Looking for compensation at multiple scales in a wetland bird community

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Abstract

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Compensatory dynamics, during which community composition shifts despite a near-constant total community size, are usually rare: synchronous dynamics prevail in natural communities. This is a puzzle for ecologists, because of the key role of compensation in explaining the relation between biodiversity and ecosystem functioning. However, most studies so far have considered compensation in either plants or planktonic organisms, so that evidence for the generality of such synchrony is limited. Here, we extend analyses of community-level synchrony to wetland birds. We analyse a 35-year monthly survey of a community where we suspected that compensation might occur due to changes in water levels, favouring birds with different habitat preferences, and potential competition. We perform both year-to-year analyses by season, using a compensation/synchrony index, as well as multiscale analyses using a wavelet-based measure, which allows for both scale- and time-dependence. We analyse synchrony both within and between guilds, with guilds defined either as tightknit phylogenetic groups or larger functional groups. We find that abundance and biomass compensation is rare, likely due to the synchronizing influence of climate (and other drivers) on birds, even after considering several temporal scales of covariation (during either cold or warm seasons, above or below the annual scale). Negative covariation in abundance at the guild or community level did only appear at the scale of a few months or several years. We also found that synchrony varies with taxonomic and functional scale: the rare cases where compensation appeared consistently in year-to-year analyses were between rather than within guilds, using functional groups. Our results suggest that abundance compensation may have more potential to emerge between broad functional groups rather than between species, as well as at relatively long temporal scales (multiple years for vertebrates), above that of the dominant synchronizing driver.

6 **Keywords:** compensation; synchrony; biodiversity; birds; time series; wavelets

27 Introduction

Density compensation occurs when individuals of a given species replace individuals of other species within a community, either because of explicit competitive processes or shifts in environmental drivers that change selection pressures (Gonzalez & Loreau, 2009). The community 30 as a whole then exhibits lower abundance variation than its constituent species (Gross et al., 31 2014): some degree of compensation or asynchrony is therefore a prerequisite to stabilization at the community level (Loreau & de Mazancourt, 2013). Understanding why environmental variation may lead to compensation is relatively easy: if species have different environmental preferences (e.g., thermal optima), and the environment changes over time, different species will be fittest at different points in time. As a consequence, relative abundances will shift over time even though the community abundance or biomass as a whole may remain relatively stable (Gonzalez & Loreau, 2009). However, the conditions for compensation to happen also depend on the particulars of the interactions between and within species in the community. 40 Compensation is particularly likely to occur when temporal environmental variation com-41 bines with a space constraint or with a strongly limiting resource, so that individuals are close to competing in a zero-sum game (sensu Hubbell, 2001 or lottery-style models, Chesson, 43 1994). When the total community size is constant over time, and the composition fluctuates, negative covariation between abundances then emerges by design (Loreau & de Mazancourt, 45 2008) since no species can increase without at least another species decreasing in abundance. Outside of this zero-sum scenario, in models where Lotka-Volterra competition is combined with temporal environmental variability, theoretical research has revealed that increased interspecific competition might not always increase species compensation (Ives et al., 1999) and might even decrease it (i.e., increase species synchrony instead, Loreau & de Mazancourt,

2008, 2013), though this depends on the fluctuation regime. Thus, in a world where total

community size varies, predicting whether compensatory dynamics can occur is intrinsically

difficult (van Klink et al., 2019).

Early investigations of the frequency of synchronous vs compensatory dynamics focused on the variance ratio, that is, the variance of the sum of the community biomass divided by the sum of the variance of the component species biomasses (Houlahan et al., 2007; 56 Gonzalez & Loreau, 2009). Unfortunately, this metric is not appropriate for communities 57 subjected to community-wide environmental forcing (Ranta et al., 2008), because a main 58 environmental driver (e.g., temperature or light) may synchronize species abundances or 59 growth rates at some temporal scale, creating large variance in community-wide biomass, in 60 spite of strongly competitive dynamics. Further research has therefore focused on specific 61 timeframes during which compensatory dynamics may be found (e.g., below the annual scale 62 at which temperature fluctuations tend to synchronize species dynamics, Vasseur et al., 2014). 63 Despite efforts to look for more meaningful temporal scales in community-level time series, 64 temporal compensation has remained surprinsingly elusive in the field (Houlahan et al., 2007; Vasseur et al., 2014); but see Morgan Ernest et al. (2008); Christensen et al. (2018). Most 66 datasets used so far to evaluate temporal compensation vs synchrony involve planktonic 67 organisms (Vasseur & Gaedke, 2007; Vasseur et al., 2014) or terrestrial plants (Bai et al., 2004; Houlahan et al., 2007; Gross et al., 2014; though see Bell et al., 2014 in fishes, Morgan Ernest et al., 2008 in mammals and van Klink et al., 2019 in beetles). Here, we take advantage of a long-term bird abundance time series in a natural reserve, with records every month for 35 years, allowing us to dig deeper into patterns of synchrony, at several temporal and taxonomic or functional scales.

