Can the biomass-ratio hypothesis predict mixed-species litter decomposition along a climatic gradient?

Antoine Tardif1,2, Bill Shipley1,*, Juliette M. G. Bloor2 and Jean-François Soussana2

1Département de Biologie, Université de Sherbrooke, Sherbrooke, QC, J1K 2R1, Canada and 2INRA, UR874, Unité de Recherche sur l’Ecosystème Prairial, F-63100 Clermont-Ferrand, France

* For correspondence. E-mail bill.shipley@usherbrooke.ca

Received: 26 July 2013 Returned for revision: 26 September 2013 Accepted: 5 December 2013 Published electronically: 29 January 2014

INTRODUCTION

Predicting ecosystem function in a changing environment is a fundamental challenge for ecologists. Ecosystems are characterized by complex interactions, both between component species and between these species and abiotic conditions. Consequently, ecosystem processes reflect the combined properties and performances of the constituent species in a changing environment; this complexity makes their study and prediction difficult in a general context.

One way to link the aggregated response of an ecosystem or a community property with the effects of each species is provided by Grime’s ‘biomass-ratio hypothesis’ (Grime, 1998). This hypothesis states that ecosystem properties are driven by the characteristics of dominant species in the community. This has been shown in various ecosystem processes (e.g. Mokany et al., 2008). The biomass-ratio hypothesis was further operationalized by Garnier et al. (2004) via community-weighted (or ‘aggregated’) means. This is formalized in Eqn (1), in which $e_i$ is the average per capita effect of species $i$ in community $c$, $p_i$ is the relative abundance of species $i$ and $E_c$ is the community-weighted effect on the ecosystem summed over all $S$ species present. The average per capita effects of each species ($e_i$) can be further decomposed into the contributions of different functional traits possessed by each species.

$$E_c = \sum_{i=1}^{S} p_i e_i$$  (1)

As required by Garnier’s version of the biomass-ratio hypothesis, this equation calculates the overall ecosystem response ($E$) as an additive function of individual species effects ($p_i e_i$). If synergistic or antagonistic interactions are present, then the biomass-ratio may not be an accurate predictor of the observed responses. If so, the biomass-ratio hypothesis would not hold because the effect of a particular species on the ecosystem would depend not only on its relative abundance, but also on the combination of species in the community.

Some studies have shown that litter decomposition rates can be correlated to functional traits such as leaf carbon and nitrogen content or leaf dry matter content (e.g. Kazakou et al., 2006; Santiago, 2007; Fortunel et al., 2009). In this context, Lavorel and Garnier (2002) and Suding et al. (2008) proposed a framework which differentiates between environmental responses to litter decomposition based on the biomass-ratio hypothesis.

Key Results Both positive and negative prediction errors occurred. Despite this, the biomass-ratio hypothesis was true as an average claim for all sites ($r = 0.91$) and for each site separately, except for the climatically intermediate site, which showed mainly synergistic deviations. Variability decreased with increasing species richness and in less favourable climatic conditions for decomposition.

Conclusions Community-weighted mean values provided good predictions of mixed-species litter decomposition, converging to the predicted values with increasing species richness and in climates less favourable to decomposition. Under a context of climate change, abiotic variability would be important to take into account when predicting ecosystem processes.

Key words: Biomass-ratio hypothesis, BMRH, climatic gradient, community-aggregated traits, community-weighted means, Festuca rubra, litter decomposition, Lolium perenne, Taraxacum officinale, Trifolium repens.
The decomposition of mixed-species plant litter, the subject of our study, provides an interesting model in this context because it constitutes an important part of the nutrient cycle and because there is already a large literature on this topic, although relatively few of these studies concern the effect of species richness in interspecific mixtures of litter (Wardle et al., 1997; reviewed by Gartner and Cardon, 2004; Hättenschwiler et al., 2005). Since litter decomposes in mixtures in most ecosystems, and since interactions occur frequently within communities, interspecific litter mixtures often show synergistic or antagonistic effects. Among the possible mechanisms for interspecific interactions, the review by Hättenschwiler et al. (2005) identified nutrient transfer, the effects of specific secondary compounds, interactions across trophic levels and improved microenvironmental conditions. Such non-additive effects have been reported in many decomposition experiments involving species mixtures (Gartner and Cardon, 2004). These non-additive interactions should therefore induce a bias in the prediction of mixed-species decomposition rates based on monoculture values (e.g. Wardle et al., 1997; Hector et al., 2000; Perez-Harguindeguy et al., 2008).

