Controversy over the decline of arthropods: a matter of temporal baseline?

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Abstract

Recently, a number of studies have reported somewhat contradictory patterns of temporal trends in arthropod abundance, from decline to increase. Arthropods often exhibit non-monotonous variation in abundance over time, making it important to account for temporal coverage in interpretation of abundance trends, which is often overlooked in statistical analysis. Combining four recently analysed datasets that led to contrasting outcomes, we first show that temporal abundance variations of arthropods are non-monotonous. Using simulations, we show non-monotony is likely to bias estimated linear abundance trends. Finally, analysing empirical data, we show that heterogeneity in estimated abundance trends is significantly related to the variation in temporal baseline of analysed time series. Once differences in baseline years, habitats and continents are accounted for, we do not find any statistical difference in estimated linear abundance trends among the four datasets. We also show that short time series produce more stochastic abundance trends than long series, making the dearth of old and long-term time series a strong limitation in the assessment of temporal trends in arthropod abundance. The lack of time series with a baseline year before global change acceleration is likely to lead to an underestimation of global change effects on biodiversity.

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**Introduction**

Over the last decades, many studies have documented a biodiversity crisis on the basis of high extinction rates (Dirzo & Raven 2003; Ceballos et al. 2015) and population losses or declines (Butchart et al. 2010; Ceballos et al. 2017), mainly in vertebrates. Worldwide erosion of biodiversity is caused by anthropogenic global change (Sala et al. 2000), i.e. a set of pressures including among others land use change, climate warming, overexploitation, or pollution (Steffen et al. 2006). In recent years, invertebrates, and especially arthropods, have been at the centre of a debate (Dornelas & Daskalova 2020; McDermott 2021) regarding the magnitude and even the directionality of the temporal trends in their abundance. Some studies showed a strong decline on the basis of standardized inventories (Hallmann et al. 2017; Seibold et al. 2019), while analyses of meta-datasets assembling heterogeneous time series evidenced a decline of terrestrial, but an increase of aquatic arthropods (Pilotto et al. 2020; van Klink et al. 2020) or no overall decline (Crossley et al. 2020). Finally, an analysis of opportunistic occurrence data revealed non-monotonic dynamics and no overall decline (Outhwaite et al. 2020).

Heterogeneity in population trends among studies is not surprising but the underlying causes need to be differentiated, particularly to tell apart abiotic/biotic factors from methodological factors. The dynamics of biodiversity changes are well-documented and their heterogeneity relatively well understood for vertebrates (Antão et al. 2020; Daskalova et al. 2020b, a; Leung et al. 2020) or for some specific functional groups of invertebrates, such as pollinators (Grab et al. 2019; Duchenne et al. 2020; Soroye et al. 2020; Millard et al. 2021). However, the heterogeneity in population trends remains poorly explained for most invertebrates. Several reasons may explain the contrasting patterns revealed by the studies involved in the arthropod decline debate. Global change pressures can vary in space among locations or ecological habitats, and species abilities to respond to environmental changes may depend on their traits or their evolutionary history (Helmus et al. 2010; Grab et al. 2019); hence spatial, ecological and taxonomic coverages are obvious sources of heterogeneity among studies and are widely discussed in the recent literature (Blowes et al. 2019; Pilotto et al. 2020). For example, some studies used global datasets with a bias towards the northern hemisphere (van Klink et al. 2020) while others considered national or even more local spatial extents (Hallmann et al. 2017). Some focused on terrestrial arthropods (Hallmann et al. 2017; Seibold et al. 2019) while others included aquatic groups (Crossley et al. 2020; van Klink et al. 2020). Finally, some studies analysed average trends by pooling multiple taxonomic groups (Hallmann et al. 2017; van Klink et al. 2020), while others reported trends for each taxonomic group (Outhwaite et al. 2020).

Differences in temporal coverage (i.e. the time period during which a taxon or community was monitored) across studies are at the core of the debate over biodiversity crisis (Pauly 1995; Cardinale et al. 2018; Loreau et al. 2022), but efforts to account for temporal coverage in statistical analysis remain limited (Cardinale et al. 2018; Didham et al. 2020; Loreau et al. 2022). Heterogeneity in temporal coverage is likely to influence the conclusions of studies assessing temporal trends in arthropod abundance for two reasons. First, there is variation in baseline years across the studies fuelling the debate: some analysed time series of identical temporal coverage and relatively old baseline year (Hallmann et al. 2017; Outhwaite et al. 2020), while others used heterogeneous datasets consisting mostly of time series with recent baseline years (Crossley et al. 2020; van Klink et al. 2020). Second, non-monotonic dynamics repeatedly reported in arthropods (Macgregor et al. 2019; Baranov et al. 2020; Høye et al. 2021; Schowalter et al. 2021), due to temporally variable environmental pressures (Baranov et al. 2020; Duchenne et al. 2020; Schowalter et al. 2021), can affect linear estimates of abundance trend beyond the original description of the shifting baseline syndrome (Fig. 1). With non-monotonic dynamics, estimated linear trends can vary from positive to negative when considering different baseline years for the same time series (e.g. Fig. 1h). As most arthropod trends were estimated assuming linear trends and available data often do not make it possible to account for all factors influencing temporal variation in abundance, it appears critical to account for differences in baseline years when making comparison within and among studies.
Figure 1: Schematic examples of how baseline year may affect estimated linear abundance trends. (a-d) Hypothetical time-series showing abundance variations over years (black) and three possible temporal coverages with contrasting baseline years: old (blue), intermediate (red) and recent (yellow). (e-h) Linear abundance trends estimated with time series starting from these three baseline years. (i-l) Pattern of estimated abundance trends against baseline year, reconstructed by estimating abundance trends over all possible baseline years. The dashed black line shows the zero-value delimiting positive (above) vs. negative (below) abundance trends.

