{\text {plot }}^{2}-n_{\text {res }} n_{\text {plot }} \sigma_{\text {pp }}^{2}\right) /\left(n_{\text {res }} n_{\text {plot }} n_{p p}\right)\end{array}\right.$ Where $M S D$ is the mean square deviation for the different sources of variability, $n_{t y p e}=2, n_{p p}$ $=200, n_{\text {plot }}=50$, and $n_{\text {res }}=10$. We then assessed the importance of spatial pattern variability in the productivity variability of each species. The sum of $\sigma_{t y p e}^{2}$ and $\sigma_{p p}^{2}$ was considered to be the overall contribution of spatial pattern to productivity variability.

## 3. Results

The results of the growth model show that, for oak, the effect of oak competition on growth is about twice higher than the effect of pine competition (see $\lambda_{\text {oak }}$ and $\lambda_{\text {pine }}$ in Table 3). The magnitude of the effect of pine competition on pine growth is close to the effect of pine competition on oak growth ( $-0.085 \mathrm{~mm} \cdot \mathrm{~m}^{-2}$ and $-0.094 \mathrm{~mm} \cdot \mathrm{~m}^{-2}$ respectively). But contrary to oak, the competition index computed on oaks has no significant effect on pine growth. The results of the simulations show that productivity in Type 2 (intimate mixture) is higher than in Type 1 (patchy mixture) for both species (Fig. 2). The difference in productivity between Type 2 and Type 1 is more pronounced for oak than for pine: $+14.7 \%$ for oak and $+11.3 \%$ for pine.

The productivity values obtained for oak and pine show some variability. If we combine the results from the two types of mixture, oak productivity varies from 0.23 to $0.36 \mathrm{~m}^{2} \cdot \mathrm{ha}^{-1} \cdot$ year $^{-1}$ (first and ninth deciles) with a coefficient of variation of 0.175 (ratio of the standard deviation to the mean). Pine productivity varies from 0.19 to $0.35 \mathrm{~m}^{2} \cdot \mathrm{ha}^{-1} \cdot \mathrm{year}^{-1}$ (first and ninth deciles) with a coefficient of variation of 0.228 . Variability in pine productivity is thus slightly higher than that of oak.

The results also show that most of the productivity variability is explained by plot effect, which represents $86 \%$ of the total variability for pine and $67 \%$ for oak (Fig. 3). The spatial pattern (type of spatial pattern + random effect in the type) explains $12 \%$ of the variability for pine and $31 \%$ for oak. The overall effect of spatial pattern on oak productivity is important. Even if the individual growth variability within a plot is high, it has little impact on the overall productivity variability (between 1 and $2 \%$ of the total variability). Variability in spatial pattern within a mixture type also has a relatively little impact, though the effect on oak productivity (5\%) is slightly higher than on pine productivity (2\%).

## 4. Discussion

### 4.1 Spatial pattern and species productivity

Spatial pattern plays a key role in the interactions between species in plant communities (Dieckmann et al. 2000). These interactions influence ecological processes involved in the species dynamics: growth, regeneration and mortality (Begon et al. 2006). Our results show that the productivity of sessile oak and Scots pine is higher in an intimate mixture (Type 2) than in a patchy mixture (Type 1). Our work has made it possible to estimate the difference in species basal area productivity between the two types of mixture. This difference was $11.3 \%$ for pine and $14.7 \%$ for oak (Fig. 2). These figures are comparable to those of Pukkala (1989) who simulated Scots pine productivity in pure stands for different spatial patterns. He found that volume productivity was $10 \%$ lower in aggregated spatial patterns compared to regular spatial patterns. Our results also show that the plot effect explains a large part of the productivity variability (Fig. 3). The plot effect, estimated with the growth model, includes several factors that affect tree growth: (i) a site effect - soil conditions vary from one plot to another and affect species productivity, (ii) an age effect - young stands have higher productivity and finally, (iii) a density effect - denser stands generally have higher productivity (Vallet and Perot 2011). The variability obtained for pine productivity is similar to that of oak productivity but is much more influenced by plot effects (Fig. 3).

