Stability of flower visitation rate over time is driven by asynchrony of pollinator abundances but does not determine the stability of plant reproductive success.

Estefanía Tobajas, Virginia Domínguez-García, Francisco P. Molina, Ignasi Bartomeus

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Abstract

- 1. The stability of ecological communities over time, often quantified using its aggregated abundance or biomass, is mainly influenced by species richness and the asynchrony of the species fluctuations. However, the relation with the stability of ecosystem functioning provided by this community is less explored, specially for functions involving several trophic levels such as plant pollination. Understanding the mechanisms driving community stability and whether this stability should be related to the stability of ecosystem functions provided is important for predicting the resilience of communities to changes in biodiversity.
- 2. Using a 5-year dataset across 13 Mediterranean shrub patches, we assessed the effect of the richness and asynchrony of pollinators as drivers of the stability of plant's visitation rate and plants reproductive success. In addition, we analyzed whether the effects of species richness and visitation rate on plant reproduction success were consistent among years.
- 3. The stability of plant's visitation rate is driven by the asynchrony of pollinator species abundances but not by the pollinator richness. In contrast, the stability of the plant reproductive success is not related to the stability of the visitation rate. A year by year analysis revels that plants are unlikely to be limited by pollinator abundances except in specific years. extreme low levels of pollinator activity
- 4. CONCLUSION

Introduction

Global environmental changes and the consequent loss of biodiversity have highlighted the need for better understanding the role of biodiversity in ecosystem function (Hong et al., 2021; Hooper et al., 2005; Lefcheck et al., 2015; Oliver et al., 2015). We have experimental and observational evidence that biodiversity enhances aggregated community-level properties, such as total biomass (Cardinale et al., 2006; Cardinale et al., 2012; Tilman et al., 2014), but the role of biodiversity for the stability of ecosystem functioning is less explored (but see Griffin et al. (2009); Grman et al. (2010)), specially for functions involving several trophic levels such as plant pollination (Ebeling et al., 2008; Garibaldi et al., 2011; Lázaro et al., 2022; Winfree & Kremen, 2009). Globally, it is estimated that at least 78% of wild plants depend to some extent on pollination by animals (Ollerton et al., 2011). Understanding the mechanisms driving pollinator community stability and how this relates to the stability of ecosystem functions and services delivered (i.e. pollination) is important for predicting changes in both community structure and ecosystem function.

Temporal stability is defined as the capacity of an ecosystem to dampen environmental perturbations over time while retaining ecological functions of interest (Ives & Carpenter, 2007; Pimm, 1984). The temporal

stability of community can be influenced by the diversity of species present and the asynchrony on the species abundance fluctuations. A higher number of species might stabilize community level functions by increasing the range of species level potential responses to environmental changes (i.e. the portfolio effect) (Hector et al., 2010; McCann, 2000; Tilman et al., 2006; Q. Xu et al., 2021), although non-significant relationships between diversity and stability have also been observed (Hooper et al., 2005; Plas, 2019). The degree of asynchrony in species abundance fluctuations is also a stabilizing mechanisms. In this sense, higher levels of asynchrony stabilize the community (Blüthgen et al., 2016; Craven et al., 2018; Ruijven & Berendse, 2007). Species asynchrony has been found to be a key factor driving positive relationships between diversity and stability (Loreau & De Mazancourt, 2008; Z. Xu et al., 2015) or even that stability is often more strongly associated with the degree of asynchrony across species than with species richness (Blüthgen et al., 2016; Huang et al., 2020). Other studies have indicated that temporal stability of the dominant species or asynchrony among these dominant species is important in maintaining the stability of ecosystems (Sasaki & Lauenroth, 2011; Valencia et al., 2020). While community temporal stability should be related to the stability of ecosystem functions and services provided by such community, there is limited evidence linking both processes on multitrophic systems.

Since the reproductive success of most plant species highly depends on pollination by animals (Ollerton et al., 2011), the stability of pollinator communities may increase the stability of plant reproductive success. A greater diversity of pollinators has been shown to increase plant reproductive success as a result of complementary in service provision arising from variation of pollinator effectiveness across space and time (Albrecht et al., 2012; Blüthgen & Klein, 2011; Fründ et al., 2010, 2013; Hoehn et al., 2008). For example, pollinator species may exhibit different ranges of temporal activity (Pisanty et al., 2016; Rader et al., 2016) and consequently may provide pollination services at different times of the day or of the season. While some studies report that sites with higher visitation rates show higher plant reproductive outputs (Garibaldi et al., 2013; Kaiser-Bunbury et al., 2017), other studies do not find such relationship (Hegland & Totland, 2012; Lundgren et al., 2013). Interestingly, studies looking at pollen limitation show that sites with higher visitation rates show no pollen limitation, indicating than above a certain threshold, changes in visitation rates are no longer relevant [Fig. 1; Cosacov et al. (2008); Williams & Winfree (2013)]. However, the occurrence and magnitude of pollen limitation can vary among years for individual populations (Ashman et al., 2004; Burd, 1994), primarily because of variation in pollinator service (Aizen & Harder, 2007).