Taxonomic and functional scales should indeed be main modulators of synchrony/compensation.
On the one hand, compensation can be high between similar and closely related species. If
two species of ducks A and B share almost the same niche, individuals from either species
experience similar competition from species A or B, and should feel the effects of other species
in the community identically. This favours priority effects (Fukami, 2015), with chance due

to movement events determining whether species A or B locally dominates, which can then provide compensation at the landscape level (Loreau et al., 2003). On the other hand, it could be argued that these two similar duck species will precisely respond in similar ways 81 to environmental variables, which tends to obfuscate compensation. Hence, more dissimilar 82 species or groups (within the same trophic level nonetheless) could exhibit more compensa-83 tion (Bai et al., 2004; Morin et al., 2014; van Klink et al., 2019) because they are more likely 84 to respond to the environment in an asynchronous manner (sensu Loreau & de Mazancourt, 85 2013). Surprisingly, such compensation between guilds has been less well explored empirically 86 than within guilds, even though there is actually some empirical evidence for compensation 87 between dissimilar guilds (e.g., Bai et al., 2004; Roscher et al., 2011; Sinclair et al., 2013; van 88 Klink et al., 2019). In this paper, we explore the level of compensation/synchrony within 89 or between guilds of a wetland bird community, along either taxonomic or functional classi-90 fications. Although a functional classification might appear intuitively more appealing, our 91 knowledge of functional traits is necessarily partial and imperfect, so that a taxonomic de-92 scription can sometimes be preferable (Clark, 2016). Our dataset is ideally suited to examine the presence of synchrony or compensation at different scales given that (i) it is a highly temporally resolved time series with respect to the species typical generation times, but it also extends well beyond generation time (timespan of 35 years) and (ii) the reserve where the data has been collected was subjected to a major management change c. 2006 (change in water levels), favouring different types of wetland birds (so that over long timescales, there is a real potential for changes in community composition).

$_{\scriptscriptstyle 100}$ Material and Methods

101 Data

The monthly time series used for the statistical analyses have been collected at the Teich 102 Ornithological Reserve, Arcachon Bay, France (44.64°N / -1.02°E), by the staff of the Teich 103 reserve, over the whole study period (1981-2016). A species list of the frequent birds is 104 provided in SI Appendix S1. The reserve comprises 120 ha of wetlands, and the counts have 105 been aggregated at the reserve scale (summed over 18 sectors where the counts are actually 106 performed, using binoculars). We use for each species the maximum observed abundance over 107 a month, which provides a "monthly snapshot" of the bird abundance, that has been used to 108 monitor the reserve since its inception. When abundance values are not reported for certain 109 species and months, we replace them by zeroes. Given the sustained observation effort (all 110 sectors are patrolled multiple times throughout the month by the staff, amateur ornithologists 111 visiting the reserve daily and communicating their findings to the reserve staff), we consider 112 that the absence of counts for a given species signals its true absence from the reserve. This 113 creates some zero abundances for rare species at the monthly scale. We have not attempted 114 to "correct" those zeroes (e.g., inferring the "missing" data with a model assuming that our 115 reserve is a subsample of a regional population) because doing so would have compromised 116 the patterns of local synchrony/compensation. However, we did check that having such 117 zeroes in the monthly time series cannot affect our conclusions (see SI). In the statistical 118 analyses, we use seasonally averaged abundances (plotted in Fig. 1), as well as the original 119 monthly data (presented in Appendix S2). We defined two seasons based on observations 120 of bird presence. We defined a 'warm season', from May to August, and a 'cold season' 121 as the months between November and February of the following year. From an ecological 122 viewpoint, this seasonal classification separates wintering birds from summer residents (some 123 of whom are breeding). This makes sense biologically because the two communities have 124

different requirements and could respond differentially to abiotic drivers. It is also useful from a more statistical perspective, as there is a partial shift in composition between the seasons, though winter and summer communities greatly overlap (i.e., species with greater abundances in the reserve in winter have also some summer residents, though these may be different individuals). The dynamics of species abundances in the Teich reserve bird community show a marked signature of seasonality (Fig. 1).

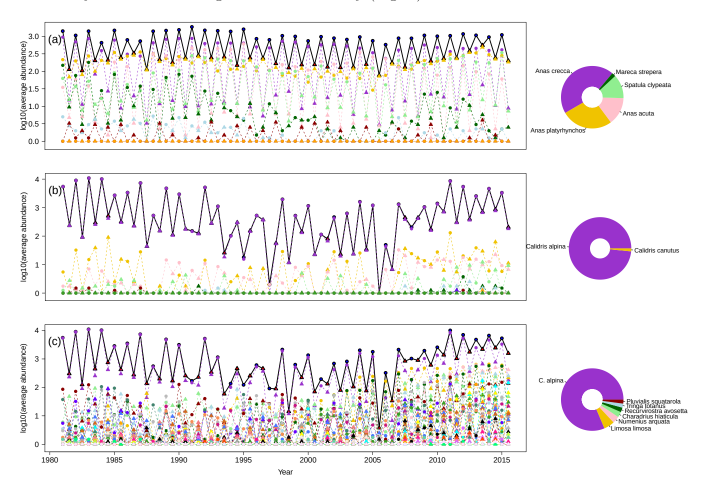


Figure 1: Time series of seasonally averaged abundance for ducks of the tribe *Anatini* (a), calidrids (b, *Calidris* genus), and all waders (c, including calidrids). The solid black lines (on top of each panel) represent the summed average abundances for each guild, dotted lines represent average abundance for each species. Circles represent the cold season and triangles, the warm season. The coloured symbols below the curves represent each species abundances, with species composition on the right side on the donut plots for the most abundant species (over 1% of relative abundance in the group considered). We added 1 to abundances before log-transforming to avoid issues with zero values.

