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More losses than gains during one century of plant biodiversity change in Germany

This is the final peer-reviewed author's accepted manuscript (postprint) of the following publication:

Published Version:

Ute Jandt, Helge Bruelheide, Florian Jansen, Aletta Bonn, Volker Grescho, Reinhard A. Klenke, et al. (2022). More losses than gains during one century of plant biodiversity change in Germany. *NATURE*, n.a., 1-11 [10.1038/s41586-022-05320-w].

Availability:

This version is available at: <https://hdl.handle.net/11585/898615> since: 2024-05-22

Published:

DOI: <http://doi.org/10.1038/s41586-022-05320-w>

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(Article begins on next page)



Helge Bruelheide, Inst. f. Geobotanik, Am Kirchtor 1, 06108 Halle

to the
handling Editor of
Nature
Dr. Armstrong

Unser Zeichen

Datum
Halle, 21.07.2022

Nature manuscript 2021-08-12657C "Plant diversity change over one century in Germany: more losers than winners"

Dear Dr. Armstrong,

thank you again for allowing us to revise the manuscript another time. We are very grateful that we could improve many further aspects of the paper. The input of the reviewers was a tremendous help, which we greatly acknowledge.

Please find the final version of the manuscript attached. Our point-by-point reply has been uploaded as a separate document.

Compared to the previous version, the main improvements are:

- We have shortened the text, rephrased the abstract and introduced hypotheses for the null models.
- We have changed the layout of Fig. 2 and 3.

We have uploaded both a clean version and version in tracking mode.

Yours sincerely, also on behalf of my co-authors

Prof. Dr. Helge Bruelheide

Senior Editor Nature, Anna Armstrong

Your manuscript entitled "Plant diversity change over one century in Germany: more losers than winners" has now been seen by referees 2 and 3. The referees remain largely happy. But in order to publish the paper we will need you to attend to all of the referees' continued concerns about presentation. No peer reviewed data should be removed altogether when making these changes.

*We are sorry that we did not seem to have met all the expectations of the reviewers.
However, we are happy that we have been given the chance to further improve the
manuscript. We believe that we now have addressed all the issues raised.*

*We have also uploaded all four "Related Manuscript Files" and made a suggestion for a
cover photograph. If the suggestion is chosen, we are prepared to describe the four time
series shown on the cover and provide the original photographs with high resolution.*

Referees' comments:

Referee #2 (Remarks to the Author):

The manuscript has improved substantially, I still like it a lot, and I only have minor comments related to writing and visual presentation of the figures. Specifically:

Visual part needs to be improved. Figures 2 and 3 still don't work, they are cluttered, confusing, with many unclear elements. What are the curly brackets for?

The curly brackets have been removed and the design of both figures has changed completely. In Fig. 2, the aggregation by species is now stated deliberately and illustrated for three plot records, with two species each. We no longer use arrows, but + and – signs to indicate decreases or increases.

Do sizes of the plants bear any meaning?

The idea was that plants with a positive and negative change in cover become larger and smaller, respectively, which obviously did not work. We now only show different icon sizes in the very left part of Fig. 2 to show that decreases and increases in cover translate into – and +. Without changing icon sizes in Fig. 3, this figure has become much simpler.

Can you shrink the plot to a printed page of Nature and will the legends be legible?

Yes, we hope so, in particular as we removed the icons on the left-hand side. However, this will finally only be seen in the proofs.

Moreover, there are at least two different fonts used in the plots, many different sizes of fonts, several levels of thickness of lines, legends are positioned at different places in different panels (compare a with b)...

Fonts now are all Arial. We have also used the same font size and refrained from using bold fonts (except for the scenarios).

..., there is a lot of redundancy with red and blue colors explained at least three times in the a-f panels, ...

We now only use colors for the different assignment of cover changes at the observation or species level. Thus, we do also not explain it repeatedly.

... and the arrows are all over the place, shifted to various positions (e.g. in Fig. 2).

The arrows have been removed.

Further, some panels are labeled by a)b)c), while others (with the plant symbols) aren't.

In Fig. 2, the labels (a and b) now also comprise the icons. However, this was not possible in Fig. 3, as here the icons refer to the whole columns.

In Fig. 2b there are “losers” and “winners” labels next to a mix of red and blue arrows, which is confusing.

We redesigned Fig. 2.

I have a hard time believing that the graphics has been edited by a graphic designer (as the authors indicate in their reply). I worry that these are things that will be hard to fix at the typesetting stage, as they relate to fundamental design/content problems. Navigating and understanding particularly Figure 3 will be a challenge, and if you combine this with very long figure captions, it’s even more challenging. The authors need to try harder to improve, clarify, and simplify the figures.

We apologize, but before we had tried really hard to illustrate Fig. 3. Now, we made a new attempt in making them as simple as possible. Here, we had a great input from iDiv’s Media & Communication department (Volker Hahn and Stefan Bernhardt).

The text is still too long at places. Some of the paragraphs in the Results, Discussion, and figure captions can be shortened, condensed, or removed. I believe that it should be possible. Particularly paragraphs on lines 214-238 and 294-327, and captions to figures 2 and 3 (some of the text there seems to repeat the results section).

We shortened both paragraphs. However, in the paragraph (previously l 214-238) we had to add the descriptions on the analyses on the effect of length of observation interval and plot sizes as request of Reviewer #3 in the last review round. To shorten the text, we moved sections of the paragraph (l 294-327) to the Supplementary Methods. This part could also be shortened because we turned the scenarios into hypotheses.

There is some confusion about the terminology referring to scale. In the intro, I’ve found “local” (lines 151, 157, 162, 180), “plot” (lines 174, 182, 186), “community” (151, 157, 159, 180, 189), “metacommunity” (164, 188), “regional scales” (168), “larger region” (163), “regional studies” (171), “sampled area” (174), “sampling area” (150), “sampling plots” (160, 162), “global scale” (144), “vegetation plot records” (174), “plot-scale” (186), “plot records” (202), and “resurveys & surveys” (172). This also happens in the rest of the paper, e.g. the discussion. There is “species level” (line 373), “species scale” (line 405), “sites” (line 369), “locations” (410), “plots” (407, 413), “coarse spatial scales” (417) and “national scale” (401) etc. In all this, it is unclear which aspects of scale the authors have in mind: Local grain? Local extent? Local support? Region grain or extent?

Metacommunity grain or extent? What is the difference between plot, site, location, survey, sampled area, and community? What is a metacommunity and how it differs from community at a regional scale? What is species level/scale? I usually roughly get what the authors have in mind, but not always. At this final stage of polishing, I suggest simplifying and clarifying this terminology as much as possible.

Thank you for pointing out our inconsistency in using terms. We have to admit that part of this happened with the many rounds of internal edits among the many co-authors. We have now harmonized the above-mentioned terms:

- “Local” is now more sparingly used. When referring to “local” and “regional”, we refer to grain, not extent, which we now make clear at the beginning of the Intro. However, we see no way to omit one of these two terms completely.
- The “community” is recorded locally. Nevertheless, we often point out that we here refer to species composition data recorded at fine grain, as this term has been also used across different grain sizes. Community composition is often observed in plots, but not exclusively

(there are many marine or freshwater communities that have been recorded plot-less). Thus, when it comes to vegetation time series, we have to mention plots.

- “vegetation-plot records” is a record made on a vegetation plot. Thus, we also use “plot record” as an abbreviation. The distinction between a “plot record” and a “plot” becomes necessary when the same plot is recorded more than once. This is common terminology in vegetation science.
- Similarly, the terms “survey” and resurvey” have concrete meanings. “Survey” is the act to record the species in a plot, “resurvey” is the act to repeat this in time.

In contrast, we removed some terms from the manuscript:

- Metacommunity: has been replaced by “regional”
- “larger region” has been replaced with “at regional extent”
- “regional studies” has been replaced by “large-grain studies”
- “sampling area” has been replaced by “sampled area”, which refers to the area of a plot that is sampled
- “global scale” has been completely removed
- “plot-scale” has been completely removed
- “species level” and “species scale” have been removed, except in the Methods when “species level” refers to taxonomic harmonization.
- “sites” has been completely removed and was partly replaced by locations.
- “coarse spatial scale” has been replaced by “coarse spatial grain”
- “national” has been replaced by “regional”

Line 165: Increase and decrease of what?

Added now “of species’ mean cover”

Lines 176-179: All this seems tangential and unnecessary.

Deleted.

Lines 209-2011: Dtto. Tangential and unnecessary.

Deleted.

Line 224: Log ratio decreased (which seems to be the current message, and which would mean a second derivative of the trend), or SR decreased as indicated by the negative log ratio?

Actually, we refer to a decrease in log ratio between species richness observed at the end and the beginning of an observation interval. Extended Data Fig. 2a shows that $\log_e(SR_{Y2}/SR_{Y1})$ was positive for time spans ≤ 2 years (that is, SR increased), while it was negative for time spans > 10 years (that is, SR decreased). Per decade, $\log_e(SR_{Y2}/SR_{Y1})$ decreased by 0.153. We now added: “indicating that more species were lost with time” (new line number 225).

Line 235: “only a marginal and” is unnecessary. If it was insignificant, it wasn’t even marginal.

Deleted.

Lines 236-237: I don’t see how no change in median indicates no change in dominance structure. You can have identical median in both time periods, but a pronounced change in the dominance, for example if increase in the dominant (high cover?) species is offset by decrease in species with low cover. In any case, what exactly do you mean by “dominance structure”?

We agree and have deleted this sentence, as we explain the discrepancy between mean and median change in cover in the following paragraph.

Line 180: Typo.

Corrected.

Lines 376-378: I don't see the direct link between homogenization and equality of change among losers and winners. Either prove/show the link mathematically, or drop the statement.

Probably, this disagreement is based on how we define homogeneity. In our opinion, homogenization is the direct consequence of redistribution of the species' cover. If decreases are distributed more equally (that is also more homogeneously) across many species and increases in cover are concentrated in few species, the latter (that is the winners) will be increasing in many communities. In consequence, the dissimilarity in species composition between these communities has to decrease (given that a quantitative dissimilarity measure is used). Mathematically, this would have to be shown by a decrease in dissimilarity, which however, is difficult to demonstrate across all plot records in our data set as many communities have no species in common. In our opinion, homogenization probably occurs within habitat types, but opening this discussion and carrying out the analysis would open a can of worms. Thus, we have decided to down-tune this statement to: "Homogenisation occurs because, across all time series, few species consistently increase in their cover, meaning that the same species are winning in many communities." (new l 372-374)

Referee #3 (Remarks to the Author):

As before, I commend the authors on their analyses and believe this paper makes a novel and important contribution by documenting long-term plant biodiversity changes in terms of cover that would have been missed by simply focusing on species richness, as most previous work has done. But while the authors have produced a strong revision of their paper, in my opinion several outstanding issues remain, and some important comments have only been partially addressed. I appreciate the new analyses that the authors have performed, and overall the analyses are appropriate and justified, and data are presented correctly, as far as I can judge. The framing of the study is also now more compelling, and I think the unique value of the dataset and insights arising from it are now harnessed more effectively. That said, I still have some concerns about the framing of the study, explained in my comments below. The Results section has also been improved, but remains difficult to read in places – again, I make specific suggestions below. Overall, while the length appears to have been reduced, the text still seems unnecessarily wordy in places (e.g. in the Results and figure captions).

Thank you for this positive assessment.

Another aspect I raised previously and has still not been resolved, in my opinion, is when results are considered to be ecologically relevant or not, and I think there could be greater transparency in the paper about this. This relates to the way the story is set-up, as mentioned in my previous review. Specifically, the authors note that due to the large sample sizes, changes in species richness can be statistically significant even if effect sizes are small, and therefore conclude that directional changes in species richness are “minor at best” (l.231). Yet on the other hand, even though the differences between winners and losers in their Gini coefficients are so small that they are imperceptible in Fig 2a, the authors note that these differences are “highly significant” (l. 245-6) and go on to interpret them. What is the reason for emphasising the ecological relevance of the latter result, but not of the species richness result? I find this troubling, since the set-up of the paper describes the paradox of changing global/regional but not local scale richness, and attempts to resolve that. Yet the paper does in fact describe significant local scale changes in richness. I

think this should be acknowledged more transparently in the paper, and not brushed off; or it should be made clear why the small differences in Gini coefficients are considered important, but not the changes in species richness, even when both are statistically significant.

We fully agree here and have now down-tuned the emphasis on significances in the Gini coefficient, too. We now write (new l 246): “While the two Gini values were highly significantly different (non-overlapping CIs even at 99.9%), their small difference might not result in ecologically meaningful effects. Nevertheless, the finding that losses in cover were more equally distributed than cover gains might point to an important ecological mechanism. If cover losses tend to occur in more uniform steps while gains result from both small and large increments, many small cover losses in a plot might be offset by a few large cover gains. The significantly different Gini coefficients show that this was at least the case in a considerable number of our change observations.”

We also agree that we too much downplayed our observation of decreasing richness. We now write at the beginning of the discussion (new l 344): “While the observed decline in species richness might be linked to the greater number of species that lost than gained cover, our analyses demonstrate that richness change at the plot level is not a necessary prerequisite for this asymmetry.” There are other places, where we also now refer to the finding of richness changes in our paper, too. However, we believe that the strength of our paper is to point to other mechanisms than richness change that can result in large-scale biodiversity change. This maybe explains why we might have too much downplayed species richness.

Specific comments

1.125 This isn't really a research question/problem statement, rather a description of a method. More importantly, I don't think this question (i.e., the discrepancy between declines of species richness at regional and global scales vs marginal changes in community species richness at local scales) is actually addressed – has there been loss of richness at a regional scale in Germany over the last century? The documentation of declines in abundance imply that regional losses will occur, but haven't necessarily done so yet (l.156). I.e., this is why the authors claim that asymmetry of losses/gains in cover could be “early warning signs” of regional species loss (l.189), which implies that regional losses haven't happened yet in Germany. For clarity, I suggest the authors think carefully about the overarching question addressed by the study and rephrase it accordingly.

Thank you for pointing out these issues. Following your advice, we have completely rewritten the Abstract. The research statement now is: “However, the majority of studies focused on species incidence and did not account for local abundance. Here, we asked whether the analysis of plant species cover changes can reveal yet unrecognized patterns of biodiversity change and provide insights into the underlying mechanisms.”

1.129 I would advise against using the word “losses” in this context, and more specifically refer to “reductions in cover” or “cover losses” throughout the paper, were relevant. “Loss” by itself implies (to me at least) that a species goes locally extinct at a location (is “lost”), which is not necessarily the case. Greater precision is important to avoid the results being misinterpreted.

This sentence has been rephrased, too. In addition, we have carefully checked all places in the manuscript, where we refer to “losses” and in most cases now write “cover losses”, except for those places where we draw conclusions on species loss as a consequence of these cover losses.

1.189 “while many species are lost from a few communities only” should be phrased more like “while rare species (those occurring in a few plots only) are lost completely” – i.e. the current formulation doesn’t necessarily imply regional species richness loss.

Thank you. We have changed the sentence accordingly.

1.203 Here a statement is missing of why these null models were developed.

We agree. As suggested we have also rephrased the null models as hypotheses (see below). We now write (new l 204): “To make sure that the observed patterns are not resulting by chance alone, we further developed null models that kept species richness constant and varied the amount and direction of change and the concentration of cover losses and gains on losers and winners. We hypothesized that the divergence in the distribution between cover losses and gains is driven by i) the proportion of species that undergo changes, ii) the ratio of increasing to decreasing species and iii) the degree to which cover losses are concentrated on a specific subset of species.”

1.205 Again, why did you do this? What are the hypotheses regarding nativity and habitat affinity? The reason for these analyses becomes clear while reading the paper but is a bit murky to the reader at this point.

We also agree with this point. We now write (new l 210): “Then, to assess whether losers and winners (i.e. those species that lost or gained cover) differed in their floristic status or habitat requirements, we analysed whether the probability of a decrease or an increase in cover depended on species being native or non-native and their habitat preference.”

1.221 “net loss of 6.04% of the species).” This should specify “per plot”, right? Although the sentence calls this “close to zero”, I would still consider a 6% loss to be concerning.

We apologize to have expressed this wrongly. A net decrease in $\log_e(SRy_2/SRy_1) = -0.062$ corresponds to an absolute decrease of 0.06 species per plot ($1 - \exp(-0.06233)$). This does not refer to the initial richness of that plot. Corrected now.

1.224-7 I don’t understand why species richness increases slightly when expressed per decade, “in consequence” of annual decreases (expressed on a log scale); perhaps I’m missing something, but please check phrasing here.

We apologize if this has not been clear enough before. As more species are lost, the longer the observation interval ($\log_e(SRy_2/SRy_1)$) decreased by 0.153 per \log_{10} years), the positive richness trend for time spans ≤ 2 years (Extended Data Fig. 2a) turns into a negative one for time spans > 10 years (Extended Data Fig. 2c). Dividing the log ratio by decade ($\log_e(SRy_2/SRy_1)$ decade $^{-1}$) reverses this trend, as, for example, then the observed positive richness trend of periods of 1 or 2 years is multiplied by 10 or 5, respectively. Please, see also our comment to reviewer #1 on the same lines.

1.228 The phrasing here, and elsewhere in the Results, is challenging to follow. E.g. “While $\log_e(SRy_2/SRy_1)$ decreased by 0.064 per \log_{10} increase in plot area” could be phrased as “While decreases in species richness were greater in larger plots”. It would greatly help readability if results were phrased in plain English wherever possible and backed-up with the statistics.

Done. We now write (new line 228): “While decreases in species richness were greater in larger plots (mean change in $\log_e(SRy_2/SRy_1)$ -0.064 per \log_{10} increase in plot area), species richness significantly decreased in all different categories of plot sizes (Extended Data Fig. 4a-4c).”

L.231 “changes in local richness are minor at best” is misleading – Fig S2 and others show there can be huge changes in local richness, but with no strong average trend towards increasing or decreasing richness. Please check phrasing here (e.g. add “directional”).

Done. We now write (new l 231): “Because of the overall very small effect sizes, we conclude that directional changes in mean local richness are minor at best, which is in accordance with preceding studies^{1,3,4}.”

1.253 there are three different ways in which this result (l. 250-254) is phrased, which all seem subtly different – it would be clearer if you could express this in one way which captures the message you wish to convey. As mentioned above, the word “losses” is misleading here, because what you mean is cover change, which might or might not entail the loss of a species from a plot.

We completely rewrote and shortened this section (new line 246): “While the two Gini values were highly significantly different (non-overlapping CIs even at 99.9%), their small difference might not result in ecologically meaningful effects. Nevertheless, the finding that losses in cover were more equally distributed than cover gains might point to an important ecological mechanism. If cover losses tend to occur in more uniform steps while gains result from both small and large increments, many small cover losses in a plot might be offset by a few large cover gains. The significantly different Gini coefficients show that this was at least the case in a considerable number of our change observations.”

L.267 Should “plants” be “plots” here? If not, then I believe it would be clearer to specify “changes in cover of individual plant species”

Replaced by “individual cover observations in plots”.

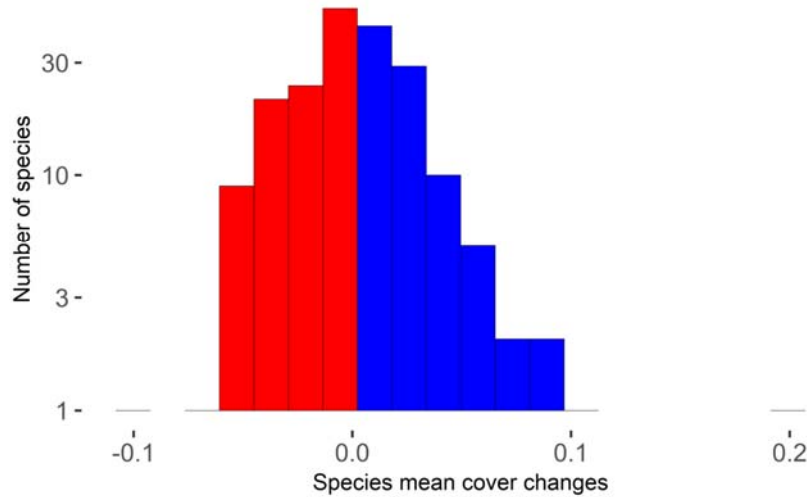
1.294 As mentioned in my previous review, this null model approach is a great addition to the paper. This paragraph remains, however, difficult to follow. I had to carefully consult Fig.3 and the supplementary methods to be able to follow it. I also mentioned previously that the rationale for the three scenarios wasn’t very clear, and this is still the case. One suggestion here would be to describe each scenario as a hypothesis to explain the empirical pattern, and then evaluate whether the null model lends support for that hypothesis or not. Currently the insights of the model are described as a series of results, and the reader is left largely to deduce the relevance of these.

We are grateful for the suggestion to turn the scenarios into hypotheses, which in our view made the approach much clearer. In addition to the graphical changes we made with Fig. 3 (see our reply to reviewer #1), we are confident that the null models now are much easier to grasp.

L.314 This result is very interesting, but can you say why non-random cover changes across species lead to inequality in the Gini coefficient in the null model – i.e. what is the mathematical/statistical phenomenon here? And what are the implications of this for the biological interpretation of divergence between losers and winners that then follows from l.328?

The mathematical phenomenon behind this outcome can be explained as follows: For hypothesis iii) we deliberately kept constant the proportion of species affected by cover change at 0.8 and the proportion of increasing species at 0.5. Thus, we could redistribute exactly the same randomly created negative cover changes as gains to the other species in the community. For these reason, the red and blue curves in Fig. 3c are absolutely identical. As the model first determined the losses and then redistributed the gains, for decreasing species this often meant extinction, for which then new species entered the community (as species richness was kept constant). Extinctions always resulted in deviation from the geometric distribution in cover declines, as the decline could never exceed the current cover of the species. Thus, declines were often smaller and more evenly spread than expected from a geometric distribution. Then, according to this scenario, the species to be selected for a

negative cover change were chosen non-randomly (i.e. in a directed way), thus, resulting in assigning these more evenly distributed declines more to the same species (please note that this was not deterministically, but simply weighted by using the species' IDs). In contrast, the gains were distributed randomly across all remaining species and new colonizers. This resulted in a larger spread, in particular of large gains, among species. This can be seen from the histogram of this scenario:



Thus, there were more winners with exceptionally high cover gains than losers with exceptionally high cover losses. This can also be seen in the extreme values and quartiles:

Min.	1st Qu.	Median	Mean	3rd Qu.	Max.
-0.0948316	-0.0050650	0.0000000	-0.0004992	0.0012849	0.1155434

We also made the focus on concentrating losses on particular species now also clearer by using the “+” and “-“ icons in Fig. 3.

Overall, we consider the way our model works in this particular aspect highly realistic. In a plot, the lower boundary of cover losses is always given by the extinction of a species in a plot. There is also a maximum that cannot be transgressed for winners, which is a relative cover of 1. However, as in nature, with the broken-stick distribution of initial cover values, there were only very few cases where species reached this upper boundary, and similar cut-off effects were observed as for declining species.

We also now added a clear interpretation of these results at new line 320: “We conclude that environmental changes that threaten specific species drive the inequality of mean cover changes of losers and winners.”

l.348 Does this result refer specifically to the null model, since you did in fact observe significant richness change? This needs to be specified here. See my major comment above about this.

As said above, we changed this section. We now write (new l 341): “Our work reconciles some issues in the debate surrounding the “biodiversity conservation paradox”¹², i.e. the discrepancy between observing a loss of species at broad scales but marginal changes in species richness at the plot scale. With support of a null model, we showed that the cover changes may affect winners and losers differently, even if plot richness does not change. While the observed decline in species richness might be linked to the greater number of species that lost than gained cover, our analyses demonstrate that richness change at the plot level is not a necessary prerequisite for this asymmetry.”

1.397-400 The wording confused me here – having greater data density in the middle of the time period could give rise to a mid-domain effect, not be a consequence of it; however, the observed pattern of greater divergence could be a consequence of the mid-domain effect, i.e. a statistical artefact. Please check the phrasing (and its implications) here.

Thank you for pointing this out. We rephrased this as follows: “Second, data density on species change observations was highest in this intermediate period, which could give rise to a mid-domain effect²⁹. In consequence, the stronger overlap of time series in the middle of the study period could have strengthened the observed trends.”

L.878 Units should be displayed in Fig1a x-axis

The unit is “number of species”. However, we do not find this helpful as the axis is expressed as $\log_e(SR_{Y2}/SR_{Y1})$. Thus, we are afraid to refrain from changing this. Any change would also affect all figures in the Extended Data, which are all carefully prepared and provide no more space on the x axes labels for additions. We hope for your understanding.

1 Cover page

2

3 Title: **Plant diversity change over one century in Germany: more losers than winners**

4

5 Article impact statement:

6 We provide a mechanistic understanding of biodiversity change using detailed time series data over
7 the past hundred years from 7,738 (semi-)permanent vegetation plots from Germany, comprising
8 1,794 vascular plant species.

9

10 Running head: **More losers than winners**

11

12

13 Keywords: Biodiversity change, Biodiversity monitoring, Gini coefficient, Temporal change, Vascular
14 plant species, Vegetation-plot resurveys

15

16 Word count: 4,424

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101 **Acknowledgments**

102 We are grateful to surveyors who recorded vegetation in the field and provided these data. We
103 acknowledge those data contributors who made their data available to us or helped in recording
104 these data: Thea Dittmann, Alexandra Erfmeier, Bernd Gerken, Kerstin Günther, Sabine Heinz,
105 Wilfried Hakes, Heike Heklau, Alfons Henrichfreise, Elisabeth Hüllbusch, Andreas Huwer, Anneke
106 Immoor, Sophie Luise Kühn, Benjamin Krause, Sebastian Leonhardt, Jennifer Reinecke, Ulrich
107 Scheidel, Immo Vollmer and Eva-Rosa Wagner. We thank Diana Bowler for her analysis of spatial
108 representativeness and Volker Hahn and Stefan Bernhard for their advice on Fig. 2 and 3. The
109 manuscript was very much improved by very detailed inputs of three anonymous reviewers and the
110 editor, Anna Armstrong. We very much appreciate the support for the strategic project sMon by the
111 German Centre for Integrative Biodiversity Research (iDiv) Halle-Jena-Leipzig, funded by the German
112 Research Foundation (DFG-FZT 118, 202548816).

113

114

115 **Author contributions**

116 U.J. and H.B. conceived the idea for the project. All authors were involved in collecting datasets,
117 developing the conceptual framework and interpreting the results. H.B. performed the statistical
118 analyses and developed the null model. U.J. and H.B. wrote the first draft of the manuscript. All
119 authors commented on and agreed with the final version of the manuscript.

120 **Competing interests**

121 The authors declare no competing interests.

122

123 Abstract

124 Recent long-term analyses of biodiversity data highlight a “biodiversity conservation
125 paradox”: biological communities show substantial species turnover over the last century^{1,2},
126 but changes in species richness are marginal^{1,3–5}. Most studies, however, focused on species
127 incidence only, neglecting changes in local abundance. Here, we asked whether the analysis
128 of plant species cover changes can reveal yet unrecognized patterns of biodiversity change
129 and provide insights into the underlying mechanisms. We compiled and analysed a dataset
130 of 7,738 (semi-)permanent vegetation plots from Germany that were surveyed between 2
131 and 54 times from 1927 to 2020 and in total comprised 1,794 vascular plant species. We
132 found that i) decrements in cover, averaged across all species and plots, occurred more
133 often than increments, ii) the number of species decreasing in cover was higher than the
134 number of those increasing, and iii) decrements were more equally distributed among losers
135 than gains among winners. Null model simulations confirmed that these trends do not
136 emerge by chance, but are the consequence of species-specific negative effects of
137 environmental changes. In the long run, these trends might result in significant species
138 losses at both local and regional scales. Summarising changes by decade shows that the
139 inequality in species mean cover changes of losers and winners diverged as early as in the
140 1960s. We conclude that species cover changes in communities convey an important but yet
141 neglected dimension of biodiversity change and should hence be more routinely considered
142 in time-series analyses.