Here, we evaluated the effect of the baseline years of time series on estimated linear trends in arthropod abundance, using the four largest datasets of arthropod time series recently analysed in studies fuelling the current debate (Dornelas et al. 2018; Crossley et al. 2020; Outhwaite et al. 2020; van Klink et al. 2020). We first characterized non-monotony in temporal variations of arthropod abundance. Then, using simulations we assessed how non-linearity can bias linear abundance trends. Finally, using a sliding baseline method on the empirical dataset aggregated here (cf. Methods, Fig. 2), we measured how non-linearity in arthropod abundance trends can produce a statistical dependency of estimated linear abundance trends on the baseline year.

Methods

The four source datasets (step 1)

We aggregated four source datasets from recent publications evaluating abundance trends in arthropods (Table S1): (i) annual occupancy estimates (the proportion of 1km² grid cells in a region occupied by a species, a proxy for abundance) at species level for a wide diversity of arthropods from Great-Britain, produced by Outhwaite et al. (2019), (ii) annual estimates of arthropod abundances mostly at species level from American Long-Term Ecological Research (LTER) sites from Crossley et al. (2020), (iii) annual estimates of arthropod abundances from the meta-analysis of van Klink et al. (2020), aggregated at the resolution of taxonomic order, and (iv) abundance time series from the BIOTIME database (Dornelas et al. 2018), mostly at the species level (Fig. S1).
Figure 2: Schematic description of data aggregation, and workflow of analyses to assess the effect of the temporal baseline on abundance trends. In Step 1, we aggregated empirical datasets from four different source datasets. Then we characterized non-monotony of abundance variations over time (Step 2), and we assessed how non-monotony can affect estimated linear abundance trends using simulations (Step 3, Goal 1). Finally, we created an array of subset datasets using a sliding baseline (Step 4) to estimate how baseline years influenced estimated linear abundance trends in empirical data (Step 5, Goal 2). Abundance trends are expressed as growth rates for all time series, allowing comparisons among common and rare species, as well as among datasets (cf. Supplementary Methods, Fig. S2 & S3). Statistical analyses were then performed accounting for habitat (aquatic vs. terrestrial), continent, data source and taxonomy.

We focused on well covered continents and habitats, using only arthropod time series from North America and Europe with information on habitat (aquatic vs. terrestrial). We homogenized taxonomy using the Global Biodiversity Information Facility (GBIF) taxonomy backbone. For data from Crossley et al. (2020) and from van Klink et al. (2020), some time series describe the temporal variations of a wide diversity of species pooled together by summing their respective abundances. For these datasets, we removed all time series with taxonomic resolution coarser than taxonomic order, except for non-insect arthropods that are often grouped at taxonomic class level in available datasets (Chilopoda, Diplododa, Collombola, Branchiopoda etc.). We retained these groups and consider them to be the same rank as taxonomic orders in the following for simplicity. Details about aggregation and filtering step are available in Supplementary Methods.

To study baseline year effects on linear abundance trends estimated from time series with consistent ending dates, we removed time series ending before 2005 (n = 14,717). We further removed the few abundance values before 1970 (n = 1,039, 0.4%) to focus on the time period when most of the data were collected. Finally we removed time series shorter than 3 years (n = 47). This led to 14,130 original time series (Table S2, Fig. S1).

Assessing the monotony of temporal variation in abundance (step 2, goal 1)

Because estimated linear abundance trends depend on the baseline year only if abundance varies non-linearly, especially non-monotonically, over time (Fig. 1), we estimated non-monotony using a Generalized Additive Model (GAM) for each original time series. We estimated the strength of non-monotony as the number of local extrema, hereafter turning points, observed in the non-linear trend predicted by the GAM, as detailed in supplementary Methods.
Simulations of temporal declines for different population dynamics (step 3, goal 1)

We used simulations to assess how non-linearity in temporal variation of abundance (i.e. population dynamics) can bias estimated linear abundance trends, depending on their baseline year. We used four different shapes of population dynamics (Fig. 4a), using four functions describing temporal variation in average population size over 42 years (cf. R script available in supplementary materials). For each shape of population dynamics, we simulated time series with different growth rates \( r \) from stable \( (r = 1) \) to declining \( (r = [0.95, 0.9, 0.85]) \). The three latter values of \( r \) correspond to declines of 5%, 10% and 15% per year, respectively. We simulated 100 abundance time series for each shape of population dynamics and growth rate. To do so, for each year, we sampled observed abundance values from a Poisson distribution with a mean equal to the average population size of the corresponding year. Then, we estimated a linear abundance trend over the entire time period (baseline year at \( t = 1 \)), as well as over truncated time series with different baseline years \( (t = 10, 20, 30) \), using a Generalized linear model (GLM) with a Poisson error structure and a log link function, and accounting for temporal autocorrelation (equation (1) below).

Generating an array of subset datasets using a sliding baseline (step 4)

To study the effect of the baseline year of time series on abundance trends estimated from the 14 130 empirical original time series, we created an array of 41 datasets, hereafter called subset datasets, corresponding to 41 different baseline years, from 1970 to 2010 by steps of one year (Fig. 2). For each of the subset datasets, time series were either truncated to start at the given baseline year, or removed if they did not include this specific baseline year. By construction, each of the original time series therefore appears several times in the array of 41 datasets, corresponding to \( n \) \( (1 \leq n \leq 41) \) truncated time series. Since time series with old baseline years are rare, the number of time series included in the subset datasets decreases with earlier baseline years.

Estimating abundance trends (step 5)

We estimated arthropod abundance trends using one GLM per truncated time series. We considered only truncated time series with at least three annual estimates of abundance, including the abundance estimate in the year used to truncate the time series.