Bird taxonomic and functional groups

The reserve is dominated by waders and waterfowl (ducks, geese and swans). These two functional groups collectively represent 68% of the total number of observed birds over the 133 years and are always present on site. Two fairly common phylogenetic groups, both in 134 abundance and occurrence, are members of the Anatini tribe (corresponding previously to 135 the Anas genus, Gonzalez et al., 2009) in ducks and members of the Calidris genus in waders. 136 Waders and ducks have different environmental preferences, with ducks (and waterfowl more 137 generally) preferring water levels allowing them to dabble (or dive for Aythini), while waders 138 usually forage on mudflats. A list of all birds found frequently in the reserve is presented 139 in Appendix S1; aside from waders and waterfowl, other common species include herons, 140 egrets and cormorants (see below). Among the fish eaters, grebes and gulls were frequently 141 counted; a few raptors were present as well. 142

To examine compensation between and within the waders and waterfowl categories, we contrasted analyses using a taxonomic classification of the species (i.e., between and within phylogenetic groups such as genera) and a functional classification of the species (26 species of waders vs 17 species of waterfowl). The waterfowl group includes all anatids (ducks, geese and swans in particular) as well as the common coot (Fulica atra, an abundant species here, which is a Rallidae but resembles a duck in morphology and foraging habits; hence its inclusion).

In addition to our main analyses on waders and waterfowl, we also "zoomed in" on a set of species that were known to exhibit potentially compensatory dynamics through competition for roosting sites: the great cormorant (*Phalacrocorax carbo*), the little egret (*Egretta garzetta*) and the grey heron (*Ardea cinerea*). The little egret and grey heron abundances were summed because of their similar requirements (i.e., they form a small functional group).

55 Statistical Analyses

56 Year-to-year analyses

We used for year-to-year analyses the synchrony index η defined by Gross *et al.* (2014), which is constructed as the mean cross-correlation between each species abundance and the summed abundances of the rest of the community (eq. 1):

$$\eta = \frac{1}{n} \sum_{i} \operatorname{Corr}(X_i, \sum_{j \neq i} X_j)$$
 (1)

where X_i is the abundance or biomass of species i in a community of n species and 160 the correlation is computed over the years. This synchrony index varies between -1 (perfect 161 compensation, total abundance is constant) and 1 (complete synchrony), while 0 represents a 162 case where all populations fluctuate independently. Contrary to other indices (e.g., Loreau & 163 de Mazancourt (2008)'s ϕ), this index is independent from the richness n of the community (or 164 more generally the number of system components) and its overall stability (Blüthgen et al., 165 2016; Hallett et al., 2016). This is particularly important here as we perform analyses at 166 different taxonomic scales, and therefore with a different n in eq. 1. All analyses performed 167 with abundance in the main text are performed with biomass in Supporting Information 168 Appendix S4. 169

We computed the synchrony index η over all available years, but separately for cold and warm seasons, using the codyn package in R (Hallett *et al.*, 2016). That is, we constructed two community-level time series of species abundances, one for the cold season and one for the warm season. To do so, we averaged monthly bird abundances, for each species, over the season duration. In follow-up analyses, we also differentiated periods before and after 2006, given that a management change occurred within the reserve in 2006. We considered both the synchrony within a given guild (e.g., among species of the *Calidris* genus) or between guilds (e.g., between the summed abundances of the 7 species of tribe *Anatini* and the sum of

the 7 Calidris species). In the latter case of between-guilds comparisons, we summed species together before seasonal averaging, to consider seasonal averages of the monthly guild-level abundance. Finally, we computed η within the community of the 60 most frequent birds.

We computed the statistical significance of the synchrony index by comparing the ob-181 served values to the distribution of η under the null hypothesis (Gouhier & Guichard, 2014), 182 which amounts to cross-correlations of value zero between species abundances (or guild-level 183 abundances, when considering taxonomic or functional groups). The challenge, in order to 184 construct such null hypothesis, is to remove all cross-correlations while keeping the exact 185 same autocorrelation in each individual time series. Therefore, for each set of time series 186 (each combination year \times season for a given community), we constructed 1000 "surrogates" 187 in which we kept auto-correlations but removed cross-correlations between time series. There 188 are multiple ways to erase cross-correlations depending on the resolution of the considered 189 community. Within guilds, we shifted the time-series (Purves & Law, 2002) while between 190 guilds (two groups only), we used a frequency-based approach (Iterative Amplitude-Adjusted 191 Fourier Transform or IAAFT, see Schreiber & Schmitz, 2000). We first explain the shift-based 192 approach: the suite of abundance values (after seasonal averaging) is displaced by a random 193 temporal lag τ , so that a value y_t is now found at $y_{t+\tau}$. At the boundary (the end of the time series), remaining points are displaced towards the beginning of the time-series, which 195 implements a toroidal shift. This method works well when comparing many times series corresponding to the multiple species. However, when computing synchrony across only two groups 197 (between guilds), spurious cross-correlations could emerge with a shift-based approach as the 198 number of possible combinations is more limited. Therefore, to test for synchrony between 199 the summed abundances of two guilds or taxonomic units, we used the more sophisticated 200 IAAFT method (Schreiber & Schmitz, 2000), which retains the frequency spectrum of the 201 time series while randomising its values. We obtained 1000 sets of randomised time series 202 for each computed synchrony index. We then compared the number of η_{H0} values which 203

exceeded or were inferior to the observed value to compute the p-value (North et al., 2002):
we use the ratio (r+1)/(n+1) where r is the number of surrogate values that are $\geq \eta_{obs}$ or $\leq \eta_{obs}$, and n is the number of surrogates. Independence of species was rejected at the
10% threshold with a Benjamini-Hochberg correction, as we compare across 2 seasons and 3
periods (all years, before 2006, after 2006), with partially overlapping data. This was found
satisfactory based on simulated data, although power is low for detecting compensation (i.e.,
the null cannot always be rejected) when only two groups are compared.