143

144

145

146 Introduction

147 Biodiversity loss is one of the most critical environmental problems^{6,7}. Globally, the
148 extinction of many taxa has been well documented^{8–10}. However, local-scale studies, that is
149 those at the fine grain of communities, do not always reflect this global trend^{2,3}, which has
150 sparked intense debates^{11–13}. The main reason for this scale discrepancy is that species
151 losses and gains through time are inherently asymmetric. At any spatial scale, it only takes
152 one individual of a new species to result in a gain, but requires the loss of all individuals of a
153 species to lead to a loss¹⁴. In consequence, at a given sampled area, the loss of all individuals
154 of one species might be compensated by single individuals of a new colonizer^{3,15}. Indeed,
155 within local communities, species turnover, not species loss, has been identified as the main
156 aspect of biodiversity change¹⁶. For example, 28% of species were found to be replaced per
157 decade in an analysis of global marine and terrestrial community data². However, except for
158 recent studies from forests^{17,18}, these analyses ignored the changes in abundance that
159 precede species turnover.

160 Time series of local communities often document the abundance of each species, an
161 information rarely available at larger scales. Yet the strength of these data has been
162 insufficiently exploited in global biodiversity assessments. In the case of plant communities,
163 the most common abundance metric is the percentage of ground covered by all individuals

164 of a species on a particular sampling plot. This allows calculating changes as percentage
165 points of cover lost or gained, which helps nd detecting declines before local extinctions
166 occur. Aggregating such cover changes across many sampling plots at regional extent allows
167 calculating the rates of decrease or increase of species' mean cover, that is, identifying losers
168 and winners. This might in turn help to understand the discrepancies in species richness
169 trends found at different spatial scales.

170 For plant species, comparisons between losers and winners have only been conducted with
171 respect to occupancy at larger grain sizes. Studies based on ~5 x 5 km grids reported
172 contrasting trends in Denmark¹⁹ or in Germany²⁰, with increasing and decreasing species
173 richness, respectively. While both studies detected an imbalance between losers and
174 winners, it is difficult to ascertain biodiversity changes at regional large-grains, as resurveys
175 at that spatial resolution often differ in sampling intensity from the initial surveys. In 5 km-
176 grid cells, species are easily overlooked, resulting in pseudo-turnover with erroneous gains
177 or losses²¹. In contrast, small-grain vegetation-plot records, ranging from a few to several
178 hundred square metres²², are usually thoroughly checked not only for species presences but
179 also for absences.

180 In vegetation science, the traditional way of analysing time series of local communities is
181 following a plot's trajectory through time and aggregating the changing occurrence or cover
182 of species in the form of plot summary metrics, such as trends in species richness, diversity
183 indices, or more sophisticated measures, such as changes in species' mean ranks or
184 abundance curves²³. This type of analysis revealed both increasing (e.g. ⁴) and decreasing
185 (e.g. ²⁴) species richness trends, and global syntheses consequently did not detect general
186 trends in community-scale species richness (e.g.^{1,3}). However, constant community-scale
187 richness may be combined with biodiversity loss at the regional scale. This might for example
188 happen when few species colonise many communities anew, while rare species (i.e. those
189 occurring in a few plots only) are lost completely. An early warning-sign of such a
190 development would be an asymmetry of cover trends across species, i.e. the fact that
191 increases in cover are concentrated in a subset of species while decreases are distributed
192 more homogeneously across many species. Indeed, a recent global analysis of local species
193 turnover has suggested that it is particularly widespread species, often non-native ones, that
194 are increasing in abundance²⁵.

195 Here, we analyse cover changes of individual species in 7,738 vegetation-plot time series,
196 spanning almost a century and a wide range of habitat types across Germany. Apart from
197 comparing the magnitude of cover decrements and increments, we tested for the inequality
198 in the distributions of cover losses and gains across all species. Therefore, we adopted the
199 Gini coefficient, a metric developed in economics to evaluate the share of incomes across
200 the inhabitants of countries²⁶. Based on the Gini-coefficients we also, we tested whether
201 cover losses were more evenly spread among losing species than cover gains were
202 distributed among winning species. We here define losers and winners by their mean change
203 in cover across all observation intervals and all plot records, which can be either negative or
204 positive. To make sure that the observed patterns are not resulting by chance alone, we
205 further developed null models that kept species richness constant and varied the amount
206 and direction of change and the concentration of cover losses and gains on losers and

207 winners. We hypothesized that the divergence in the distribution between cover losses and
 208 gains is driven by i) the proportion of species that undergo changes, ii) the ratio of increasing
 209 to decreasing species and iii) the degree to which cover losses are concentrated on a specific
 210 subset of species. Then, to assess whether losers and winners (i.e. those species that lost or
 211 gained cover) differed in their floristic status or habitat requirements, we analysed whether
 212 the probability of a decrease or an increase in cover depended on species being native or
 213 non-native and their habitat preference. Finally, we assessed the temporal dynamics of
 214 cover losses and gains and asked whether they occurred at the same point in time.

215

216 Results

217 The 7,738 vegetation-plot time series covered the period from 1927 to 2020 (Extended Data
 218 Fig. 1). Plot richness change, calculated as the log ratio of species richness (SR) at the end
 219 and the beginning of the observation time interval (Fig. 1a), varied more than tenfold in
 220 absolute numbers. Even though we observed a significant decrease in species richness over
 221 time, the estimated effect sizes were close to zero (mean $\log_e(SR_{Y2}/SR_{Y1}) = -0.062$,
 222 corresponding to a mean net loss of 0.06 species per plot). There was a tendency of shorter
 223 observation intervals having significant increases and longer observation intervals significant
 224 decreases in species richness (Extended Data Fig. 2a-2c). On average, $\log_e(SR_{Y2}/SR_{Y1})$
 225 decreased by 0.153 per \log_{10} years ($p < 0.001$ according to a mixed model), indicating that
 226 more species were lost with time. In consequence, the change in species richness was also
 227 close to zero but significantly positive when richness change was expressed per decade
 228 (mean $\log_e(SR_{Y2}/SR_{Y1})$ decade $^{-1} = +0.062$, Extended Data Fig. 3). While decreases in species
 229 richness were greater in larger plots (mean change in $\log_e(SR_{Y2}/SR_{Y1})$ -0.064 per \log_{10}
 230 increase in plot area), species richness significantly decreased in all different categories of
 231 plot sizes (Extended Data Fig. 4a-4c). Because of the overall very small effect sizes, we
 232 conclude that directional changes in mean local richness are minor at best, which is in
 233 accordance with preceding studies^{1,3,4}. Similarly, the effect sizes for Shannon's index of
 234 diversity, Pielou's index of evenness and the change in the species rank abundance curve (as
 235 a measure of curve-change²³) were significantly negative, but of small magnitude (Extended
 236 Data Fig. 5a, b, c).

237 Across all plots, there were 458,311 change observations, i.e. species x plot records x time
 238 interval combinations. There were more negative ($n = 172,252$) than positive ($n = 166,554$)
 239 observations, and on average, decrements were larger than increments (4.05 and 3.97
 240 percentage points, respectively, according to a t-test ($p=0.003$, $df=338,187$). For each
 241 interval, species change was assessed as change in per cent cover, expressed as percentage
 242 points. Across all observations, the values of both negative and positive cover changes were
 243 not evenly distributed, which is illustrated by the Lorenz curves (Fig. 2a) and the
 244 corresponding Gini coefficients. Gini coefficients of 0.712 (95% confidence intervals (CIs)
 245 0.710 and 0.714) and 0.718 (CIs 0.717 and 0.721) were obtained for negative and positive
 246 change observations, respectively. While the two Gini values were highly significantly
 247 different (non-overlapping CIs even at 99.9%), their small difference might not result in
 248 ecologically meaningful effects. Nevertheless, the finding that losses in cover were more

equally distributed than cover gains might point to an important ecological mechanism. If cover losses tend to occur in more uniform steps while gains result from both small and large increments, many small cover losses in a plot might be offset by a few large cover gains. The significantly different Gini coefficients show that this was at least the case in a considerable number of our change observations. Moreover, cover changes also depended on interval length. Cover decreased significantly more in longer than in shorter observation intervals (by -0.042 percentage points per \log_{10} interval length, $p < 0.001$ according to a mixed model) as well as in larger than in smaller plots (by -0.14 percentage points per \log_{10} area, $p < 0.001$).

Across all intervals independent of interval length, there were more losers than winners per plot, with an average difference of 0.407 species (CIs 0.246 and 0.569, Extended Data Fig. 5d), which corresponds to the observed decrease in plot richness (Fig. 1a). Despite on average larger decrements than increments and fewer winners than losers in plots there was a significant increase of 2.5 percentage points in mean cover of all the species in a plot across all plot records (Extended Data Fig. 5e). In contrast, we observed a (insignificant) decrease of 0.7 percentage points in median cover (Extended Data Fig. 5f). These opposing directions of changes in mean and median cover are the direct consequence of a higher inequality of increments compared to decrements.

While these changes of individual cover observations in plots are so subtle that they may only be detectable in large datasets, they add up when species mean changes are calculated. Out of the total 1,794 vascular plant species in our study, there were 41% more losers than winners, with 1011 and 719 species, respectively. In consequence, the median across all species' mean cover changes was significantly negative (-0.063 percentage points, CIs -0.089 and -0.035, $p < 0.001$, Extended Data Fig. 6). Mean cover changes of species did not depend on their overall frequency in the dataset (regression of species' mean cover change on \log_{10} frequency, $p=0.601$). Decreases in species mean cover were also consistent with respect to observation interval length (Extended Data Fig. 2d-f) and plot size (Extended Data Fig. 4d-f). In all analyses, there were not only more losers than winners, the amounts of cover losses and gains were also not distributed equally within both groups. This is demonstrated by Lorenz curves, which in Fig. 2b are based on mean cover changes per species. The Gini coefficients for species with mean negative (0.692, CIs 0.660 and 0.718) and positive changes (0.778, CIs 0.720 and 0.816) differed by almost 0.1. The larger Gini coefficient for winners indicates that there were a few winners that gained disproportionately more mean cover than others, while the mean cover losses among losers were more equally distributed. Comparing Fig. 2a and 2b shows that two factors contributed to the inequality of biodiversity change. First, decreases occurred in smaller and more equal cover changes than gains. Second, the gains were concentrated in fewer winning species, while the cover losses were distributed among more losers.

These results also hold when rare species were excluded from the calculations. Fig. 1b shows the histogram of mean cover changes of the 578 species for which at least 100 time interval observations were available. Here, the change in mean cover was -0.165 percentage points ($p < 0.001$), showing that species mean losses in cover were significantly higher than species mean gains. In other words, there was a redistribution of species: fewer species increased in

292 dominance and frequency while more species decreased in cover and potentially
293 disappeared locally.

294 To understand the factors that determine the divergence in Gini coefficients between
295 decreases and increases in cover and to disentangle those from possible species richness
296 effects, we performed a series of null model simulations to test three different hypotheses.
297 We hypothesized that the divergence in the distribution between cover losses and gains is
298 driven by i) the proportion of species that undergo changes, ii) the ratio of increasing to
299 decreasing species and iii) the degree to which cover losses are concentrated on a specific
300 subset of species (Fig. 3, for further explanations and a graphical illustration see
301 Supplementary Methods). In all null models, species richness of each plot was kept constant
302 to avoid confounding effects of richness change, and only cover changes were redistributed
303 among losers and winners (for details see Methods). In contrast to hypothesis i), the
304 divergence in the distribution between cover losses and gains did not depend on the
305 proportion of species that undergo change. While subjecting more species to cover changes
306 increased the Gini coefficients for both negative and positive change observations (Fig. 3a),
307 this did not propagate to the species mean change values (Fig. 3d). We could confirm
308 hypothesis ii) which posited that the ratio of increasing to decreasing species drives the
309 divergence in the Gini coefficients of decrements and increments (Fig. 3b). Inequality is
310 higher for the kind of change that is more frequent. However, in the empirical data the
311 increments were more unequal (Fig. 2a) although they are less frequent. As a corollary, the
312 observed divergence of inequality is unlikely to be a mere consequence of the absolute
313 number of losses and gains. There was also support for hypothesis iii). Concentrating losses
314 on a specific subset of species did not affect the inequality of decrements and increments
315 across all species (Fig. 3c). However, it resulted in mean cover losses of losers being more
316 evenly distributed than the cover gains of winners (Fig. 3f), like in the empirical data (Fig.
317 2b). This pattern was not obtained by the other two model simulations. Neither was the
318 proportion of species affected by cover change (Fig. 3d), nor had the proportion of
319 increasing species (Fig. 3e) significant impacts on the divergence of the Gini coefficients
320 between losers and winners. We conclude that environmental changes that threaten specific
321 species drive the inequality of mean cover changes of losers and winners.

322 To determine the identity of losers and winners we focused on the 578 species with at least
323 100 time-interval observations, of which 161 showed significant differences in cover losses
324 and gains across all plots (binomial test at $p < 0.05$, with Holm correction, Fig. 4). Among
325 these 161 species with a directional change, native species decreased and neophytes
326 increased more often than expected by chance (at $p < 0.05$). Comparing the species' habitat
327 affinities revealed that significant decreases occurred among species of mires and spring
328 fens (level 1 EUNIS habitat Q), grasslands (R) and arable land (V), while forest species (T)
329 increased more often than expected by chance.

330

331 The times when cover losses and gains occurred were highly species-specific, as can be
332 inferred from the temporal course of the Gini coefficients for the 1011 losers and 719
333 winners (Fig. 5). Changes started to be more unequally distributed among winners than
334 among losers as early as in the 1960s. Since then, inequality of both gains and losses in cover

335 increased, with cover gains always being significantly more unevenly distributed among
 336 winners than losses among losers until 2010.

337

338

339 Discussion

340 Our work reconciles some issues in the debate surrounding the “biodiversity conservation
 341 paradox”¹², i.e. the discrepancy between observing a loss of species at broad scales but
 342 marginal changes in species richness at the plot scale. With support of a null model,
 343 we showed that the cover changes may affect winners and losers differently, even if plot
 344 richness does not change. While the observed decline in species richness might be linked to
 345 the greater number of species that lost than gained cover, our analyses demonstrate that
 346 richness change at the plot level is not a necessary prerequisite for this asymmetry. Overall,
 347 we found a higher number of losers than winners at the scale of Germany. This depends on
 348 two phenomena. First, cover losses were more evenly distributed than gains at the
 349 community scale. Second, cover losses and gains were concentrated in different species.

350 Finding 41% more losers than winners nationwide might even be considered a conservative
 351 estimate for Germany’s low to mid elevations, and certainly underestimates total plant
 352 biodiversity change. On the one hand, our study also includes plots in the alpine region,
 353 where positive richness changes have been described²⁷. On the other hand, our work suffers
 354 of most of the shortcomings noted in other studies on local time series^{13,28}, including the
 355 lack of spatial representativeness, varying lengths of observation intervals and a bias
 356 towards habitats that are least affected by human activities (see Extended Data Fig. 7 and 8).
 357 For instance, time series are usually discontinued in case of substantial land-use change,
 358 such as when a (semi-)natural habitat is converted into agricultural or urban land (one of the
 359 few exceptions was Hundt 2001²⁹, included in our analysis). In consequence, it is not
 360 surprising that the predicted 30% of local species extinctions due to land conversion³⁰
 361 remain mostly unnoticed in vegetation-plot time series like ours. We do not want to address
 362 all the criticism brought forward on calculating biodiversity change from local time
 363 series^{13,28}, which we think is mostly justified. Yet, we note that our time series covered about
 364 half the number of vascular plant species that occur in Germany, including rare habitats
 365 which often harbour rare plant species. This means that even if the spatial
 366 representativeness is incomplete for entire Germany, the representativeness at the level of
 367 individual species is unprecedented and grants robustness to our results.

368 Our results show that minor asymmetries of cover losses and gains in communities sum up
 369 when being aggregated by species, potentially hinting at population declines and extinctions
 370 at larger spatial extents. This is in agreement with trends observed across Germany^{20,31–33},
 371 including biotic homogenisation³⁴, which was already put forward in earlier studies on time
 372 series^{1,3}, but had not yet been properly tested¹³. Homogenisation occurs because, across all
 373 time series, few species consistently increase in their cover, meaning that the same species
 374 are winning in many communities. This supposedly results in a decreasing dissimilarity
 375 between communities. Other studies that analysed species changes conform to our finding
 376 of a prevalence of losers over winners, such as from Denmark³⁵, the UK³⁶ or Germany²⁰.

377 Although neophytes were more frequently increasing than decreasing, confirming global
378 observations^{37,38}, most winners were native species, as has been reported already for
379 German forest communities^{39–41}. Similarly, the habitat affinities of declining species being
380 concentrated in mires, grasslands and arable land reflect both the trends revealed by
381 Germany's Red List of vascular plants⁴² and floristic mapping programmes²⁰.

382 Our time series also provide important temporal information on species losses and gains.
383 The strongest asymmetry between cover losses and gains occurred between the end of the
384 1960s and the beginning of the 21st century, indicating rapid species turnover, most likely as
385 a result of substantial land-use changes (e.g. ⁴³). All systematic monitoring programmes on
386 vegetation, however, started only after year 2000, and thus, cannot provide information on
387 the second half of the 20th century. Our findings confirm the early warnings from the first
388 Red Lists in Germany⁴⁴ as well as estimated richness changes from floristic mapping
389 programmes when intervals between 1997–2017 were compared to the ones between 1960
390 and 1987²⁰. However, these results have to be interpreted with great caution for several
391 reasons. First, it is probable that later time series were established at locations and habitats
392 in the focus of nature conservation efforts, thus receiving a more favourable management
393 than the average landscape. Second, data density on species change observations was
394 highest in this intermediate period, which could give rise to a mid-domain effect²⁸. In
395 consequence, the stronger overlap of time series in the middle of the study period could
396 have strengthened the observed trends. We note, however, that early inequalities in cover
397 losses and gains at the plot scale will ultimately result in species extinctions at the regional
398 scale, representing another aspect of extinction debt⁴⁵.

399 Overall, we consider the overarching analysis of numerous local vegetation-plot time series
400 an important piece of evidence in the ongoing attempts to assess biodiversity change and
401 understand the underlying mechanisms. We demonstrated that changes in species cover
402 within communities are a neglected source of information when assessing biodiversity
403 change at large spatial extents. We advocate therefore the compilation of further existing
404 community time series worldwide, especially from vegetation plots of which few have
405 already been mobilised in global databases, such as BioTime⁴⁶. Compared to temporal
406 database analyses⁴⁷ and meta-analyses⁴⁸, repeated observations on the same locations
407 represent the most sensitive strategy for analysing temporal vegetation changes⁴⁹. However,
408 a careful quality control is a key prerequisite for this type of analysis¹³. In particular,
409 aggregating changes across different communities by species rather than aggregating
410 changes per plot requires much more attention for harmonising different taxonomies to
411 prevent pseudo-turnover²¹. With appropriate care taken, plot time series of community data
412 across larger regions should form a crucial backbone in future biodiversity monitoring.
413 Unravelling the temporal taxonomic turnover at the community scale^{1,2} allows insights into
414 the mechanisms of species losses and gains that monitoring at coarser spatial grains, such as
415 floristic mapping at grid sizes of several kilometres, alone cannot provide.

416

417

418 **Methods**419 **Data compilation**

420 We compiled as many long-term repeated vegetation-plot records from Germany as we
421 could access, including data from published studies, as well as results from grey literature
422 and conservation assessments. The data includes 92 projects (Extended Data Table 1, for a
423 description of the data see ref⁵⁰).

424 The different steps of data preparation and analysis are summarised in Extended Data Fig. 9.
425 Within each project, the plot resurvey ID indicates which plot observations from different
426 times were made on the same plot or set of plots at the same site, allowing their comparison
427 between different points in time. Plot resurvey IDs either refer to a single plot that was
428 repeatedly visited, which was either permanently marked (using poles, magnets etc.) or was
429 semi-permanent, e.g. provided with exact coordinates or other ways of descriptions of the
430 exact locality. In some cases, when the exact locations were not precise, resurveys used
431 several plots to match one previous plot, resulting in a one-to-many relationship. In this
432 case, all plot records received the same plot resurvey ID and all plot records for the same
433 point in time were combined. There were also resurveys with sets of plots at a site that
434 could not be matched by single plots but only by another set of plots, resulting in a many-to-
435 many relationship. Such resurveys were done to compare a particular community at a
436 particular site at two points in time, each represented by a set of plots, which then all
437 received the same plot resurvey ID. Accordingly, all plot records for the same point in time
438 were combined by averaging the species cover values and then treated as a single
439 observation. Some of our studies included experimental treatments with different
440 management of habitats (e.g. abandonment or establishment of grazing, succession and
441 disturbance). To exclude treatments that are not representative for biodiversity change in
442 Germany, from these studies we included only the control plots⁵¹, plots that reflected the
443 ambient land use at the site⁵², that were unfenced⁵³ or were subjected to continuous
444 grazing⁵⁴. Finally, 7,738 unique plot resurvey IDs remained, comprising a total of 23,641
445 vegetation-plot records, ranging from 1927 to 2020. We retrieved coordinates for all
446 locations (longitude and latitude), either from the original sources or by looking up plot
447 locations from maps. The duration and survey times of each project are shown in Extended
448 Data Fig. 1. Since different projects used different cover scales, we converted cover into per
449 cent, following the default conversion of the Turboveg 2 program⁵⁵. For example, for the
450 seven-grade Braun-Blanquet scale the transformation was r + 1 2 3 4 5 → 1% 2% 3% 13%
451 38% 63% 88%, respectively.

452 The location of all plots of all projects is displayed in Extended Data Fig. 7. We assigned the
453 individual plot locations to the grid cells of the quadrants of German ordnance maps
454 ("MTBQ," 0°10' × 0°6', approximately 5.6 km × 5.9 km in the centre of Germany), and tested
455 whether the grid cells analysed differed from those without observations with respect to
456 population density, road density, urban cover, cropland cover and protected areas. This
457 clearly revealed that the sampled grid cells were not representative for the whole area of
458 Germany. Surprisingly, they showed significantly higher human population densities, road
459 densities and urban cover, while cover of cropland and the amount of protected area was
460 lower, which indicates that many time series were made in regions with higher human
461 pressures. Our time series were also biased with respect to habitat types. This was
462 illustrated by assigning all plot records of the time series to EUNIS classes, using the expert
463 system EUNIS-ESy⁵⁶ and the corresponding R code⁵⁷. Each time series was assigned to the

464 habitat type by using the earliest plot record that resulted in level 3 EUNIS classification
 465 (Extended Data Fig. 8). While the time series covered 92 of the approximately 150 EUNIS
 466 habitat types encountered in Germany, most of the 23,641 plot records came from
 467 grasslands (level 1 EUNIS habitat R, n=14,849, 62.8%), followed by forests and other wooded
 468 lands (T, n= 5,440, 23%). In contrast, arable land, which makes up more than 36% of the land
 469 cover in Germany, was only represented by 816 plot records (V, vegetated man-made
 470 habitats, 3.5%).

471

472 **Taxonomic harmonisation**

473 All projects were linked to the standardized species list German SL 1.3⁵⁸. The nomenclature
 474 for vascular plants followed the concepts of Wisskirchen et al.⁵⁹, with additional
 475 aggregations to higher taxonomic levels according to German SL 1.3⁵⁸. As some authors
 476 recorded subspecies and other infraspecific taxa, species were aggregated at the species
 477 level, using vegdata⁶⁰. Some closely related species that, from our experience, were often
 478 mistaken in the field were merged at the aggregate or genus level. Species aggregates were
 479 also used when different taxon names of the same aggregate occurred in different projects,
 480 to prevent that the same taxon might appear under different taxon names. The taxon name
 481 harmonisation was a crucial step in our approach, as our aim was to assess species cover
 482 changes across projects (Fig. 1). We used our own R code to merge taxon names and the
 483 notation of the ESy expert system⁵⁶ to protocol all steps. The species harmonisation forms
 484 the first section of the ESy system and shows which taxon names were aggregated under the
 485 name of a broader taxonomic concept (Extended Data Table 2). In addition, within single
 486 projects, we used customised aggregations when the same taxa were reported at different
 487 taxonomic levels at different points in time in the same plot resurvey IDs (Extended Data
 488 Table 3). For example, while in all but one year of a time series of a specific plot *Orchis*
 489 *militaris* was reported, only one year reported *Orchis* spec. at the genus level. Unaccounted
 490 for, such a leap between taxonomic levels within a time series would result in incorrect
 491 species change observations. To avoid losing the predominating information at the species
 492 level by aggregating all records to *Orchis*, we assumed that the taxon was also *Orchis*
 493 *militaris* in that particular year. If more than one taxon occurred in previous years, we
 494 equally distributed the cover among those taxa. This happened for example when a record
 495 was taken late in spring when the two species *Anemone nemorosa* and *A. ranunculoides*
 496 could no longer be distinguished.

497 The percentage cover values of the same aggregated taxon name as well as those of taxa
 498 occurring in different layers of the same plot were merged, assuming a random overlap of
 499 their cover values and making sure that the combined cover values cannot exceed 100%⁵⁶.
 500 We removed bryophytes and lichens using the vegdata package in R⁶⁰.
 501 Finally, the original list of 3,280 taxon names that included bryophytes and lichens was
 502 reduced to 1,794 taxon names of vascular plants. In the following, for the sake of simplicity,
 503 we refer to these taxon names as species.

504

505 **Temporal change analysis**

506 Instead of fitting trends for individual time series, different intervals of the same time series
 507 were treated as separate change observations. This was achieved by separating all records
 508 into 458,311 plot resurvey triplets, i.e. ID x species x time interval observations, where the
 509 interval designated two subsequent observations between year 1 and year 2 for the start

510 and end of the interval, respectively. Separating a time series in its different intervals avoids
 511 the problem of establishing a baseline against which the changes are being compared^{13,28}.

512 Temporal change analysis at the plot level

513 At the plot level, the triplets were aggregated into plot resurvey ID x time interval
 514 combinations (in total n=13,987). With a total of 7,738 plot resurvey IDs, this corresponds to
 515 on average 1.81 resurvey intervals per plot resurvey ID. This means that, on average, a time
 516 series had about three observation events. While most plot resurvey IDs were only repeated
 517 once (one interval, n=6,006), 798 had two intervals, 213 had three intervals and 721 had
 518 four or more intervals. The longest time series comprised 54 intervals (Uwe Wegener,
 519 montane Harz meadows⁶¹). For each interval and plot resurvey ID, we calculated the change
 520 in species richness (SR), Shannon's index of diversity and Pielou's index of evenness. In
 521 addition, we calculated the change in the rank abundance curves, using the formula for
 522 curve-change in ref²³. The change in rank abundance reflects the area between the two rank
 523 abundance curves for the later and earlier observation. Rank abundance curves are
 524 constructed by plotting the species' cumulative relative cover (ranging from 0 to 1) against
 525 the species' ranks in cover values, calculating ranks from highest to lowest cover and then
 526 dividing the ranks by the maximum rank (with scaled ranks ranging from 0 to 1).
 527 Furthermore, we calculated the number of species with decreases and increases in cover as
 528 well as mean and median cover across all species in a plot record.