Considering this potential source of error, one could expect that predictions based on the biomass-ratio hypothesis would be incorrect since they implicitly assume that there are no interactions between species. However, some studies (Perez-Harguindeguy et al., 2008; Pakeman et al., 2011) showed that the use of the biomass-ratio hypothesis can still be accurate since the observed values did not produce any overall bias relative to predicted values. How can a hypothesis that assumes the absence of interspecific interactions provide accurate predictions of a phenomenon in which such interactions are common? Our study aims to explain such an apparent paradox.

Based on the ‘idiomsyncratic hypothesis’ of Lawton (1994), an ‘idiomsyncratic annulment’ hypothesis (Tardif and Shipley, 2013) was recently proposed to explain why the biomass-ratio hypothesis often provides good predictions of mixed-species litter decomposition despite the common presence of interactions between species. It assumes that interactions exist, but, being equally likely to be both positive or negative, they cancel each other out, leading to a global average rate around the predicted value. Furthermore, increasing the number of species increases the number of potential interactions which more fully cancel each other out, leading to a better prediction, i.e. through a decrease of the variation in observed decomposition rates between mixtures having the same number of species (Tardif and Shipley, 2013). However, Tardif and Shipley’s (2013) empirical demonstration was conducted for tree species under controlled conditions, and so might not be applicable to ecosystems or communities in situ. Indeed, many environmental factors (e.g. temperature, humidity, soil pH, differing decomposer communities) affect litter decomposition rates (e.g. Berg et al., 1993; Aerts, 1997; Murphy et al., 1998; Cornwell et al., 2008; Zhang et al., 2008; Bontti et al., 2009) and environmental variability could strongly affect species interactions in mixture.

In particular, if the abiotic conditions drive deviations in a particular direction (synergistic or antagonistic), and thus unbalance the equilibrium between positive and negative deviations, then the assumption of symmetrical bias around zero required by idiomsyncratic annulment would be false. Moreover, as more limiting abiotic conditions reduce the overall rate of decomposition, a reduction of the variability between mixtures would be expected.

Among the abiotic conditions known to affect decomposition rates, mean annual temperature (MAT) and mean annual precipitation (MAP) are often used in litter decomposition experiments (e.g. Epstein et al., 2002; Zhang et al., 2008; Bontti et al., 2009). We therefore concentrate on variation in MAT and MAP. Thus, under natural conditions, the effects of abiotic environmental variability (i.e. climate) represent a second source of error in the prediction of decomposition rates, especially in species mixtures. We expect that abiotic conditions limiting to decomposition will reduce the potential variability of decomposition rates between species. In our study, we tested the ‘idiomsyncratic annulment’ hypothesis on abiotic variability, applied to the decomposition of grassland litter. We measured both within- and between-site variability in decomposition, the latter based on a correlated gradient of both temperature and precipitation, which is described in more detail below.

Here, we report a litterbag experiment along a climatic gradient to examine the effects of the species richness and the abiotic conditions on litter decomposition. Specifically, we tested (1) whether the observed specific decomposition rates in mixed-species litters are equal to the community-weighted average decomposition rates between species. However, some studies (Perez-Harguindeguy et al., 2008; Pakeman et al., 2011) showed that the use of the biomass-ratio hypothesis can still be accurate since the observed values did not produce any overall bias relative to predicted values. How can a hypothesis that assumes the absence of interspecific interactions provide accurate predictions of a phenomenon in which such interactions are common? Our study aims to explain such an apparent paradox.