211 Wavelet analyses

In addition to the time-domain analyses above, we performed wavelet analyses at multiple 212 temporal scales, ranging from a month to several years. Wavelet analyses provide information 213 on community synchrony for a given temporal scale or frequency, as well as a given location 214 in time along the time series. This was done at the whole community level, including the 60 215 most frequent bird species, and for the rich wader and waterfowl communities, as well as the 216 group formed by the great cormorant, grey heron and little egret. All wavelet analyses take 217 as input the monthly time series data. Based on the work by Keitt (2008) and follow-up by Vasseur et al. (2014), we used the wavelet modulus ratio to measure the synchrony between 219 time series 220

$$\rho(t,s) = \frac{\int_{-\infty}^{+\infty} \frac{1}{\sqrt{2\pi}} e^{-\frac{1}{2}(\frac{\tau-t}{s})^2} |\sum_i w_i(\tau,s)| d\tau}{\int_{-\infty}^{+\infty} \frac{1}{\sqrt{2\pi}} e^{-\frac{1}{2}(\frac{\tau-t}{s})^2} \sum_i |w_i(\tau,s)| d\tau}$$
(2)

where $w_i(t,s)$ is the continuous Morlet wavelet transform of species i at time t for scale s, and $|\cdot|$ is the modulus of the complex number. The numerator considers the total abundance variation $|\sum_i w_i(\tau,s)|$ at a given temporal scale s and location in time τ , while the denominator considers a weighted sum of the fluctuation amplitude of each species $(\sum_i |w_i(\tau,s)|)$.

The Gaussian weights in the numerator and denominator ensure that $\rho(s,t)$ is specific to

scale s and time t. This index ρ is close to 0 when species (or compartments) compensate and reaches 1 when they are synchronous (Keitt, 2008). Significance of high and low values of ρ were evaluated using a 10% overall level. The null hypothesis was constructed using the IAAFT algorithm (Schreiber & Schmitz, 2000), using 1000 surrogate time series, and computing of the corresponding ρ values for each one (similar to Cazelles et al., 2014). The robustness of the wavelet approach to the presence of exactly zero values is tested in SI Appendix S6. Appendices S7 and S8 further test the ability of ρ to identify compensation or synchrony in cases of skewed species abundance distribution, either in the mean or in the amplitude of temporal variation.

All datasets and statistical analyses are available in a GitHub repository https://github.com/fbarraquand/BirdTimeSeries_Teich and stored at Zenodo [will be done for the final version] (Picoche, Aluome & Barraquand, 2020). Finally, we want to highlight a conceptual issue worth keeping in mind: both ρ and η indicate synchrony when reaching one, but such synchrony should be understood as the reciprocal of compensation rather than exactly synchronized peaks and troughs for all species (i.e., phase synchrony). Unlike phase synchrony, compensation and community-level synchrony depend on the distribution of abundance variation within the community. In the limit case where a single species abundance fluctuates more than all others combined, compensation may not even be reachable, since variation in the abundance of that dominant species cannot be offset by changes in numbers of other species. Only when species densities have commensurate temporal variability will the concepts of community-level synchrony and phase synchrony exactly match.

Results

$_{ t 48}$ Synchrony within phylogenetic or functional groups

Using a taxonomic classification of the community, focusing on the genera Calidris and tribe 249 Anatini (formerly Anas) as two key examples of taxonomic units with contrasted preferences, 250 within-genus synchrony dominates year-to-year analyses for the two seasons (Fig. 2). Us-251 ing functional groups (waders and waterfowl), synchrony within functional groups was also 252 prominent. The Gross et al. (2014) synchrony indices are indeed mostly positive, and always 253 positive whenever significantly different from the null hypothesis of no temporal correlation between species. Therefore, there is no compensation within guilds (Fig. 2a and b) across 255 years, for the two seasons. This matches the patterns obtained within the entire wetland bird 256 community (Fig. 3a): synchrony dominates when abundances are computed at the species 257 level. 258

For the cold season, abundances within *Calidris* and *Anatini* display opposite changes in synchrony values in response to the management change in 2006, with species within *Anatini* becoming less synchronous over time, although we should mention that these changes are not statistically significant. For the warm season, the management change, which consisted of lowering the water levels, created little change in communities of species within the *Anatini* and *Calidris*: they are all synchronous.

Even though there is no widespread community-wide or genus-wide compensation across years (separating the two seasons), there could be compensation at finer temporal scales, e.g. a month or two, or coarser scales, over several years. Such compensation could also occur at specific time intervals instead of throughout the whole time series, a time-dependency that wavelet analyses allow to reveal. When we consider the wavelet modulus ratio (Fig. 4), that is, a time-varying and scale-dependent strength of synchrony, we can see that there is synchrony even at a fine temporal scale throughout most of the time series. However,

post-2006, there seems to be a possibility for episodic compensation on a temporal scale of approximately 2-4 months, for both waders and waterfowl. There could also be withinguild compensation at scales of 5 years, approximately post-2000 for waders and pre-2005 for waterfowl. Waterfowl synchrony trends likely influence whole-community trends (Fig. 3).