529 For all change metrics calculated at the plot level, we calculated log response ratios of the
 530 metric at time Y2 divided by that at time Y1, except for the change in rank abundance curves
 531 and losses and gains, where we used the difference between area and number of species,
 532 respectively. To assess the impact of plot size on the change of species richness, we tested
 533 the effect of $\log_{10}(\text{surface area in m}^2)$ on $\log_e(\text{SR}_{Y2}/\text{SR}_{Y1})$. Additionally, we analysed the
 534 distribution of plot records with respect to $\log_e(\text{SR}_{Y2}/\text{SR}_{Y1})$ separately for small ($> 25 \text{ m}^2$),
 535 medium-size (25 m^2) and large plots ($> 25 \text{ m}^2$). A similar analysis was carried out for testing
 536 the effect of the observation length (\log_{10} interval length in years) on the change of species
 537 richness and analysing the distribution of plot records separately for short (≤ 2 years),
 538 medium (> 2 and ≤ 10 years) and long observation intervals (> 10 years). We additionally
 539 expressed richness change per decade (mean $\log_e(\text{SR}_{Y2}/\text{SR}_{Y1})$ decade⁻¹). The departure of
 540 effect sizes and differences from 0 in all these analyses were assessed with mixed effects
 541 models, using the time-series ID as random factor, thus taking into account the non-
 542 independence of intervals from the same time series. As there were 13,987 plot resurvey ID
 543 x time interval combinations, the test statistics tended to be significant, even when the
 544 mean of the test metric was close to zero. We used mixed models to calculate confidence
 545 intervals using Wald-test approximation⁶².

546

547 Temporal change analysis by species

548 In total there were 458,311 plot resurvey ID x species x time interval combinations, for which
 549 the difference in cover for every species k and time interval m was calculated as $\Delta\text{cover}_{k,m} =$
 550 $\text{cover}_{k,m,Y2} - \text{cover}_{k,m,Y1}$ and expressed as percentage points. Similar to our analyses for the
 551 change of species richness, we also tested the effect of $\log_{10}(\text{surface area in m}^2)$ and of

552 observation length (\log_{10} interval length in years) on $\Delta\text{cover}_{k,m}$, using mixed effects models
 553 with the time-series ID as random factor.

554 Here, Y_2 and Y_1 refer to the end and the start year of an interval, defined as the two nearest
 555 points in a time series.

556 To compare the distribution of cover changes across all species, we considered positive and
 557 negative cover change observations separately ($n= 184,678$ and $192,162$ time interval
 558 observations, respectively). We then sorted the cover changes in each category (positive or
 559 negative cover changes) according to increasing absolute values and plotted the cumulative
 560 sums of cover changes against the proportion of observations in each category, thus
 561 obtaining a Lorenz curve. We calculated the unweighted Gini coefficient for each category,
 562 according to⁶³ and using the bias correction implemented in the DescTools package⁶⁴:

$$563 G_{\text{cover}} = \frac{\sum_i^n \sum_j^n |\Delta\text{cover}_i - \Delta\text{cover}_j|}{2 \sum_i^n \sum_j^n \Delta\text{cover}_j} \frac{n}{n-1}$$

564 with Δcover_i and Δcover_j being cover changes of change observations i and j in plots,
 565 irrespective of species, and n the total number of change observations. G_{cover} is calculated
 566 separately for negative or positive change observations, using either only all negative or
 567 positive change observations. The Gini coefficient is a measurement of inequality in
 568 distribution²⁶, given as a value between 0 and 1 with 0 indicating a perfectly equal
 569 distribution.

570 Across all plot resurvey IDs, there were 458,311 species x time interval combinations with a
 571 value for cover change. For species comparisons, we aggregated cover changes by species
 572 across all plot resurvey IDs and intervals. We counted the number of positive, zero or
 573 negative cover changes per species and subjected them to an exact binomial test, using the
 574 stats package. We adjusted the significance levels for multiple testing using Holm correction.
 575 When showing changes by species in graphs (Fig. 4), we confined the list to those species
 576 with $p < 0.05$ after Holm correction and with ≥ 100 time interval observations ($n=161$). To
 577 compare the distribution of cover changes among all species, we calculated the mean cover
 578 change per species, expressed as percentage points in cover. As the cover changes were
 579 highly dependent on species and many species occurred only rarely in the time series, we
 580 tested the probability to increase with a non-parametric exact binomial test. We assigned
 581 the floristic status native, archaeophyte and neophyte (the latter two being exotic species
 582 arriving in Germany before or after 1492, respectively) to these 161 species, using the
 583 BIOLFLOR database⁶⁵. We assigned species to their preferred habitat, using the level 1
 584 habitats of the EUNIS habitat classification⁶⁶. This was achieved by assigning all 225,606
 585 vegetation plots in the German Reference Vegetation Database⁶⁷ to EUNIS classes, using the
 586 expert system EUNIS-ESy⁵⁶ and the corresponding R code⁵⁷. We then calculated the affinity
 587 of the 161 species with a significant change to each of the 150 EUNIS classes that occurred in
 588 Germany, using the Φ coefficient of association^{68,69}. Then, the habitat preference of a
 589 species was defined as the EUNIS class to which the species had the highest Φ coefficient.
 590 For further analysis, we used the highest hierarchy of the EUNIS system (level 1). To assess
 591 which categories of floristic status and EUNIS habitat level 1 preference departed from the
 592 expected probability to increase, which is 0.5, we scaled the probability response to -1 to 1

593 and calculated linear models without intercept. Additionally, we tested whether mean cover
 594 changes of species depended on their overall frequency in the dataset and analysed subsets
 595 of species based on different interval lengths and plot sizes in which the species occurred,
 596 using the same categories of interval lengths and plots sizes as used for analysing species
 597 richness.

598 We calculated the Gini coefficient for inequality of changes, separately for species with
 599 negative and positive mean cover changes (that is losers and winners), respectively. The Gini
 600 coefficient based on species means was also calculated using the DescTools package⁶⁴ and is
 601 defined as:

$$602 G_{\overline{cover}} = \frac{\sum_i^N \sum_j^N |\overline{\Delta cover}_i - \overline{\Delta cover}_j|}{2 \sum_i^N \sum_j^n \overline{\Delta cover}_j} \frac{N}{N-1}$$

603 with $\overline{\Delta cover}_i$ and $\overline{\Delta cover}_j$ being the mean cover changes of species i and j and N the total
 604 number of species. $G_{\overline{cover}}$ is calculated separately for losers or winners, using either only all
 605 negative or positive species mean cover changes. Applied in this way, the Gini coefficient
 606 $G_{\bar{x}}$ indicates that either the losses or the gains in cover were not distributed equally among
 607 species. To assess the significance in the difference between the Gini coefficients of losers
 608 and winners, we calculated 95% confidence intervals from bootstrapping, using percentiles,
 609 bias correction and 1,000 replicates. For analysing the temporal course of the distribution of
 610 cover changes in the groups of winners and losers, we calculated the Gini coefficient $G_{\bar{x}}$ as
 611 described above using a moving window of five years, using only records from 1945 onwards
 612 because of data scarcity before this date. Cover changes of all resurvey ID x species x time
 613 interval combinations were aggregated by species and year for all years that fell into a
 614 window of five years. In every window, 300 species were resampled by chance, repeated
 615 100 times, and Gini coefficients were calculated separately for all decrements and
 616 increments of the means of these 300 resampled species. Temporal trends with confidence
 617 intervals were calculated from the Gini coefficients from these 100 runs.

618 Null model scenarios

619 To assess the mechanisms that might drive the inequality of cover changes among losers vs.
 620 winners, we set up a simple model, serving as a theoretical null expectation (see illustration
 621 in Supplementary Methods). Corresponding to our data analysis, the null model was not
 622 spatially explicit. In contrast to previously developed null models, such as in¹, our aim was
 623 also not to model stochastic colonization or extinction but stochastic changes in cover, which
 624 to our knowledge had not been attempted before. Extinction only happened when cover
 625 decreased below zero, and then, was exactly counterbalanced by colonization. In this way
 626 we kept species richness constant, which contrasts null models such as in¹. Our null model
 627 also differs from traditional null models in community ecology which reshuffle cover values
 628 across communities and/or species^{70–72} but do not allow for random decrease and increase
 629 in cover.

630

631 Simulating random communities

632 We simulated random communities and subjected them to different scenarios. First, we
 633 created a pool of 200 species with frequencies randomly drawn from a log-normal
 634 distribution, using the rlnorm function in R (meanlog = 1.5, sdlog = 1.2). Summing up all
 635 frequencies resulted in a total of 1,810 occurrences. We then drew random species richness

values for 100 communities from a normal distribution, varying the mean and standard deviation to obtain the same total number of occurrences (1,810), which was achieved by using a Gaussian distribution with mean=19.13 and sd=9 species). We chose these parameters in a way to be similar to the richness distribution of our empirical data set (mean=23.4, sd=13.7). Finally, cover values were randomly assigned to the species in each community according to a broken-stick distribution⁷³, using the drbs function of the sads package⁷⁴, which resulted in a sum of a total cover of 100% in each community.

644 *Introducing cover change: three scenarios*

We then introduced different types of change to this random community, using three different scenarios. In all scenarios species richness was kept constant, which reflected our own findings and those in refs^{1,3–5}. However, we allowed species turnover by replacing species that – owing to randomly introduced decreases – had cover values <0. Newly colonising species were randomly selected from the pool of 200 species, with the drawing probability weighted by the species' frequency. In the scenarios 1 and 2, this made sure that the species frequency distribution in the species pool remained constant (except for random noise). Species decreases in cover were introduced by varying three parameters, which corresponded to the three scenarios in which these parameters were varied: 1) the proportion of species affected by cover change in a community (to simulate different rates of turnover in community composition), 2) the proportion of species with increase in cover among those species affected by change (to simulate differences in the distribution of cover losses and gains, irrespective of species) and 3) the identity of the species to decrease in cover (to simulate that cover losses and gains might be concentrated in certain species). Decrements were either assigned randomly or according to the descending ID of the species, which resulted in species with higher ID values being more frequently selected for losing cover than other species.

In each community, according to these parameters, species were randomly chosen that underwent a decrease. The cover of all decreasing species in each community was summed up and redistributed according to a geometric distribution. For example, in a community of 24 species where 50% of all species were selected to change in cover and 50% of those were subjected to decrease in cover, the summed cover of these six species was redistributed (but randomly assigned) to the same six species as 0.125, 0.0625, 0.03125, 0.015625, 0.0078125 and 0.0078125. Note that the smallest change occurred twice to result in a sum of 0.25. If the decrease in cover assigned to a species was larger than the current cover of that species, its cover became 0 and the species was replaced as described above. The actually applied decrements were then assigned to the species that – according to the given parameters – were selected for increase. The number of increasing species also comprised the newly colonising species in a community. If the number of decreasing and increasing species was the same, the exactly same cover changes of decreasing species were randomly assigned as increments to the increasing species, taking the decrements and changing their sign. In this case, the absolute values of all increments and decrements across all communities were exactly the same, and thus, could not result in differences in the equality of their distribution. If the number of decreasing species was higher than that of increasing species, each two randomly chosen decrements were combined until the number of required increments was reached. Vice versa, if the number of decreasing species was lower than that of increasing species, randomly selected decrements were divided by 2 until the number of required increments was reached. In the latter two cases, the equality of the distribution of decrements was no longer the same as that of the increments.

684 For all scenarios, we measured the inequality of increments and decrements by the Gini
685 coefficient as described above. As in the empirical data, we calculated the Gini coefficient i)
686 across all cover changes, separately for increments and decrements, but irrespective of
687 species, and ii) on species-aggregated mean values of increments and decrements.

688

689 The analyses were calculated in R version 4.0.3 using the packages stats, foreign, reshape2,
690 data.table, tidyverse, Hmisc, sads and DescTools. Graphs were produced with the packages
691 ggplot2, egg and vcd.

692

693 **Data availability**

694 All data is available as data paper⁵⁰ and available at <https://doi.org/10.25829/idiv.3508-c17blk> under the terms specified by CC BY 4.0.

696 A current version of the data paper submitted to Scientific Data has been uploaded. As the
697 Nature paper might be published before acceptance, we have also submitted the data paper
698 to bioRxiv (also uploaded).

700 [Please note that the DOI link is not yet activated, which will happen around July 22nd,
701 2022. In the meantime you can already access the metadata via
702 <https://idata.idiv.de/ddm/Data>ShowData/3508?version=0> and the full dataset here:
<https://cloud.uni-halle.de/s/wei1ljqnq2Wet0A>

703 This part marked in yellow will then be deleted from the paper]

704

705 **Code availability**

706 The R code to retrieve resurvey ID x species x time interval combinations and to calculate the
707 results of this paper is provided in Supplementary Code 1 and on <https://github.com/idiv->
708 biodiversity/ReSurveyGermany_Analysis, the R code to produce the null models in
709 Supplementary Code 2 and on <https://github.com/idiv->
710 biodiversity/ReSurveyGermany_null_models.

711

712

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878

879

880 **Captions:**

881

882 Fig. 1 | Temporal change in a) plant species richness of plots and b) mean cover change of
883 species. The black dashed line shows zero change, while the red solid line in a) shows the
884 mean change of richness and in b) the species' mean change in cover in percentage points.
885 a) Comparisons of species richness (SR) in plots recorded at subsequent points in time
886 ($n=13,987$). An effect size of ± 0.69 corresponds to double or half the initial number of
887 species, while an effect size of ± 2.3 indicates tenfold or one tenth of the initial number of
888 species. Estimated mean overall effect size was -0.062 according to a mixed effects model
889 ($p=2 \times 10^{-16}$, $df=5,310$) with a 95% confidence interval between -0.071 and -0.053 . b)
890 Comparisons of species' mean cover changes between subsequent records. Only species
891 with at least 100 observations of change ($n=578$) were included. The estimated overall mean
892 of the species' mean cover change was -0.165 ± 0.089 according to a t-test ($p=3.1 \times 10^{-4}$,
893 $df=577$).

894

895 Fig. 2 | Lorenz curves for cover decreases (red) and increases (blue), reported in percentage
896 points across the whole observation period from 1927 to 2020 a) across all change
897 observations irrespective of species ($n= 172,252$ and $166,554$ observations of decrease and
898 increase, respectively) and b) aggregated by species, by averaging all change observations
899 from a) across all plots and time interval ($n= 1011$ and 719 losers and winners, i.e. species
900 with negative and positive mean cover change, respectively). The icons on the left illustrate
901 these two types of aggregation of cover changes for six change observations, from which
902 each three decreased (-) or increased (+). In a) these decreases are sorted by sign (- or +), in
903 b) they are averaged by species, defining losers and winners, exemplified here as one species
904 each with negative or positive mean change, respectively. The Lorenz curves show the
905 cumulative amount of cover decrease and increase (added in order of their ascending
906 absolute values) as a function of the cumulative number of change observations (expressed
907 as proportion of the total number of observations). The diagonal black line indicates the
908 theoretical curve that would result if all observed changes would be equal in size. The Gini
909 coefficient, a measure of inequality, is the area between this diagonal line and the actual
910 Lorenz curve divided by the entire area under the diagonal line. Thus, 0 and 1 indicate
911 maximal equality and inequality, respectively. The differences between the Gini coefficients
912 in both graphs were significant at $p=0.05$ but the confidence intervals in a) are so small that
913 they are invisible in the graph.

914

915 Fig. 3 | Lorenz curves based on null model simulations of changes in species cover. In all
916 simulations, the number of species per community was kept constant, but species turnover
917 (extinctions and colonisations) was allowed (for details see Methods and Supplementary
918 Methods). The graphs in the top row (a, b, c) show cover changes by plot ID x species x time
919 interval (corresponding to Fig. 2a), separated into negative (red) and positive (blue) change
920 observations. The bottom row of graphs (d, e, f) show mean cover changes per species
921 (corresponding to Fig. 2b), separated into species with average cover increase and decrease

922 (losers in red and winners in blue. The columns correspond to ourcr three hypotheses. In
923 each panel, the blue and red Gini coefficients next to each other refer to the same scenario;
924 * indicates a statistically significant difference in the Gini coefficient between the two Gini
925 coefficients from the same scenario, n.s. not significant (all at p=0.05).

926

927 Fig. 4 | Probability of increase in cover for the 161 species with a significantly negative or
928 positive change (binomial test at p <0.05, with Holm correction) and at least 100 change
929 observations. Decreasing species are those with a probability to increase of <0.5, and thus,
930 increase less often than expected by chance, and their names are plotted to the bottom of
931 the y-axis at 0.5, while the names of increasing species are plotted to the top. The colours of
932 taxon names show their floristic status, with black, purple and pink for native, archaeophyte
933 and neophyte, respectively. The bar colour indicates the species' affinity to level 1 EUNIS
934 habitats⁶⁶ and the error bars indicate the 95% CIs. The three most declining and increasing
935 species are illustrated by name and photographs. Plant photographs were obtained from
936 www.floraweb.de.

937

938 Fig. 5 | Temporal course of the Gini coefficient, calculated by a moving window approach
939 with a window width of five years, separately for losers (species with mean cover losses, red)
940 and winners (species with mean cover gains, blue) in this time window. The coloured lines
941 show the mean values of 100 resampling events of 300 species each per time window, while
942 the confidence bands show the standard deviations across these samples. Non-overlapping
943 confidence bands indicate significantly different Gini coefficients between losers and
944 winners. The increasing Gini coefficients indicate increasingly unequal distribution of cover
945 changes with time.

946

947 **Supplementary Figures**

948 Extended Data Fig. 1 | Temporal coverage of the 92 projects included in the study. The coloured lines
 949 indicate the start and the end of a project, black diamonds show in which years surveys were made.
 950 Resurvey type refers to either studies that were repeated within a particular community across a site
 951 without attempts to match plots (community comparison), or were carried out on matched plots,
 952 which were either permanently marked or relocated from exact descriptions (semi-permanent). The
 953 lower graph shows the number of times a particular year was included in any of the time series.

954

955 Extended Data Fig. 2 | Impact of the observation interval length on the temporal change of species
 956 richness (SR) in plot records (a-c) and mean cover change of species (d-f). The responses are shown
 957 separately for short (≤ 2 years), medium (> 2 and ≤ 10 years) and long observation intervals (> 10
 958 years). The black dashed line shows zero change, while the red solid line in a) - c) shows the mean
 959 change of richness and in d) – f) the species' median change in cover in percentage points. According
 960 to a mixed effects model estimated mean overall effect size was in a) $+0.025$ ($p=3.9 \times 10^{-9}$, $df=4,142$),
 961 b) $+0.007$ ($p=0.093$, $df=3,903$) and c) -0.150 ($p<2 \times 10^{-16}$, $df=8,612$). In d) – f) plot Interval comparisons
 962 of the mean of all cover changes per species between time points Y1 and Y2 of the start and end
 963 year, respectively, are shown on an axis with a sign*square root-transformation. According to an
 964 exact binomial test estimated overall median of cover change was in d) 0 (95 per cent confidence
 965 interval 0 and 0.007), e) -0.02 (CI -0.02 and 0) and f) -0.26 (CI -0.53 and 0.002).

966

967 Extended Data Fig. 3 | Temporal change of plot records expressed per decade. Interval comparisons
 968 of species richness (SR) in plot records between time points Y1 and Y2 of the start and end year,
 969 respectively, and divided by the length of the interval in decades $((Y2-Y1)*10)$ ($n=13,987$). Estimated
 970 overall effect size was $+0.062$ according to a mixed effects model ($p=1.8 \times 10^{-7}$) with a 95%
 971 confidence interval between $+0.039$ and $+0.086$.

972

973 Extended Data Fig. 4 | Impact of plot surface area on the temporal change of species richness (SR) in
 974 plot records (a-c) and mean cover change of species (d-f). The responses are shown separately for
 975 small ($> 25 m^2$), medium-size ($25 m^2$) and large plots ($>25 m^2$). The black dashed line shows zero
 976 change, while the red solid line in a) - c) shows the mean change of richness and in d) – f) the species'
 977 median change in cover in percentage points. According to a mixed effects model estimated mean
 978 overall effect size was in a) -0.03 ($p=0.064$, $df=487$), b) -0.031 ($p=1.55 \times 10^{-13}$, $df=4,204$) and c) -0.095
 979 ($p<2 \times 10^{-16}$, $df=9,124$). In d) – f) plot Interval comparisons of the mean of all cover changes per
 980 species between time points Y1 and Y2 of the start and end year, respectively, are shown on an axis
 981 with a sign*square root-transformation. According to an exact binomial test estimated overall
 982 median of cover change was in d) -0.017 (95 per cent confidence interval -0.065 and -0.001), e) $-$
 983 0.019 (CI -0.043 and -0.006) and f) -0.26 (CI -0.134 and -0.050).

984

985 Extended Data Fig. 5 | Temporal change of plot records. The histograms show the interval
 986 comparisons of plot records between time points Y1 and Y2 of the start and end year, respectively.
 987 The black dashed line shows the zero change, while the red solid line shows the mean change as
 988 predicted from a mixed effects model. a) Change in Shannon's index of diversity (H). Estimated mean
 989 effect size for H -0.055 ($p=2.2 \times 10^{-16}$, $df=5,462$, 95% confidence interval -0.064 and -0.047). b)
 990 Change in Pielou's index of evenness (E). Estimated mean effect size for E -0.019 ($p=2.6 \times 10^{-16}$, 95%

991 confidence interval -0.024 and -0.015). c) Difference in the area under the rank abundance curves.
 992 Estimated mean difference -0.143 ($p=0.00211$, 95% confidence interval -0.194 and -0.091). d)
 993 Difference in the number of cover gains and losses. Estimated mean difference -0.407 ($p=7.9 \times 10^{-7}$,
 994 95% confidence interval -0.569 and -0.246). e) Change in mean cover of all the species in a plot (in
 995 per cent covered ground). Estimated mean effect size for mean cover $+0.025$ ($p=1.0 \times 10^{-10}$, 95%
 996 confidence interval $+0.018$ and $+0.033$). f) Change in median cover of all the species in a plot (per
 997 cent of covered ground). Estimated mean effect size for median cover -0.007 ($p=0.2984$, 95%
 998 confidence interval -0.021 and $+0.007$).

999

1000 Extended Data Fig. 6 | Temporal change in mean cover change of all species in percentage points.
 1001 Plot Interval comparisons of the mean of all cover changes per species between time points Y1 and
 1002 Y2 of the start and end year, respectively, shown on an axis with a sign*square root-transformation.
 1003 The black dashed line shows the zero change, while the red solid line shows the median change in
 1004 cover across all species. All species in the dataset were included ($n=1,794$). Estimated overall median
 1005 of cover change was -0.0625 (95 per cent confidence interval -0.089 and -0.035) and significantly
 1006 different from zero according to an exact binomial test ($p<0.001$).

1007

1008 Extended Data Fig. 7 | Map of plot locations of all plots of all projects ($n=23,641$). One or several
 1009 plots are summarised under the same plot resurvey ID ($n=7,738$). Note that the more complete
 1010 coverage of Bavaria resulted from including the grassland monitoring Bavaria which started in 2002⁷⁵.