Based on the ‘idiomsyncratic hypothesis’ of Lawton (1994), an ‘idiomsyncratic annulment’ hypothesis (Tardif and Shipley, 2013) was recently proposed to explain why the biomass-ratio hypothesis often provides good predictions of mixed-species litter decomposition despite the common presence of interactions between species. It assumes that interactions exist, but, being equally likely to be both positive or negative, they cancel each other out, leading to a global average rate around the predicted value. Furthermore, increasing the number of species increases the number of potential interactions which more fully cancel each other out, leading to a better prediction, i.e. through a decrease of the variation in observed decomposition rates between mixtures having the same number of species (Tardif and Shipley, 2013). However, Tardif and Shipley’s (2013) empirical demonstration was conducted for tree species under controlled conditions, and so might not be applicable to ecosystems or communities in situ. Indeed, many environmental factors (e.g. temperature, humidity, soil pH, differing decomposer communities) affect litter decomposition rates (e.g. Berg et al., 1993; Aerts, 1997; Murphy et al., 1998; Cornwell et al., 2008; Zhang et al., 2008; Bontti et al., 2009) and environmental variability could strongly affect species interactions in mixture.  

In particular, if the abiotic conditions drive deviations in a particular direction (synergistic or antagonistic), and thus unbalance the equilibrium between positive and negative deviations, then the assumption of symmetrical bias around zero required by idiomsyncratic annulment would be false. Moreover, as more limiting abiotic conditions reduce the overall rate of decomposition, a reduction of the variability between mixtures would be expected.

Among the abiotic conditions known to affect decomposition rates, mean annual temperature (MAT) and mean annual precipitation (MAP) are often used in litter decomposition experiments (e.g. Epstein et al., 2002; Zhang et al., 2008; Bontti et al., 2009). We therefore concentrate on variation in MAT and MAP. Thus, under natural conditions, the effects of abiotic environmental variability (i.e. climate) represent a second source of error in the prediction of decomposition rates, especially in species mixtures. We expect that abiotic conditions limiting to decomposition will reduce the potential variability of decomposition rates between species. In our study, we tested the ‘idiomsyncratic annulment’ hypothesis on abiotic variability, applied to the decomposition of grassland litter. We measured both within- and between-site variability in decomposition, the latter based on a correlated gradient of both temperature and precipitation, which is described in more detail below.

Here, we report a litterbag experiment along a climatic gradient to examine the effects of the species richness and the abiotic conditions on litter decomposition. Specifically, we tested (1) whether the observed specific decomposition rates in mixed-species litters are equal to the community-weighted average decomposition rates between species.

**MATERIALS AND METHODS**

**Study sites**

The experiment was conducted in three sites distributed along correlated altitudinal–climatic gradients near Clermont-Ferrand in central France: Laqueuille (45°38′33″N, 2°44′11″E, 1045 m a.s.l.), Theix (45°43′22″N, 3°1′21″E, 880 m a.s.l.) and Crouël (45°46′32″N, 3°8′48″E, 340 m a.s.l.). These sites have a contrasting climate. Laqueuille is the coolest (MAT = 8.0 °C) and wettest (MAP = 1100 mm), Theix is intermediate (MAT = 8.7 °C and MAP = 830 mm) and Crouël is hottest (MAT = 12.0 °C) and driest (MAP = 570 mm, with a drought in August). The climatic data were provided by CLIMATIK, an agroclimatic database (MétéoFrance and INRA convention). However, this experimental design was not made to test a single climatic factor such as temperature or precipitation, but rather to test a gradient from more favourable to more limiting abiotic conditions for decomposition.

**Construction of litter bags**

The experiment involved a set of four herb species: common dandelion (Taraxacum officinale L., F.H. Wigg), white clover (Trifolium repens L.), perennial ryegrass (Lolium perenne L.) and red fescue (Festuca rubra L.). The species were chosen in...
order to maximize the differences in leaf functional traits known to affect decomposition rates, especially leaf dry matter content (Kazakou et al., 2006; Fortunel et al., 2009; Pakeman et al., 2011) among the species present in our sites; leaf dry matter content values from Grime et al. (2007) are 137 (T. officinale), 179 (T. repens), 212 (L. perenne) and 267 mg g⁻¹ (F. rubra). Litter of the four herbaceous species was gathered in autumn 2009 by cutting the senescent leaves of several individuals between vegetation and decomposition. We first plotted

\[ \ln \left( \frac{M_i(t)}{M_i(0)} \right) = a_i + b_i t + c_i t^2 + \varepsilon_{ij} \]  