### 4.2 Influence of spatial and growth interactions

Intra- and interspecific competition are crucial to understand the effect of mixture on forest productivity and forest dynamic (Kelty 2006; Forrester et al. 2006). As in the study of Perot et al. (2010), our results showed that, for both species, intraspecific competition had a more negative effect on growth than interspecific competition (see parameters $\lambda_{\text {oak }}$ and $\lambda_{\text {pine }}$ in Table 3). Oak had little impact on pine growth probably because pines had a greater girth than oaks
on average (Table 1). The light interception by the pine foliage is lower than the light interception by the oak foliage (Balandier et al. 2006; Sonohat et al. 2004). This may help to explain that in our oak model, the interspecific competition was lower than the intraspecific competition. The two species involved have different light requirements but also different root distribution patterns (Brown 1992). The complementarity in nutrient and water use could also contribute to explain why intraspecific competition was more severe than interspecific competition. The local competition and the spatial features of each mixture type help to explain the results of this work. Two spatial features vary simultaneously between Types 1 and 2: the intraspecific pattern and the interspecific pattern. Ripley's function and inter-type function (Ripley 1977; Lotwick and Silverman 1982; Perot and Picard 2012) can be used to characterized and compare these two dimensions. On average in the patchy mixture (Type1), there are more oaks around an oak tree than in the intimate mixture (Type 2) (see L functions at 10 m for oak in Fig. 4). Consequently, the competition index $I C_{o a k}$ is higher, on average, in Type 1 than in Type 2. In contrast, in Type 1 mixture, there are fewer pines on average around an oak tree than in Type 2 (see inter-type functions at 10 m in Fig. 4). Consequently, the competition index $I C_{\text {pine }}$ is lower, on average, in Type 1 than in Type 2. In addition, the parameters of the growth model must be examined. Parameters $\lambda_{\text {oak }}$ and $\lambda_{\text {pine }}$ (Table 3) show that oak competitors $\left(I C_{o a k}\right)$ have a more negative effect on oak growth than do pine competitors $\left(I C_{\text {pine }}\right)$ ( $\lambda_{\text {oak }}$ is more negative than $\lambda_{\text {pine }}$ ). In the intimate mixture (Type 2 ) there are more pines around oaks than in the patchy mixture (Type 1) and pines are less competitive than oaks. This explains why oak productivity is higher, on average, in the intimate mixture than in the patchy mixture. For one particular simulation, the final result is complex because productivity depends on both intra- and interspecific competition (estimated through parameters $\lambda_{\text {oak }}$ and $\lambda_{\text {pine }}$ and also on intra- and interspecific spatial patterns. Variability in the spatial pattern of a mixture type thus explains why oak productivity in Type 2 is not always
higher than in Type 1 (Fig. 2). The reasoning is similar for pine but the result is easier to analyze because there is no interspecific competition parameter in the individual growth model. For pine, productivity depends only on the intraspecific spatial pattern.

### 4.3 Influence of species assemblage and stand age

The effect of spatial pattern on species productivity in mixed stands should depend on species assemblage. In our study, both oak and pine were favored in the intimate mixture because, for both species, intraspecific competition was more severe than interspecific competition. Other authors have also shown that interspecific competition was lower than intraspecific competition (e.g. Forrester and Smith 2012), while some studies have shown the opposite in some conditions (e.g. Pretzsch et al . 2010). Intensity of interactions may also change with species assemblages. Further works involving other species are therefore necessary to generalize our results. Moreover, for tree species, the competition relationship between species may depend on stand developmental stage (Filipescu and Comeau 2007; Cavard et al. 2011). Pine is a fast growing species compared to oak (Duplat and Tran-Ha 1997; Perot et al. 2007). In young stage, pine is probably more competitive than oak. Consequently, oak productivity could be favored by a patchy mixture at an earlier stage. In addition, Getzin et al. (2006) showed that interspecific competition is less intense at older stages than at younger stages, probably due to the spatial sharing of resources. In our study, this would explain why the mixture type had less impact on pine productivity than on oak productivity, and why pine is more influenced by plot effects (site, age, density) than oak.

## Conclusion

Our study is innovative in that we worked on a mature mixed forest. For such complex forests, models and simulations can provide interesting quantitative results that would be difficult to obtain through experimentation. The two mixture types that we tested are realistic
oak-pine mixtures found in central France (Ngo Bieng et al. 2006). Our results show that their spatial differences are contrasted enough to have an impact on the productivity of both species in the mixture. From a practical point of view, our work shows the interest of favoring intimate mixtures in mature oak-pine stands to optimize tree species productivity. Oak is the species that benefits most from this type of management. In order to achieve more general results, further work is needed to determine the change in competition between oak and pine over time.