1011

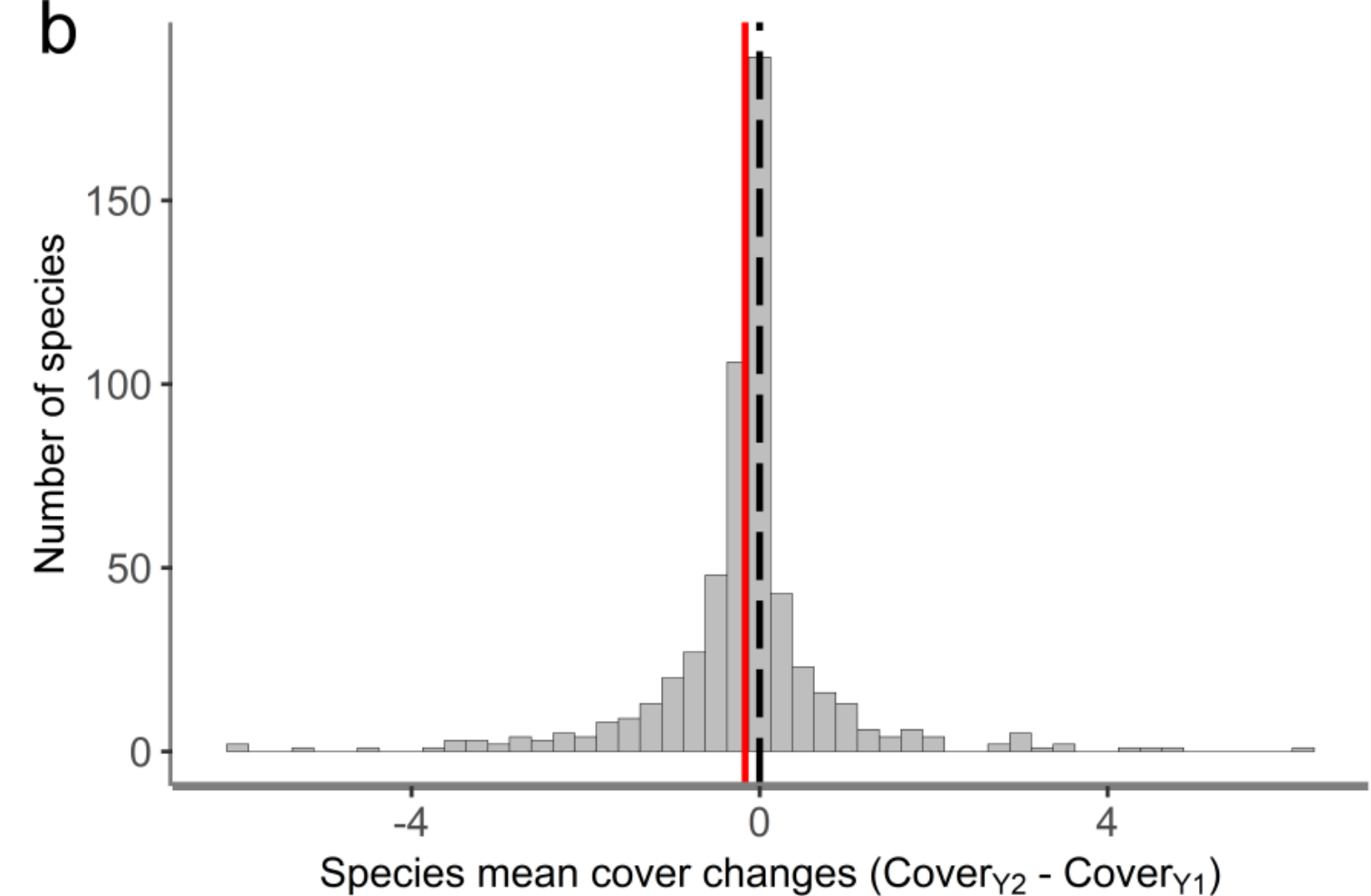
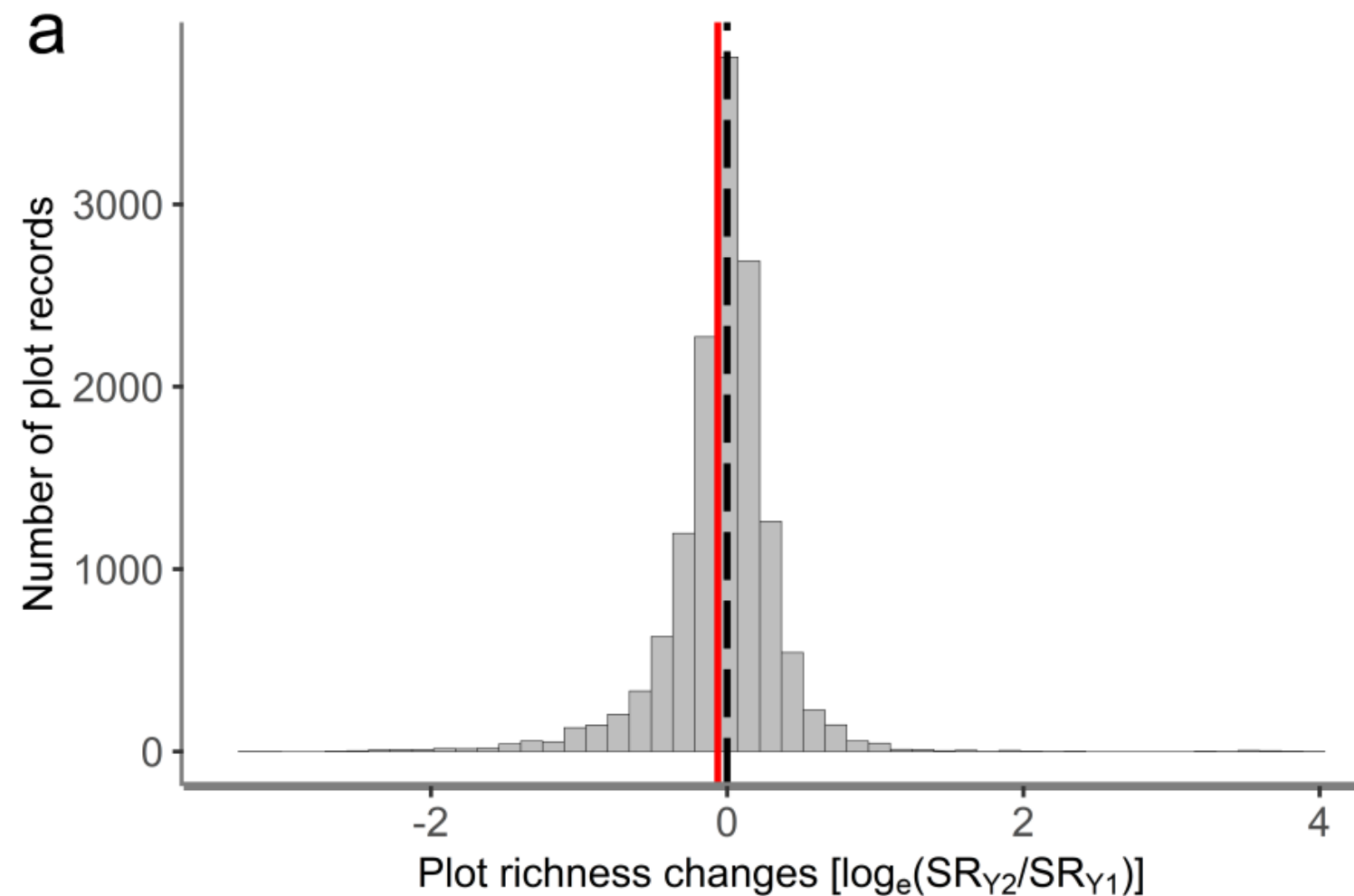
1012 Extended Data Fig. 8 | Assignment of time series plot records to EUNIS habitat types. Each time
 1013 series was assigned to the habitat type by using the earliest plot record that matched with the level 3
 1014 EUNIS classification. The classification was based on the EUNIS-ESy expert system⁵⁶ using the R code
 1015 implementation⁵⁷. ?: plots not assigned to any level 3 EUNIS habitat type, +: assigned to more than
 1016 one level 3 EUNIS habitat type, A: Marine habitats, C: Inland surface waters, H: Inland sparsely
 1017 vegetated habitats or devoid of vegetation, N: Coastal habitats, Q: Wetlands, R: Grasslands and lands
 1018 dominated by forbs, mosses or lichens, S: Heathlands, scrub and tundra, T: Forests and other wooded
 1019 land, V: Vegetated man-made habitats, including arable land. Labels for EUNIS habitats were only
 1020 printed at the top of the corresponding bar section when the number of assigned records was ≥ 150 .

1021

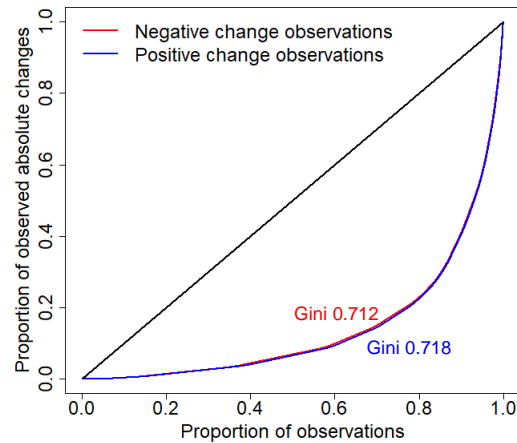
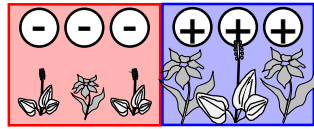
1022 Extended Data Fig. 9 | Steps of data preparation and analysis.

1023

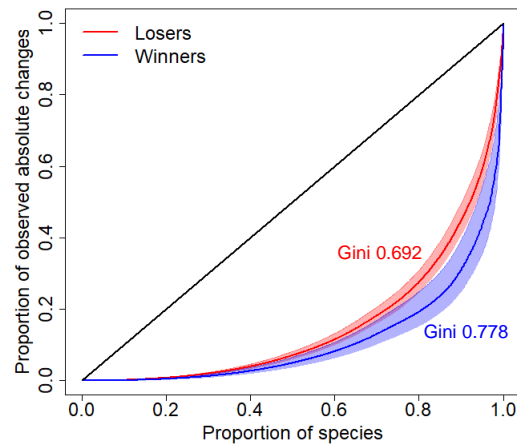
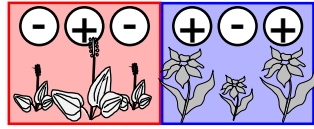
1024



a Change observations by
Plot ID x species x time interval



b Change observations
aggregated by species
(across all plots
and time intervals)



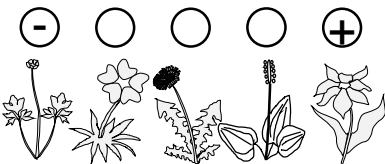
Hypotheses: The divergence in the distribution between cover losses and gains is driven by ...

i) ...the proportion of species that undergo changes

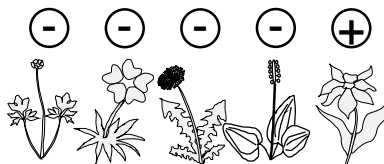
ii) ...the ratio of increasing to decreasing species

iii) ...the degree to which cover losses are concentrated on a specific subset of species

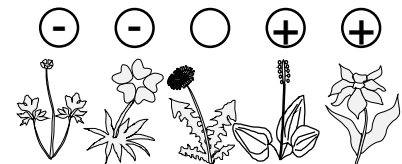
Proportion of species with cover changes 0.4



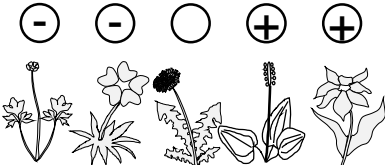
Proportion of increases in all changes 0.2



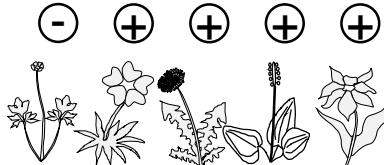
Random species selection



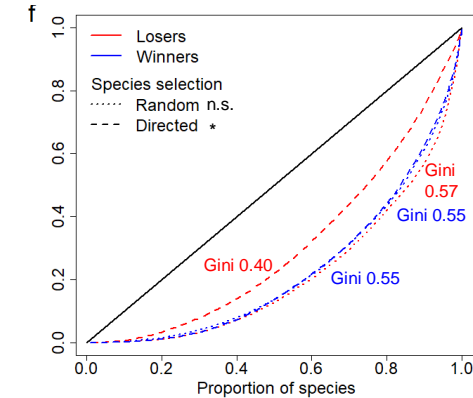
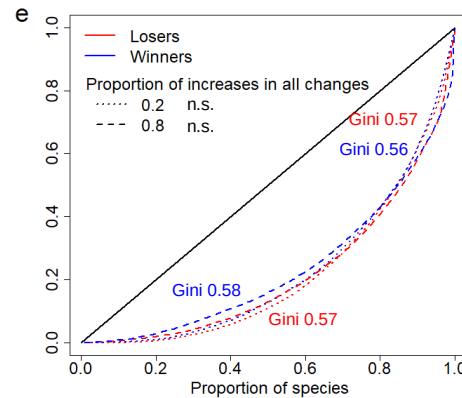
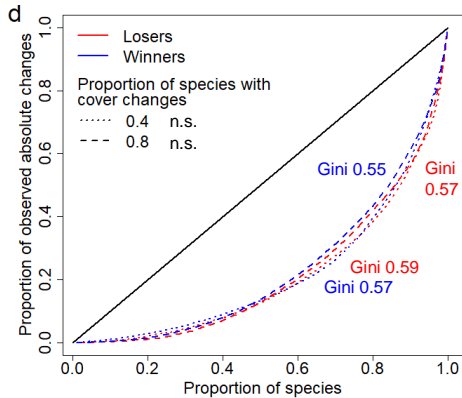
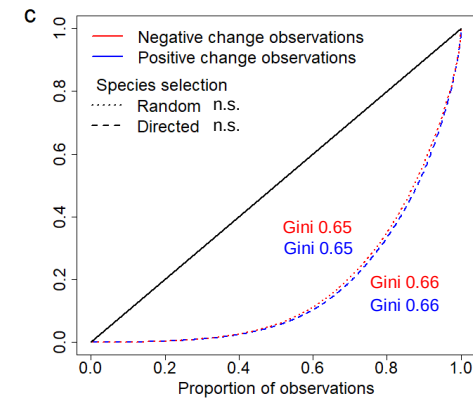
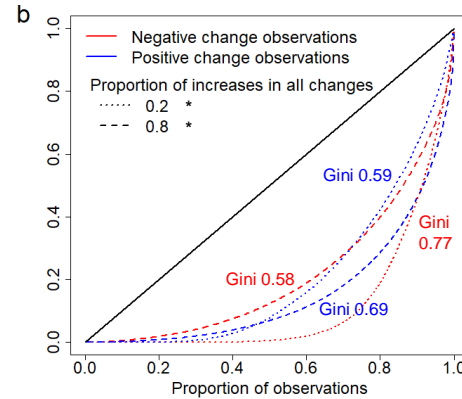
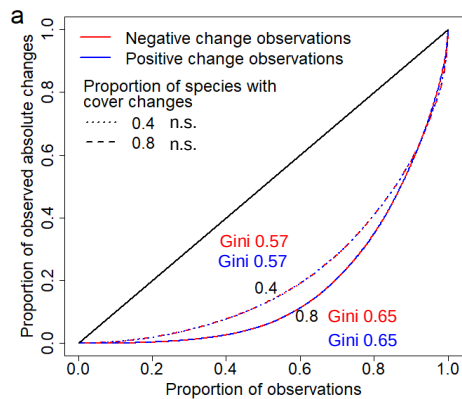
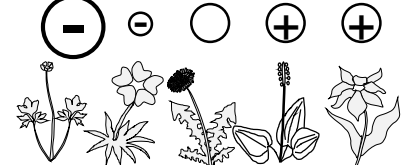
Proportion of species with cover changes 0.8

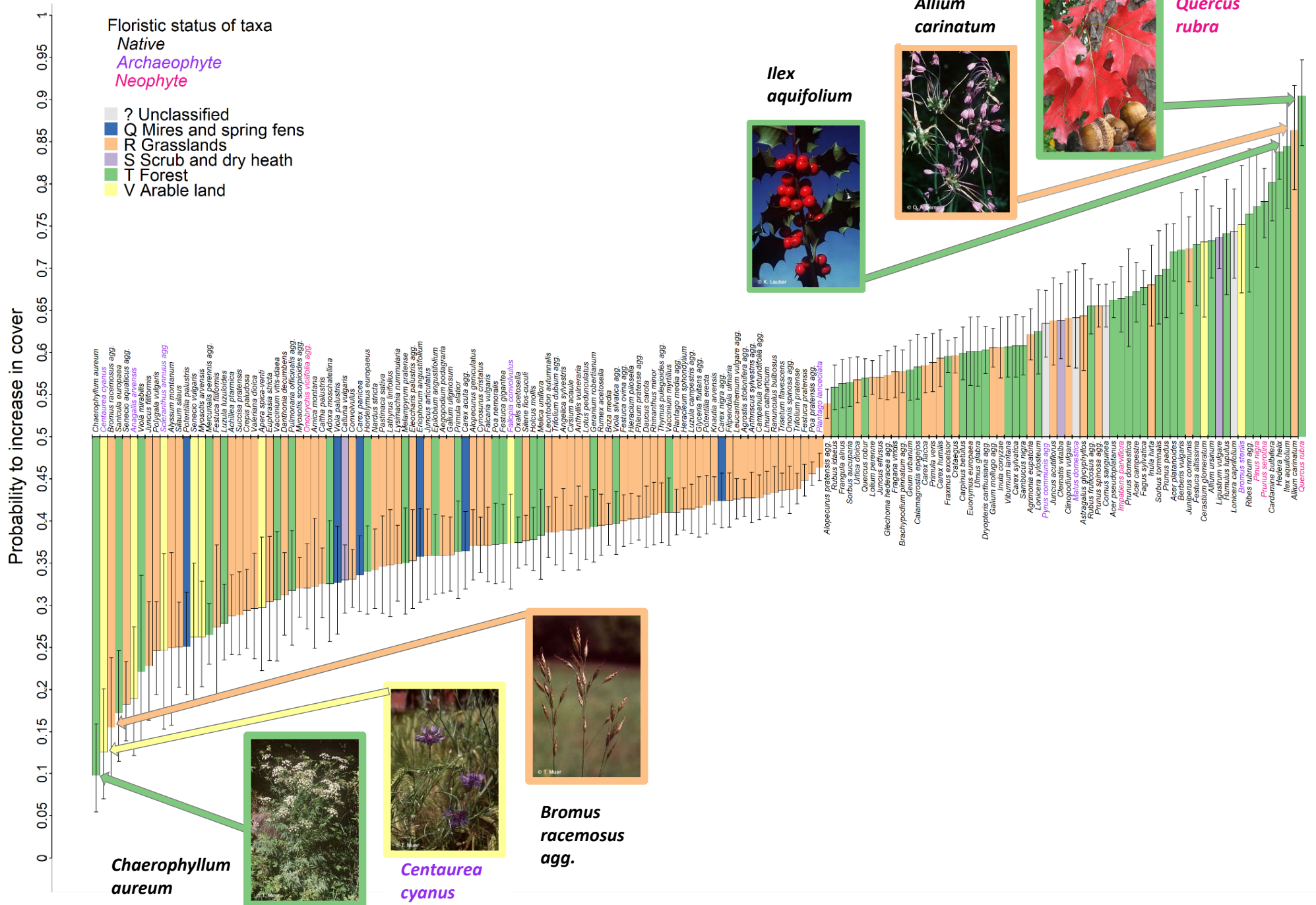


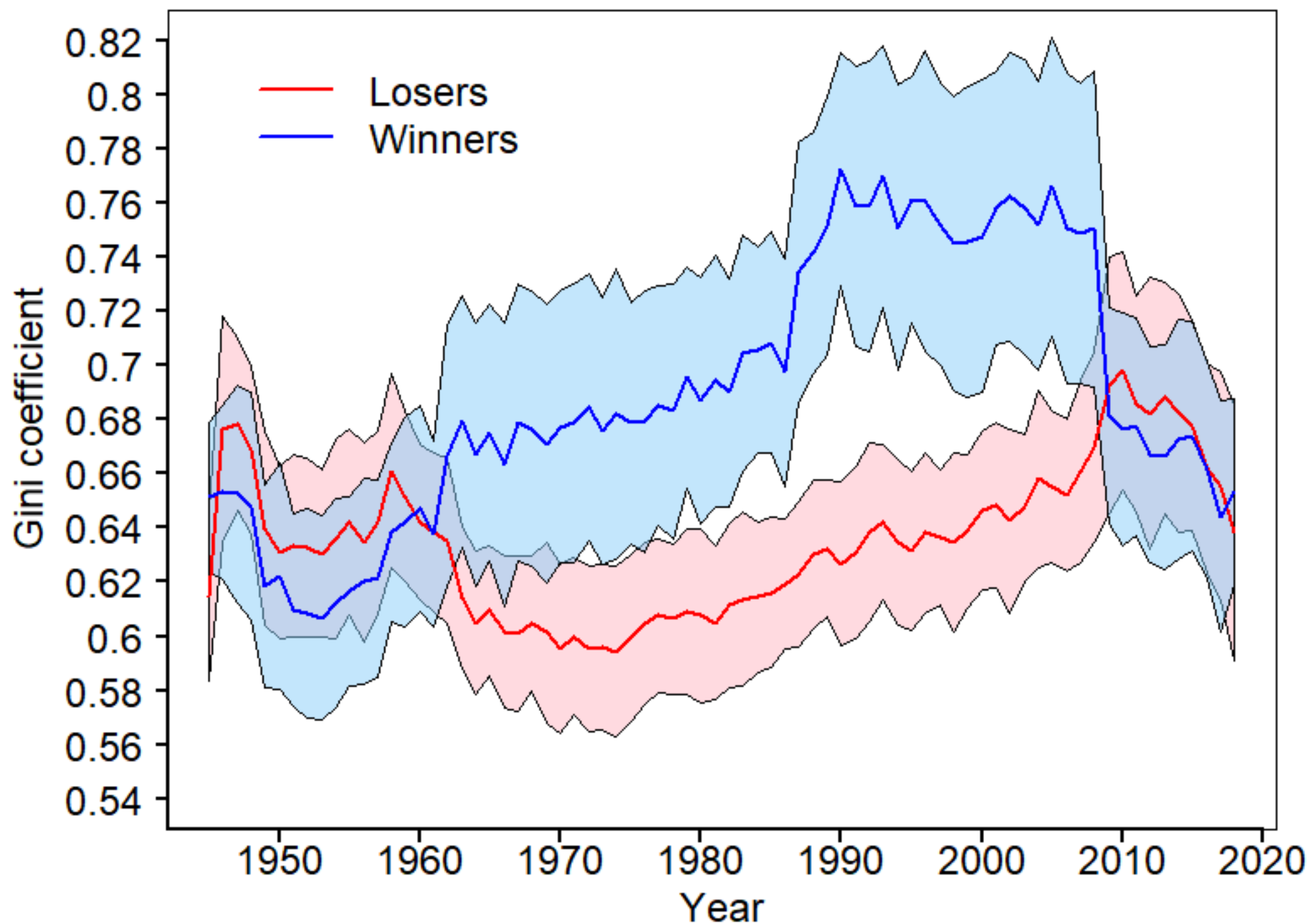
Proportion of increases in all changes 0.8