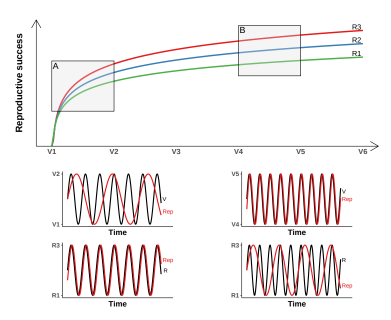


Figure 1. Conceptual figure showing the general relationship between visitation rate of pollinators and plant reproductive success to different species richness. As the number of visits from pollinators increases, the plant reproductive success is expected to increase until than above a certain threshold, changes in visitation rates are no longer relevant to plant reproductive success.

Here, we use a 5-year dataset collected for a total of 12 plant species across 13 Mediterranean woodland patches but to study the role of pollinator richness and asynchrony as drivers of stability of its visitation rates we use the 5 most represented plant species. Further, we evaluate whether more stable pollinator communities foster a more stable plant reproductive success along years, measuring both the stability of fruit and seed set. We expect that the stability of visitation rates will increase with higher richness and asynchrony of pollinators. In addition, a greater stability of plant reproductive success is expected with higher stability of visitation rate. However, given the lack of a clear relationship between pollinator and plant reproductive output stability, we further analyzed the variability on the yearly effect of pollinator richness and visitation rate on the reproductive success of the plants.

Material and method

Our study was conducted in the southwest of the Iberian Peninsula (Huelva and Sevilla - Spain). We selected 13 sites in Mediterranean shrub ecosystems that ranged from sites with little to no disturbances to more modified ones with crops and urban areas nearby. These sites are similar in the plant community composition and the most abundant plant species are known to be attractive to flower visitors and to depend on pollinators to maximize its reproduction (e.g. dominated by Cistus crispus, Cistus ladanifer, Cistus salviifolius, Halimium halimifolium, and Lavandula pedunculata). Each site was surveyed at least 7 times per year during the flowering season from 2015 to 2019 (from March to June). Within each site, we selected 3-4 individuals belonging to 3-9 plant species (mean \pm standard error: 5.92 ± 0.54 plant species per site). We performed focal observations where all the floral visitors (from now on referred to as pollinators) that contacted with their flowers were registered. Only pollinators that could not be identified in the field were captured, stored, and identified to species level in the laboratory. For each plant individual, we counted the total number of flowers observed in each survey and at the end of the season, we recorded the number of fruits developed and the number of seeds per fruit in a subsample of 3 fruits. Focal observations had a duration of five minutes per individual in 2015-2016 and three minutes from 2017 to 2019. Hence, for each plant individual, the sampling effort was standardized in number of visits per flower per minute. To that end, we divided the total visits observed by the number of flowers observed and by the total observation time (time observation multiplied by number of survey per year). Pollinator richness is calculated as the number of pollinator species observed visiting the plant, fruit set as the proportion of fruits developed, and the seed set as the mean number of seeds per fruit for each plant individual sampled per site.

Stability

We analyze five plant species (Cistus crispus, Cistus ladanifer, Cistus salviifolius, Halimium halimifolium, and Lavandula pedunculata) that were recorded in at least 5 sites in a minimum of two years per site (show mean sites / years per plant). The stability of pollinator visitation rates was quantified as the inverse of the coefficient of variation (mean/SD) for each plant species in each plot following Tilman, 1999. To examine the role of richness and synchrony of pollinator species abundances as drivers of stability of visitation rate, we calculate the richness as the total number of pollinator species that visit each plant species in each plot. The synchrony index for each plant species in the plot was calculated following Loreau and de Mazancourt (2008), as:

$$\phi = \frac{\sigma_{x_T}^2}{(\sum_{i=1}^S \sigma_{x_i})^2}$$

where $\sigma_{x_T}^2$ is the variance of the community abundance of each plot and σ_{x_i} is the standard deviation of species i abundance in a community with S species. This synchrony measurement varies from zero, when fluctuations are perfectly asynchronous, to one, when fluctuations are perfectly synchronous (Loreau & De Mazancourt, 2008). Finally, we express asynchrony as $1 - \phi$. We also calculated the stability of fruit proportion and of the seed number for each plant species in each plot.

Statistical analysis

To discover the main drivers of visitation rate stability on the five plant species studied, we carried out two linear mixed models using as response variable the stability of visitation rate and as explanatory variables pollinator asynchrony and richness, respectively. Similarly, linear mixed model was used to test the influence of pollinator richness on asynchrony of pollinator species abundances. In all cases, site and plant species were included as random factor. For a deeper understanding of the processes stabilizing visitation rate, we also analyzed the response for each plant species separately.