To obtain comparable abundance trends among the various sources, expressed in the same unit, we used a model structure that allows the estimation of growth rates. To do so we used GLMs with Poisson error structure with a log link function for count data, from van Klink et al. (2020), Crossley et al. (2020) and BIOTIME, GLMs with a binomial error structure with a logit link function for occupancy estimates from Outhwaite et al. (2019), and GLMs with a gaussian error structure with a log link function for density estimates from BIOTIME. Trends estimated with a log or logit link functions are expressed as the logarithm of a growth rate (see Supplementary Methods, Fig. S2 & S3), allowing comparisons between common and rare species, but also among datasets. Therefore, this approach with appropriate link functions gives the same importance to rare and to common species, in contrast to classic standardization of abundance by mean and standard deviations, which biases average trends by giving more weight to species with lower inter-annual variability in abundance (Fig. S4).

The GLM used for each truncated time series explains the abundance estimates of each year \( i \) by a Poisson or Binomial distribution, of parameter \( \lambda_i \) and \( p_i \) respectively, which depend on a linear year effect (\( \beta \)):

\[
\log(\lambda_i) \text{ or } \logit(p_i) = \alpha + \beta \times year_i + \varphi_i + e_i
\]

where \( \alpha \) is the intercept, \( \varphi_i \) is a temporal random walk of order one \((\varphi_1 \sim \mathcal{N}(\varphi_{i-1}, \sigma_{\varphi}^2))\), with \( \varphi_1 = 0 \) to account for temporal autocorrelation and \( e_i \) an error term. We fitted these GLMs using the INLA R package (Rue et al. 2009).

Evaluating the importance of the effect of baseline year in arthropod abundance trends (goal 2)

Pooling together the slopes of the year effect from the linear models presented above, across the 41 subset datasets, we obtained 192,244 abundance trends. We removed abundance trends estimated from time series with a single non-zero yearly estimate of abundance. We did so because growth rates estimated from such time series are likely to be extreme (i.e. strongly negative or positive) if this positive abundance...
estimate is by chance at the end or at the beginning of the truncated time series (Fig. S5). We thus kept the 175,796 abundance trends derived from truncated time series with ≥2 non-zero abundance estimate.

We examined the effect of baseline year used to truncate the original time series on abundance trends. To this end, we used a Bayesian linear mixed-effects model explaining abundance trends with spatial and temporal variables. Since we do not expect a linear relationship between abundance trends and baseline year (Fig. 1), the effect of baseline year was modelled as a polynomial of order three. We added a continent effect, a habitat effect, and all two- and three-way interactions between these effects and the polynomial baseline year effect (equation (2)), because recent results suggest that terrestrial and aquatic arthropods exhibit differences in their abundance trends and because population trends can vary over space (Outhwaite et al. 2020; van Klink et al. 2020). To assess whether differences among source datasets (Crossley et al. 2020; Outhwaite et al. 2020; van Klink et al. 2020) persist after spatio-temporal variables have been taken into account, we included a dataset effect, with four levels.

We also accounted for pseudo-replication in species belonging to the same taxonomic orders, as well as for the fact that several truncated time series are obtained from the same original time series, by adding a random effect of taxonomic order and time series ID on the intercept. Data from van Klink et al. (2020) and Crossley et al. (2020) originated from multiple sites and different data sources. Thus, we accounted for this structure by adding a random site effect nested in a random data source effect. For data from Outhwaite et al. (2019), the data source corresponds to the groups used to calculate occupancy estimates in the original dataset and the site corresponds to the country coverage of the data for each group (UK or GB), both extracted from the Online-only Table 1 (Outhwaite et al. 2019).

The linear mixed-effect model is thus the following:

\[ AT_{jchsdli} = \alpha_{ch} + \beta_{s} + \varphi_{bch} + \theta_{1o} + \theta_{2d} + \theta_{3i} + \theta_{4t} + \epsilon_{jchsdli} \]  

where \( AT_{jchsdli} \) is the abundance trend of truncated time series \( j \) with baseline \( b \), from continent \( c \), habitat \( h \), dataset \( s \), order \( o \), data source \( d \), site \( l \), from original time series \( i \). \( \alpha_{ch} \) is the intercept for all combinations of continent \( c \) and habitat \( h \), while \( \beta_{s} \) is the effect of the source dataset \( s \). \( \varphi_{bch} \) is a baseline effect that depends on baseline \( b \), continent \( c \) and habitat \( h \). It is modelled as a temporal random walk of order one (\( \varphi_{bch} \sim \mathcal{N}(\varphi_{(b-1)c}, \sigma_{0}^{2}) \), with \( \varphi_{1c} = 0 \)). The \( \theta \)s denote random effects on the intercept: \( \theta_{1o} \) for taxonomic order, \( \theta_{2d} \) for data source, \( \theta_{3i} \) for site and \( \theta_{4t} \) for time series ID. \( \epsilon_{jchsdli} \) is an error term following \( \mathcal{N}(0, \sigma^{2}) \).

To account for the fact that our response variable is estimated, and thus each value has an associated standard error (\( sde_{j} \)), we modelled the residual variance as a function of this error. In addition, since the four source datasets have different taxonomic scopes or different spatial scales, we expect that residual variance (\( \sigma^{2} \)) will be strongly structured by the source dataset. Finally, the baseline year also affects the number of abundance estimates in time series, which is expected to affect the stochasticity of abundance trends (Bahat et al. 2021). We modelled the dependence of variance of the residuals (\( \sigma^{2} \)) on standard error associated to truncated trends of time series \( j \left( AT_{j} \right) \), on source dataset \( s \) and on the number of years with data in each truncated time series (\( ny \)):

\[ \log(\sigma^{2}_{js}) = \alpha_{s} + \beta_{1s} \times n_{y} + \beta_{2s} \times n_{y}^{2} + \beta_{3} \times \log(\text{sde}_{j}) \]  

where \( \alpha_{s} \) is the intercept, which depends on source dataset \( s \), and \( \beta_{1s} \) and \( \beta_{2s} \) the polynomial effects of the number of years with data in truncated time series \( j \), for each dataset, on the residual variance \( \sigma^{2}_{s,j} \). \( \beta_{3} \) is the effect of standard error associated to \( AT_{j} \) on the residual variance \( \sigma^{2}_{s,j} \). Parameters of the two models, one for the mean (equation (2)) and one for the residual variance (equation (3)), are estimated simultaneously. On this model (equation (2)), we estimated the variance of abundance trends explained by each random or fixed effect, as the ratio of its variance on the sum of all these variances plus the variance of the residuals. Priors used are detailed in the R script available in supplementary material. The model was
fitted using the R2jags R package (Su & Yajima 2012), with 3 chains using 60,000 iterations with a burnin of 50,000 and a thin rate of 3, which was enough to reach convergence (all parameters with Rhat<1.1).