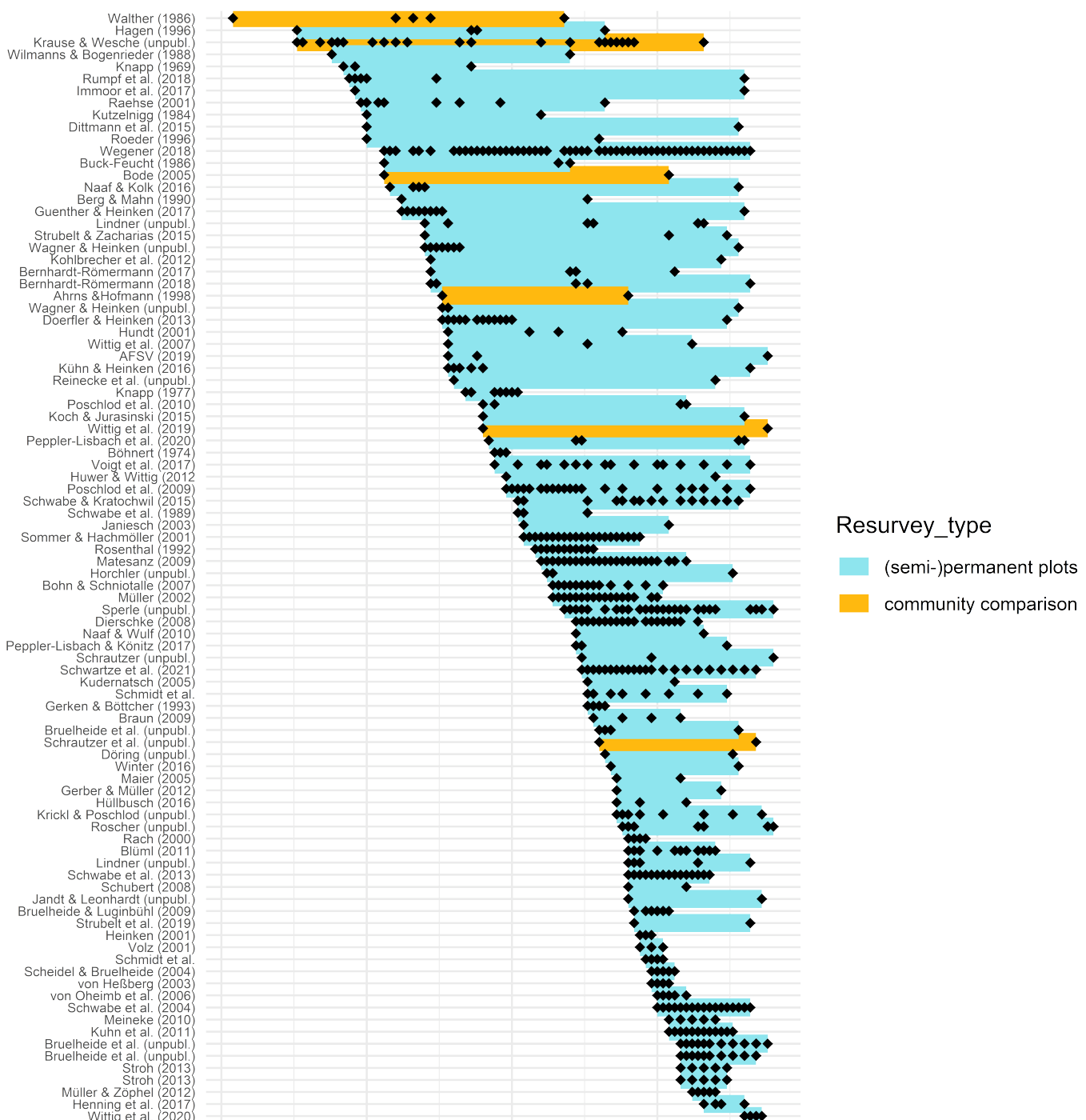


Directed species selection

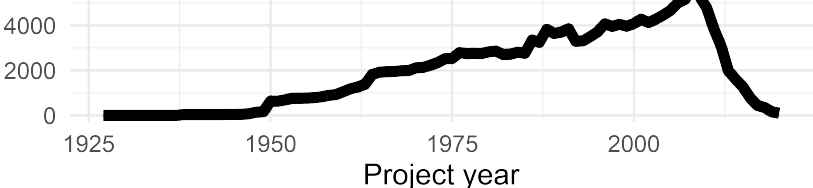


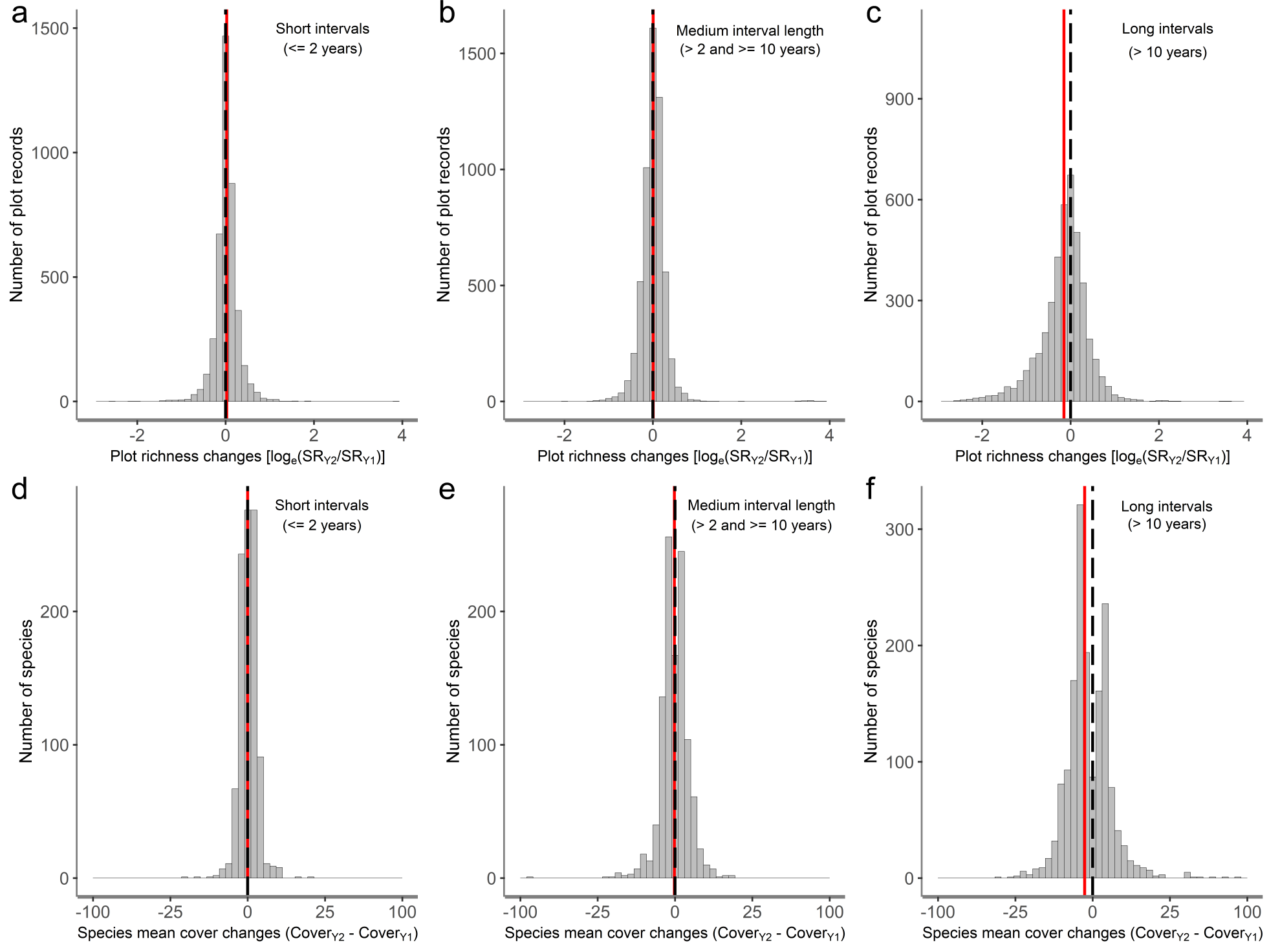






Number of time series



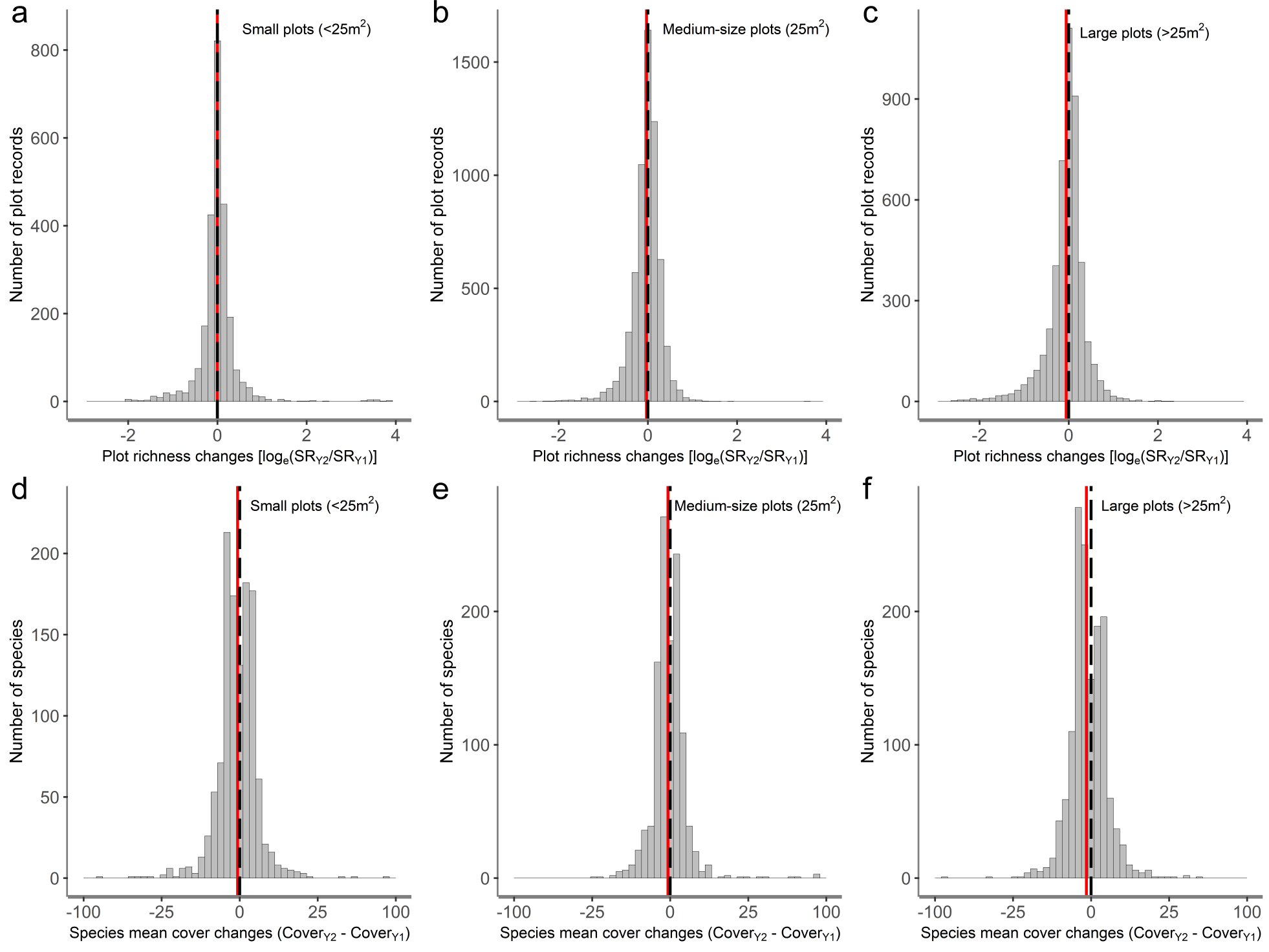


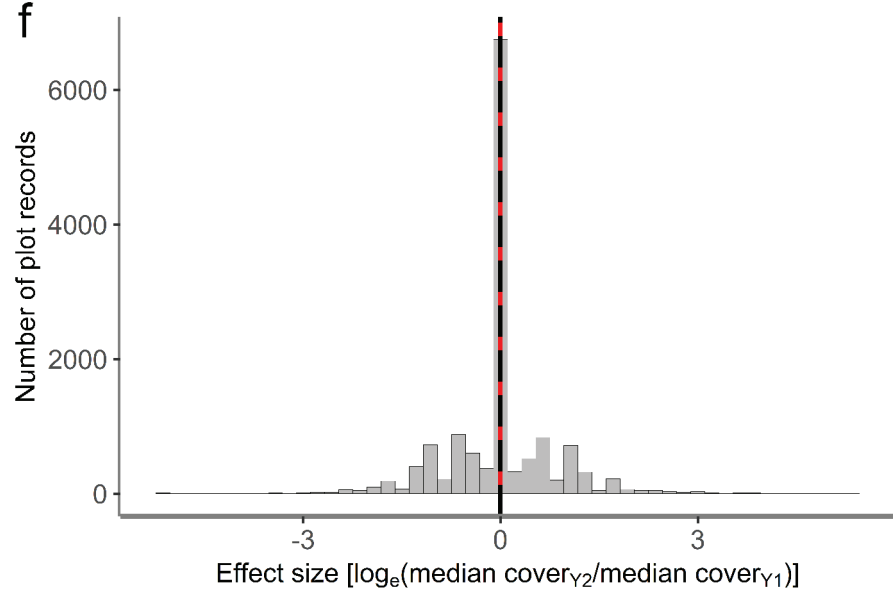
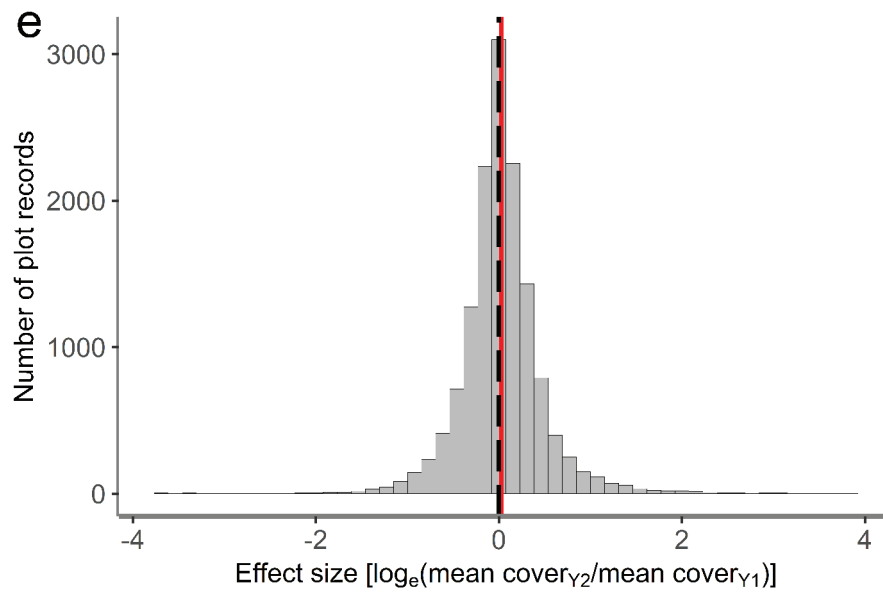
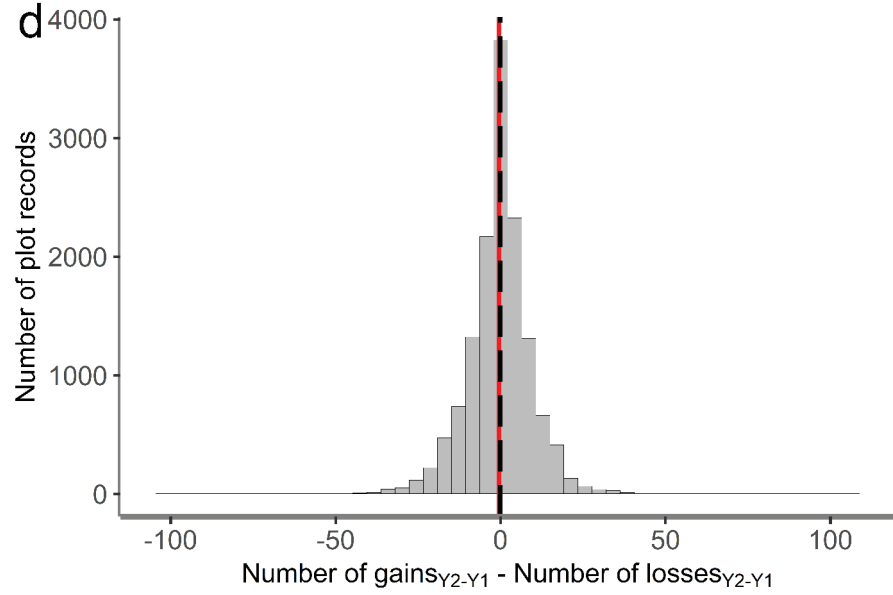
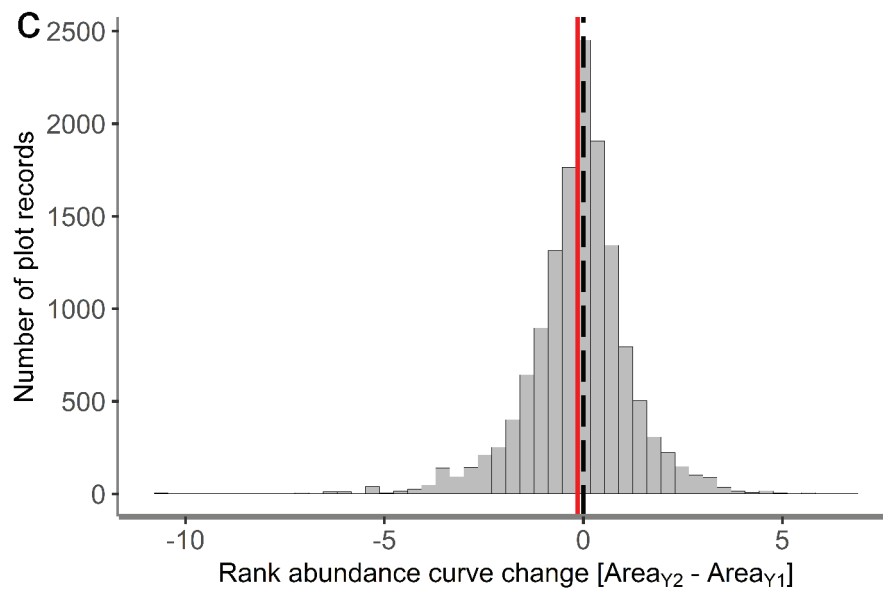
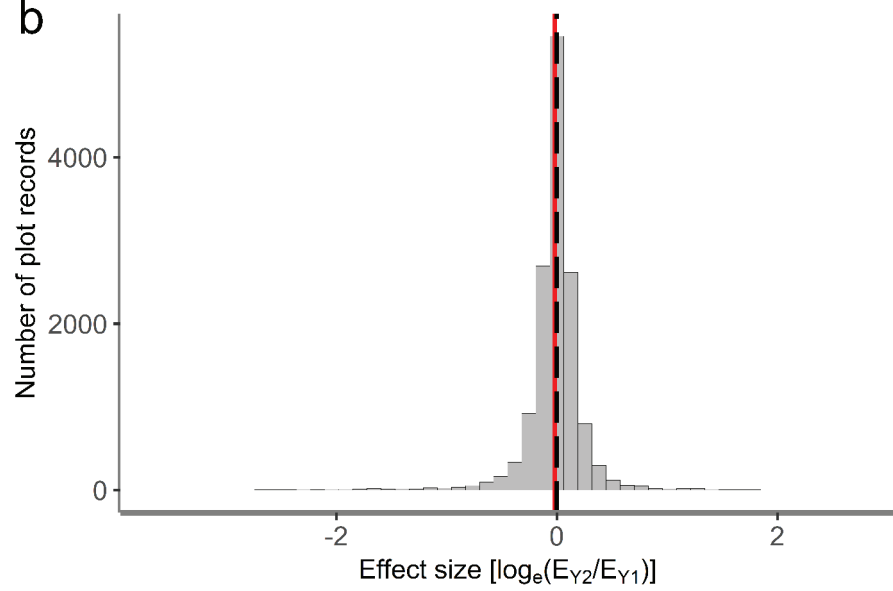
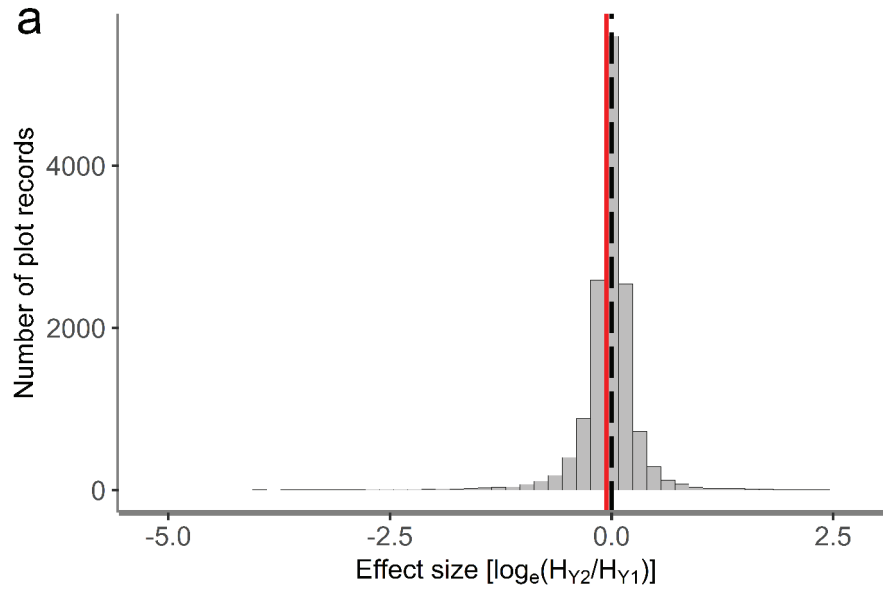
Number of plot records

1500
1000
500
0

-2 0 2
Effect size [$\log_e(\text{SR}_{Y2}/\text{SR}_{Y1}) \text{ decade}^{-1}$]







Number of species

200

100

0

-25

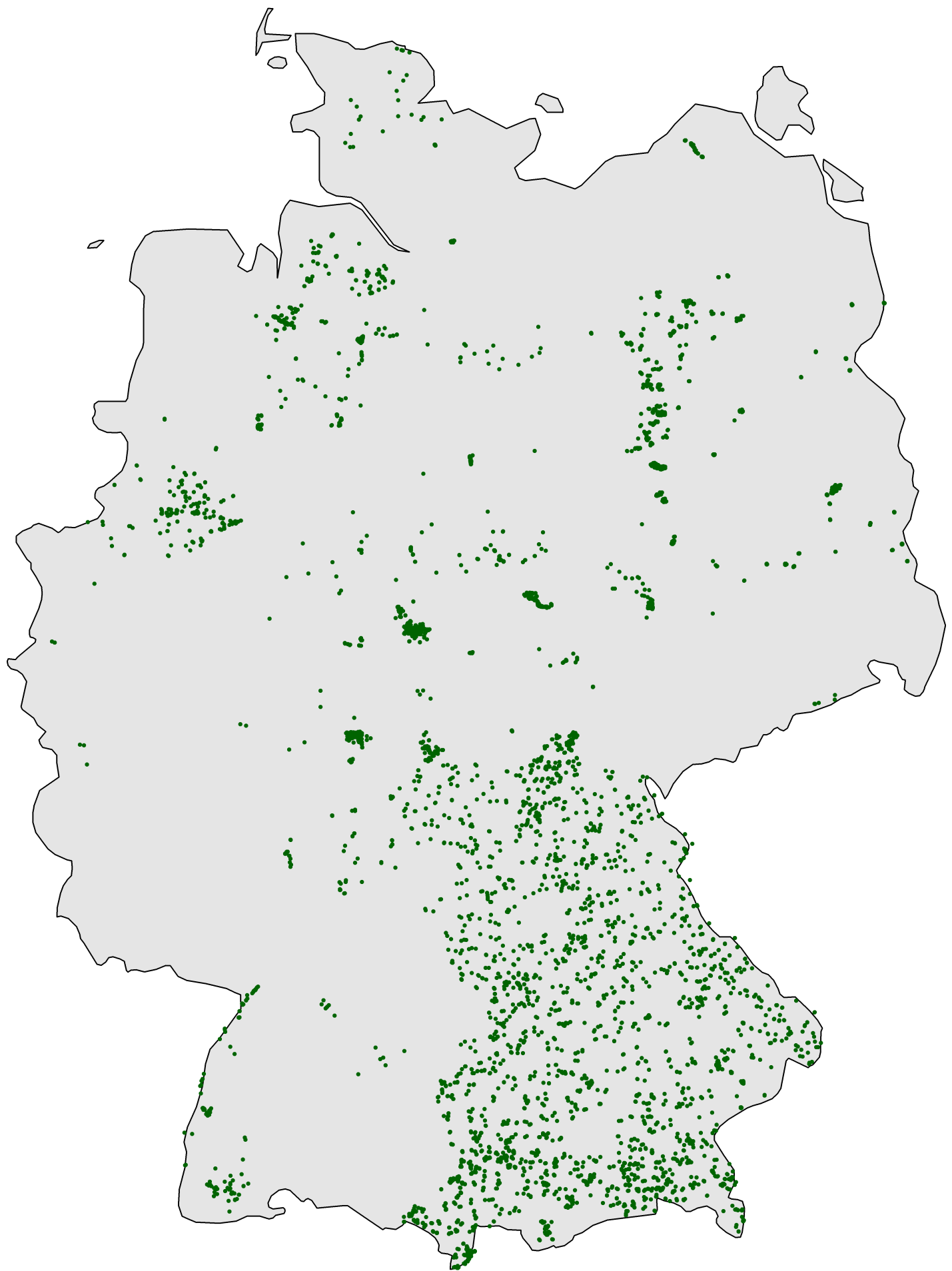
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25

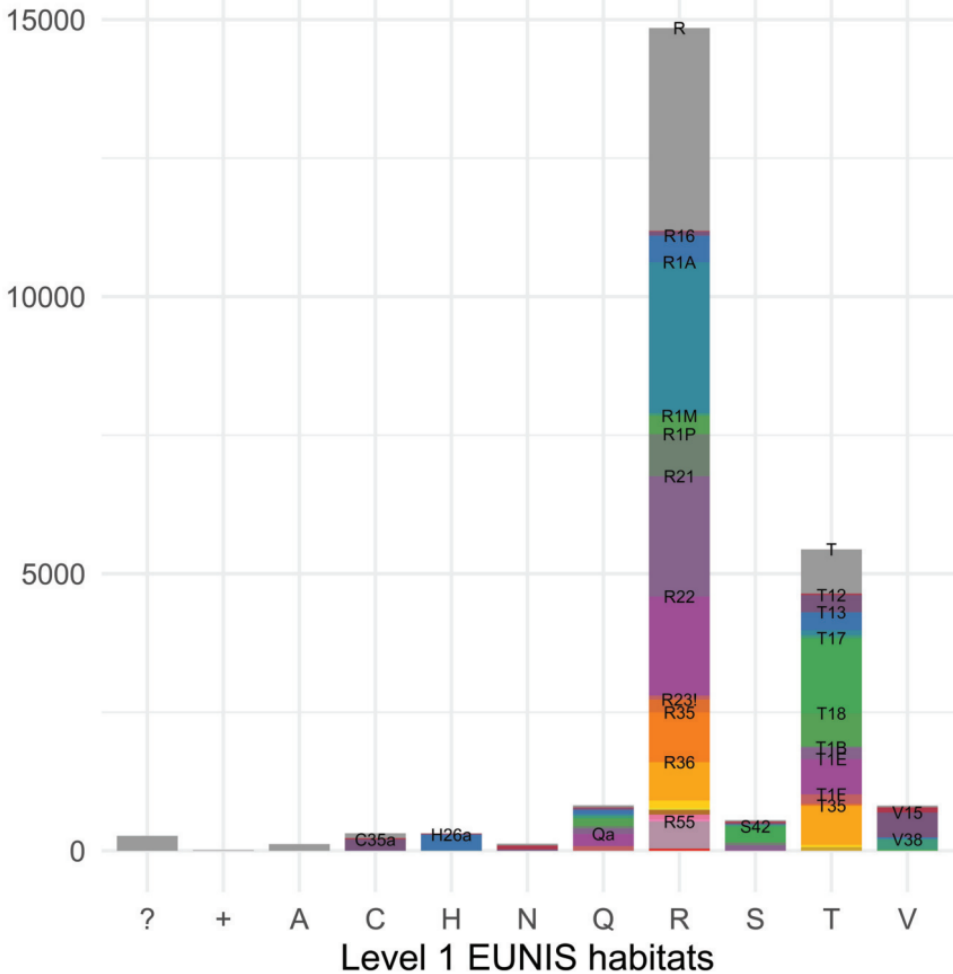
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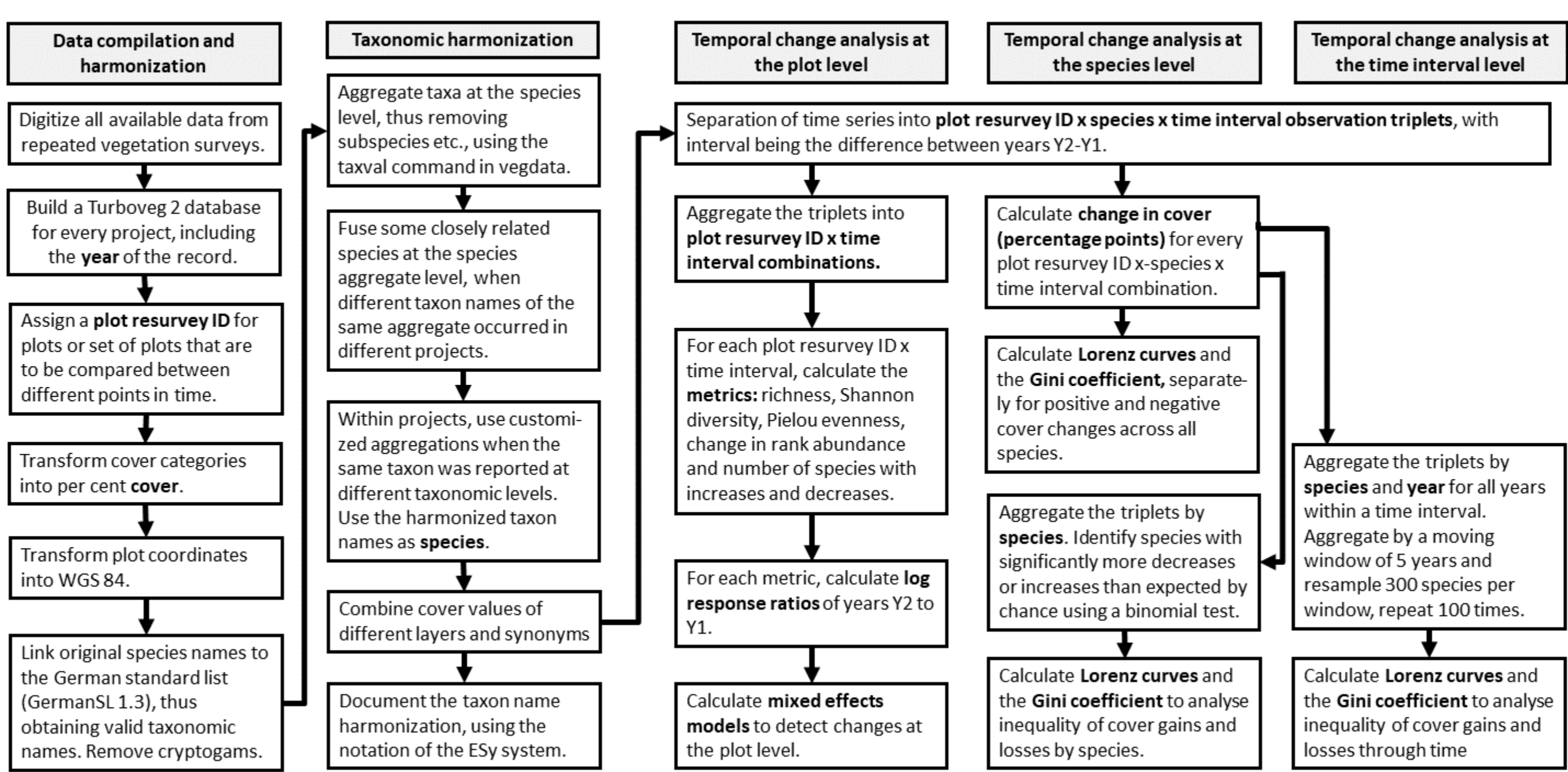
Species mean cover changes ($\text{Cover}_{Y2} - \text{Cover}_{Y1}$)





Number of records assigned





1 **Supplementary Tables**

2
3 Jandt, U., Bruelheide, H., Jansen, J., Bonn, A., Grescho, V., Klenke, R., Sabatini, F.M.,
4 Bernhardt-Römermann, M., Blüml, V., Dengler, J., Diekmann, M., Dörfler, I., Döring, U.,
5 Dullinger, S., Haider, S., Heinken, T., Horchler, P., Kuhn, G., Lindner, M., Metze, K., Müller, N.,
6 Naaf, T., Peppler-Lisbach, C., Poschlod, P., Roscher, C., Rosenthal, G., Rumpf, S., Schmidt, W.,
7 Schrautzer, W., Schwabe, A., Schwartze, P., Sperle, T., Stanik, N., Storm, C., Voigt, W.,
8 Wegener, U., Wesche, K., Wittig, B., Wulf, M. (2022): Plant diversity change over one century
9 in Germany: more losers than winners. – Nature

10
11

12
13 Extended Data Table 1 | List of all projects included in this study. PROJECT_ID: internal
14 reference number. EUNIS habitat types of time series were assigned to the habitat type by
15 using the earliest plot record that resulted in level 3 EUNIS classification. The classification
16 was based on the EUNIS-ESy expert system⁵⁶ using the R code implementation⁵⁷. When a
17 project included several habitat types, they are shown in decreasing numbers of plot
18 records. Code for habitat types are ?: plots not assigned to any level 3 EUNIS habitat type, +:
19 assigned to more than one level 3 EUNIS habitat type, A: Marine habitats, C: Inland surface
20 waters, H: Inland sparsely vegetated habitats or devoid of vegetation, N: Coastal habitats, Q:
21 Wetlands, R: Grasslands and lands dominated by forbs, mosses or lichens, S: Heathlands,
22 scrub and tundra, T: Forests and other wooded land, V: Vegetated man-made habitats,
23 including arable land.