To study whether the stability of the visitation rate influenced in the stability of the plant reproductive success, we run linear mixed models using as response variables the stability of fruit proportion or the stability of seed number and as explanatory variable the visitation rate stability. Site and plant species were included as factor random.

Finally, we further analyzed the variability on the yearly effect of pollinator richness and visitation rate on the reproductive success of the plants. For this we employed all plant species sampled and we analyzed the interactive effect between richness and year, and between visitation rate and year on fruit proportion using binomial generalized linear mixed model, and on the number of seeds employing linear mixed model. P values were obtained using the type III Anova (package: car) The number of seeds per fruit was centered and scaled per plant species to allow meaningful comparisons across species. For all models, we included plant species identity nested within site and site as random effects because of multiple individuals of the same plant species are measured at each site.

All the analyses of this study were conducted in R software (v. 4.0.2; R Core Team, 2022) and the lme4 (Bates et al., 2015) and ggplot2 (Wickham, 2016) packages. DHARMa package was employed for checking all models (Harting 2017)

Results

Stability of visitation rate and its drivers: richness and asynchrony

Stability of visitation rates ranged from 0.64 to 9.24 (mean \pm SE: 1.928 \pm 0.232), whereas pollinator richness ranged from 2 to 20 species (8.128 \pm 0.686) and the pollinator asynchrony varied from 0 to 0.998 (0.754 \pm 0.032). There is a strong positive relationship between stability of visitation rates and asynchrony of pollinator species abundances with greater visitation rate stability when pollinator asynchrony increases (Estimate \pm standard error: 2.903 \pm 1.134; p = 0.017) (Fig 2A), whereas stability of visitation rate was not influenced by the pollinator richness (-0.001 \pm 0.063; p = 0.986) (Fig 2B). We also found a positive relationship between asynchrony and pollinator richness (0.020 \pm 0.005, p = 0.001) (Fig 2C). Furthermore, we observed one data point has a high leverage on the stability analysis (*C.crispus*, Fig 2A-B), when we remove this point, although the relationship is not significant, the tendency is still positive (0.968 \pm 0.745; p = 0.215).

When we analysed each plant species separately, we observed positive relationship between asynchrony of pollinators and stability of visitation rate for *C. crispus*, *H. halimifoliun* and *L. pedunculata*, weak relationship for *C. salviifolious* and a weak negative relationship for *C.ladanifer*. For the relationship with pollinator richness, *C. crispus*, *C. ladanifer* and *C. salviifolius* showed a negative relationships wile *H. halimifolium* and *L. pedunculata* showed positive relationships, explaining the overall lack of relationship across all five plants (Supplementary material-Fig 1).

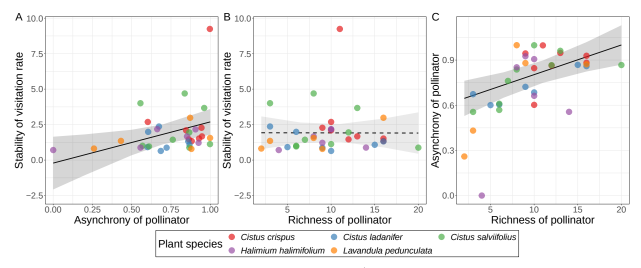


Figure 2.Relationships between stability of visitation rate (calculated as the inverse of the coefficient of variation) and asynchorny of pollinators (Loreau and de Mazancourt's index) (A), stability of visitation rate and pollinator richness (B), and asynchorny of pollinators and richness (C). Dots represent the different plant species included in the analysis. Significant models are representing with solid line.

Stability of visitation rate and stability of the plant reproductive success

Stability of fruit proportion varied from 2.62 to 120.66 (mean \pm SE: 19.806 \pm 3.615), whereas the stability of seed number ranged 1.28 to 55.86 (7.018 \pm 1.441). When we analyzed whether the stability of the pollinator's visitation rate affected the stability of the plant reproductive success, we did not find a clear relationship when analyzing the fruit proportion (Estimate \pm standard error: -1.547 \pm 2.611; p=0.558) (Fig 3A). However, contrary to expected, the stability of seed number did show a negative relationship with the stability of pollinator's visitation rate (-1.887 \pm 0.959; p=0.061) (Fig 3B). Although this tendency is influenced by a high value in the stability of seed number of *H. halimifolium* (Fig 3B) since when we removed this point the relationship disappears (0.561 \pm 0.535; p=0.306).

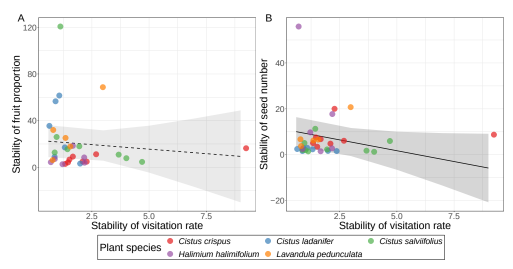


Figure 3. Relationship between stability of plant reproductive succes: fruit proportion (A) and seed number (B) and the stability of visitation rate. Dots represent the different plant species included in the analysis. Significant models are representing with solid line.