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## 7. Tables

Table 1: Dendrometric characteristics of the nine plots used for growth models (Orléans Forest, France). BA = basal area; Other = other broadleaf tree species; $D=$ mean diameter at a height of $130 \mathrm{~cm} ;$ Age = mean age of the cored trees at a height of $\mathbf{1 3 0} \mathbf{c m} ; \mathbf{H o}=$ dominant height. Only the height of the sample trees was measured. The dominant height was estimated with a measure of the dominant diameter and a height-diameter relationship fitted for each species and each plot using the sample trees; $\mathbf{P P}=\mathbf{t y p e}$ of spatial pattern, $1=$ patchy mixture, 2 = intimate mixture, 3 = intermediate type with cluster of pines and oaks randomly scattered; For diameters and ages, values represent the mean with the standard deviation in parentheses.

| Plot | Area (ha) | $\begin{gathered} \mathrm{BA}_{\text {oak }} \\ \left(\mathrm{m}^{2} . \mathrm{ha}^{-1}\right) \\ \hline \end{gathered}$ | $\begin{gathered} \mathrm{BA}_{\text {pine }} \\ \left(\mathrm{m}^{2} . \mathrm{ha}^{-1}\right) \\ \hline \end{gathered}$ | $\begin{gathered} \mathrm{BA}_{\text {other }} \\ \left(\mathrm{m}^{2} . \mathrm{ha}^{-1}\right) \\ \hline \end{gathered}$ | $\begin{gathered} \mathrm{BA}_{\text {total }} \\ \left(\mathrm{m}^{2} . \mathrm{ha}^{-1}\right) \\ \hline \end{gathered}$ | $\begin{aligned} & \mathrm{D}_{\text {oak }} \\ & (\mathrm{cm}) \\ & \hline \end{aligned}$ | $\begin{aligned} & \mathrm{D}_{\text {pine }} \\ & (\mathrm{cm}) \end{aligned}$ | Age $_{\text {oak }}$ | Age $_{\text {pine }}$ | $\begin{gathered} \begin{array}{c} \mathrm{Ho}_{\text {oak }} \\ (\mathrm{m}) \end{array} \\ \hline \end{gathered}$ | $\begin{gathered} \mathrm{Ho}_{\text {pine }} \\ (\mathrm{m}) \\ \hline \end{gathered}$ | PP |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| P108 | 0.80 | 9.6 | 19.8 | 1.4 | 30.8 | 17.7 (6.74) | 36.2 (5.31) | 68 (4.3) | 66 (2.5) | 22.3 | 23.0 | 2 |
| P178 | 1.00 | 16.5 | 10.0 | 1.5 | 28.0 | 21.5 (10.49) | 36.5 (7.56) | 78 (4.6) | 77 (1.8) | 21.1 | 22.1 | 1 |
| P184 | 0.75 | 10.9 | 12.0 | 2.1 | 25.1 | 17.5 (8.88) | 36.3 (7.76) | 71 (8.6) | 68 (4.2) | 21.9 | 20.8 | 3 |
| P216 | 0.50 | 11.2 | 12.1 | 0.9 | 24.1 | 17.0 (6.39) | 27.8 (7.6) | 52 (2.8) | 50 (2.2) | 18.8 | 19.0 | 2 |
| P255 | 1.00 | 12.6 | 10.5 | 1.1 | 24.2 | 17.8 (7.54) | 31.7 (6.25) | 69 (5.9) | 62 (4.6) | 20.1 | 19.7 | 2 |
| P534 | 0.50 | 12.2 | 19.6 | 1.0 | 32.7 | 16.6 (6.54) | 37.4 (6.5) | 59 (2.3) | 83 (3.2) | 22.1 | 22.5 | 2 |
| P563 | 0.50 | 13.6 | 11.9 | 0.2 | 25.7 | 25.1 (10.12) | 35.6 (4.58) | 70 (3.1) | 69 (2.3) | 24.5 | 23.0 | 2 |
| P57 | 1.00 | 11.2 | 11.4 | 0.4 | 23.0 | 16.7 (6.36) | 34.3 (6.41) | 67 (7.1) | 62 (3.1) | 20.4 | 21.2 | 1 |
| P78 | 0.70 | 14.7 | 16.5 | 1.0 | 32.2 | 20.1 (7.48) | 42.2 (8.79) | 62 (5.2) | 112 (17.5) | 21.8 | 25.6 | 2 |

Table 2 Parameters in the point process models. $n c l_{s p}=$ number of aggregates for species $s p ; r l_{s p}=$ radius of aggregates for species $s p ;$ dreg $=$ distance of regularity which corresponds to the minimum distance allowed between pines; drep $=$ repulsion distance between oaks and pines; und = oak understory; dattr = distance of intraspecific attraction between understory oaks and canopy oaks.