(2)

as a function of time, where \( M_i(t) \) is the mass of the litter mixture \( i \) in litterbag \( j \) at time \( t \), and \( M_i(0) \) is the initial mass of litter mixture \( i \) (1.5 ± 0.1 g) in litterbag \( j \). As we observed non-linear trends, we did not use the simple negative exponential model of Olson (1963) (i.e. linear on a semi-log scale) but instead we modelled the ln-transformed proportional mass loss as a quadratic function of time [Eqn (2)], which gave a better empirical fit to our data and where \( a_i, b_i \) and \( c_i \) are coefficients of the equation for mixture \( i \), \( t \) is the harvest date (days) and \( \varepsilon_{ij} \) is the deviation of mixture \( i \) in litterbag \( j \) from its predicted value.

\[ k_i(t) = b_i t + 2c_i t \]

(3)

Of course, this quadratic function makes no biological sense if extrapolated beyond our data, since it would eventually result in a mass gain. As in Tardif and Shipley (2013), we used mixed models, because the mass loss depends on both fixed (time) and random (the error between litterbags of a given mixture) effects.

Obtaining observed decomposition rates, and those predicted by the biomass-ratio hypothesis. A first derivative as a function of time of Eqn (2) provided \( \dot{k}_i(t) \), i.e. the observed decomposition rates of mixture \( i \) at each time \( t \) [Eqn (3)].

\[ \dot{k}_i(t) = \frac{\sum_{n=1}^{S} p_{in} b_n(t)}{\sum_{n=1}^{S} p_{in}} \]

where \( p_{nj} = 1/S \) is the initial mass proportion of species \( n \) in the mixture \( i \), and \( b_n \) is the empirically estimated (observed) specific decomposition rate of species \( n \) in mixture \( i \) when measured in monoculture at time \( t \). These values were obtained by calculating the community-weighted values of \( b_i \) and \( c_i \) for mixture \( i \), i.e.

\[ \hat{b}_i = \frac{\sum_{n=1}^{S} p_{in} b_n}{\sum_{n=1}^{S} p_{in}} \quad \text{and} \quad \hat{c}_i = \frac{\sum_{n=1}^{S} p_{in} \varepsilon_n}{\sum_{n=1}^{S} p_{in}} \]

Substituting these into Eqn (3). We thus obtained Eqn (4), with the predicted decomposition rate of mixture \( i \) at time \( t \).

\[ \dot{k}_i(t) = \hat{b}_i t + 2\hat{c}_i t \]

(4)

Testing the mass-ratio hypothesis. The biomass-ratio hypothesis assumes that the mean observed decomposition of a mixture within a site is equal to the values predicted from a community-weighted approach [Eqn (5a)], with \( \varepsilon_{ij} \) representing the residual variation around a single mixture.

\[ \ln \left( \frac{M_i(t)}{M_i(0)} \right) = \hat{a}_i + \hat{b}_i t + \hat{c}_i t^2 + \varepsilon_{ij} \]  

(5a)
To test this hypothesis, two new variables were created for each mixture by multiplying the community-weighted values of each \( b_i \) and \( c_i \) by time and its square, respectively. For each site, we then fitted a mixed-model regression following Eqn (5b) but also including the litter mixtures as the between-group random variable. If the biomass-ratio hypothesis is strictly true, then the estimated coefficients of \( \beta \) and \( \gamma \) are both equal to 1 and there will be no significant between-mixture variation associated with these parameters. If the biomass-ratio hypothesis is true on average, but not for every mixture, then the estimated coefficients of \( \beta \) and \( \gamma \) are both equal to 1 but there will be significant between-mixture variation associated with these parameters. If the biomass-ratio hypothesis is not true even on average, then one or both of \( \beta \) and \( \gamma \) will be significantly different from 1.