The effect of pollinators on reproductive success depends on the year

For analyses whether the variability of pollinator richness and visitation rate across years affected to the reproductive success of plants we employed total plant species sampled. When we analysed the fruit proportion, we found that the interaction between year and pollinator richness, and year and visitation rate affected the fruit proportion (richness and year: $\chi 2 = 28.976$, df = 4, p = <0.0001; visitation rate and year: $\chi 2 = 40.691$, df = 4, p = <0.0001). Furthermore, we observed a significant effect of year by itself, not so of pollinator richness or visitation rate. Although we observed a general saturation of the proportion of fruits developed with most species consistently over 80% of fruit set, in years like 2017 we found that a greater richness and visitation rate increased the fruit proportion. We also observed a positive relationship between richness and fruit proportion in 2019. However in 2018, the relationship between richness and fruit proportion was negative (Fig 3). esto requiere un poco más de story telling. Por ejemplo, estos años son años altos o bajos en polinizadores?

For the seed number, we found an effect of year in both models in addition to the interaction between visitation rate and year ($\chi 2 = 9.113$, df = 4, p = 0.058) but not the interaction between pollinator richness and year ($\chi 2 = 2.592$, df = 4, p = 0.628). We observed that a higher visitation rate increased the seed number in 2019 (Supplementary material-Fig 2).

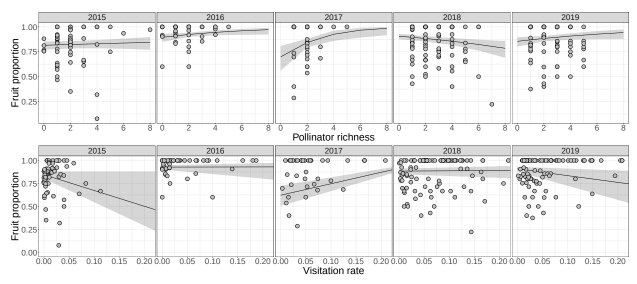


Figure 4. Interaction effect between richness and year (top row), and visitation rate and year (bottom row) on fruit proportion. For the visualization purpose, we omit in the visitation rate (x-axis) the values outside of the 95%.

Discussion

We show that pollinator's visitation rate stability increase with higher asynchrony of pollinators species abundances and that greater species richness only increase stability of visitation rate indirectly, via enhanced species asynchrony. However, contrary to expectations, we found no relationship between the stability of visitation rates and the stability of plant reproductive success. This is because in our study system, plant reproduction is not pollinator limited in most years, with most plants reaching maximum reproductive success even in the sites with lower pollinator abundances. However, this do not imply that pollinators are not important or that are always in enough numbers to disregard them, because for specific years the relationship between pollinator activity and plant's reproductive output kicks in as important seria ideal decir quales... los que tienen poco polinizador? Los secos, quizas? When and where pollinators are limiting is the key question then. [esta frase a lo mejor puede ser una conclusing sentence del paper, por ahora la dejo aquí]

While drivers of temporal stability have been well study in plant communities (Craven et al., 2018; Gross et al., 2014; Kigel et al., 2021; Lepš et al., 2018; Ruijven & Berendse, 2007; Sasaki et al., 2019; Valencia et al., 2020), relatively few studies have evaluated how this same drivers can affect community stability in mobile organisms such as pollinators (Ebeling et al., 2008; Lázaro et al., 2022; Senapathi et al., 2021). A first clear outcome of our analysis is that asynchrony of species abundances is a key driver of visitation rate stability. This stability-asynchrony relationship has been found by different theoretical and empirical studies (Blüthgen et al., 2016; Kigel et al., 2021; Lepš et al., 2018; Mazancourt et al., 2013; Sasaki et al., 2019; Valencia et al., 2020). For example, Lázaro et al. (2022) also found for plants, pollinators and their interactions that more asynchronous communities being more temporal stable. The fact that species asynchrony increases the temporal stability of the visitation rate is likely because a decrease in the abundance of one pollinator species is compensated by an increase of the abundance of another. [mirate los appaers de Winfree sobre esto que lo explica muy bien y explica mecanismos. On the other hand, species richness is considered one of the strongest drivers of community stability [in plants?] (Tilman et al., 2006; de Mazancourt et al., 2013; Hallett et al., 2014), although Valencia et al. (2020) have reported that this relation is not nearly as strong [where?]. Pollinator studies have observed a positive relationship between richness and stability of what? in crops?? (Ebeling et al 2008; Senapathi et al 2021), however, our results show that richness was positively associated with asynchrony, which might positively affect the stability of visitation rate, but indirectly. This most likely reflects niche variation among species that affect their relative performance over time in a temporally variable environment (Yachi & Loreau, 1999; Gross et al., 2014; Craven et al 2018).