**Figure 3:** Empirical time series of arthropod abundance are non-monotonous, leading to unstable abundance trends over a gradient of baseline years. (a) Distribution of the number of original time series as a function of the number of turning points (a proxy for non-monotony, cf. Methods) and of the number of years with data in the corresponding time series. (b) Variation in direction of abundance trends (positive vs. negative) of truncated time series, as a function of the number of years with data in the corresponding time series. Truncated time series are from the same original time series which is truncated for every possible baseline year (cf. Methods, Fig. 2). Boxplots represent minimum and maximum values (bottom and top of vertical lines), first and third quartiles (Q1 and Q3, bottom and top of boxes) and median (thick horizontal lines); colours indicate sample size (number of original time series). Points with values outside of the range [Q1-1.5(Q3-Q1), Q3+1.5(Q3-Q1)] are considered as outliers and represented as full circles.

**Results**

The number of turning points per empirical time series increases with the number of years with data in the time series, indicating that the vast majority of population trends are non-monotonous when time series are long enough (Fig. 3a). As expected from Figure 1, the strength of the non-monotony, measured as the number of turning points per time series, affects trend estimation. Non-monotonous abundance trends are characterized by multiple changes in trend direction over time, such that estimated linear trends may have opposite signs depending on the baseline year considered as the start of the time series (Fig. 3b). This pattern is consistent across the four source datasets (Fig. S6).
Figure 4: Non-linear population dynamics can bias estimated abundance trends. (a) Raw patterns of population dynamics when the population is stable, i.e. with a growth rate \( r = 1 \), or when the population declines by 5\% (\( r=0.95 \)), 10\% (\( r=0.9 \)) or 15\% (\( r=0.85 \)) each year. (b) Scaled patterns of population dynamics shown in (a) but using a log10 y-axis scale illustrating that the red curve corresponds to linear dynamics (cf. Supplementary methods, Fig. S2). (c) Estimated values of log(\( r \)) as a function of the shape of the population dynamics and of the baseline year used to calculate the population trend. Boxplots have the same meaning as in Figure 3. (d) Estimated values of log(\( r \)) as a function of the shape of the population dynamics and of the baseline year used to truncate the time series. In (c) and (d) the dashed horizontal line shows the value of the logarithm of the true (simulated) growth rate. Boxplots have the same meaning as in Figure 3. The right panel of (d) without outliers is represented in Figure S7.
Using simulated population dynamics (Fig. 4a), we show that similar rates of declines lead to different estimated growth rates, depending on the shape of the population dynamics (Fig. 4a-c). Assuming linearity to estimate population trends from non-linear dynamics can induce a strong bias when the growth rate departs from stability ($r=1$), by either overestimating or underestimating the genuine simulated trends (Fig. 4c). As expected, truncating the time series towards more recent baseline years increases the bias in the estimated growth rate, except in simulations with linear population dynamics (Fig. 4d). In addition to the magnitude of the bias, uncertainty (i.e. dispersion of values around the median) also increases when truncating the time series towards more recent baseline years (Fig. 4d). Short and recent time series tend to produce extreme estimated growth rates, which can even be opposite to the simulated long-term decline (Fig. 4d). These results suggest that not accounting for non-linearity in population trends can induce strong biases, and that short-term and recent time series cannot be used to infer long-term population change, even if population dynamics are linear, due to the strong uncertainties on estimated values.

When looking for general patterns in all empirical truncated time series together, statistical analyses show that average linear abundance trends estimated from the truncated time series strongly depend on the baseline year used for truncation, in interaction with habitats and continents (Fig. 5). This suggests that the non-linearity in temporal variations of abundance (Fig. 3b) leads to a strong dependence of the Arthropod population trends to the considered period (Fig. 5a). Results show a higher uncertainty in North America likely because time series with old baseline years are scarce for this region (Fig. 5b). This highlights that data are missing to estimate long-term abundance changes of Arthropods, and that recent data cannot help to fill this blank because of non-linearity.

**Figure 5: Effect of the baseline year on average abundance trends.** (a) Relationships between the baseline year of times series and estimated abundance trends (log of the growth rate) relative to the abundance trends estimated with the oldest baseline of the dataset (1970), for each continent and habitat. Error bars are 95% confidence intervals. (b) Number of truncated time series used for each baseline value.

Importantly, for the two source datasets including short and long time series (*i.e.* those from Outhwaite *et al.* and van Klink *et al.*, Fig. 6a), the residual variance of the model strongly decreases with the number of years with data in time series. This is consistent with our previous results based on simulations of truncated time series (Fig. 4c), and suggests that the stochasticity in abundance trends estimated from
short time series is much greater than that from long time series. For data from Crossley et al. and BIOTIME, the relationship between the residual variance in abundance trends and the number of years with data is strongly parabolic. However, those datasets are strongly biased towards recent and short time series (Fig. 6a), prompting caution in interpreting this signal.