**Testing the hypothesis of idiosyncratic annulment.** Idiosyncratic annulment occurs when the biomass-ratio hypothesis is not strictly true, i.e. when the ecosystem response (mixed-species litter decomposition) is not simply the effect contributed by each species weighted by its relative abundance, but when the differences between the observed effect and that predicted by the biomass-ratio hypothesis tend to cancel out more completely as the species richness of the community increases. We tested this in two ways.

To test this hypothesis, we first used Monte Carlo simulations to describe the probability distribution of the standard deviation of mixed-species litter decomposition rates. Next, the standard deviation of the observed deviations between observed and predicted (CWM) decomposition rates was regressed on the number of species per mixture (two, three or five), and a permutation test was used (Manly, 1997) to compare slopes, which were assumed to be independent of the species richness of the mixture. Simulations were done in R version 2.12.0 (R Development Core Team, 2010; see also Tardif and Shipley, 2013).

### RESULTS

**A quadratic function for modelling the dynamics of litter decomposition**

Visual inspection of ln(mass loss) over time suggested a non-linear response. We therefore fitted a linear mixed model regressing the natural logarithm of observed mass ratio (remaining mass divided by initial mass) as a function of ‘site’, ‘time’, ‘time2’ and interactions (as fixed factors), and ‘species mixture’ nested in ‘time’ and ‘time2’ (as random terms). An analysis of variance (ANOVA) identified highly significant \((P < 0.001)\) effects associated with all fixed effects factors: sites, harvest times, squared time and interactions with site. In particular, adding a quadratic term for time significantly improved the model \((F\text{-value} = 127.72, P\text{-value} < 0.001)\). As significant differences between sites were also found, we thus used a quadratic model to model the dynamic of decomposition, for each site separately. This implies a varying decomposition rate over time.

**Empirical patterns of mass loss**

Mass loss during the experimental period ranged from 16 to 86 % across mixtures (Fig. 1). There was an initial rapid rate of mass loss, which then decreased with time (Fig. 1). An initial examination of the data also showed a much stronger variability in the site of Laqueuille (the coolest and wettest site) than in Theix (the intermediate site) and Crouël (the hottest and driest site).

<table>
<thead>
<tr>
<th>Site</th>
<th>Remaining mass (ln scale)</th>
<th>Time (weeks)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laqueuille</td>
<td>0</td>
<td>0 – 20</td>
</tr>
<tr>
<td>Theix</td>
<td>-2</td>
<td>0 – 25</td>
</tr>
<tr>
<td>Crouël</td>
<td>-1.5</td>
<td>0 – 25</td>
</tr>
</tbody>
</table>

**Fig. 1.** The natural logarithm of the observed remaining mass of leaf litter as a proportion of initial mass, plotted against time (weeks) from the beginning of the experiment, for each litterbag and each of three sites (open circles). Dotted grey and solid black lines represent the quadratic function of each mixture (mono- and polycultures) and their mean estimate, respectively.
The biomass-ratio hypothesis

Our first hypothesis was that the observed specific decomposition rates in mixed-species litter mixtures were equal to the community-weighted values \((k_{\text{CWM}})\) obtained from monocultures (i.e. the biomass-ratio hypothesis), within each site. For a given site, the biomass-ratio hypothesis is strictly true if neither of the slope coefficients \(\beta\) and \(\gamma\) [Eqn (5b)] differs significantly from 1 and if there is also no significant between-mixture variance associated with them. The biomass-ratio hypothesis is true as an average claim but not true in every litter mixture if there is a significant between-mixture variance associated with either \(\beta\) or \(\gamma\). Our results showed a significant between-mixture variance of the two fixed slopes \(\beta\) and \(\gamma\) for each of the tested sites in any site \((P = 0.002, <0.001\text{ and } <0.001\text{ for Crouël, Theix and Laqueuille, respectively})\). Notably, the fixed slopes estimates did not differ from unity in both Laqueuille and Crouël (Fig. 2), supporting the biomass-ratio hypothesis on average but not in every mixture. However, in Theix, \(\beta\) and \(\gamma\) were both different from unity. In this site, the biomass-ratio hypothesis was therefore rejected (Fig. 2).