Despite the stability is crucial to the sustainability of ecosystem functions, relatively few studies have evaluated the stability of the pollinator visits frequency on pollination services, and those who did used agricultural systems [right?] (Garibaldi et al. 2011, Lázaro and Alomar 2019, Hünicken et al 2021, Bishop et al 2022). The majority of these studies were focused on crops, however, to our knowledge, how the stability of pollinator community affect to the reproductive success of the wild plant community remaining largely unknown. In pollinator-dependent crops have been observed a greater yield stability with the pollinator diversity and the stability of visitation rate (Garibaldi et al. 2011, Hünicken et al 2021), however in our study system (wild plants), we did not find this relationship or even observed a negative relationship between seed number stability and visitation rate stability.

Supplementary material

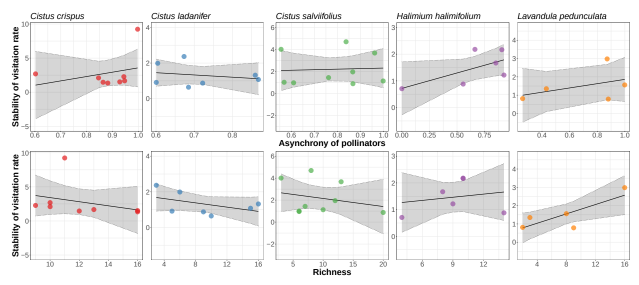


Figure 1. Changes in the stability of visitation rate with the asynchrony (top row) and richness (botton row) are shown for each plant species

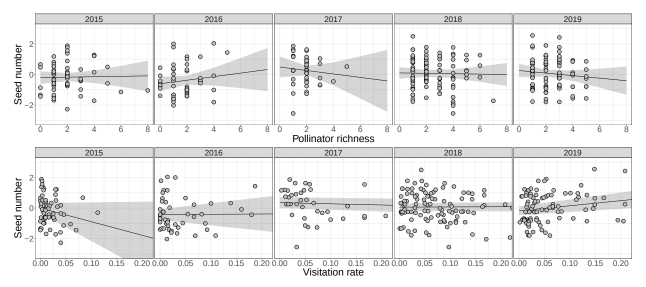


Figure 2. Interaction effect between richness and year (top row), and visitation rate and year (bottom row) on seed number. For the visualization purpose, we omit in the visitation rate (x-axis) the values outside of the 95%.

References

Aizen, M. A., & Harder, L. D. (2007). Expanding the limits of the pollen-limitation concept: effects of pollen quantity and quality. *Ecology*, 88(2), 271–281.

Albrecht, M., Schmid, B., Hautier, Y., & Müller, C. B. (2012). Diverse pollinator communities enhance plant reproductive success. *Proceedings of the Royal Society B: Biological Sciences*, 279(1748), 4845–4852. https://doi.org/10.1098/rspb.2012.1621

Ashman, T. L., Knight, T. M., Steets, J. A., Amarasekare, P., Burd, M., Campbell, D. R., Dudash, M. R.,