PROJECT_ID	Project Name	Reference	EUNIS habitat type
77	AFSV (2019)	Arbeitsgemeinschaft Forstliche Standorts- und Vegetationskunde (AFSV) (2019) Nordwest-Eifel - Standorte, Waldgesellschaften, Nutzungen gestern und heute. Exkursionsführer der AFSV-Tagung 2019 in der Nordwesteifel. Verlag Dr. Kessel, Remagen-Oberwinter.	T17
1	Ahrns & Hofmann (1998)	Ahrns, C. & Hofmann, G. (1998) Vegetationsdynamik und Florenwandel im ehemaligen mitteldeutschen Waldschutzgebiet "Hainich" im Intervall 1963 - 1995. Hercynia N.F. 31: 33-64.	T17
4	Berg & Mahn (1990)	Berg, C. & Mahn, E.-G. (1990) Anthropogene Vegetationsveränderungen der Strassenrandvegetation in den letzten 30Jahren-die Glatthaferwiesen des Raumes Halle Saale. Tuexenia 10: 185-195.	R22, R
2	Bernhardt-Römermann (2017)	(Echinger Lohe)	T1F, T1E, T13, T
3	Bernhardt-Römermann (2018)	(Sonneberg)	T35, T32, T1F, T18, T17, T, S42, R57, R, ?

5	Blüml (2011)	Blüml, V. (2011) Langfristige Veränderungen von Flora und Vegetation des Grünlandes in der Dümmerniederung (Niedersachsen) unter dem Einfluss von Naturschutzmaßnahmen. Dissertation, Universität Bremen.	R55, R36, R35, R21, R, Qb, Q52, Q51
6	Bode (2005)	Bode, F. (2005) Subrezenter Vegetations- und Landschaftswandel im Südschwarzwald. Dissertation, Albert-Ludwigs-Universität Freiburg.	S42
92	Bohn & Schniotalle (2007)	Bohn, U. & Schniotalle, S. (2007) Hochmoor-, Grünland- und Waldrenaturierung im Naturschutzgebiet "Rotes Moor", Hohe Rhön 1981 - 2001: Ergebnisse 20-jähriger wissenschaftlicher Begleituntersuchungen im Rahmen und im Anschluss an ein E+E-Vorhaben des Bundes. Bundesamt für Naturschutz, Bonn.	T12, S92, R37, R35, R23!, R23, R1M, Qa, Q24, Q22, ?
78	Böhnert (1974)	Böhnert, W. (1978) Ökologische Untersuchungen auf den Kreidesandsteinhöhen der Harslebener Berge bei Quedlinburg. Naturschutz und naturkundliche Heimtforschung in den Bezirken Halle und Magdeburg 15 (2): 11-23	S42, R
7	Braun (2009)	Braun, W. (2009) Ein Niedermoor wächst über Hochmoortorf. Ber. d. Bayerischen Botanischen Gesellschaft 79: 127-146. München.	Qb, Qa, Q25, Q24, H26a
88	Bruelheide & Luginbühl (2009)	Bruelheide, H. & Luginbühl, U. (2009) Peeking at ecosystem stability: making use of a natural disturbance experiment to analyze resistance and resilience. Ecology 90 (5): 1314-1325.	T1F, T1E, T17, T, H25
17	Bruelheide et al. (unpubl.)	(Bergwiesen)	R55, R37, R35, R23!, R22, R1M, R, Q24, ?
90	Bruelheide et al. (unpubl.)	(Preuk)	T35, T1B, T, Sa
91	Bruelheide et al. (unpubl.)	(Schiessplatz)	T35, R
8	Buck-Feucht (1986)	Buck-Feucht, G. (1986) Vergleich alter und neuer Vegetationsaufnahmen im Forstbezirk Kirchheim unter Teck. Mitteilungen des Vereins für forstliche Standortskunde und Forstpflanzenzüchtung (32): 43-49.	T1E, T18, T17, T13, T
76	Dierschke (2008)	Dierschke, H. (2008) Dynamik und Konstanz an naturnahen Flussufern -27 Jahre Dauerflächenuntersuchungen am Oderufer (Harzvorland). Braunschweiger Geobotanische Arbeiten 9: 119-138.	R55

10	Dittmann et al. (2015)	Dittmann, T., Heinken, T. & Schmidt, M. (2018) Die Wälder von Magdeburgerforth (Fläming, Sachsen-Anhalt) – eine Wiederholungsuntersuchung nach sechs Jahrzehnten. <i>Tuexenia</i> 38: 11-42.	T35, T1E, T1B, T18, T17, T15, T13, T12, T
11	Doerfler & Heinken (2013)	Vegetation change of wet and moist forests in NE German nature reserves	T3M, T3J, T35, T1E, T17, T16, T15, T13, T12, T, S92, Qb, Qa, Q53, +, ?
12	Döring (unpubl.)	Feuchtwälder in den Landkreisen Diepholz und Uelzen; Wiederholungsaufnahmen von PD Dr. Wilfried Hakes [NW-FVA] im Rahmen des Forschungsvorhabens "NaLaMa-nT" erhoben	T1E, T16, T15, T13, T12, T
62	Gerber & Müller (2012)	Gerber, L. & Müller, F. (2012) Flora und Vegetation den Naturschutzgebietes Am Galgenteich Altenberg. Berichte der Arbeitsgemeinschaft sächsischer Botaniker N.F. 21: 65-123.	S42, S32, R1M, R
13	Gerken & Böttcher (unpubl.)	Abschlußbericht, unpubl.	V38, R22, R1A, R, H26a, ?
15	Günther et al. (2021)	Günther, K., Schmidt, M., Quitt, H. & Heinken, T. (2021): Veränderungen der Waldvegetation im Elbe-Havelwinkel von 1960 bis 2015. <i>Tuexenia</i> 41: 53-85.	T3M, T35, T1B, T16, T15, T13, T12, T, S92, S42
16	Hagen (1996)	Hagen, T. (1996) Vegetationsveränderungen in Kalk-Magerrasen des Fränkischen Jura. Laufener Forschungsbericht 4. Bayerische Akademie f. Naturschutz u. Landschaftspflege, Laufen, Salzach. 218 S.	R1A, R16, R13
79	Heinken (2001)	Heinken, A. (2001) Vegetationsentwicklung von Auengrünland nach Wiederüberflutung. Diss. Math.-Nat. Fak. Humboldt-Universität Berlin. 161 S.	R36, R22, R21, R, Qb, Q52, Q51
43	Heinrich, Marstaller & Voigt (2012)	Eine Langzeitstudie zur Sukzession in Halbtrockenrasen - Strukturwandlung in einer Dauerbeobachtungsfläche im Naturschutzgebiet "Leutratal und Cospoth" bei Jena (Thüringen). Artenschutzreport Jena 30: 1-80.	T36, T1H, T19, T17, T, Sa, S35, R51!, R22, R1A, R16, H26a, ?
68	Henning et al. (2017)	Henning, K.; Lorenz, A.; von Oheimb, G.; Härdtle, W.; Tischew, S. Year-round cattle and horse grazing supports the restoration of abandoned, dry sandy grassland and heathland communities by supressing Calamagrostis. <i>Journal for Nature Conservation</i> 40: 120-130.	V38, S42, R1P, R1A, N19, N15
54	von Heßberg (2003)	von Heßberg, A. (2003) Landschafts- und Vegetationsdynamik entlang renaturierter Flussabschnitte von Obermain und Rodach. Dissertation, Universität Bayreuth.	V15, T13, S91, R55, R21, H25

74	Horchler (unpubl.)	Horchler, P., Henrichfreise, A. Vollmer, I. (2013) Wiederholungsaufnahme von 54 Auenwald-Vegetationsaufnahmen am Oberrhein.	T1H, T1F, T1E, T13, T11, T, Sa, S35, R55, Q52, C23, ?
19	Hüllbusch et al. (2016)	Hüllbusch, E., Brandt, L.M., Ende, P. & Dengler, J. (2016) Little vegetation change during two decades in a dry grassland complex in the Biosphere Reserve Schorfheide-Chorin (NE Germany). <i>Tuexenia</i> 36: 395-412.	R22, R1P, R1B, R1A, R, N15!!
20	Hundt (2001)	Hundt, R. (2001) Ökologisch-geobotanische Untersuchungen an den mitteldeutschen Wiesengesellschaften unter besonderer Berücksichtigung ihres Wasserhaushaltes und ihrer Veränderung durch die Intensivbewirtschaftung. Mitteilungen aus dem Biosphärenreservat Rhön/Thüringen. 3. Monographie. 366 S.	V37, R55, R37, R36, R35, R23!, R22, R1M, R1A, R, Q53, Q43, +
48	Huwer & Wittig (2012)	unpublished data for: Huwer, A. & Wittig, R. (2012) Changes in the species composition of hedgerows. <i>Tuexenia</i> 32: 31-53. Göttingen.	T1E, T1B, T13, T12, T11, T, Sa, S37, ?
21	Immoor et al. (2017)	Immoor, A., Zacharias, D., Müller, J. & Diekmann, M. (2017) A re-visitation study (1948–2015) of wet grassland vegetation in the Stedinger Land near Bremen, North-western Germany. <i>Tuexenia</i> 37: 271-288.	R36, R35, R21, R, Qb
23	Jandt & Leonhardt (unpubl.)	(Kyffhäuser)	S42, R1BI, R1B, R1A, R16, R13
22	Janiesch (2003)	Janiesch, P. (2003) Vegetationsökologische Untersuchungen in einem Erlenbruchwald im nördlichen Münsterland - 25 Jahre im Vergleich. Abhandlungen aus dem Westfälischen Museum für Naturkunde: Vegetation und Fauna in Westfalen (ed Westfälisches Museum für Naturkunde), pp. 71-80, Münster.	T15, T13
70	Knapp (1969)	Knapp, R. (1969) Änderungen in der Vegetation Hessischer Gebirge in den letzten Jahrzehnten. Mitteilungen der Floristisch-Soziologischen Arbeitsgemeinschaft N.F. 14: 274-286	R35, R22, R1M, R
71	Knapp (1977)	Knapp, R. (1977) Dauerflächen-Untersuchungen über die Einwirkung von Haustieren und Wild während trockener und feuchter Zeiten in Mesobromion- Halbtrockenrasen in Hessen. Mitteilungen der Floristisch-Soziologischen Arbeitsgemeinschaft N.F. 19: 269-274	R1A
87	Koch & Jurasiński (2015)	Koch, M. & Jurasiński, G. (2015) Four decades of vegetation development in a percolation mire complex following intensive drainage and abandonment. <i>Plant Ecology & Diversity</i> 8: 49-60	T16, T12, T, R55, R35, Qb, Qa, Q53, Q52, Q51, Q43, Q42, Q41, Q24, C35a, ?

59	Kohlbrecher et al. (2012)	unpublished data for: Kohlbrecher, C., Wesche, K., Hilbig, W., Leuschner, C., Meyer, S. (2012) Veränderungen der Segetalvegetation am Kyffhäusergebirge in den letzten 50 Jahren. Landschaftspflege und Naturschutz in Thüringen 49: 1-9.	V15
24	Krause & Wesche (2011)	unpublished data for: Krause, B., Culmsee, H., Wesche, K., Bergmeier, E. & Leuschner, C. (2011) Habitat loss of floodplain meadows in north Germany since the 1950s. Biodiversity and Conservation 20 (11): 2347-2364.	R22, R21, R1P, R, C35a, A25c
84	Krickl & Poschlod (unpubl.)	(calcareous grasslands SW-Germany)	R1A, R16, H26a
65	Kudernatsch (2005)	Kudernatsch, T. (2005) Auswirkungen der globalen Erwärmung auf die Vegetation alpiner Kalk-Magerrasen im Nationalpark Berchtesgaden. Dissertation TU München, Department für Ökologie, Fachgebiet Geobotanik Weihenstephan. 151 S.	S22, R44, R43!, R
25	Kühn & Heinken (2017)	unpublished data for: Kühn, S.L., Heinken, T. (2017): Vegetationsveränderungen im NSG Bredower Forst im Verlauf von 50 Jahren – Analyse anhand historischer Vegetationsaufnahmen. Natursch. Landschaftspfl. Brandenbg. 26(4): 4-16.	T35, T1E, T1B, T18, T17, T, ?
14	Kuhn et al. (2011)	unpublished data for: Kuhn, G.; Heinz, S.; Meyer, F. (2011) Grünlandmonitoring Bayern, Ersterhebung der Vegetation 2002 - 2008. LfL Schriftenreihe Bayerische Landesanstalt für Landwirtschaft 3.	V38, V37, V15, V, S22, R55, R44, R37, R36, R35, R23!, R23, R22, R21, R1M, R1A, R16, R, Qb, Q53, Q52, Q51, Q43, Q41, C35d, ?
26	Kutzelnigg (1984)	Kutzelnigg, H. (1984) Veränderungen der Ackerwildkrautflora im Gebiet um Moers/Niederrhein seit 1950 und ihre Ursachen. Tuexenia (4): 81-102.	V37, V15
58	Lindner (unpubl.)	(Hechtmoor)	T1B, T16, T, Sb, S92, S41, R, Qb, Qa, Q51, Q42, Q25, Q24, Q21, ?
60	Lindner (unpubl.)	(Süderbrarup)	R55, R52, R22, R1M, R
27	Maier (2005)	Maier, M. (2005) Untersuchung zur Entwicklung von Flora und Fauna in einem Feuchtwiesenschutzgebiet (Naturschutzgebiet Bornhorster Huntewiesen). Diplomarbeit Landschaftsökologie an der Carl von Ossietzky Universität Oldenburg.	R36, R35, Qb, Q53, Q51
28	Matesanz (2009)	Matesanz, S., Brooker, R.W., Valladares, F. & Klotz, S. (2009) Temporal dynamics of marginal steppic vegetation	R16

		over a 26-year period of substantial environmental change. Journal of Vegetation Science 20 (2): 299-310.	
55	Meineke & Menge (2010)	Meineke, T., & Menge, K. (2010) Wirkungskontrolle zum PROFIL Kooperationsprogramm Naturschutz, Teilbereich Besondere Biotoptypen im FFH-Gebiet Bergwiesen und Wolfsbachtal bei Hohegeiß. Im Auftrag des Niedersächsischen Landesbetriebs für Wasserwirtschaft Küsten- und Naturschutz, Hannover.	R23!, R21, R
63	Müller & Zöphel (2012)	Müller, F. & Zöphel, B. (2012) Bestandssituation, Biologie und Ökologie von Gentianella lutescens im Osterzgebirge. Berichte der Arbeitsgemeinschaft sächsischer Botaniker N.F. 21: 139-184	R23!
29	Müller (2002)	Müller, N. (2002) Auswertung der Langzeituntersuchungen von Dauerflächen im Augsburger Stadtgebiet zur Renaturierung von Lechhainen. Ber. Bayer. Landesamt Umweltschutz (Hrsg.): 97 S.	V38, V, R55, R37, R22, R1A, R16, N15, H26a
30	Naaf & Kolk (2016)	unpublished data for: Naaf, T. & Kolk, J. (2016) Initial site conditions and interactions between multiple drivers determine herb-layer changes over five decades in temperate forests. For Ecol Manag 366: 153-165.	T1H, T1E, T1B, T18, T17, T15, T13, T12, T, Q51, ?
31	Naaf & Wulf (2010)	Naaf, T. & Wulf, M. (2010) unpublished data for: Habitat specialists and generalists drive homogenization and differentiation of temperate forest plant communities at the regional scale. Biol Conserv 143: 848-855.	T1F, T1E, T17, T13, T12, T, Sa
32	Peppler-Lisbach & Könitz (2017)	Peppler-Lisbach, C. & Könitz, N. (2017): Vegetationsveränderungen in Borstgrasrasen des Werra-Meißner-Gebietes (Hessen, Niedersachsen) nach 25 Jahren – Tuexenia 37: 201-228.	S42, R37, R22, R21, R1M, R
44	Peppler-Lisbach et al. (2020)	Peppler-Lisbach, C, Stanik, N, Könitz, N, Rosenthal, G. (2020) Long-term vegetation changes in <i>Nardus</i> grasslands indicate eutrophication, recovery from acidification, and management change as the main drivers. Applied Vegetation Science 23: 508-521. https://doi.org/10.1111/avsc.12513	Sb, Sa, S42, R37, R35, R23!, R23, R22, R1M, R
83	Poschlod et al. (2009)	unpublished data for: Poschlod, P., Schreiber, K.-F., Mitlacher, K., Römermann, C. & Bernhardt-Römermann, M. (2009): Entwicklung der Vegetation und ihre naturschutzfachliche Bewertung. In: Schreiber, K.-F., Brauckmann, H.-J., Broll, G., Krebs, S. & Poschlod, P. (Hrsg.): Landschaftspflege und Naturschutz im Extensivgrünland. 30 Jahre Offenhaltungsversuche Baden-Württemberg. – Naturschutz-Spectrum Themen 97: 243-288.	T1E, R22, R21, R1M, R1A, N19

85	Poschlod et al. (2010)	unpublished data for: Poschlod, P., Kos, M., Roauer, S., Seemann, A., Wiesmann, O., Zeltner, G., Kohler, A. (2006) Long-term monitoring in rivers of south Germany since the 1970ies - macrophytes as indicators for the assessment of water quality. In: Müller, F., Baessler, C., Schubert, H. & Klotz, S. (Eds.): Long-term ecological research. Between Theory and Application. Berlin: Springer, pp. 189-199.	Qb, Q52, Q51, C23, C22b, ?
64	Rach (2000)	Rach, C. (2000) Charakterisierung von Renaturierungsprozessen in Bruchwäldern - Ökologische Untersuchungen in zwei Landschaftsräumen Nordwestdeutschlands. Dissertation Fachbereich Biologie, Geo- und Umweltwissenschaften Universität Oldenburg. 201 S.	T15, T12, T
36	Raeħse (2001)	Raeħse, S. (2001) Veränderungen der hessischen Grünlandvegetation seit Beginn der 50er Jahre am Beispiel ausgewählter Tal- und Bergregionen Nord- und Mittelhessens. Kassel, University Press GmbH. 222 S.	V15, S42, R55, R37, R36, R35, R22, R21, R1M, R1A, R, Qb, Q53, Q51, N19
37	Reinecke et al. (unpubl.)	unpublished data for: Reinecke, J., Klemm, G., Heinkein, T. (2014): Vegetation change and homogenization of species composition in temperate nutrient-deficient Scots pine forests after 45 yr. <i>J. Veg. Sci.</i> 25: 113-121.	T35, T1B, T, S42, R54, ?
80	Roeder et al. (1996)	Röder, H., Fischer, A., Klöck, W. (1996) Waldentwicklung auf Quasi-Dauerflächen im Luzulo-Fagetum der Buntsandsteinrhön (Forstamt Mittelsinn) zwischen 1950 und 1990. <i>Forstw. CB1.</i> 115, 321-335	T35, T1H, T18, T, S42, R, ?
86	Roscher (unpubl.)	(East Thuringia)	R51!, R22, R1A, R16, R
38	Rosenthal (1992)	Rosenthal, G. (1992) Erhaltung und Regeneration von Feuchtwiesen. Vegetationsökologische Untersuchungen auf Dauerflächen. <i>Diss. Bot.</i> 182: 1-283. Berlin, Stuttgart.	V15, R55, R36, R35, R, Q51
67	Rumpf et al. (2018)	unpublished data for: Rumpf, S.; Dullinger, S. (2018) Range dynamics of mountain plants decrease with elevation. <i>PNAS</i> 115(8):1848-1853.	Sb, S42, S22, R56, R55, R44, R43, R41, R23!, R21, R, H32c, H26b, H24, H23
39	Scheidel & Bruelheide (2004)	Scheidel, U. & Bruelheide, H. (2004) Versuche zur Beweidung von Bergwiesen im Harz. <i>Hercynia N.F.</i> 37: 87-101	S32, R35, R, ?
33	Schmidt et al.	Garbitz, D. (1990): Vegetation und Standortsbedingungen im Naturwald "Staufenberg". Dipl.-Arb. Syst.-Geobot. Institut, Universität Göttingen. Melcher, S. (1999): Flora und Vegetation im Naturwald "Großer Staufenberg" (Forstamt Walkenried, Revier	T3M, T35, T1F, T1E, T18, T17, T, Sa, +, ?

		<p>Staufenberg). Dipl.-Arb. Institut f. Waldbau, Abt. I, Universität Göttingen.</p> <p>Mölder, A., Streit, M., Schmidt, W. (2014): When beech strikes back: How strict nature conservation reduces herb-layer diversity and productivity in Central European deciduous forests. <i>Forest Ecology and Management</i> 319: 51-61.</p> <p>Kohls, K. (1994): Geobotanische Untersuchungen in Wäldern des Forstamtes Sellhorn (Lüneburger Heide). Dipl.-Arb. Syst.-Geobot. Institut, Universität Göttingen.</p> <p>Albrecht, B. (2000): Vegetationskundliche Untersuchungen im Naturwaldreservat "Meninger Holz" unter besonderer Berücksichtigung der Vegetationsentwicklung. Dipl.-Arb. FG Naturschutz, FB Biologie, Universität Hamburg.</p> <p>Happe, E. (1995): Vegetation und Standortsverhältnisse im Naturwald und Naturschutzgebiet "Totenberg" (Bramwald). Dipl.-Arb. Institut f. Waldbau, Abt. I, Universität Göttingen.</p> <p>Fischer, C., Parth, A., Schmidt, W. (2009): Vegetationsdynamik in Buchen-Naturwäldern. Ein Vergleich aus Süd-Niedersachsen. <i>Hercynia</i> 42: 45-68.</p>	
34	Schmidt et al.	<p>Kompa T., Schmidt, W. (2005): Buchenwald-Sukzession nach Windwurf auf Zechstein-Standorten des südwestlichen Harzvorlandes. <i>Hercynia</i> N.F. 38: 233-261.</p> <p>Schmidt, W. (2002): Die Naturschutzgebiete Hainholz und Staufenberg am Harzrand – Sukzessionsforschung in Buchenwäldern ohne Bewirtschaftung. <i>Tuexenia</i> 22: 151-213.</p> <p>Schmidt, W., Heinrichs, S. (2012) 13 Jahre nach dem Sturm - Vegetationsentwicklung im Buchen-Naturwald "Königsbuche" (südwestliches Harzvorland, Niedersachsen). <i>Hercynia</i> 45: 81-110.</p>	V39, T1F, T1E, T18, T17, T13, T12, T, Sa, S32, R57, R55, R, ?
73	Schrautzer (unpubl.)	Resurvey Eidertal	R55, R35, Q53, Q52
72	Schrautzer et al. (unpubl.)	Resurvey of Härdtle - Beckmann	T17, T13, T12, T
40	Schubert (2008)	Schubert, R. (2008) Vegetationsdynamik in einigen Naturschutzgebieten Sachsen-Anhalts. <i>Mitteilungen florist. Kart. Sachsen-Anhalt</i> (Halle 2008) 13: 53-75.	V38, T, S42, S38, R55, R35, R22, R1A, R, Q51, ?
42	Schwabe & Kratochwil (2015)	Schwabe, A. & Kratochwil, A. (2015) Pflanzensoziologische Dauerflächen-Untersuchungen im Bannwald "Flüh" (Südschwarzwald) unter besonderer	T18, T, S42, R52

		Berücksichtigung der Weidfeld-Sukzession. standort.wald 49: 5-49	
41	Schwabe et al. (1989)	Schwabe, A., Kratochwil, A. & Bammert, J. (1989) Sukzessionsprozesse im aufgelassenen Weidfeld-Gebiet des "Bannwald Flüh" (Südschwarzwald) 1976-1988 - Mit einer vergleichenden Betrachtung statistischer Auswertungsmethoden. Tuexenia 9: 351-370. Göttingen.	S42, R52, R1M, R, ?
89	Schwabe et al. (2004)	(Grundaufnahmen) unpublished data for: Schwabe, A.; Zehm, A., Nobis, M., Storm, C., Suess, K. (2004) Auswirkungen von Schaf-Erstbeweidung auf die Vegetation primär basenreicher Sand-Ökosysteme. Ber. NNA 1:/2004: 39-54.	V38, V37, V34, V, R1P, R1B, R1A, R13, R, N15!!
57	Schwabe et al. (2013)	unpublished data for Schwabe, A., Suess, K., Storm, C. (2013) What are the long-term effects of livestock grazing in steppic sandy grassland with high conservation value? Results from a 12-year field study. Tuexenia 33: 189-212. Göttingen.	V38, R1P, R1B, R1A, R16, R13, R11, R
69	Schwartz et al. 2021	Schwartz, P., Birkner, L., Velbert, F. & Hölzel, N. (2021) Vielfalt durch extensive Grünlandnutzung. – 30 Jahre Dauermonitoring auf unterschiedlich bewirtschafteten Feuchtgrünlandflächen. Natur in NRW, 1/2021, 16-21. Paderborn. part of unpublished data for: Poptcheva, K., Schwartz, P., Vogel, A., Kleinebecker, T. & Hölzel, N. (2009) Changes in wet meadow vegetation after 20 years of different management in a field experiment (North-West Germany). Agriculture, Ecosystems & Environment, 134 (1-2), 108–114.	R55, R37, R36, R35, R21, R, Q53
61	Sommer & Hachmöller (2001)	Sommer, S. & Hachmöller, B. (2001) Auswertung der Vegetationsaufnahmen von Dauerbeobachtungsflächen auf Bergwiesen im NSG Oelsen bei varierter Mahd im Vergleich zur Brache. Berichte der Arbeitsgemeinschaft sächsischer Botaniker N.F. 18: 99-135	R23!
9	Sperle (unpubl.)		V39, V11, T3K, T12, Sb, S42, S38, R56, R55, R37, R35, R22, R1A, R, Qb, Qa, Q51, Q42, Q41, Q25, Q24, ?
45	Stroh (2013)	Stroh, H.-G. (2013) Wiederholung der vegetationskundlichen Erfassungen an den Dauerbeobachtungsflächen der Kalkmagerrasen des Altendorfer Berges. Bericht für das Jahr 2012. Im Auftrag des Niedersächsischen Landesbetriebs für Wasserwirtschaft Küsten- und Naturschutz, Hannover.	R1A, ?
46	Stroh (2013)	Stroh, H.-G. (2013) Wiederholung der vegetationskundlichen Erfassungen an den	Sa, R1A, ?

		Dauerbeobachtungsflächen der Kalkmagerrasen der Weper und des Gladebergs. Zwischenbericht für das Jahr 2012. Im Auftrag des Niedersächsischen Landesbetriebs für Wasserwirtschaft Küsten- und Naturschutz, Hannover.	
47	Strubelt & Zacharias (2015)	Strubelt, I., Diekmann, M. & Zacharias, D. (2015) Langzeitmonitoring der Vegetation über 52 Jahre im Hartholzauenwald (<i>Querco-Ulmetum minoris</i> Issler 1924) im Haseder Busch (Landkreis Hildesheim). Braunschweiger Geobotanische Arbeiten 11: 173-247	T3M, T1F, T1E, T17, T13, T
66	Strubelt et al. (2019)	Strubelt, I., Diekmann, M., Peppler-Lisbach, C., Gerken, A. & Zacharias, D. (2019) Vegetation changes in the Hasbruch forest nature reserve (NW Germany) depend on management and habitat type. Forest Ecology and Management, 444, 78–88.	T1E, T1B, T18, T17, T13, T
56	Volz (2001)	Volz, H. (2001) Vegetationskundliches Monitoring im NSG Lange Rhön Gebiet Leitgraben. Bearbeitungszeitraum 2001. Gutachten Umweltplanung Volz, Giessen, 62 S.	R57, R35
35	von Oheimb et al. (2006)	v. Oheimb, G., Eischeid, I., Finck, P., Grell, H., Härdtle, W., Mierwald, U., Riecken, U., Sandkühler, J. (2006) Halboffene Weidelandschaft Höltigbaum. Perspektiven für den Erhalt und die naturverträgliche Nutzung von Offenlandlebensräumen. Naturschutz und Biologische Vielfalt 36: 1-280. Landwirtschaftsverlag Münster.	V39, T13, T, R55, R52, R36, R35, R22, R21, R1P, R1M, R, Qb, Q53, Q52, N15, C35a, ?
50	Wagner & Heinken (unpubl.)		T3M, T35, T1H, T1E, T1B, T18, T13, T, S42, R, ?
51	Wagner & Heinken (unpubl.)		T35, T1F, T1E, T1B, T18, T17, T13, T12, T, R55, R51, R1M, R, ?
49	Walther (1986)	Walther, K. (1986) Die Vegetation des Maujahn 1984. Wiederholung der vegetationskundlichen Untersuchung eines wendländischen Moores. Tuexenia 6: 145-193. Göttingen.	T1B, R37, R35, R1P, Q51
52	Wegener (2018)	Wegener, U. (2018) Vegetationswandel des Berggrünlands nach Untersuchungen von 1954 bis 2016. Wege zur Erhaltung der Bergwiesen. Abh. und Ber. aus dem Museum Heineanum 11: 35-101.	R23
53	Wilmanns & Bogenrieder (1988)	Wilmanns, O. & Bogenrieder, A. (1986) Veränderungen der Buchenwälder des Kaiserstuhls im Laufe von vier Jahrzehnten und ihre Interpretation - pflanzensoziologische Tabellen als Dokumente. Abhandlungen aus dem westfälischen Museum für Naturkunde 48(2): 55-80	T18, T, ?