At the Laqueuille and Crouël sites, the measured decomposition rates in mixtures showed no systematic bias and good correlations with respect to the predicted values based on the independently measured monoculture values and the biomass-ratio hypothesis (Fig. 3). Both positive and negative deviations were observed relative to the predicted rates, but a general 1:1 relationship exists even though there is no statistical necessity for this relationship. However, the site of Theix showed mainly synergistic deviations (decomposition rates with higher absolute values than those predicted by the CWM) which contradict the biomass-ratio hypothesis, even as an average claim.

The ‘idiosyncratic annulment’ hypothesis

The variability of observed decomposition rates, measured as the standard deviation in \(k\)-values between mixtures having the same number of species, decreased with increasing number of species in the mixture (from one to three species, Fig. 4). Although results at Theix and Crouël showed similar variability with two- and three-species mixtures, the decreasing variability was clearest for the coolest and wettest site (Laqueuille), at all levels of species richness.

We then used Monte Carlo simulations to determine if the observed variability of the \(k\)-values was less than expected based purely on the fact that CMWs are sums of random variables. Using 20,000 Monte Carlo simulations for each, we compared the measured range of \(k\)-values for litters having the same number of species with the range of \(k\)-values expected, assuming that this variation was due solely to sampling variation arising from different species compositions, as explained in the Materials and Methods. In both Laqueuille and Theix, for two- and three-species mixtures and at all times, the variation of observed values was consistent with the values of Monte Carlo simulations (e.g. at 18 weeks, Laqueuille, \(P = 0.813, 0.177\); Theix, \(P = 0.440, 0.785\) for the two- and three-species mixtures,
respectively). In Crouël, however, at 18 weeks for both two- and three-species mixtures and 28 weeks for two-species mixtures, the variations of observed values were significantly lower than the values of Monte Carlo simulations (e.g. \( P = 0.001 \) and 0.012 for the two- and three-species mixtures, respectively, at 18 weeks). Thus, the decrease in variability of decomposition rates between mixtures having the same number of species was consistent with mathematical expectations across sites and harvest dates, except at two harvest dates at Crouël; here the significantly lower variability indicates a biological effect.

To test further the idiosyncratic annulment sensu stricto, permutation tests were applied to determine if the observed variability significantly decreases as the number of species in the mixtures increases, at a given time. Analyses based on 50,000 permutations could not detect a significant decrease in the variability of the prediction errors with increase of species richness (e.g. at 18 weeks, \( P = 0.538, 0.462 \) and 0.621 for Laqueuille, Theix and Crouel, respectively).

**The climate effect**

The mean decomposition rates of Laqueuille (the coldest and wettest site) and Crouël (the hottest and driest site) were similar at the beginning of the experiment, then diverged (Fig. 5A). The site of Laqueuille showed the most favourable conditions for decomposition on average, with the highest mean \( k \)-values calculated over mixtures and dates. The climatically intermediate site of Theix had the slowest mean decomposition rate at the beginning of the experiment, but then increased to reach a rate similar to that of Laqueuille at the end of the experiment.

Figure 5B shows that the variability of observed decomposition rates was largest at the coolest and wettest site (Laqueuille) and lowest at the warmest and driest site (Crouel) at all times during the experiment. Within each site, we observed a U-shaped curve with two distinct and opposite patterns centred around 10–18 weeks (i.e. the fourth and the fifth harvest time; June 22 and August 17, respectively). During the first part of the experiment (from 0 to 10–18 weeks), the variability (standard deviation of the observed \( k \)-value between mixtures at each site) decreased to reach a minimum in July–August, then increased during the second part of the experiment (from 10–18 to 28 weeks). Before this shift, the observed variability of decomposition rates between mixtures (within sites) decreased with slowing rates over time. After the shift, observed variability increased even as decomposition rates continued to decrease. The fastest decomposing site (Laqueuille, the coolest and wettest site) then showed the highest level of variability, and the slowest decomposing site (Crouël, the hottest and driest site) showed the lowest variability.