Although statistically significant, the effects of baseline year explain only a small fraction of the total variation in abundance trends (Fig. 6b). Overall, random effects controlling for methodological issues, spatial and taxonomic heterogeneity explain 56% of the variation in abundance trends (Fig. 6b). A high proportion of this variation in abundance trends is explained by the random effect of time series ID, controlling for the artificial dependencies among abundance trends generated by the truncation procedure (cf. Methods). Most of the remaining variation in abundance trends is explained by taxonomic order and local site random effects, highlighting that abundance trends are strongly heterogeneous among clades and among sites. This questions the relevance of global estimates and stresses the need to carefully identify the drivers of such heterogeneity. Some groups, such as Trombidiformes (mites), Orthoptera, Collembola and Isopoda, exhibit more positive average abundance trends while other groups, such as Archaeognatha (jumping bristletails), Hymenoptera, Coleoptera, Dermaptera and Blattodea are associated with more negative trends (Fig. S8). Strikingly, once the various sources of heterogeneity are controlled for, we do not find any significant difference among the four source datasets from which the abundance estimates were extracted (Fig. 6c).

![Figure 6: Variation in abundance trends is strongly structured by methodological effects. (a) Predicted (lines) and observed (points) residual variance of the model for each source dataset. Predictions are from equation (3). The histogram at the top shows the distribution of truncated time series along the x-axis, per dataset. (b) Variation in abundance trends explained by the model effects, and residual variation. (c) Black dots show the effect (± CI95%) of the source dataset on abundance trends, relatively to the reference level (BIOTIME).](image-url)
Discussion

Accounting for differences in baseline years contributes to settling the debate on arthropod decline

Our analysis shows that the non-monotony of empirical abundance time series induces a strong dependency of estimated abundance trends to the baseline year, as well as a strong uncertainty in abundance trends from short time series. Our findings therefore bring statistical support to the fact that most of the available data regarding arthropods, which are biased towards recent and short time series, are not appropriate to extrapolate long-term population trends, as suggested recently in discussions over vertebrates and arthropods population trends (Daskalova et al. 2021; Loreau et al. 2022; Mehrabi & Naidoo 2022).

We show that baseline year is statistically linked to abundance trends, and we highlight that average linear abundance trends, if they make any sense, should be interpreted in the light of the temporal window covered by the analyzed time series. By statistically assessing the effect of shifting baseline, we provide a general picture of how abundance trends change as function of baseline year, which may help to reposition the findings of past and future studies in a broader context, and hopefully to make between-studies comparison easier. Importantly, once we controlled average temporal trends of studies by the various sources of heterogeneity we did not find any statistical difference among source datasets, suggesting that the contradiction among recent results regarding arthropod decline comes from methodological issues related to temporal coverage as well as taxonomical and geographical bias.

The importance to define temporal baseline and use common spatial yardsticks when evaluating temporal change was previously emphasized in the wider context of biodiversity decline (Lotze & Worm 2009; Mihoub et al. 2017; Cardinale et al. 2018; Stouffer et al. 2021), but here we bring further statistical support to this caveat. Indeed, although the importance of comparing results with common baselines is well known, the baseline effect is rarely explicitly accounted for in quantitative analysis (but see Macgregor et al. (2019) for an example), in contrast to other sources of heterogeneity in abundance trends (e.g. space and taxonomy). Although van Klink et al. (2020) tested the effect of the starting year by truncating their time series (see Fig. 3 of van Klink et al. 2020), they did not formally test for a baseline effect: discarding data earlier than a given baseline threshold, from 1960 to 2005, did not result in constraining the baseline years to be equal across all time series. Since their dataset was mainly composed of time series with a baseline year after 1990, discarding data before a giving baseline threshold did not affect the overall distribution of baseline years much, except for recent thresholds (post-1990).

We also show that shorter time series exhibit much more stochasticity (i.e. higher residual variance) than long term series, which increases the uncertainty of results from these short-term time series that are commonly used in assessing arthropod abundance trends (Seibold et al. 2019; Crossley et al. 2020; van Klink et al. 2020). In other words, our results show that short-term series need to be much more replicated than long-term time series to reach the same level of confidence in the results. This is consistent with previous trends assessment regarding moths in Great-Britain (Macgregor et al. 2019) and more generally with the fact that arthropod abundance trends estimated from short time series are strongly sensitive to year to year variations (Daskalova et al. 2021). Here we show that uncertainty decreases exponentially with the length of time series, highlighting the importance to maintain existing biodiversity monitoring schemes. Obtaining long enough time series is critical for assessing reliable abundance trends, which echoes hot and recent debates about abundance trends in vertebrates (Leung et al. 2022; Loreau et al. 2022).

Limitations and future challenges

In addition to the dramatic lack of data for some regions of the world, our study suggests that available arthropod abundances suffer from temporal limitations that should be carefully kept in mind when assessing their trends. First, the significant effects of baseline year and study area (continent and local site) suggest that the estimation of the arthropod abundance trends suffers from large uncertainties, mainly due to the lack of historical data. Here, 1970 is the oldest baseline we can tackle with the data at hand, but it cannot be considered as a reference before the rise in global change pressures (Mihoub et al. 2017), leading to a likely underestimation of global change effects on arthropod abundance. Since we cannot go back in time to sample biodiversity, this issue will remain difficult to solve. Museum collections and other sources of historical data could help fill this gap, although extracting reliable information from such data is still challenging (Isaac et al. 2014; Bartomeus et al. 2019; Duchenne et al. 2020; Outhwaite et al. 2020).
Whether or not scientists can manage to obtain the necessary data and apply relevant methods to effectively turn back the clock, our analysis stresses the critical need to maintain long-term monitoring and secure appropriate archiving of related data (Millar & Searcy 2019).