81	Winter (unpubl.)	Winter, R. (2016) Flora und Vegetationsentwicklung der Sukzessionsfläche "Wildnis am Bunker Valentin" in der Bremer Weseraue bei Farge. Bachelorthesis Hochschule Bremen, Studiengang Technische und angewandte Biologie, 61 S.	T, S42, Q51
18	Wittig et al. (2007)	unpublished data for: Wittig, B., Waldmann, T., Diekmann, M. (2007) Veränderungen der Grünlandvegetation im Holtumer Moor über vier Jahrzehnte. <i>Hercynia</i> N.F. 40: 285-300.	V38, V37, V15, V11, V, T16, T15, T13, T12, S92, S41, R55, R52, R37, R36, R35, R22, R21, R, Qa, Q53, Q51, Q22
75	Wittig et al. (2019)	Wittig, B., Müller, J., Mahnke-Ritoff, A. (2019) Talauen-Glatthaferwiesen im Verdener Wesertal (Niedersachsen). <i>Tuexenia</i> 39: 249-265. Göttingen	R22
82	Wittig et al. (2020)	Wittig, B., Müller, J., Quast, R., Miehlich, H. (2020) <i>Arnica montana</i> in Calluna-Heiden auf dem Schießplatz Unterlüß (Niedersachsen). <i>Tuexenia</i> 40: 131-146. Göttingen.	S42, R1M

24

25

26 Extended Data Table 2 | List of all taxa that were harmonized across all projects. The format
 27 of the list follows the rules of the ESy system⁵⁶. The taxon names that were aggregated
 28 below a broader concept name are indented using five blanks. The number to the right
 29 shows the German SL 1.3⁵⁸ number for each taxon.

30	Achillea atrata agg.	18
31	Achillea atrata	19
32	Achillea atrata agg.	18
33	Achillea millefolium agg.	27
34	Achillea millefolium	31
35	Achillea millefolium agg.	27
36	Achillea millefolium subsp. collina	20096
37	Achillea millefolium subsp. millefolium	32
38	Achillea pannonica	34
39	Achillea setacea	36
40	Acinos arvensis	49
41	Acinos arvensis	49
42	Calamintha acinos	976
43	Satureja acinos	23760
44	Aconitum lycoctonum	14242
45	Aconitum lycoctonum	14242
46	Aconitum lycoctonum subsp. vulparia	20209
47	Aconitum vulparia	68
48	Adonis aestivalis	76
49	Adonis aestivalis	76
50	Adonis aestivalis var. citrinus	27247
51	Agrimonia eupatoria	99
52	Agrimonia eupatoria	99
53	Agrimonia eupatoria subsp. eupatoria	100
54	Agrostis canina agg.	120
55	Agrostis canina	121
56	Agrostis coarctata	20180
57	Agrostis stricta	122
58	Agrostis vinealis	20684
59	Agrostis capillaris	20178
60	Agrostis capillaris	20178
61	Agrostis capillaris subsp. oreophila	7086
62	Agrostis tenuis	130
63	Agrostis vulgaris	20179
64	Agrostis stolonifera agg.	127
65	Agrostis alba var. stolonifera	27181
66	Agrostis gigantea	128
67	Agrostis stolonifera	129
68	Agrostis stolonifera agg.	127
69	Agrostis stolonifera subsp. stolonifera	6550
70	Aira caryophyllea	26920
71	Aira caryophyllea	26920
72	Aira caryophyllea subsp. caryophyllea	20218
73	Alchemilla conjuncta agg.	154
74	Alchemilla conjuncta agg.	154
75	Alchemilla hoppeana	166
76	Alchemilla nitida	10027
77	Alchemilla fissa agg.	177
78	Alchemilla fallax	179
79	Alchemilla fissa	180
80	Alchemilla fissa agg.	177
81	Alchemilla hybrida agg.	190
82	Alchemilla flabellata	194
83	Alchemilla glaucescens	195
84	Alchemilla hybrida agg.	190
85	Alchemilla mollis	200

86	<i>Alchemilla acutiloba</i> var. <i>mollis</i>	20237
87	<i>Alchemilla vulgaris</i> agg.	209
88	<i>Alchemilla acutiloba</i> var. <i>stellata</i>	11694
89	<i>Alchemilla crinita</i>	219
90	<i>Alchemilla glabra</i>	235
91	<i>Alchemilla micans</i>	20278
92	<i>Alchemilla monticola</i>	245
93	<i>Alchemilla subcrenata</i>	265
94	<i>Alchemilla vulgaris</i>	26467
95	<i>Alchemilla vulgaris</i> agg.	209
96	<i>Alchemilla vulgaris</i> auct.	20289
97	<i>Alchemilla xanthochlora</i>	273
98	<i>Alisma plantago-aquatica</i> agg.	275
99	<i>Alisma gramineum</i>	276
100	<i>Alisma lanceolatum</i>	277
101	<i>Alisma plantago-aquatica</i>	278
102	<i>Alisma plantago-aquatica</i> agg.	275
103	<i>Allium scorodoprasum</i>	12756
104	<i>Allium rotundum</i>	310
105	<i>Allium scorodoprasum</i>	12756
106	<i>Allium scorodoprasum</i> subsp. <i>rotundum</i>	20326
107	<i>Allium senescens</i>	6583
108	<i>Allium montanum</i>	294
109	<i>Allium senescens</i>	6583
110	<i>Allium senescens</i> subsp. <i>montanum</i>	20321
111	<i>Allium senescens</i> var. <i>montanum</i>	12712
112	<i>Alopecurus pratensis</i> agg.	334
113	<i>Alopecurus pratensis</i>	336
114	<i>Alopecurus pratensis</i> agg.	334
115	<i>Alyssum alyssoides</i>	345
116	<i>Alyssum alyssoides</i>	345
117	<i>Alyssum calycinum</i>	20351
118	<i>Alyssum montanum</i>	350
119	<i>Alyssum montanum</i>	350
120	<i>Alyssum montanum</i> subsp. <i>gmelinii</i>	26457
121	<i>Alyssum montanum</i> subsp. <i>montanum</i>	26456
122	<i>Anagallis arvensis</i>	393
123	<i>Anagallis arvensis</i>	393
124	<i>Anagallis arvensis</i> subsp. <i>arvensis</i>	394
125	<i>Anemone narcissiflora</i>	434
126	<i>Anemonastrum narcissiflorum</i>	20423
127	<i>Anemone narcissiflora</i>	434
128	<i>Anthoxanthum aristatum</i>	20071
129	<i>Anthoxanthum aristatum</i>	20071
130	<i>Anthoxanthum puelii</i>	466
131	<i>Anthoxanthum odoratum</i> agg.	463
132	<i>Anthoxanthum alpinum</i>	464
133	<i>Anthoxanthum odoratum</i>	465
134	<i>Anthoxanthum odoratum</i> agg.	463
135	<i>Anthriscus sylvestris</i> agg.	469
136	<i>Anthriscus sylvestris</i>	473
137	<i>Anthriscus sylvestris</i> agg.	469
138	<i>Anthriscus sylvestris</i> subsp. <i>alpestris</i>	20460
139	<i>Anthriscus sylvestris</i> subsp. <i>sylvestris</i>	20463
140	<i>Anthyllis vulneraria</i>	477
141	<i>Anthyllis vulneraria</i>	477
142	<i>Anthyllis vulneraria</i> subsp. <i>carpatica</i>	479
143	<i>Anthyllis vulneraria</i> subsp. <i>pseudovulneraria</i>	14783
144	<i>Aphanes inexspectata</i>	494
145	<i>Aphanes inexspectata</i>	494
146	<i>Aphanes microcarpa</i>	6554
147	<i>Aquilegia vulgaris</i> agg.	507
148	<i>Aquilegia atrata</i>	508

149	<i>Aquilegia vulgaris</i>	510
150	<i>Arabis bellidifolia</i>	20511
151	<i>Arabis bellidifolia</i> subsp. <i>bellidifolia</i>	20512
152	<i>Arabis glabra</i>	520
153	<i>Arabis glabra</i>	520
154	<i>Turritis glabra</i>	24405
155	<i>Arabis hirsuta</i> agg.	521
156	<i>Arabis hirsuta</i>	523
157	<i>Arabis nemorensis</i>	20072
158	<i>Arctium minus</i>	13702
159	<i>Arctium minus</i>	13702
160	<i>Arctium minus</i> agg.	547
161	<i>Lappa communis</i>	14348
162	<i>Arenaria serpyllifolia</i> agg.	563
163	<i>Arenaria serpyllifolia</i>	13703
164	<i>Arenaria serpyllifolia</i> agg.	563
165	<i>Armeria maritima</i>	581
166	<i>Armeria elongata</i>	577
167	<i>Armeria maritima</i>	581
168	<i>Armeria maritima</i> agg.	575
169	<i>Armeria maritima</i> subsp. <i>elongata</i>	20584
170	<i>Armeria maritima</i> var. <i>elongata</i>	20590
171	<i>Armeria vulgaris</i>	26026
172	<i>Artemisia campestris</i> agg.	594
173	<i>Artemisia campestris</i>	596
174	<i>Artemisia campestris</i> agg.	594
175	<i>Artemisia campestris</i> subsp. <i>campestris</i>	6466
176	<i>Artemisia vulgaris</i> agg.	616
177	<i>Artemisia vulgaris</i>	618
178	<i>Artemisia vulgaris</i> agg.	616
179	<i>Arum maculatum</i> agg.	622
180	<i>Arum maculatum</i>	625
181	<i>Arum maculatum</i> agg.	622
182	<i>Asplenium ruta-muraria</i>	674
183	<i>Asplenium ruta-muraria</i>	674
184	<i>Asplenium ruta-muraria</i> subsp. <i>ruta-muraria</i>	676
185	<i>Aster lanceolatus</i> agg.	15106
186	<i>Aster lanceolatus</i>	693
187	<i>Aster parviflorus</i>	15085
188	<i>Atriplex prostrata</i> agg.	20706
189	<i>Atriplex calotheca</i>	756
190	<i>Atriplex hastata</i>	20693
191	<i>Atriplex hastata</i> agg.	755
192	<i>Atriplex latifolia</i>	760
193	<i>Atriplex prostrata</i>	762
194	<i>Atriplex sagittata</i>	754
195	<i>Atriplex nitens</i>	20702
196	<i>Aurinia saxatilis</i>	777
197	<i>Alyssum saxatile</i>	354
198	<i>Betonica officinalis</i>	826
199	<i>Betonica officinalis</i>	826
200	<i>Stachys officinalis</i>	24165
201	<i>Betula alba</i> agg.	99014
202	<i>Betula pendula</i>	829
203	<i>Betula verrucosa</i>	20786
204	<i>Betula pubescens</i>	830
205	<i>Betula pubescens</i>	830
206	<i>Betula pubescens</i> subsp. <i>carpatica</i>	6472
207	<i>Betula pubescens</i> subsp. <i>pubescens</i>	6471
208	<i>Bidens frondosa</i>	834
209	<i>Bidens frondosa</i>	834
210	<i>Bidens melanocarpa</i>	20788
211	<i>Bidens tripartita</i>	836

212	<i>Bidens tripartita</i>	836
213	<i>Bidens tripartita</i> subsp. <i>tripartita</i>	838
214	<i>Biscutella laevigata</i>	844
215	<i>Biscutella laevigata</i>	844
216	<i>Biscutella laevigata</i> subsp. <i>kernerii</i>	6652
217	<i>Bistorta officinalis</i>	27744
218	<i>Bistorta officinalis</i>	27744
219	<i>Persicaria bistorta</i>	23029
220	<i>Polygonum bistorta</i>	4420
221	<i>Bistorta vivipara</i>	20801
222	<i>Bistorta vivipara</i>	20801
223	<i>Persicaria vivipara</i>	23045
224	<i>Polygonum viviparum</i>	4437
225	<i>Bolboschoenus maritimus</i>	852
226	<i>Bolboschoenus maritimus</i>	852
227	<i>Schoenoplectus maritimus</i>	23820
228	<i>Bothriochloa ischaemum</i>	854
229	<i>Andropogon ischaemum</i>	20417
230	<i>Bothriochloa ischaemum</i>	854
231	<i>Dichanthium ischaemum</i>	21538
232	<i>Brachypodium pinnatum</i> agg.	862
233	<i>Brachypodium pinnatum</i>	863
234	<i>Brachypodium pinnatum</i> agg.	862
235	<i>Brachypodium rupestre</i>	864
236	<i>Bromus arvensis</i>	883
237	<i>Bromus arvensis</i>	883
238	<i>Bromus arvensis</i> subsp. <i>arvensis</i>	7191
239	<i>Bromus hordeaceus</i> agg.	895
240	<i>Bromus hordeaceus</i>	896
241	<i>Bromus hordeaceus</i> agg.	895
242	<i>Bromus hordeaceus</i> subsp. <i>hordeaceus</i>	897
243	<i>Bromus mollis</i>	901
244	<i>Bromus racemosus</i> agg.	908
245	<i>Bromus commutatus</i>	909
246	<i>Bromus racemosus</i>	910
247	<i>Bromus ramosus</i> agg.	911
248	<i>Bromus benekenii</i>	912
249	<i>Bromus ramosus</i>	913
250	<i>Bromus ramosus</i> agg.	911
251	<i>Callitricha palustris</i> agg.	999
252	<i>Callitricha obtusangula</i>	1003
253	<i>Callitricha palustris</i>	1004
254	<i>Callitricha palustris</i> agg.	999
255	<i>Callitricha stagnalis</i>	1006
256	<i>Campanula rotundifolia</i> agg.	1057
257	<i>Campanula rotundifolia</i>	1072
258	<i>Campanula rotundifolia</i> agg.	1057
259	<i>Campanula scheuchzeri</i>	1073
260	<i>Cardamine bulbifera</i>	20944
261	<i>Cardamine bulbifera</i>	20944
262	<i>Dentaria bulbifera</i>	1896
263	<i>Cardamine heptaphylla</i>	20951
264	<i>Dentaria heptaphyllos</i>	91015
265	<i>Cardamine pratensis</i> agg.	1105
266	<i>Cardamine dentata</i>	20945
267	<i>Cardamine palustris</i>	1109
268	<i>Cardamine pratensis</i>	15133
269	<i>Cardamine pratensis</i> agg.	1105
270	<i>Cardaminopsis arenosa</i>	1114
271	<i>Arabis arenosa</i>	20509
272	<i>Cardaminopsis arenosa</i>	1114
273	<i>Cardaminopsis halleri</i>	1116
274	<i>Arabis halleri</i>	20518

275	Cardaminopsis halleri	1116
276	Cardaminopsis petraea	1117
277	Arabis hispida	20526
278	Carduus nutans agg.	1140
279	Carduus nutans	1143
280	Carduus nutans agg.	1140
281	Carex acuta agg.	26775
282	Carex acuta	20989
283	Carex acuta subsp. tricostata	6685
284	Carex gracilis	1230
285	Carex arenaria agg.	1159
286	Carex arenaria	1160
287	Carex ligerica	1161
288	Carex pseudobrizoides	1162
289	Carex atrata agg.	1164
290	Carex atrata	14018
291	Carex atrata agg.	1164
292	Carex elata	1204
293	Carex elata	1204
294	Carex elata subsp. elata	1205
295	Carex flacca	1216
296	Carex flacca	1216
297	Carex glauca	21026
298	Carex flava agg.	1219
299	Carex demissa	20011
300	Carex flava	1220
301	Carex flava agg.	1219
302	Carex flava var. oederi	25010
303	Carex lepidocarpa	1222
304	Carex oederi	1223
305	Carex serotina	21070
306	Carex tumidicarpa	1225
307	Carex viridula	6693
308	Carex muricata agg.	1249
309	Carex divulsa	1250
310	Carex leersiana	1251
311	Carex muricata	6687
312	Carex muricata agg.	1249
313	Carex muricata subsp. muricata	6689
314	Carex pairae	1252
315	Carex spicata	1253
316	Carex nigra agg.	1254
317	Carex fusca	21024
318	Carex nigra	1256
319	Carex nigra agg.	1254
320	Carex ornithopoda agg.	1262
321	Carex ornithopoda	1263
322	Carex ornithopoda agg.	1262
323	Carex ovalis	21055
324	Carex leporina	1240
325	Carex ovalis	21055
326	Carex vulpina agg.	1301
327	Carex otrubae	1302
328	Carex vulpina	1303
329	Carex vulpina agg.	1301
330	Carex × elytroides	10122
331	Carex × elytroides	10122
332	Carex acuta × nigra	90526
333	Carex × figertii	90596
334	Carex davalliana × dioica	90549
335	Carex × xanthocarpa	10137
336	Carex flava × hostiana	90562
337	Carlina acaulis	1306

338	Carlina acaulis	1306
339	Carlina acaulis subsp. simplex	1308
340	Carlina vulgaris	1312
341	Carlina vulgaris	1315
342	Carlina vulgaris subsp. vulgaris	21094
343	Castanea sativa	1323
344	Castanea sativa	1323
345	Castanea vesca	21097
346	Caucalis platycarpos	1329
347	Caucalis lappula	21108
348	Caucalis platycarpos	1329
349	Centaurea jacea	1347
350	Centaurea angustifolia	21122
351	Centaurea jacea	1347
352	Centaurea jacea agg.	24981
353	Centaurea jacea subsp. amara	21132
354	Centaurea jacea subsp. angustifolia	1348
355	Centaurea nigra	26577
356	Centaurea nemoralis	21138
357	Centaurea nigra	26577
358	Centaurea nigra subsp. nemoralis	1370
359	Centaurea scabiosa	1390
360	Centaurea scabiosa	1390
361	Centaurea scabiosa subsp. scabiosa	1397
362	Centaurea stoebe	25004
363	Centaurea maculosa subsp. rhenana	21134
364	Centaurea rhenana	21146
365	Centaurea stoebe	25004
366	Centaurium erythraea	1406
367	Centaurium erythraea	1406
368	Centaurium erythraea subsp. erythraea	1407
369	Centaurium umbellatum	21156
370	Centaurium littorale	1409
371	Centaurium littorale	1409
372	Centaurium minus	7104
373	Cerastium arvense	1431
374	Cerastium arvense	1431
375	Cerastium arvense subsp. arvense	1432
376	Cerastium fontanum agg.	1449
377	Cerastium caespitosum	21178
378	Cerastium fontanum	1450
379	Cerastium fontanum agg.	1449
380	Cerastium fontanum subsp. triviale	21187
381	Cerastium holosteoides	1451
382	Cerastium triviale	25269
383	Cerastium vulgare	21213
384	Cerastium pumilum agg.	1460
385	Cerastium glutinosum	1461
386	Cerastium pallens	21198
387	Cerastium pumilum	1462
388	Cerastium pumilum agg.	1460
389	Ceratocapnos claviculata	21221
390	Ceratocapnos claviculata	21221
391	Corydalis claviculata	1670
392	Cerinthe glabra	1473
393	Cerinthe alpina	21227
394	Chaerophyllum hirsutum	26947
395	Chaerophyllum hirsutum	26947
396	Chaerophyllum hirsutum agg.	1490
397	Chaerophyllum villarsii	1493
398	Chamaecytisus ratisbonensis	1504
399	Chamaecytisus ratisbonensis	1504
400	Cytisus ratisbonensis	21484

401	<i>Chamaecytisus supinus</i>	1506
402	<i>Chamaecytisus supinus</i>	1506
403	<i>Cytisus supinus</i>	21486
404	<i>Chamaespartium sagittale</i>	1509
405	<i>Chamaespartium sagittale</i>	1509
406	<i>Genista sagittalis</i>	21992
407	<i>Genistella sagittalis</i>	2615
408	<i>Chenopodium album</i> agg.	1514
409	<i>Chenopodium album</i>	1515
410	<i>Chenopodium album</i> agg.	1514
411	<i>Chenopodium strictum</i> subsp. <i>striatiforme</i>	6482
412	<i>Cirsium acaule</i>	1556
413	<i>Cirsium acaule</i>	1556
414	<i>Cirsium acaulon</i>	21327
415	<i>Cirsium heterophyllum</i>	1569
416	<i>Cirsium helenioides</i>	21329
417	<i>Cirsium heterophyllum</i>	1569
418	<i>Cirsium vulgare</i>	1579
419	<i>Cirsium lanceolatum</i>	21330
420	<i>Cirsium vulgare</i>	1579
421	<i>Cirsium × rigens</i>	26068
422	<i>Cirsium acaule</i> × <i>oleraceum</i>	90817
423	<i>Clinopodium vulgare</i>	1593
424	<i>Calamintha clinopodium</i>	978
425	<i>Consolida regalis</i>	1627
426	<i>Consolida regalis</i>	1627
427	<i>Delphinium consolida</i>	21510
428	<i>Convolvulus</i>	60816
429	<i>Convolvulus</i>	60816
430	<i>Convolvulus arvensis</i>	1632
431	<i>Conyza canadensis</i>	1638
432	<i>Conyza canadensis</i>	1638
433	<i>Erigeron canadensis</i>	21687
434	<i>Crataegus</i>	61165
435	<i>Crataegus curvisepala</i>	1697
436	<i>Crataegus laevigata</i> agg.	1701
437	<i>Crataegus laevigata</i>	1701
438	<i>Crataegus lindmanii</i>	1699
439	<i>Crataegus monogyna</i>	1707
440	<i>Crataegus monogyna</i> subsp. <i>monogyna</i>	1708
441	<i>Crataegus monogyna</i> subsp. <i>nordica</i>	1709
442	<i>Crataegus monogyna</i> var. <i>monogyna</i>	90937
443	<i>Crataegus oxyacantha</i>	21400
444	<i>Crataegus rhipidophylla</i>	26677
445	<i>Crataegus × macrocarpa</i>	1705
446	<i>Crataegus × media</i>	50021
447	<i>Crataegus calycina</i>	6733
448	<i>Crepis bocconi</i>	21408
449	<i>Crepis pontana</i>	1735
450	<i>Cruciata laevipes</i>	1766
451	<i>Cruciata laevipes</i>	1766
452	<i>Galium cruciata</i>	21963
453	<i>Cystopteris fragilis</i> agg.	1825
454	<i>Cystopteris alpina</i>	21473
455	<i>Cystopteris fragilis</i>	1827
456	<i>Cytisus nigricans</i>	21483
457	<i>Cytisus nigricans</i>	21483
458	<i>Lembotropis nigricans</i>	3333
459	<i>Cytisus scoparius</i>	1837
460	<i>Cytisus scoparius</i>	1837
461	<i>Sarothamnus scoparius</i>	5245
462	<i>Dactylis glomerata</i> agg.	1842
463	<i>Dactylis</i>	397

464	<i>Dactylis aschersoniana</i>	21487
465	<i>Dactylis glomerata</i>	1843
466	<i>Dactylis glomerata</i> agg.	1842
467	<i>Dactylis glomerata</i> subsp. <i>glomerata</i>	13464
468	<i>Dactylis polygama</i>	1846
469	<i>Dactylorhiza maculata</i> agg.	1852
470	<i>Dactylorhiza fuchsii</i>	1853
471	<i>Dactylorhiza fuchsii</i> × <i>maculata</i>	90986
472	<i>Dactylorhiza maculata</i>	1857
473	<i>Dactylorhiza maculata</i> agg.	1852
474	<i>Dactylorhiza majalis</i> agg.	1861
475	<i>Dactylorhiza majalis</i>	1862
476	<i>Dactylorhiza majalis</i> agg.	1861
477	<i>Dactylorhiza majalis</i> subsp. <i>majalis</i>	1866
478	<i>Dactylorhiza traunsteineri</i>	1871
479	<i>Dactylorhiza</i> × <i>carnea</i>	13760
480	<i>Dactylorhiza maculata</i> × <i>incarnata</i>	90995
481	<i>Danthonia decumbens</i>	1874
482	<i>Danthonia decumbens</i>	1874
483	<i>Danthonia decumbens</i> subsp. <i>decumbens</i>	1876
484	<i>Sieglungia decumbens</i>	24019
485	<i>Daucus carota</i>	1886
486	<i>Daucus</i>	61271
487	<i>Daucus carota</i>	1886
488	<i>Daucus carota</i> subsp. <i>carota</i>	6753
489	<i>Deschampsia cespitosa</i> agg.	1903
490	<i>Deschampsia cespitosa</i>	1904
491	<i>Deschampsia cespitosa</i> agg.	1903
492	<i>Deschampsia flexuosa</i>	20725
493	<i>Aira flexuosa</i>	20226
494	<i>Avenella flexuosa</i>	783
495	<i>Deschampsia flexuosa</i>	20725
496	<i>Dianthus gratianopolitanus</i>	1934
497	<i>Dianthus caesius</i>	21522
498	<i>Dianthus gratianopolitanus</i>	1934
499	<i>Dianthus superbus</i>	1949
500	<i>Dianthus superbus</i>	1949
501	<i>Dianthus superbus</i> subsp. <i>superbus</i>	1952
502	<i>Digitalis grandiflora</i>	1960
503	<i>Digitalis ambigua</i>	21540
504	<i>Dipsacus fullonum</i>	1980
505	<i>Dipsacus fullonum</i>	1980
506	<i>Dipsacus sylvestris</i>	20021
507	<i>Draba aizoides</i> agg.	2000
508	<i>Draba aizoides</i>	2001
509	<i>Draba aizoides</i> agg.	2000
510	<i>Drosera longifolia</i>	21588
511	<i>Drosera anglica</i>	2025
512	<i>Drosera longifolia</i>	21588
513	<i>Dryopteris carthusiana</i> agg.	2030
514	<i>Dryopteris</i> × <i>deweverti</i>	50029
515	<i>Dryopteris carthusiana</i>	2032
516	<i>Dryopteris carthusiana</i> × <i>dilatata</i>	91118
517	<i>Dryopteris carthusiana</i> agg.	2030
518	<i>Dryopteris dilatata</i>	2033
519	<i>Dryopteris expansa</i>	2031
520	<i>Dryopteris spinulosa</i>	21604
521	<i>Dryopteris filix-mas</i> agg. s. l.	94728
522	<i>Dryopteris affinis</i>	2038
523	<i>Dryopteris filix-mas</i>	2037
524	<i>Dryopteris filix-mas</i> agg.	2035
525	<i>Eleocharis palustris</i> agg.	2087
526	<i>Eleocharis palustris</i>	2091