- Johnston, M. O., Mazer, S. J., Mitchell, R. J., Morgan, M. T., & Wilson, W. G. (2004). Pollen limitation of plant reproduction: ecological and evolutionary causes and consequences. *Ecology*, 85(9), 2408–2421.
- Blüthgen, N., & Klein, A.-M. (2011). Functional complementarity and specialisation: The role of biodiversity in plant-pollinator interactions. *Basic and Applied Ecology*, 12(4), 282–291. https://doi.org/10.1016/j.baae.2010.11.001
- Blüthgen, N., Simons, N. K., Jung, K., Prati, D., Renner, S. C., Boch, S., Fischer, M., Hölzel, N., Klaus, V. H., Kleinebecker, T., Tschapka, M., Weisser, W. W., & Gossner, M. M. (2016). Land use imperils plant and animal community stability through changes in asynchrony rather than diversity. *Nature Communications*, 7. https://doi.org/10.1038/ncomms10697
- Burd, M. (1994). Bateman's principle and plant reproduction: the role of pollen limitation in fruit and seed set. *The Botanical Review*, 60(4), 373–425.
- Cardinale, B. J., Duffy, J. E., Gonzalez, A., Hooper, D. U., Perrings, C., Venail, P., Narwani, A., Mace, G. M., Tilman, D., Wardle, D. A., & Kinzig, A. P. (2012). Biodiversity loss and its impact on humanity. Nature, 486(7401), 59–67. https://doi.org/10.1038/nature11148.Access
- Cardinale, B. J., Srivastava, D., Duffy, J. E., Wright, J. P., Downing, A. L., Sankaran, M., & Jouseau, C. (2006). Effects of biodiversity on the functioning of trophic groups and ecosystems. *Nature*, 443(7114), 989–992.
- Cosacov, A., Nattero, J., & Cocucci, A. A. (2008). Variation of pollinator assemblages and pollen limitation in a locally specialized system: The oil-producing Nierembergia linariifolia (Solanaceae). *Annals of Botany*, 102(5), 723–734. https://doi.org/10.1093/aob/mcn154
- Craven, D., Eisenhauer, N., Pearse, W. D., Hautier, Y., Isbell, F., Roscher, C., Bahn, M., Beierkuhnlein, C., Bönisch, G., Buchmann, N., Byun, C., Catford, J. A., Cerabolini, B. E. L., Cornelissen, J. H. C., Craine, J. M., De Luca, E., Ebeling, A., Griffin, J. N., Hector, A., ... Manning, P. (2018). Multiple facets of biodiversity drive the diversity-stability relationship. Nature Ecology and Evolution, 2(10), 1579–1587. https://doi.org/10.1038/s41559-018-0647-7
- Ebeling, A., Klein, A.-M., Schumacher, J., Weisser, W. W., & Tscharntke, T. (2008). How does plant richness affect pollinator richness and temporal stability of flower visits? *Oikos*, 117(12), 1808–1815. https://doi.org/10.1111/j.1600-0706.2008.16819.x
- Fründ, J., Dormann, C. F., Holzschuh, A., & Tscharntke, T. (2013). Bee diversity effects on pollination depend on functional complementarity and niche shifts. *Ecology*, 94(9), 2042–2054. https://doi.org/10.1890/12-1620.1
- Fründ, J., Linsenmair, K. E., & Blüthgen, N. (2010). Pollinator diversity and specialization in relation to flower diversity. Oikos, 119(10), 1581–1590. https://doi.org/10.1111/j.1600-0706.2010.18450.x
- Garibaldi, L. A., Steffan-Dewenter, I., Kremen, C., Morales, J. M., Bommarco, R., Cunningham, S. A., Carvalheiro, L. G., Chacoff, N. P., Dudenhöffer, J. H., Greenleaf, S. S., Holzschuh, A., Isaacs, R., Krewenka, K., Mandelik, Y., Mayfield, M. M., Morandin, L. A., Potts, S. G., Ricketts, T. H., Szentgyörgyi, H., ... Klein, A. M. (2011). Stability of pollination services decreases with isolation from natural areas despite honey bee visits. *Ecology Letters*, 14(10), 1062–1072. https://doi.org/10.1111/j.1461-0248.2011.01669.x
- Garibaldi, L. A., Steffan-Dewenter, I., Winfree, R., Aizen, M. A., Bommarco, R., & Cunningham, S. A., ... & Bartomeus, I. (2013). Wild Pollinators Enhance Fruit Set of Crops Regardless of Honey Bee Abundance. *Science*, 339, 1608–1611. https://doi.org/10.1126/science.1230200
- Griffin, J. N., O'Gorman, E. J., Emmerson, M. C., Jenkins, S. R., Klein, A.-M., Loreau, M., & Symstad, A. (2009). Biodiversity and the stability of ecosystem functioning. In *Biodiversity, ecosystem functioning, and human wellbeing: An ecological and economic perspective.* (pp. 78–93). https://doi.org/10.1093/acprof
- Grman, E., Lau, J. A., Schoolmaster, D. R., & Gross, K. L. (2010). Mechanisms contributing to stability in ecosystem function depend on the environmental context. *Ecology Letters*, 13(11), 1400–1410. https://doi.org/10.1111/j.1461-0248.2010.01533.x
- Gross, K., Cardinale, B. J., Fox, J. W., Gonzalez, A., Loreau, M., Wayne Polley, H., Reich, P. B., & Ruijven, J. van. (2014). Species richness and the temporal stability of biomass production: A new analysis of recent biodiversity experiments. *American Naturalist*, 183(1), 1–12. https://doi.org/10.1086/673915
- Hector, A., Hautier, Y., Saner, P., Wacker, L., Bagchi, R., Joshi, J., Scherer-Lorenzen, M., Spehn, E. M., Bazeley-White, E., M.Weilenmann, Caldeira, M. C., Dimitrakopoulos, P. G., Finn, J. A., Huss-Danell, K., Jumpponen, A., & Loreau, M. (2010). General stabilizing effects of plant diversity on grassland