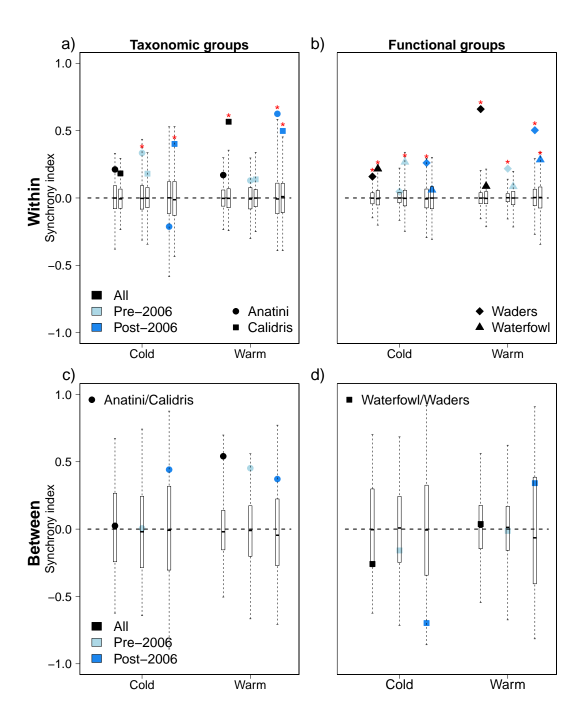


Figure 2: Gross' synchrony index (η) as a function of the season (cold and warm seasons), calculated within (top, a-b) and between (bottom, c-d) groups. The groups considered were different taxonomic groups (Anatini, Calidris, left a-c) or functional groups (waders vs waterfowl, right b-d). The index was computed in each panel on the whole dataset (black) or using two periods: before and after 2006 (light and dark blue), the year of the change in water level management. Boxplots indicate the distribution of η under the null hypothesis (independent species) and filled symbols correspond to the observed values. Red stars correspond to synchrony values significantly different from the null model, at the 10% threshold with a Benjamini-Hochberg correction.

We thus find contrasted results regarding the effect of the management change on synchrony within guilds or within the whole bird community, depending on the type of analyses.
Year-to-year analyses yield unclear results for both guilds. At shorter (one or two months)
and longer (five years) timescales though, wavelet analyses show that the management change
may decrease synchrony and even promote compensation.

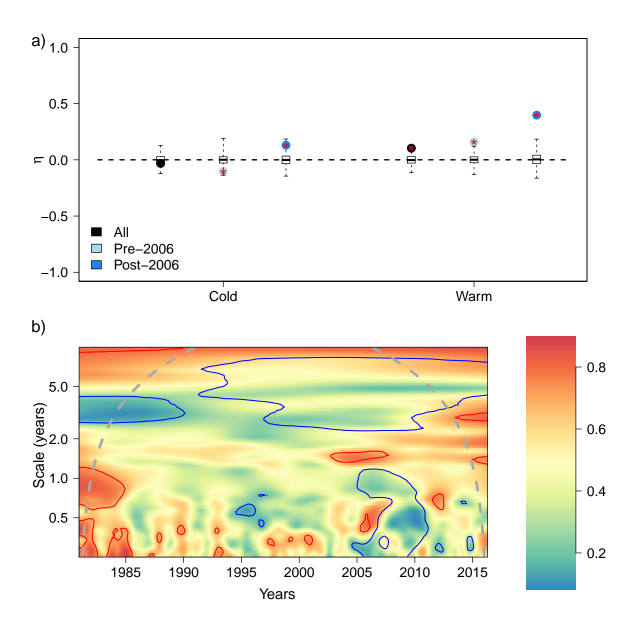


Figure 3: Synchrony indices for the whole community of frequently observed birds. Panel a) presents yearly synchrony (η) for both seasons and b) the wavelet modulus ratio (ρ) . The latter index scales from 0 (compensation, blue color) to 1 (synchrony, red color). Red and blue lines respectively delineate regions of significantly lower and higher synchrony than the null model (independently fluctuating species, but conserving their original Fourier spectrum), at the 10% level.

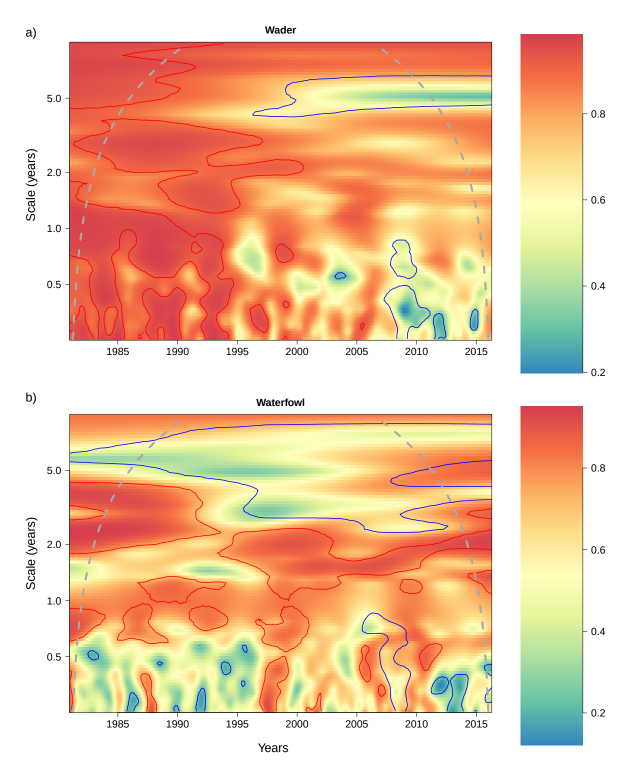


Figure 4: Wavelet modulus ratio (ρ) for a) the wader community and b) the waterfowl community. The index ρ scales from 0 (compensation, blue color) to 1 (synchrony, red color). Red and blue lines respectively delineate regions of significantly lower and higher synchrony than the null model (independently fluctuating species, but conserving their original Fourier spectrum), at the 10% level.