Second, we used the same datasets as previous studies, so our analysis does not introduce new elements to assess the reliability of these abundance trends with respect to potential biases related to spatial and taxonomic coverage or data quality (Desquilbet et al. 2020). For example, some of the time series included in van Klink et al.’s and Crossley et al.’s datasets were produced by experimental studies manipulating environmental conditions likely to influence abundance trends (Desquilbet et al. 2020, 2021), leading to criticisms about the use of these time series to assess temporal trends (Cardinale et al. 2018). Similarly, using only or mostly data from research stations, such as Long-Term Ecological Research sites, could bias estimated abundance trends upward as these locations are often partially protected from disturbances. Our results show that local site strongly explains heterogeneity in abundance trends, which is consistent with the strong influence of local changes in environmental conditions on arthropod abundance trends (Seibold et al. 2019). This has consequences for the interpretation of global trends obtained from a non-representative sample of sites. This potential bias stresses the need for standardized protocols to monitor arthropod abundance in numerous sites, representative of the areas covered by different habitats and land-uses, to handle the diversity of anthropic pressures on biodiversity, some remaining restricted to particular areas while others apply widely over space. Monitoring schemes based on citizen sciences are one way to tackle this challenge (van Swaay et al. 2008; Jeliazkov et al. 2016), since they can produce protocoled or semi-protocoled datasets over a large set of habitats and landscapes, over seasons and years, at national or even continental scales. Citizen science monitoring schemes are often recent, but would be of considerable help to ensure long-term monitoring of species abundances, should they be maintained in the future.

Moreover, by expressing abundance trends as growth rates, we gave the same importance to rare species as to common species, which could be debated. Decline of extremely rare species could be poorly informative and less likely to affect ecosystem functioning than decline of common species, while rare species can exhibit extreme abundance trends (Fig. S9), thus affecting average abundance trends in a non-negligible way. On the other hand, rare species can contribute greatly to some biodiversity metrics, such as species richness, phylogenetic diversity or functional diversity, making it relevant to weigh them similarly as common species for some purposes. Consistent with previous comments (O’Hara & Kotze 2010; Desquilbet et al. 2021), we show that transforming abundance counts with log(x+1) before statistical analysis of the data, as done by Crossley et al. 2020 and van Klink et al. 2020, instead of using model structures adapted to the data (i.e. GLM instead of linear models), can introduce an asymmetrical bias, flattening the abundance trends of rare species (Fig. S2). In a similar way, standardizing data by dividing by standard deviation also strongly biases the relative values of abundance trends among species (Fig. S4), stressing the need to explicitly test the effect of any transformations performed on the data.

Finally, the large variation in abundance trends across sites, taxonomic groups, habitats and continents brings into question the relevance of producing global multitaxon linear trends. Global multitaxon trends are likely to be disconnected from the ecological causes and consequences of biodiversity changes. Losses in one place or one taxonomic group cannot be balanced by gains in another place or taxonomic group. Losses and gains can have contrasting ecological and evolutionary consequences, that need to be assessed at relevant ecological scales, e.g. at community level or for a given functional group. Moreover, our results show that variation in arthropod abundance over time is non-linear and sometimes non-monotonous. This suggests that the use of linear analyses is inadequate, despite being the most straightforward and used analysis, and should at least always be associated to the temporal coverage of the data. Disconnecting arthropod decline assessments from temporal yardsticks can affect the understanding of published results making them apparently contradictory. This is particularly important for topics of interest for the general public such as arthropod decline, as it could lead to undermining trust in science (Dornelas & Daskalova 2020).

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underlying their work available, and for building step by step our knowledge about the way biodiversity changes over time. We also thank an anonymous reviewer from a previous submission for his/her insightful comments and opinions on that manuscript. The simulations were performed at the HPCaVe centre at Sorbonne Université. François Duchenne was funded by the European Research Council (ERC) under the European Union’s Horizon 2020 research and innovation program (grant agreement N° 787638). A preprint version of this article (https://doi.org/10.1101/2022.02.09.479422) has been peer-reviewed and recommended by Peer Community In Ecology (https://doi.org/10.24072/pci.ecology.100098).

Data, scripts and codes availability

All data used here were publicly available (cf. Methods). Scripts used for analyses are available in Supplementary materials (https://doi.org/10.1101/2022.02.09.479422) and here: https://github.com/f-duchenne/Controversy-over-the-decline-of-arthropods.

Supplementary material

Supplementary material are available online: https://doi.org/10.1101/2022.02.09.479422

Conflict of interest disclosure

The authors declare no competing interests.

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References


Supplementary Information for:

Controversy about the decline of arthropods: a matter of temporal baseline?

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This part includes:

Supplementary Methods

Fig. S1 to 9

Table S1 & S2
Supplementary Methods

Aggregating datasets

For data from Outhwaite et al. (2019), we focused on occupancy estimates labelled at Great-Britain level, which can in fact correspond to two different spatial coverages (United Kingdom or Great-Britain). Since for this dataset occupancy estimates are modelled, not observed, we excluded occupancy estimates that converged poorly in their analyses (Rhat<1.2): this yielded 138,095 occupancy estimates for 3,496 species, from 1970 to 2015, which correspond to 57% of the arthropod occupancy estimates at Great-Britain level from the original dataset.

For data from Crossley et al. (2020) (https://datadryad.org/stash/dataset/doi:10.5061/dryad.cc2fz645) we used the file “External_Database_S1_PerSpecies_Abundance_LTER.annotated.csv.” After data filtering, we obtained 78,187 annual abundance estimates from 1943 to 2019 with a taxonomic resolution varying from species to order levels. This corresponds to 94% of the abundance estimates from the original dataset.

For data from van Klink et al. (2020) we used the file “aax9931-Van-Klink-SM-CORRECTED-Data-S1.txt.” Some data from American LTER sites and from BIOTIME could be duplicated with time series from Crossley et al. (2020) and from BIOTIME respectively, but because van Klink et al. aggregated data at taxonomic order (Fig. S1), we considered them as different. We also removed biomass data to keep abundance data only. We restricted our analysis to time series from North America and Europe as other locations were poorly represented. After data filtering, we obtained 26,785 annual abundance estimates from 1932 to 2018 with a taxonomic resolution at order level. This corresponds to 43% of the abundance estimates from the original dataset.