**DISCUSSION**

The complexity of ecosystems makes it difficult to predict their process rates from information on their component parts. One suggested way of linking the properties of component species to ecosystem processes is to use community-weighted values of species-level traits (Garnier et al., 2004), a proposal derived from Grime’s biomass-ratio hypothesis (Grime, 1998). Our main objective was to determine to what degree CWM values provide accurate predictions of ecosystem processes. Specifically, can monoculture decomposition rates, weighted by the relative abundance of each species in a mixed-species litter, be used to predict mixed-species decomposition rates in the field?

Here, we focused on two of the main sources of errors occurring in such predictions: interactions between species causing deviations from the linear sums predicted by community-weighted values and temporal variation in the abiotic environment (climate) causing correlated responses in such deviations.

Specifically, we tested the following hypotheses: (1) the decomposition rates of mixed-species litter agree with Grime’s biomass-ratio hypothesis, (2) deviations from the predicted CWMs tend to cancel out as the number of species in the mixture increases (i.e. idiosyncratic annulment); and (3) the variation of decomposition rates between litter mixtures decreases as the abiotic conditions of the site become more limiting.

This study reveals contrasting results within and across sites. All sites considered, the CWM values of mixture decomposition rates based on abundance-weighted monoculture values predicted the actual mixture decomposition rates well (\( r = 0.91 \))
and with no bias (Fig. 3). Thus, at this scale, the biomass-ratio hypothesis provides a good quantitative description of the mixed-species response. However, the results differ between sites. In the two climatically extreme sites (Laqueuille, the coolest and wettest; and Crouël, the hottest and driest), good support was found for the biomass-ratio hypothesis on average even though this was not true in every mixture. Furthermore, there was no statistical difference (Fig. 2) between values predicted by the biomass-ratio hypothesis and that actually observed. Indeed, we found abundant evidence of both positive and negative deviations in litter mixtures from the expected decomposition rates using CWMs, but such deviations were symmetrically distributed around zero. As a result, the average differences between observed and predicted rates of mixtures were approximately zero. This is consistent with results conducted under constant environmental conditions (Tardif and Shipley, 2013).

However, in the climatically intermediate site (Theix), the biomass-ratio hypothesis was not true, even on average (Fig. 2), and only 34% of the variance in the observed decomposition rates was explained by the community-weighted values in this site. The equilibrium between positive and negative deviations from predictions that we found in Laqueuille and Crouël was not present in Theix, where most interactions were synergistic (Fig. 3).

Considering the similarity of the responses in the two other sites and under controlled conditions, this result from the intermediate site is surprising. Given that our experiment under variable environmental conditions involved only three sites, it is difficult to identify a causal explanation for this contrasted behaviour in Theix. The same soil, species mixtures and sources of litter were used at all three sites. The same decomposer community existed in all three sites at the beginning of the experiment, although this probably changed over the course of the experiment. The most obvious difference between the sites was in the abiotic conditions.

Our results show that the biomass-ratio hypothesis can lead to biased predictions within particular sites due to correlated responses between mixtures in a particular direction, but still produce unbiased and relatively accurate predictions at a larger scale involving different sites. This is similar to what was found by Tardif and Shipley (2013): CWMs can both over- and underestimate observed rates within a given species mixture but, taken over many mixtures, they provide unbiased predictions. Therefore, it is essential to consider the scale of the study when using CWMs to predict decomposition rates.

The species richness of a mixture is also an important factor when predicting the global decomposition rate. As already found for example in Keith et al. (2008), Perez-Harguindeguy et al. (2008) and Tardif and Shipley (2013), the variability of observed decomposition rates in mixtures decreased as the number of species increased (Fig. 4). However, the magnitude of this decrease varied between sites. A part of this effect is a necessary mathematical consequence of (1) the reduction of variation of weighted random variables compared with the values of the component variables and (2) the covariance occurring when different mixtures contain species in common (Tardif and Shipley, 2013). However, beyond this purely statistical effect, we had expected a biological one, caused by a combinatorial increase in the number of possible species interactions with an increasing species richness of the litter (Tardif and Shipley, 2013), which we called ‘idiosyncratic annulment’. This biological effect would decrease the variability of the prediction errors even below that expected due to purely statistical reasons. We did not observe this except for at Crouël (the hottest and driest site) after 10–18 weeks of the decomposition experiment. Using permutation tests to test the idiosyncratic annulment sensu stricto, we could not detect any additional decrease in variance with increasing species richness of the litter mixture. This is consistent with our previous results in Tardif and Shipley (2013), observed under controlled conditions. Either the idiosyncratic annulment effect does not exist or it is too weak to be detected. Despite this, the decreasing variability of observed decomposition rates with an increasing number of species due to purely statistical effects still makes the predictions using CWMs more accurate in litter mixtures containing more species.