Synchrony between phylogenetic or functional groups

More easily interpretable results can be found when we examine synchrony vs compensation
between functional groups (Fig. 2d). Since we consider only two functional or phylogenetic
groups, the Gross et al. (2014) index reduces to a simple correlation between two groups.

Anatini and Calidris are positively correlated in the warm season (for all periods), and have
unclear correlations during the cold season (Fig. 2c). In contrast, waders and waterfowl are
negatively correlated during the cold season and positively correlated during the warm season
(Fig. 2d). Although the negative correlation is not statistically significant, it is consistent
for both pre- and post-2006 periods.

Synchrony in a small module with known competition

Compensation could be expected upon visual inspection of the time series of the two groups 291 formed by cormorant on the one hand, and little egret plus grey heron (summed as a small 292 functional group) on the other hand (Fig. 5, though see SI Appendix S3 for alternative 293 representations). However, we see on Fig. 6 that synchrony is in fact the rule around the 294 annual scale and below, when considering the wavelet modulus ratio. We wondered if the 295 patterns in Fig. 5 were caused by the use of a log scale, but we found that in fact the 296 correlation was higher rather than lower on the log scale (Appendix S3). However, over long 297 temporal scales (~ 8 years) we observe consistent compensation, which could correspond to 298 the slow change in composition observed within this small community module, that was 299 already visible on the abundance time series plot (Fig. 5). There is some statistically 300 significant compensation over shorter timescales as well, but only at very specific times. The 301 absence of marked compensation at short temporal scales may be an inevitable consequence 302 of the difference in the amplitude of temporal variation between the two groups (Appendix 303 S8), as opposite annual phases for the two time series can be observed before 2000 (Fig. 5).

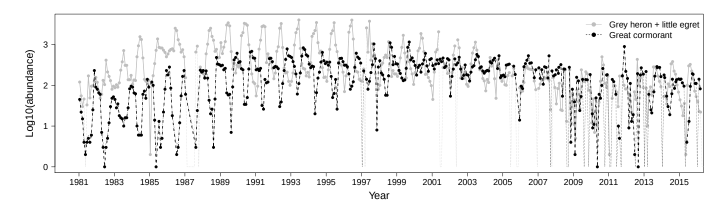


Figure 5: Time series of great cormorant abundance (dash-dotted black line), as well as summed abundances of grey heron and little egret (solid grey line).

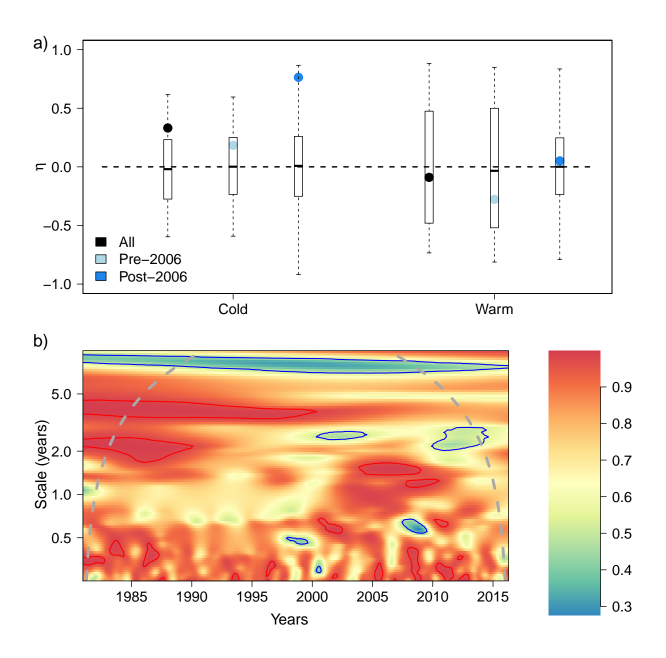


Figure 6: Synchrony analyses of the group formed by cormorant vs egret and heron. Panel a) presents yearly synchrony (η) for both seasons and b) the wavelet modulus ratio (ρ) . The latter index scales from 0 (compensation, blue color) to 1 (synchrony, red color). Red and blue lines respectively delineate regions of significantly lower and higher synchrony than the null model (independently fluctuating species, but conserving their original Fourier spectrum), at the 10% level.

Discussion

Between-species compensation was not found across years (for two separate seasons), syn-306 chrony between species being the rule. In other words, there was no widespread "functional 307 compensation" (sensu Gonzalez & Loreau 2009) within genera or guilds in year-to-year anal-308 yses of cold and warm seasons. Yet, summing the abundances of species within a guild and 309 comparing these total abundances of contrasted guilds, it was possible to find compensation 310 across years, during the cold season corresponding to wintering birds (although the null hy-311 pothesis of no correlation could not be rejected); that is, there was compensation between guilds. These results are robust to using biomass in place of abundance (SI Appendix S4). A 313 zoom on a module of three species with known competition also revealed clear compensation 314 at scales ≈ 8 years. We elaborate below on these findings.