For BIOTIME data we extracted all data, and we kept only time series regarding arthropods from terrestrial and freshwater habitats, from North America and Europe. We removed LTER data to avoid duplicated time series with Crossley et al. (2020). We inferred zeros on a given plot for a given species for a given year when the plot has data for at data from the same taxonomic order than the focal species for the given year. Then we removed time series with only zero abundance values. After data filtering, we obtained 79,250 annual abundance estimates from 1898 to 2016 with a taxonomic resolution varying from species to order levels.

Assessing monotony of abundance variation over time

We estimated non-monotony of temporal abundance variations using a Generalized Additive Model (GAM) for each original time series, using the mgcv R package (Wood 2017). These GAMs have a Poisson error structure (with a log link function) for abundance data from van Klink et al. (2020), Crossley et al. (2020) and BIOTIME, or a gaussian error structure for logit transformed occupancy estimates from Outhwaite et al. (2019). We used a smooth (spline) effect of the year. For time series sampled at different periods, we added a spline effect of the period penalized by a ridge penalty, to model it as a random effect. To avoid non-identifiable models, the dimension of the basis used to represent the year smooth term is constrained to be smaller than to the number of years (duration) of the time series, minus one if it is sampled over different sampling periods, with a maximum basis value of 10.

Then extracting the polynomial effect of the year on abundance, we assessed the non-monotony of this polynomial effect as the number of local maximums and minimums (i.e. turning points) observed.

Comparing abundance trends among species

A common problem when comparing abundance trends over many species is that estimated trends are not easily comparable among species, especially between rare and common species, since abundance trends depend on initial abundance. Indeed, two species, a rare and an abundant one, with an abundance shifting in a year from 10 to 15 individuals and from 10,000 to 15,000 individuals respectively, are both growing with a rate of 50% per year. However, measuring these abundance variations on the number of individuals will give a gain of 5 individuals for the first one against a gain of 5,000 individuals for the second species. Demographic effects measured on rough measure of abundance (number of individuals, occupancy probability, etc.) depend on the level of abundance, hiding the decline of rare species (Fig. S2 & S3). Note that standardizing abundances of each species by subtracting the mean and dividing by standard deviation does not solve this problem in a proper way, since it does not allow to estimate growth rate but
can instead create artefacts by increasing the importance of stable species with low inter-annual variation in their abundance, relatively to species declining/increasing but with high inter-annual variations in abundance (Fig. S4).

This dependency of abundance trends to initial abundance can be overcome by expressing population trends in terms of growth rates, a multiplicative factor which correspond to the growth of the population. In a time-discrete system, a growth rate > 1 is associated with increasing abundance over time, a growth rate equals to 1 corresponds to a constant abundance over time, while a growth rate < 1 is associated with decreasing abundance over time. Such measure allows to compare trends among species, regardless of their initial abundance, but is cannot be estimated directly using classic linear models. Indeed, the demography of a population with growth rate \( r \), as shown in Fig. S6, can be expressed by the following geometric progression:

\[
A_t = A_{t=0} \times r^t
\]

where \( A_t \) is the abundance at time \( t \), \( A_{t=0} \) is the initial abundance (intercept), while \( r \) is the growth rate. The problem is that this product cannot be estimated using classic linear models which, by definition, estimate linear functions (\( y = b + ax \)). Here, we transformed occupancy estimates using a \( \logit \) function and abundance counts using a \( \log \) function to linearize geometric changes, then allowing to estimate growth rates using linear models. If we study the logarithm of the abundance instead of the abundance, then the function describing the population demography becomes linear:

\[
\log (A_t) = \log(A_{t=0} \times r^t)
\]

\[
\log (A_t) = \log(A_{t=0}) + \log(r^t)
\]

\[
\log (A_t) = \log(A_{t=0}) + t \times \log (r)
\]

If we set \( a = \log (r) \) and \( b = \log(A_{t=0}) \), we have:

\[
\log (A_t) = b + at
\]

Thus, by regressing linearly \( \log(abundance) \) against time we can estimate the logarithm of the growth rate, which is a measure of abundance trend independent of the initial abundance (i.e., the rarity) of species (Figure S6). Since \( \log(1) = 0 \), then the sign and the value of the estimated slope indicate the direction and the magnitude of the abundance trend, respectively. However, since \( \log(0) \) is not defined and since we have zero abundance values, we need to use a model structure which allows zero abundances. Usually, studies use \( \log (A_t + \varepsilon) \) instead of \( \log (A_t) \), choosing \( \varepsilon = 1 \) arbitrarily (Crossley et al. 2020; van Klink et al. 2020), while the value of \( \varepsilon \) will strongly affect the result by breaking the linearization of the geometric progression (Fig. S2).

Here, to avoid this arbitrary transformation, we used a Generalized linear model (GLM) with a Poisson error structure, using a \( \log \) link function, which allows to estimate \( \log (A_t) = b + at \) by considering zero abundance data.

Regarding the occupancy probabilities, the same logic applies but we have to account for the fact that the proportion of grid cells occupied by the species is bounded between 0 and 1. So the population dynamic can be view as a logistic regression with a carrying capacity \( (K) \) of 1:

\[
\frac{dP_t}{dt} = r^*P_t \left(1 - \frac{P_t}{K}\right)
\]

where \( r^* \) is the growth rate of a time-continuous system (0 = stable, negative = decline, positive = increase). Thus, we have:

\[
P_t = \frac{KP_0e^{-r^*t}}{K + P_0(e^{-r^*t} - 1)} = \frac{K}{1 + \frac{K-P_0}{P_0}e^{-r^*t}}
\]

If we set \( K = 1 \), we have:

\[
P_t = \frac{1}{1 + \frac{1-P_0}{P_0}e^{-r^*t}}
\]

If we logit transform \( P_t \), we get:

\[
\log \left( \frac{P_t}{1-P_t} \right) = \log(P_t) - \log(1-P_t) = -\log \left( \frac{K-P_0}{P_0} \right) + r^*t
\]

Switching from \( r^* \) to \( r \), so from continuous to discrete time we get:

\[
\log \left( \frac{P_t}{1-P_t} \right) = \log \left( \frac{K-P_0}{P_0} \right) + t \times \log (r)
\]
where $r$ is the growth rate of a discrete time system ($1 = \text{stable, below 1 = decline, above 1 = increase}$). If we set $a = \log (r)$ and $b = -\log \left(\frac{K-e_0}{e_0}\right)$, we have:

$$\log \left(\frac{P_t}{1-P_t}\right) = b + at$$

Thus, by regressing linearly $\logit(\text{occupancy probability})$ against time we can estimate, as previously, the logarithm of the growth rate, which is a measure of abundance trend independent of the initial abundance (i.e. the rarity) of species (Figure S3).