- productivity through population asynchrony and overyielding. Ecology, 91(8), 2213–2220.
- Hegland, S. J., & Totland, Ø. (2012). Interactions for pollinator visitation and their consequences for reproduction in a plant community. *Acta Oecologica*, 43, 95–103. https://doi.org/10.1016/j.actao.2012.06.002
- Hoehn, P., Tscharntke, T., Tylianakis, J. M., & Steffan-Dewenter, I. (2008). Functional group diversity of bee pollinators increases crop yield. *Proceedings of the Royal Society B: Biological Sciences*, 275(1648), 2283–2291. https://doi.org/10.1098/rspb.2008.0405
- Hong, P., Schmid, B., De Laender, F., Eisenhauer, N., Zhang, X., Chen, H., Craven, D., De Boeck, H. J., Hautier, Y., Petchey, O. L., Reich, P. B., Steudel, B., Striebel, M., Thakur, M. P., & Wang, S. (2021). Biodiversity promotes ecosystem functioning despite environmental change. *Ecology Letters*, 25(2), 555–569. https://doi.org/10.1111/ele.13936
- Hooper, D. U., Chapin, F. S., Ewel, J. J., Hector, A., Inchausti, P., Lavorel, S., Lawton, J. H., Lodge, D. M., Loreau, M., Naeem, S., Schmid, B., Setälä, H., Symstad, A., Vandermeer, J. H., & Wardle, D. A. (2005). Effects of biodiversity on ecosystem functioning: a consensus of current knowledge. *Ecological Monographs*, 75(1), 3–35. https://doi.org/10.1890/04-0922
- Huang, M., Liu, X., & Zhou, S. (2020). Asynchrony among species and functional groups and temporal stability under perturbations: Patterns and consequences. *Journal of Ecology*, 108(5), 2038–2046. https://doi.org/10.1111/1365-2745.13418
- Ives, A. R., & Carpenter, S. R. (2007). Stability and diversity of ecosystems. Science, 317(5834), 58–62. https://doi.org/10.1126/science.1133258
- Kaiser-Bunbury, C. N., Mougal, J., Whittington, A. E., Valentin, T., Gabriel, R., Olesen, J. M., & Blüthgen, N. (2017). Ecosystem restoration strengthens pollination network resilience and function. *Nature*, 542(7640), 223–227. https://doi.org/10.1038/nature21071
- Kigel, J., Konsens, I., Segev, U., & Sternberg, M. (2021). Temporal stability of biomass in annual plant communities is driven by species diversity and asynchrony, but not dominance. *Journal of Vegetation Science*, 32(2), 1–12. https://doi.org/10.1111/jvs.13012
- Lázaro, A., Gómez-Martínez, C., González-Estévez, M. A., & Hidalgo, M. (2022). Portfolio effect and asynchrony as drivers of stability in plant–pollinator communities along a gradient of landscape heterogeneity. *Ecography*, 1–14. https://doi.org/10.1111/ecog.06112
- Lefcheck, J. S., Byrnes, J. E. K., Isbell, F., Gamfeldt, L., Griffin, J. N., Eisenhauer, N., Hensel, M. J. S., Hector, A., Cardinale, B. J., & Duffy, J. E. (2015). Biodiversity enhances ecosystem multifunctionality across trophic levels and habitats. *Nature Communications*, 6. https://doi.org/10.1038/ncomms7936
- Lepš, J., Májeková, M., Vítová, A., Doležal, J., & Bello, F. de. (2018). Stabilizing effects in temporal fluctuations: management, traits, and species richness in high-diversity communities. *Ecology*, 99(2), 360–371. https://doi.org/10.1002/ecy.2065
- Loreau, M., & De Mazancourt, C. (2008). Species synchrony and its drivers: Neutral and nonneutral community dynamics in fluctuating environments. *American Naturalist*, 172(2). https://doi.org/10. 1086/589746
- Lundgren, R., Lázaro, A., & Totland, Ø. (2013). Experimental pollinator decline affects plant reproduction. Journal of Pollination Ecology, 11(7), 46–56.
- Mazancourt, C. de, Isbell, F., Larocque, A., Berendse, F., De Luca, E., Grace, J. B., Haegeman, B., Wayne Polley, H., Roscher, C., Schmid, B., Tilman, D., Ruijven, J. van, Weigelt, A., Wilsey, B. J., & Loreau, M. (2013). Predicting ecosystem stability from community composition and biodiversity. *Ecology Letters*, 16(5), 617–625. https://doi.org/10.1111/ele.12088
- McCann, K. S. (2000). The diversity-stability debate. Nature, 405(May).
- Oliver, T. H., Heard, M. S., Isaac, N. J. B., Roy, D. B., Procter, D., Eigenbrod, F., Freckleton, R., Hector, A., Orme, C. D. L., Petchey, O. L., Proença, V., Raffaelli, D., Suttle, K. B., Mace, G. M., Martín-López, B., Woodcock, B. A., & Bullock, J. M. (2015). Biodiversity and Resilience of Ecosystem Functions. Trends in Ecology and Evolution, 30(11), 673–684. https://doi.org/10.1016/j.tree.2015.08.009
- Ollerton, J., Winfree, R., & Tarrant, S. (2011). How many flowering plants are pollinated by animals? Oikos, 120, 321-326. https://doi.org/10.1111/j.1600-0706.2010.18644.x
- Pimm, S. L. (1984). The complexity and stability of ecosystems. Nature, 315(6021), 635-636.
- Pisanty, G., Afik, O., Wajnberg, E., & Mandelik, Y. (2016). Watermelon pollinators exhibit complementarity in both visitation rate and single-visit pollination efficiency. *Journal of Applied Ecology*, 53(2), 360–370.