316 Synchrony within or between guilds

Given that we compare the level of synchrony/compensation within guilds (with many 317 species) and between guilds (with only a handful of groups), we checked in Appendix S5, 318 using the dynamical model of Gross et al. (2014), if changing the number of "compartments" 319 (n) in the index η could affect its value. It did not have marked effects, unless the num-320 ber of compartments is equal to 2, in which case significance is hard to achieve and some 321 compensatory dynamics can be missed with weak environmental response. Additionally, we 322 found – still using this dynamical model – that if two guilds respond in opposite ways to 323 a shared environmental driver, the stronger the response of growth rates to the driver, the lesser the compensation indicated by η at the whole community level. An intuitive explanation of this modelling result is that when there are two groups and many species within a 326 group, a stronger forcing homogeneizes the dynamics within a group as much as it creates 327 differences between groups. This might explain the low levels of compensation that we found 328

in our empirical dataset, at the overall wetland bird community level (Fig. 3), in spite of
the clear presence of two guilds (waders and waterfowl) reacting in opposite way to a shared
driver (here, water levels). Analyses at several taxonomic/functional scales are therefore warranted to be conclusive about compensation, which mirrors what was suggested by earlier
plant studies (e.g., Bai et al., 2004). Future case studies with more than two main functional
groups may be instructive, to challenge the generality of our findings.

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We used correlation between the summed abundances of closely related species (species within the *Anatini* tribe vs species within the *Calidris* genus) or the summed abundances of functionally similar species (waders vs waterfowl) to uncover compensation. The functional group classification produced some compensation between guilds while the taxonomic classification did not, despite the contrasted habitat preferences of these two phylogenetic groups. Using functional groups produced more logical results, although as we stressed above, due to the low power of the tests, the null hypothesis of no compensation at the yearly scale was still plausible as well.

We expected to see compensation at the "functional group scale" for both cold and warm 343 seasons. The separation of seasons allowed to differentiate summer residents (some of whom may be breeding) and wintering birds, in order to remove the overwhelming influence of the seasonal migratory cycle. In both of those seasons though, we had reason to expect waders and waterfowl to have different environmental preferences. Instead, waders and waterfowl were found to correlate negatively only during the cold (wintering) season. A simple explanation is that the reserve might be closer to its carrying capacity for these species in winter, 349 so that space is limited and increases in one functional group are compensated by decreases 350 in the other. The dominant species in each guild (Fig. 1), such as C. alpina for waders and 351 A. crecca for waterfowl, are migratory species which are much more abundant in winter than 352 summer in that area, which adds to the plausibility of the reserve reaching carrying capac-353 ity. Of course, the space constraint should not be taken too literally: birds are obviously

mobile and do forage outside of the reserve (e.g., waders moving to the nearby Arcachon bay mudflats), but there are costs to those movements (energetics, mortality risk due to nearby hunting) which make the reserve a very attractive wintering site where birds both rest and forage to some degree. Packing even more birds over its 120 ha may just not be feasible, so that increases in one guild result in decrease in the other. Compensation might therefore be easier to detect during the cold season because the study area is "filled", and it is not detected in our warm season (May to August) because there are less birds overall.

It may be better to say that we detected "compensation" rather than "compensatory 362 dynamics" between bird species (Gonzalez & Loreau, 2009), if compensatory dynamics is 363 thought to result from births and deaths, i.e., population dynamics. Indeed, the observed 364 long-term changes in species composition (more waders, proportionally less waterfowl; Ap-365 pendix S2) is likely due to an increased inflow of birds preferring low water levels (waders), 366 and outflow of birds preferring high water levels (waterfowl), under an overall space constraint 367 (at least in winter, as we explained above). Bird settlement decisions for both winter and 368 spring/summer seasons are the proximal causes of bird species composition in the reserve, 369 rather than population dynamics. However, it would be incorrect to conclude that because 370 the local compensation in winter that we found results from bird behaviour, it is disconnected 371 from regional-scale community dynamics: which species are present in the reserve - safe from 372 hunting - affects ultimately their survival and reproductive success, which then feeds back 373 into the regional-scale community dynamics.

Effect of the change in management on synchrony

Although we performed a first set of analyses using the whole time series, we have also performed year-to-year analyses pre- and post-2006. The reason for these additional analyses is that a marked change in management occurred around 2006, after which the water levels were lower. Separating pre-/post-2006 and comparing to the previous analyses allows to

disentangle the effect of the "normal" dynamics from the effect of this management change.

Pre- and post-2006 analyses showed very little differences with whole time series analyses

for either the warm or cold season. However, in the wavelet modulus ratio analyses, we see

at monthly or 5-year timescales more compensation after 2006 for waders; this could reflect

that the community is becoming saturated with waders. The effects of disturbances on the

level of synchrony or compensation are likely idiosyncratic: for instance, Keitt (2008) found

increased synchrony after disturbance while van Klink et al. (2019) found no clear effect.