| Species | Tree position | Type of spatial pattern | Parameters in the point process model |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | $\begin{gathered} n c l_{\text {pine }} \\ \left(\mathrm{ha}^{-1}\right) \end{gathered}$ | $\begin{gathered} r c l_{\text {pine }} \\ (\mathrm{m}) \end{gathered}$ | dreg <br> (m) | $\begin{aligned} & \text { ncl oak } \\ & \left(\text { ha }^{-1}\right) \end{aligned}$ | $\begin{gathered} r_{c} l_{\text {oak }} \\ (\mathrm{m}) \end{gathered}$ | drep <br> (m) |
| Oak and pine | Canopy | Type 1 (Patchy mixture) | 13 | 18 | 5 | 7 | 17 | 18 |
|  |  |  | $\begin{gathered} n c l_{\text {pine }} \\ \left(\mathrm{ha}^{-1}\right) \end{gathered}$ | $r c l_{\text {pine }}$ (m) | dreg <br> (m) | drep <br> (m) | $p$ |  |
| Oak and pine | Canopy | Type 2 (Intimate mixture) | 38 | 8 | 10 | 4 | 0.15 |  |
|  |  |  | $\begin{aligned} & n c l_{u n d} \\ & \left(\mathrm{ha}^{-1}\right) \end{aligned}$ | $\begin{gathered} r c l_{\text {und }} \\ (\mathrm{m}) \end{gathered}$ | dattr <br> (m) | drep <br> (m) |  |  |
| Oak | Understory | Type 1 and Type 2 | 37 | 12 | 52 | 2 |  |  |

Table 3 Parameter estimates of the spatially explicit individual growth model (see Eq. 1).

|  |  | Parameter estimates |  |  |  |  | Model statistics |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Intercept $\alpha$ (mm) | $\begin{aligned} & \text { girth } \\ & \beta \\ & \left(\mathrm{mm} \cdot \mathrm{~cm}^{-1}\right) \end{aligned}$ | $C I_{o a k}$ <br> $\lambda_{\text {oak }}$ <br> (mm.m ${ }^{-2}$ ) | $\begin{aligned} & C I_{\text {pine }} \\ & \lambda_{\text {pine }} \\ & \left(\mathrm{mm} \cdot \mathrm{~m}^{-2}\right) \end{aligned}$ | $\delta^{\text {a }}$ | RSE | df | AIC |
| Oak | Estimates | 3.335 | 0.126 | -0.196 | -0.094 | 0.526 | 1.013 | 218 | 1196 |
|  | Std. error | 1.202 | 0.018 | 0.042 | 0.024 |  |  |  |  |
|  | P -value | 0.006 | <0.001 | <0.001 | <0.001 |  |  |  |  |
|  | $\sigma_{\text {plot }}$ | 2.036 | 0.048 |  |  |  |  |  |  |
| Pine | Estimates | 2.711 | 0.0654 |  | -0.0855 | 0.621 | 0.838 | 258 | 1413 |
|  | Std. error | 1.054 | 0.0094 |  | 0.0241 |  |  |  |  |
|  | P -value | 0.011 | <0.001 |  | $<0.001$ |  |  |  |  |
|  | $\sigma_{\text {plot }}$ |  | 0.0145 |  |  |  |  |  |  |

[^0]Table 4 Dendrometric features of the initial stand used in simulations (stand area = $\mathbf{1} \mathbf{h a}$ ). For girth, the value in parentheses corresponds to the standard deviation.

|  | Number of trees |  |  | Girth (cm) |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
| Species | Canopy | Understory | mean | min. | max. |  |
| Oak | 284 | 208 | $53.2(23.2)$ | 23 | 129 |  |
| Pine | 237 | 2 | $97.1(21.0)$ | 33 | 160 |  |

## 8. Figure captions

a) Patchy mixture (Type 1)
b) Intimate mixture (Type 2)


- Oak (canopy)
$\Delta$ Pine (canopy)
- Oak (understory)
- Pine (understory)

25 meters

Fig. 1 a) Patchy mixture (Type 1) simulated with the point process models; b) Intimate mixture (Type 2) simulated with the point process models.


Fig. 2 Productivity comparison between the two mixture types for oak and pine.

Oak
Decomposition of the productivity variability


Pine
Decomposition of the productivity variability


Fig. 3 Decomposition of the productivity variability for oak and pine following Equation
(3). The different sources of variability are: type of mixture (Type), spatial point pattern within the type (PP), plot random effect (Plot), and tree random effect (Tree).


Fig. 4 L function and intertype L function calculated with 1000 simulations of Type 1 and Type 2 mixtures. For the intraspecific $L$ function, $L(r)$ less than 0 indicates spatial regularity, $L(\mathbf{r})$ greater than 0 indicates spatial aggregation. For the intertype $L$ function, $L(r)$ less than 0 indicates spatial repulsion between the two species, $L(r)$ greater than 0 indicates spatial attraction between the two species.


[^0]:    ${ }^{\mathrm{a}} \delta$ is the parameter of the variance model (see Eq. 2).