Our third hypothesis was that the variation between the decomposition rates measured in the litter mixtures decreases as the abiotic conditions of the site become more limiting to decomposition, i.e. when the average measured decomposition rates at the site are lower.

Our results supported this hypothesis, when comparing the responses both (1) across the sites differing in average conditions of temperature and precipitations and (2) within each site as the conditions became more or less favourable to decomposition (cf. figure of temperatures and precipitations in Supplementary Data Fig. S1).

First, the coldest and wettest site (Laqueuille) showed the most favourable conditions for litter decomposition along the gradient (Fig. 5A), with the highest mean k-values across the whole experiment (all mixtures considered) and also having the greatest variability in decomposition rates between litter mixtures. In contrast, the hottest and driest site (Crouël) had the poorest conditions for litter decomposition, the lowest mean k-values across the whole experiment (all mixtures considered) and also the least variability in decomposition rates between litter mixtures.

Then, as shown in Fig. 5B, we also found a clear effect of the climate on the variability of the observed k-values within sites: the variability decreased with the hotter and drier climate occurring in July–August, especially for the site of Crouël (cf. figure of temperatures and precipitations in Supplementary Data Fig. S1). Increased precipitation and/or decreasing temperatures during the autumn were associated with a rise in variability of decomposition rates between litter mixtures (especially in Theix and Laqueuille). This reduction in the variability of decomposition rates between litter mixtures at a given site and time that was induced by a hotter and drier climate was presumably due to a reduction in the activity of decomposers under such conditions which may have restricted the strength of the interactions between the species in the litter.

A consequence of our results is that we can expect better predictions using the CWM under a climate less favourable to decomposition. As proposed in Tardif and Shipley (2013), the spatial scale will also be a determinant in the prediction of species mixture decomposition rates because increasing spatial scales have higher species richness which would decrease the variability between different species mixtures in the litter. Considering the differences of structure (e.g. spatial pattern of species distribution) and species richness between communities (or biomes), we expect better predictions in communities (or
biomes) with higher species richness, for a given spatial scale (e.g. temperate grasslands vs. temperate forest, or tropical vs. temperate forest). However, even at larger spatial scales, CWMs of monoculture decomposition rates would be expected to have lower predictive accuracy in climates more favourable to decomposition.

Our results are also important in the current context of climate change and should be included in the predictive ecosystem models. Indeed, IPPC predictions for the next decades of this study area (central France region) imply warming and summer droughts (IPCC, 2001). If our results are more generally true, then such warmer and drier conditions would result in better predictions of litter decomposition rates using the CWM.

Beyond the example of litter decomposition and as previously suggested in Tardif and Shipley (2013), these results support the idea that the biomass-bias hypothesis, operationalized as CWMs, could offer the opportunity to predict ecosystem processes at larger spatial scales and in a changing environment. These initial results are encouraging, but other studies must be conducted to extrapolate to other plant types and climates. Also, it would be interesting to study the variations of functional diversity of the involved species, as it might be an important source of error in the predictions.

SUPPLEMENTARY DATA

Supplementary data are available online at www.aob.oxfordjournals.org and consist of Figure S1: further details concerning Supplementary data are available online at www.aob.oxfordjournals.org.

ACKNOWLEDGEMENTS

We thank David Shipley, Alexandre Salcedo and the technical staff of Crouël, Theix and Laqueuille for technical assistance, and Dr Katja Klumpp for revisions and helpful comments. We also thank the Centre for Forest Studies (CEF) in Quebec, Canada, and INRA Grassland Ecosystem Research Unit (UREP), France, for their collaboration. This research was funded by the Natural Sciences and Engineering Research Council (NSERC) of Canada.

LITERATURE CITED