So, here we used a GLM with a binomial error structure, using a logit link function, which allows to estimate $\logit (P_t) = b + at$.

References


**Figure S1**: Taxonomic distribution of time series. (a) Number of species across taxonomic groups and (b) source datasets, as a function of their taxonomic resolution.
Figure S2: Representation of a geometric decline over time. (a) Abundance, measured as number of individuals, against time, for two species declining with a rate of 20% per year (growth rate = 0.8). Zooming on a part on the bottom of the y-axis, we can see that both species decline with the same rate, but since one is rare and the other is common, variations in number of individuals are much larger for the common species. (b,c,d) represents the same dynamics as in a, but applying the transformation \( \log(x + \varepsilon) \) to the abundance, with three values of \( \varepsilon \).
Figure S3: Representation of a geometric decline over time. (a) Occupancy probability, measured as the proportion of grid cell occupied by a species on a given area, against time, for two species declining at a growth rate of 0.8, as in figure S2. Zooming on a part on the bottom of the y-axis, we can see that both species decline with the same rate, but since one is rare and the other is common, variations in number of individuals is much larger for the common species. (b) represents the same dynamics as in a, but when applying the transformation logit(x) to the occupancy probability. On the logit scale we can see that declines are linear and identical (same slopes) between both species, and can easily be estimated with a linear model.
Figure S4: Simulated data showing how standardization can affect abundance trends. (a) abundance values over time and associated abundance trends, for one strongly declining species, with high inter-annual variation in abundance (green), and for another species slightly declining, with low inter-annual variation in abundance (purple). (b) shows the same data as in (a) but after standardization of abundance values within each time series (minus mean and then divided by standard deviation). In (b) abundance trends are calculated on standardized data, giving similar trends among species while in fact the green species is declining more than the purple one. Because purple species exhibit low inter-annual variability in abundance, the absolute value of its trend is artefactually increased by the standardization.
Figure S5: Truncated time series with only one non-zero yearly estimate of abundance produce extreme value of abundance trends. Density distribution of the absolute value of abundance trends, log(growth rate), as function of the number of non-zero yearly estimate of abundance contained in the truncated time series (1 vs >1). To preserve readability the x-axis is square-root transformed.
Figure S6: Same figure as in the main text (Figure 3b), but for each source dataset. Proportion of abundance trends from truncated time series with different directions (positive vs. negative), as a function of the number of turning points in the corresponding original time series. Boxplots represent minimum and maximum values (bottom and top of vertical lines), first and third quartiles (Q1 and Q3, bottom and top of boxes) and median (thick horizontal lines); colours indicate sample size (number of original time series). Points with values outside of the range \([Q1-1.5(Q3-Q1), Q3+1.5(Q3-Q1)]\) are considered as outliers and represented as full circles.
Figure S7: **Estimated growth rate when a decline of 15%/year is simulated.** This is the same plot than in the left panel of Figure 4d, but without the outliers to improve readability of average biases. The dashed horizontal line shows the value of the logarithm of the true (simulated) growth rate.
Figure S8: Random effects of taxonomic order on abundance trends. The dots show the mean of the posterior distribution while the error bars show the confidence interval at 95% (quantile 2.5% and 97.5%).
Figure S9: Estimated abundance trends of truncated time series as a function of average abundance. Abundance trends (log growth rate) as a function of the average abundance of the original time series (measured on the original abundance scale), per source dataset and habitat.
Table S1: Description of the 4 datasets used.

<table>
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<tr>
<th>Dataset's name</th>
<th>Kind of abundance estimate</th>
<th>Original spatial scope</th>
<th>Spatial scope used here</th>
<th>Taxonomical resolution of time-series</th>
<th>Original temporal coverage</th>
<th>Temporal coverage used here</th>
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<tr>
<td>Outhwaite et al. 2019</td>
<td>Annual occupancy estimate (i.e. the proportion of 1km² grid cells in a region occupied by a species, a proxy for abundance)</td>
<td>Great-Britain or United-Kingdom or region (Wales, England, Scotland, Northern Ireland) levels</td>
<td>Time-series at Great-Britain level</td>
<td>Species level</td>
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<td>1970-2015</td>
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<td>Crossley et al. 2019</td>
<td>Local annual abundance count</td>
<td>Local time series spread across the USA</td>
<td>Local time series spread across the USA</td>
<td>Species level mostly (cf. Figure S1)</td>
<td>1943-2019</td>
<td>1970-2019</td>
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<tr>
<td>van Klink et al. 2020</td>
<td>Local annual abundance count, aggregated from literature</td>
<td>Local time series spread across the world</td>
<td>Local time series spread across North America and Europe</td>
<td>Order level</td>
<td>1925-2018</td>
<td>1970-2018</td>
</tr>
<tr>
<td>BIOTIME database</td>
<td>Local annual abundance count</td>
<td>Local time series spread across the world</td>
<td>Local time series spread across North America and Europe</td>
<td>Species level</td>
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<td>1970-2018</td>
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Table S2: Number of original time series across datasets, continents and habitats, at the end of the filtering process.

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<th>aquatic</th>
<th>terrestrial</th>
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