- https://doi.org/10.1111/1365-2664.12574
- Plas, F. van der. (2019). Biodiversity and ecosystem functioning in naturally assembled communities. Biological Reviews, 94(4), 1220–1245. https://doi.org/10.1111/brv.12499
- Rader, R., Bartomeus, I., Garibaldi, L. A., Garratt, M. P. D., Howlett, B. G., Winfree, R., Cunningham, S. A., Mayfield, M. M., Arthur, A. D., Andersson, G. K. S., Bommarco, R., Brittain, C., Carvalheiro, L. G., Chacoff, N. P., Entling, M. H., Foully, B., Freitas, B. M., Gemmill-Herren, B., Ghazoul, J., ... Woyciechowski, M. (2016). Non-bee insects are important contributors to global crop pollination. Proceedings of the National Academy of Sciences of the United States of America, 113(1), 146–151. https://doi.org/10.1073/pnas.1517092112
- Ruijven, J. van, & Berendse, F. (2007). Contrasting effects of diversity on the temporal stability of plant populations. *Oikos*, 116(8), 1323–1330. https://doi.org/10.1111/j.2007.0030-1299.16005.x
- Sasaki, T., & Lauenroth, W. K. (2011). Dominant species, rather than diversity, regulates temporal stability of plant communities. *Oecologia*, 166(3), 761–768. https://doi.org/10.1007/s00442-011-1916-1
- Sasaki, T., Lu, X., Hirota, M., & Bai, Y. (2019). Species asynchrony and response diversity determine multifunctional stability of natural grasslands. *Journal of Ecology*, 107(4), 1862–1875. https://doi.org/10.1111/1365-2745.13151
- Senapathi, D., Fründ, J., Albrecht, M., Garratt, M. P. D., Kleijn, D., Pickles, B. J., Potts, S. G., An, J., Andersson, G. K. S., Bänsch, S., Basu, P., Benjamin, F., Bezerra, A. D. M., Bhattacharya, R., Biesmeijer, J. C., Blaauw, B., Blitzer, E. J., Brittain, C. A., Carvalheiro, L. G., ... Klein, A. M. (2021). Wild insect diversity increases inter-annual stability in global crop pollinator communities. *Proceedings of the Royal Society B: Biological Sciences*, 288(1947). https://doi.org/10.1098/rspb.2021.0212
- Tilman, D., Isbell, F., & Cowles, J. M. (2014). Biodiversity and ecosystem functioning. *Annual Review of Ecology, Evolution, and Systematics*, 45, 471–493. https://doi.org/10.1146/annurev-ecolsys-120213-091917
- Tilman, D., Reich, P. B., & Knops, J. M. H. (2006). Biodiversity and ecosystem stability in a decade-long grassland experiment. *Nature*, 441(7093), 629–632. https://doi.org/10.1038/nature04742
- Valencia, E., Bello, F. de, Galland, T., Adler, P. B., Lepš, J., E-Vojtkó, A., Klink, R. van, Carmona, C. P., Danihelka, J., Dengler, J., Eldridge, D. J., Estiarte, M., García-González, R., Garnier, E., Gómez-García, D., Harrison, S. P., Herben, T., Ibáñez, R., Jentsch, A., ... Götzenberger, L. (2020). Synchrony matters more than species richness in plant community stability at a global scale. Proceedings of the National Academy of Sciences of the United States of America, 117(39), 24345–24351. https://doi.org/10.1073/pnas.1920405117
- Williams, N. M., & Winfree, R. (2013). Local habitat characteristics but not landscape urbanization drive pollinator visitation and native plant pollination in forest remnants. *Biological Conservation*, 160, 10–18. https://doi.org/10.1016/j.biocon.2012.12.035
- Winfree, R., & Kremen, C. (2009). Are ecosystem services stabilized by differences among species? A test using crop pollination. *Proceedings. Biological Sciences / The Royal Society*, 276(1655), 229–237. https://doi.org/10.1098/rspb.2008.0709
- Xu, Q., Yang, X., Yan, Y., Wang, S., Loreau, M., & Jiang, L. (2021). Consistently positive effect of species diversity on ecosystem, but not population, temporal stability. *Ecology Letters*, 24(10), 2256–2266. https://doi.org/10.1111/ele.13777
- Xu, Z., Ren, H., Li, M. H., Van Ruijven, J., Han, X., Wan, S., Li, H., Yu, Q., Jiang, Y., & Jiang, L. (2015). Environmental changes drive the temporal stability of semi-arid natural grasslands through altering species asynchrony. *Journal of Ecology*, 103(5), 1308–1316. https://doi.org/10.1111/1365-2745.12441