387 Synchrony in a small module with known competition

We now zoom in on the cormorant-heron-egret module, for which we knew beforehand that 388 competition for resting and roosting sites in the summer season occurs between, on the one hand, great cormorants, and on the other hand, little egrets and grey herons (C. Feigné, 390 pers. obs.). Abundance time series suggested some negative correlation, but it was not 391 found in year-to-year analyses for which synchrony (or an absence of relation) dominates. Instead, we find that compensation mostly occurs on a scale of 8 years, much above the 393 annual scale, which is a likely consequence of the slow shift in frequencies of cormorants and 394 little egrets / grey herons. The reason why we do not find a compensation at the montly to 395 annual scale pre-2000 in spite of some opposition of annual phases may be related to the large 396 difference in the amplitude of short-term temporal variation between the two groups. When 397 one functional group or species dominates the temporal variation, as shown in SI Appendix 398 S8, its dominance of temporal variation can forbid the occurrence of compensation since by 399 definition no increase in the numbers of the species that fluctuate less may compensate for 400 the decreases in the species that fluctuate more (and vice versa). 401

¹² Conclusion and perspectives for theory

Overall, our results suggest to search for compensation more often between rather than within 403 functional groups, and over relatively long timescales, above the typical temporal autocorrelation of the dominant driver (e.g., above 5 years if the main driver is a seasonal climate). 405 This rejoins the recent findings of van Klink et al. (2019) who found that increased func-406 tional differences between species tend to decrease synchrony in beetles, as well as earlier 407 results of Bai et al. (2004) on negative covariation of plant functional groups. Our suggestion 408 goes against calls to search for compensation within closely related species but at very short 409 timescales (Vasseur & Gaedke, 2007; Gonzalez & Loreau, 2009), below the timescale of the 410 main synchronizing seasonal environmental driver, in order to filter out precisely its synchro-411 nizing effect. Searching for compensation at temporal scales below the seasonal abiotic driver 412 (e.g., temperature) was partly motivated by studies on plankton whose population dynamics 413 are usually much faster than the dominant abiotic driver, with short generation times, so 414 that the effects of competition may be manifest at the scale of a few weeks or months. 415

In theory, we could have expected compensation to manifest also at the smallest temporal 416 scale of our survey (monthly). Indeed, the community dynamics in our case are driven by the 417 movements and settlement decisions of birds, reacting to perceived food and space availability, 418 rather than by births and deaths directly. Such behavioural dynamics can certainly be much 419 faster than bird population dynamics, and could operate at the scale of weeks or months. 420 However, such compensation due to short-term movements was not observed except perhaps in some years. We suspect that because many species share common abiotic drivers (e.g., disturbances due to nearby hunting, local climatic conditions) fluctuating even within a single season, their dynamics can be synchronized by these drivers at monthly temporal scales. It 424 is noteworthy that even in planktonic systems, the temporal scale of compensation has often 425 been found to be well above that of the forcing driver (Keitt, 2008; Brown et al., 2016). Thus 426 our findings reinforce previous suggestions to search for compensation over relatively long timescales (several years for vertebrates or plants).

The attractor of community dynamics, i.e., the shape of community trajectories in phase 429 space, seems to be more or less an annual cycle here: the dominant species fluctuate season-430 ally, but even though there are shifts in some species dynamics, no abundant species seem to 431 exhibit violent multi-year oscillations. If we had to describe our community mathematically, 432 a dynamical model with a stable fixed point forced by seasonality and some noise would prob-433 ably be appropriate. This mild fluctuation scenario somehow contrasts with the dynamics of 434 other communities, such as insect pests, that have quite often multi-year cycles (on top of 435 seasonal cycles, for multivoltine species), with possibly strong indirect interactions between 436 similar species mediated by predators and parasitoids (Murdoch et al., 2003). In the latter 437 context of internally-generated variability ("Endogenous compensatory cycles" in Gonzalez 438 & Loreau, 2009), compensation may be more likely: Klapwijk et al. (2018) recently reported 439 only transient synchrony between species of moths, that typically exhibit such multi-year 440 fluctuations. 441

In many ways, searching for abundance compensation using biodiversity time series data 442 is searching for needles in a haystack: only some specific temporal and functional/taxonomic scales allow to see compensation whilst numerous confounding factors make the community co-vary positively at all other scales (Vasseur et al., 2014). When a common species fluctuates 445 much more than the rest, this can also lessen or forbid compensation. Thus, although the knowledge of specific biological mechanisms increasing the densities of some species at the expense of others can help, synchrony will likely dominate community-level time series data 448 for closely related species, even in species that compete strongly (Ranta et al., 2008; Loreau & de Mazancourt, 2008). This is true even in cases of known mechanisms of competition 450 for space or shifts in community composition due to abiotic changes affecting differentially 451 species preferences, as in this study. We therefore suggest that "zooming out" functionally 452 (considering summed abundances of dissimilar functional groups) and temporally (consider-453

ing temporal scales well above the periodicity of the dominant abiotic driver) may often be the best strategy to see the compensation that will inevitably manifest at some scales, if the community-level biomass is to be maintained within bounds in the long run.

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Data accessibility

All the code and data used for analyses are available at https://github.com/fbarraquand/
BirdTimeSeries_Teich and archived at Zenodo [will be done for the final version], DOI:XXXXX (Picoche, Aluome & Barraquand, 2020).

469 Authors' contributions

FB, LC and CF designed the original project. CF coordinated the data collection and provided knowledge on functional groups. CA, FB and CP standardized the bird abundance database and performed exploratory analyses. Final statistical analyses were designed by FB and CP, and coded mostly by CP. FB and CP led the article writing, with inputs from all authors.

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