527	<i>Eleocharis palustris</i> agg.	2087
528	<i>Eleocharis uniglumis</i>	2094
529	<i>Scirpus palustris</i>	23856
530	<i>Elymus arenosus</i>	27779
531	<i>Elymus repens</i> subsp. <i>arenosus</i>	6543
532	<i>Elymus athericus</i>	27782
533	<i>Elymus pungens</i>	27905
534	<i>Elymus caninus</i>	20145
535	<i>Agropyron caninum</i>	103
536	<i>Elymus caninus</i>	20145
537	<i>Roegneria canina</i>	23598
538	<i>Elymus repens</i>	27778
539	<i>Agropyron repens</i>	27914
540	<i>Agropyron repens</i> subsp. <i>caesium</i>	6541
541	<i>Elymus repens</i>	27778
542	<i>Elymus repens</i> subsp. <i>repens</i>	27781
543	<i>Elytrigia repens</i>	21639
544	<i>Triticum repens</i>	24393
545	<i>Empetrum nigrum</i> agg.	2103
546	<i>Empetrum nigrum</i>	2105
547	<i>Empetrum nigrum</i> agg.	2103
548	<i>Epilobium angustifolium</i>	2113
549	<i>Chamaenerion angustifolium</i>	21237
550	<i>Epilobium angustifolium</i>	2113
551	<i>Epilobium ciliatum</i>	21642
552	<i>Epilobium adenocaulon</i>	2109
553	<i>Epilobium ciliatum</i>	21642
554	<i>Epilobium tetragonum</i>	2126
555	<i>Epilobium tetragonum</i>	2126
556	<i>Epilobium tetragonum</i> subsp. <i>lamyi</i>	2127
557	<i>Epilobium tetragonum</i> subsp. <i>tetragonum</i>	2128
558	<i>Epipactis atrorubens</i>	2130
559	<i>Epipactis atropurpurea</i>	21653
560	<i>Epipactis atrorubens</i>	2130
561	<i>Epipactis rubiginosa</i>	21664
562	<i>Epipactis helleborine</i> agg.	2131
563	<i>Epipactis helleborine</i>	2134
564	<i>Epipactis latifolia</i>	21660
565	<i>Equisetum fluviatile</i>	2143
566	<i>Equisetum fluviatile</i>	2143
567	<i>Equisetum limosum</i>	21674
568	<i>Equisetum × mildeanum</i>	91305
569	<i>Equisetum pratense</i> × <i>sylvaticum</i>	91302
570	<i>Erica carnea</i>	21680
571	<i>Erica carnea</i>	21680
572	<i>Erica herbacea</i>	2163
573	<i>Erigeron acris</i>	2167
574	<i>Erigeron acris</i>	2167
575	<i>Erigeron acris</i> subsp. <i>acris</i>	2168
576	<i>Erigeron annuus</i>	2178
577	<i>Erigeron annuus</i>	2178
578	<i>Erigeron strigosus</i>	21695
579	<i>Erigeron glabratus</i>	21689
580	<i>Erigeron glabratus</i>	21689
581	<i>Erigeron polymorphus</i>	2186
582	<i>Erodium cicutarium</i> agg.	2197
583	<i>Erodium cicutarium</i>	2199
584	<i>Erodium cicutarium</i> agg.	2197
585	<i>Erophila verna</i>	12356
586	<i>Draba verna</i>	21586
587	<i>Erophila verna</i>	12356
588	<i>Erophila verna</i> agg.	2205
589	<i>Erophila verna</i> subsp. <i>verna</i>	21715

590	<i>Erysimum maschallianum</i>	2229
591	<i>Erysimum durum</i>	21726
592	<i>Erysimum odoratum</i>	2234
593	<i>Erysimum erysimoides</i>	21727
594	<i>Erysimum odoratum</i>	2234
595	<i>Euphorbia verrucosa</i>	2309
596	<i>Euphorbia brittingeri</i>	21740
597	<i>Euphorbia verrucosa</i>	2309
598	<i>Euphrasia officinalis</i>	13816
599	<i>Euphrasia officinalis</i>	13816
600	<i>Euphrasia officinalis</i> subsp. <i>rostkoviana</i>	26254
601	<i>Euphrasia rostkoviana</i>	2332
602	<i>Euphrasia rostkoviana</i> agg.	2329
603	<i>Fallopia convolvulus</i>	2359
604	<i>Fallopia convolvulus</i>	2359
605	<i>Polygonum convolvulus</i>	23228
606	<i>Festuca brevipila</i>	26591
607	<i>Festuca brevipila</i>	26591
608	<i>Festuca duriuscula</i>	21820
609	<i>Festuca trachyphylla</i>	2406
610	<i>Festuca filiformis</i>	21824
611	<i>Festuca capillata</i>	21813
612	<i>Festuca filiformis</i>	21824
613	<i>Festuca tenuifolia</i>	2405
614	<i>Festuca ovina</i> agg.	2385
615	<i>Festuca guestfalica</i> et <i>ovina</i>	91404
616	<i>Festuca guestfalica</i>	6483
617	<i>Festuca lemanii</i>	94730
618	<i>Festuca ovina</i>	2396
619	<i>Festuca ovina</i> s. l.	91415
620	<i>Festuca ovina</i> subsp. <i>ovina</i>	91418
621	<i>Festuca ovina</i> var. <i>firmula</i>	21845
622	<i>Festuca ovina</i> agg.	2385
623	<i>Festuca ovina</i> subsp. <i>glaucha</i>	10352
624	<i>Festuca pallens</i>	2397
625	<i>Festuca cinerea</i>	2387
626	<i>Festuca glauca</i>	2392
627	<i>Festuca glaucina</i>	91400
628	<i>Festuca pallens</i>	2397
629	<i>Festuca pallens</i> * <i>glaucina</i>	91420
630	<i>Festuca pallens</i> subsp. <i>pallens</i>	6792
631	<i>Festuca quadriflora</i>	21852
632	<i>Festuca pumila</i>	2416
633	<i>Festuca quadriflora</i>	21852
634	<i>Festuca rubra</i> agg.	2417
635	<i>Festuca heteromalla</i>	7346
636	<i>Festuca nigrescens</i>	2420
637	<i>Festuca nigrescens</i> subsp. <i>nigrescens</i>	91414
638	<i>Festuca rubra</i>	2421
639	<i>Festuca rubra</i> agg.	2417
640	<i>Festuca rubra</i> subsp. <i>commutata</i>	21856
641	<i>Festuca rubra</i> subsp. <i>fallax</i>	13510
642	<i>Festuca rubra</i> subsp. <i>rubra</i>	2425
643	<i>Festuca rubra</i> var. <i>genuina</i>	21863
644	<i>Festuca trichophylla</i>	2426
645	<i>Festuca rupicola</i>	2402
646	<i>Festuca rupicola</i>	2402
647	<i>Festuca sulcata</i>	27166
648	<i>Festuca valesiaca</i> subsp. <i>sulcata</i>	10354
649	<i>Festuca valesiaca</i>	13712
650	<i>Festuca pulchra</i>	13523
651	<i>Festuca valesiaca</i>	13712
652	<i>Festuca violacea</i> agg.	2441

653	<i>Festuca norica</i>	2443
654	<i>Festuca violacea</i> agg.	2441
655	<i>Filipendula vulgaris</i>	2462
656	<i>Filipendula hexapetala</i>	21900
657	<i>Filipendula vulgaris</i>	2462
658	<i>Frangula alnus</i>	2472
659	<i>Frangula alnus</i>	2472
660	<i>Rhamnus frangula</i>	23535
661	<i>Fumana procumbens</i>	2486
662	<i>Fumana procumbens</i>	2486
663	<i>Fumana vulgaris</i>	21921
664	<i>Galeopsis ladanum</i> agg.	2520
665	<i>Galeopsis angustifolia</i>	2521
666	<i>Galeopsis ladanum</i>	2522
667	<i>Galeopsis tetrahit</i> agg.	2526
668	<i>Galeopsis bifida</i>	2527
669	<i>Galeopsis tetrahit</i>	2528
670	<i>Galeopsis tetrahit</i> agg.	2526
671	<i>Galium aparine</i> agg.	2532
672	<i>Galium aparine</i>	2533
673	<i>Galium aparine</i> agg.	2532
674	<i>Galium spurium</i>	2534
675	<i>Galium glaucum</i>	2542
676	<i>Asperula glauca</i>	20637
677	<i>Galium glaucum</i>	2542
678	<i>Galium mollugo</i> agg.	2548
679	<i>Galium album</i>	2549
680	<i>Galium album</i> subsp. <i>album</i>	2550
681	<i>Galium mollugo</i>	2555
682	<i>Galium mollugo</i> agg.	2548
683	<i>Galium palustre</i>	2564
684	<i>Galium elongatum</i>	2563
685	<i>Galium palustre</i>	2564
686	<i>Galium palustre</i> agg.	2561
687	<i>Galium palustre</i> subsp. <i>elongatum</i>	21974
688	<i>Galium palustre</i> subsp. <i>palustre</i>	21975
689	<i>Galium pusillum</i> agg.	2569
690	<i>Galium anisophyllum</i>	2570
691	<i>Galium pumilum</i>	2572
692	<i>Galium pusillum</i> agg.	2569
693	<i>Galium valdepilosum</i>	2577
694	<i>Galium rotundifolium</i>	2579
695	<i>Galium scabrum</i>	21982
696	<i>Galium saxatile</i>	2585
697	<i>Galium harcynicum</i>	2543
698	<i>Galium saxatile</i>	2585
699	<i>Galium verum</i> agg. s. l.	94733
700	<i>Galium × pomeranicum</i>	2599
701	<i>Galium verum</i>	2600
702	<i>Galium verum</i> agg.	2598
703	<i>Galium verum</i> subsp. <i>verum</i>	24929
704	<i>Galium wirtgenii</i>	2601
705	<i>Gentiana acaulis</i> agg.	91530
706	<i>Gentiana acaulis</i>	2616
707	<i>Gentiana clusii</i>	2624
708	<i>Gentianella ciliata</i>	2656
709	<i>Gentiana ciliata</i>	22041
710	<i>Gentianella ciliata</i>	2656
711	<i>Gentianopsis ciliata</i>	22040
712	<i>Gentianella germanica</i> agg.	2657
713	<i>Gentiana germanica</i>	22044
714	<i>Gentianella aspera</i>	2659
715	<i>Gentianella germanica</i>	2662

716	<i>Gentianella germanica</i> agg.	2657
717	<i>Gentianella germanica</i> subsp. <i>germanica</i>	7352
718	<i>Gentianella lutescens</i>	2665
719	<i>Geum × intermedium</i>	50040
720	<i>Geum rivale</i> × <i>urbanum</i>	91547
721	<i>Glechoma hederacea</i> agg.	2710
722	<i>Glechoma hederacea</i>	2711
723	<i>Glechoma hederacea</i> agg.	2710
724	<i>Globularia punctata</i>	2718
725	<i>Globularia bisnagarica</i>	2718
726	<i>Globularia elongata</i>	22079
727	<i>Globularia punctata</i>	2718
728	<i>Globularia vulgaris</i>	26117
729	<i>Globularia willkommii</i>	22080
730	<i>Glyceria fluitans</i> agg.	2719
731	<i>Glyceria declinata</i>	2720
732	<i>Glyceria fluitans</i>	2721
733	<i>Glyceria fluitans</i> agg.	2719
734	<i>Glyceria notata</i>	26260
735	<i>Helianthemum nummularium</i>	26596
736	<i>Helianthemum nummularium</i>	26596
737	<i>Helianthemum nummularium</i> agg.	2774
738	<i>Helianthemum nummularium</i> subsp. <i>obscurum</i>	22127
739	<i>Helianthemum nummularium</i> subsp. <i>ovatum</i>	22128
740	<i>Helianthemum ovatum</i>	2780
741	<i>Helictotrichon pratense</i>	20728
742	<i>Avena pratensis</i>	20717
743	<i>Avenochloa pratensis</i>	788
744	<i>Avenula pratensis</i>	20732
745	<i>Helictotrichon pratense</i>	20728
746	<i>Helictotrichon pubescens</i>	20729
747	<i>Avena pubescens</i>	20718
748	<i>Avenochloa pubescens</i>	789
749	<i>Avenula pubescens</i>	20733
750	<i>Helictotrichon pubescens</i>	20729
751	<i>Helictotrichon versicolor</i>	20730
752	<i>Avenula versicolor</i>	20734
753	<i>Hepatica nobilis</i>	2815
754	<i>Anemone hepatica</i>	20422
755	<i>Hepatica nobilis</i>	2815
756	<i>Hepatica triloba</i>	10391
757	<i>Herniaria glabra</i>	2832
758	<i>Herniaria glabra</i>	2832
759	<i>Herniaria glabra</i> subsp. <i>glabra</i>	2833
760	<i>Hieracium angustifolium</i>	25621
761	<i>Hieracium glaciale</i>	2885
762	<i>Hieracium caespitosum</i>	2866
763	<i>Pilosella caespitosa</i>	12173
764	<i>Hieracium lactucella</i>	2899
765	<i>Hieracium auricula</i>	2857
766	<i>Hieracium lactucella</i>	2899
767	<i>Pilosella lactucella</i>	12147
768	<i>Hieracium murorum</i>	25659
769	<i>Hieracium murorum</i>	25659
770	<i>Hieracium sylvaticum</i>	2942
771	<i>Hieracium pilosella</i>	2923
772	<i>Hieracium pilosella</i>	2923
773	<i>Pilosella officinarum</i>	14992
774	<i>Hieracium piloselloides</i>	2924
775	<i>Hieracium piloselloides</i>	2924
776	<i>Pilosella piloselloides</i>	12226
777	<i>Hieracium rothianum</i>	25654
778	<i>Hieracium setigerum</i>	12219

779	<i>Hieracium subramosum</i>	10562
780	depr. <i>Hieracium subramosum</i>	94848
781	<i>Hippocrepis emerus</i>	21379
782	<i>Coronilla emerus</i>	1655
783	<i>Hornungia petraea</i>	2993
784	<i>Hornungia petraea</i>	2993
785	<i>Hutchinsia petraea</i>	22188
786	<i>Hypericum maculatum</i> agg.	3027
787	<i>Hypericum × desetangssii</i>	3029
788	<i>Hypericum maculatum</i>	20046
789	<i>Hypericum maculatum</i> agg.	3027
790	<i>Hypericum maculatum</i> subsp. <i>maculatum</i>	20047
791	<i>Hypericum perforatum</i>	3036
792	<i>Hypericum perforatum</i>	3036
793	<i>Hypericum perforatum</i> subsp. <i>angustifolium</i>	22208
794	<i>Hypericum tetrapterum</i>	3042
795	<i>Hypericum quadrangulum</i>	22214
796	<i>Hypericum tetrapterum</i>	3042
797	<i>Inula conyzae</i>	22227
798	<i>Inula conyza</i>	3069
799	<i>Inula conyzae</i>	22227
800	<i>Jasione laevis</i>	3117
801	<i>Jasione perennis</i>	22240
802	<i>Juncus alpinus</i>	22243
803	<i>Juncus alpinoarticulatus</i>	91853
804	<i>Juncus alpinus</i>	22243
805	<i>Juncus articulatus</i>	3136
806	<i>Juncus articulatus</i>	3136
807	<i>Juncus supinus</i>	22278
808	<i>Juncus bufonius</i> agg.	3140
809	<i>Juncus bufonius</i>	3141
810	<i>Juncus bufonius</i> agg.	3140
811	<i>Juncus compressus</i> agg.	3147
812	<i>Juncus compressus</i>	3148
813	<i>Juncus gerardii</i>	3149
814	<i>Juncus trifidus</i>	26603
815	<i>Juncus monanthos</i>	3165
816	<i>Juncus trifidus</i> subsp. <i>monanthos</i>	22281
817	<i>Juniperus communis</i>	3168
818	<i>Juniperus communis</i>	3168
819	<i>Juniperus communis</i> subsp. <i>communis</i>	3170
820	<i>Kobresia myosuroides</i>	22299
821	<i>Elyna myosuroides</i>	2102
822	<i>Kobresia myosuroides</i>	22299
823	<i>Koeleria pyramidata</i> agg.	3233
824	<i>Koeleria</i>	61398
825	<i>Koeleria cristata</i>	22305
826	<i>Koeleria gracilis</i>	22312
827	<i>Koeleria macrantha</i>	3235
828	<i>Koeleria pyramidata</i>	3236
829	<i>Koeleria pyramidata</i> agg.	3233
830	<i>Lamium galeobdolon</i> agg.	22337
831	<i>Galeobdolon argentatum</i>	21942
832	<i>Galeobdolon luteum</i>	20026
833	<i>Galeobdolon luteum</i> agg.	91498
834	<i>Galeobdolon montanum</i>	20027
835	<i>Lamiastrum galeobdolon</i>	3256
836	<i>Lamiastrum montanum</i>	3257
837	<i>Lamium argentatum</i>	24905
838	<i>Lamium galeobdolon</i>	22338
839	<i>Lamium galeobdolon</i> agg.	22337
840	<i>Lamium montanum</i>	22340
841	<i>Larix decidua</i>	3272

842	<i>Larix decidua</i>	3272
843	<i>Larix europaea</i>	10606
844	<i>Lathyrus linifolius</i>	3301
845	<i>Lathyrus linifolius</i>	3301
846	<i>Lathyrus montanus</i>	22390
847	<i>Ledum palustre</i>	3328
848	<i>Ledum palustre</i>	3328
849	<i>Rhododendron tomentosum</i>	28144
850	<i>Leontodon helveticus</i>	3346
851	<i>Leontodon helveticus</i>	3346
852	<i>Leontodon pyrenaicus</i> subsp. <i>helveticus</i>	7365
853	<i>Leontodon hispidus</i>	3347
854	<i>Leontodon hastilis</i>	25306
855	<i>Leontodon hispidus</i>	3347
856	<i>Leontodon hispidus</i> subsp. <i>hastilis</i>	22413
857	<i>Leontodon hispidus</i> subsp. <i>hispidus</i>	6837
858	<i>Leontodon saxatilis</i>	3354
859	<i>Leontodon saxatilis</i>	3354
860	<i>Leontodon taraxacoides</i>	22424
861	<i>Leucanthemum atratum</i> agg.	3375
862	<i>Leucanthemum atratum</i> agg.	3375
863	<i>Leucanthemum halleri</i>	3378
864	<i>Leucanthemum vulgare</i> agg.	3381
865	<i>Chrysanthemum leucanthemum</i>	21299
866	<i>Leucanthemum ircutianum</i>	3384
867	<i>Leucanthemum maximum</i>	3386
868	<i>Leucanthemum vulgare</i>	3387
869	<i>Leucanthemum vulgare</i> agg.	3381
870	<i>Linaria vulgaris</i> agg.	3429
871	<i>Linaria vulgaris</i>	3433
872	<i>Linaria vulgaris</i> agg.	3429
873	<i>Lindernia dubia</i>	3434
874	<i>Gratiola anagallidea</i>	13832
875	<i>Linum perenne</i> agg.	3454
876	<i>Linum austriacum</i>	3455
877	<i>Linum leonii</i>	3458
878	<i>Linum perenne</i>	26993
879	<i>Lithospermum arvense</i>	3475
880	<i>Buglossoides arvensis</i> subsp. <i>sibthorpiana</i>	20947
881	<i>Lithospermum arvense</i>	3475
882	<i>Lotus corniculatus</i> agg.	3505
883	<i>Lotus corniculatus</i>	3508
884	<i>Lotus corniculatus</i> agg.	3505
885	<i>Lotus corniculatus</i> subsp. <i>corniculatus</i>	22509
886	<i>Lotus corniculatus</i> var. <i>ciliatus</i>	25841
887	<i>Lotus tenuis</i>	3510
888	<i>Lotus pedunculatus</i>	22517
889	<i>Lotus pedunculatus</i>	22517
890	<i>Lotus uliginosus</i>	3516
891	<i>Luzula campestris</i> agg.	3530
892	<i>Luzula campestris</i>	3531
893	<i>Luzula campestris</i> agg.	3530
894	<i>Luzula campestris</i> subsp. <i>multiflora</i>	22525
895	<i>Luzula multiflora</i>	11817
896	<i>Luzula pallens</i>	25388
897	<i>Luzula sudetica</i>	3536
898	<i>Luzula luzuloides</i>	3541
899	<i>Luzula albida</i>	3526
900	<i>Luzula luzuloides</i>	3541
901	<i>Luzula luzuloides</i> subsp. <i>rubella</i>	14424
902	<i>Luzula sylvatica</i>	3547
903	<i>Luzula sylvatica</i>	3547
904	<i>Luzula sylvatica</i> subsp. <i>sylvatica</i>	3549

905	<i>Lycopersicon esculentum</i>	3558
906	<i>Lycopersicon esculentum</i>	3558
907	<i>Solanum lycopersicum</i>	24088
908	<i>Lycopus europaeus</i>	3562
909	<i>Lycopus europaeus</i>	3562
910	<i>Lycopus europaeus</i> subsp. <i>europaeus</i>	3563
911	<i>Malus sylvestris</i> agg.	3580
912	<i>Malus sylvestris</i>	3582
913	<i>Malus sylvestris</i> agg.	3580
914	<i>Matricaria</i>	60765
915	<i>Chamomilla</i>	67025
916	<i>Matricaria</i>	60765
917	<i>Matricaria discoidea</i>	3597
918	<i>Chamomilla suaveolens</i>	21249
919	<i>Matricaria discoidea</i>	3597
920	<i>Matricaria recutita</i>	21248
921	<i>Chamomilla recutita</i>	21247
922	<i>Matricaria recutita</i>	21248
923	<i>Medicago sativa</i> agg.	3616
924	<i>Medicago</i> × <i>varia</i>	3620
925	<i>Medicago falcata</i>	3617
926	<i>Medicago sativa</i>	11820
927	<i>Medicago sativa</i> agg.	3616
928	<i>Melampyrum pratense</i>	3638
929	<i>Melampyrum pratense</i>	3638
930	<i>Melampyrum pratense</i> subsp. <i>pratense</i>	13853
931	<i>Melica ciliata</i> agg.	3643
932	<i>Melica ciliata</i>	3644
933	<i>Melica ciliata</i> var. <i>nebrodensis</i>	13015
934	<i>Melica nutans</i> agg.	3648
935	<i>Melica nutans</i>	3650
936	<i>Melica nutans</i> agg.	3648
937	<i>Melica picta</i>	3651
938	<i>Mentha spicata</i> agg.	3676
939	<i>Mentha longifolia</i>	3677
940	<i>Mentha suaveolens</i>	3680
941	<i>Mercurialis perennis</i> agg.	3691
942	<i>Mercurialis ovata</i>	3692
943	<i>Mercurialis perennis</i>	3694
944	<i>Minuartia hybrida</i>	3720
945	<i>Alsine tenuifolia</i>	25807
946	<i>Minuartia rubra</i>	22674
947	<i>Minuartia fastigiata</i>	3715
948	<i>Minuartia rubra</i>	22674
949	<i>Minuartia verna</i>	26607
950	<i>Minuartia verna</i>	26607
951	<i>Minuartia verna</i> subsp. <i>gerardii</i>	22675
952	<i>Minuartia verna</i> subsp. <i>hercynica</i>	6499
953	<i>Molinia caerulea</i> agg.	3756
954	<i>Molinia</i>	61408
955	<i>Molinia arundinacea</i>	3757
956	<i>Molinia caerulea</i>	3758
957	<i>Monotropa hypopitys</i> agg.	3762
958	<i>Monotropa hypopithecia</i>	3763
959	<i>Monotropa hypopitys</i>	3764
960	<i>Monotropa hypopitys</i> agg.	3762
961	<i>Montia fontana</i>	3765
962	<i>Montia fontana</i>	3765
963	<i>Montia fontana</i> subsp. <i>chondrosperma</i>	3767
964	<i>Mycelis muralis</i>	3781
965	<i>Lactuca muralis</i>	22329
966	<i>Mycelis muralis</i>	3781
967	<i>Myosotis arvensis</i>	3782

968	<i>Myosotis arvensis</i>	3782
969	<i>Myosotis arvensis</i> subsp. <i>arvensis</i>	3783
970	<i>Myosotis discolor</i>	3785
971	<i>Myosotis discolor</i>	3785
972	<i>Myosotis versicolor</i>	22757
973	<i>Myosotis ramosissima</i>	3792
974	<i>Myosotis hispida</i>	22735
975	<i>Myosotis ramosissima</i>	3792
976	<i>Myosotis scorpioides</i> agg.	20051
977	<i>Myosotis caespitosa</i>	22731
978	<i>Myosotis laxa</i>	6500
979	<i>Myosotis laxiflora</i>	3788
980	<i>Myosotis multiflora</i>	12098
981	<i>Myosotis nemorosa</i>	3789
982	<i>Myosotis nemorosa</i> subsp. <i>brevisetacea</i>	22741
983	<i>Myosotis palustris</i>	22743
984	<i>Myosotis palustris</i> agg.	3786
985	<i>Myosotis palustris</i> s. l.	15315
986	<i>Myosotis scorpioides</i>	3791
987	<i>Myosotis scorpioides</i> agg.	20051
988	<i>Myosotis scorpioides</i> subsp. <i>scorpioides</i>	12104
989	<i>Myosotis stricta</i>	3804
990	<i>Myosotis micrantha</i>	22740
991	<i>Myosotis stricta</i>	3804
992	<i>Myosotis sylvatica</i> agg.	3793
993	<i>Myosotis alpestris</i>	3794
994	<i>Myosotis decumbens</i>	3795
995	<i>Myosotis sylvatica</i>	3802
996	<i>Nigritella nigra</i> agg.	3851
997	<i>Nigritella nigra</i>	7167
998	<i>Nigritella nigra</i> agg.	3851
999	<i>Odontites luteus</i>	3863
1000	<i>Euphrasia lutea</i>	21768
1001	<i>Odontites luteus</i>	3863
1002	<i>Orthanthella lutea</i>	3863
1003	<i>Odontites vernus</i> agg.	12023
1004	<i>Odontites ruber</i>	13749
1005	<i>Odontites vernus</i>	3866
1006	<i>Odontites vernus</i> agg.	12023
1007	<i>Odontites vulgaris</i>	3867
1008	<i>Onobrychis viciifolia</i> agg.	3906
1009	<i>Onobrychis arenaria</i>	3907
1010	<i>Onobrychis sativa</i>	93791
1011	<i>Onobrychis viciifolia</i>	3912
1012	<i>Ononis spinosa</i> agg.	3920
1013	<i>Ononis procurrens</i>	22831
1014	<i>Ononis repens</i>	3922
1015	<i>Ononis repens</i> subsp. <i>procurrens</i>	3923
1016	<i>Ononis spinosa</i>	3925
1017	<i>Ononis spinosa</i> agg.	3920
1018	<i>Ononis spinosa</i> subsp. <i>maritima</i>	22837
1019	<i>Ophrys holoserica</i>	3955
1020	<i>Ophrys fuciflora</i>	22852
1021	<i>Ophrys holoserica</i>	3955
1022	<i>Ophrys insectifera</i>	3956
1023	<i>Ophrys insectifera</i>	3956
1024	<i>Ophrys muscifera</i>	22859
1025	<i>Oreopteris limbosperma</i>	22909
1026	<i>Dryopteris montana</i>	21597
1027	<i>Lastrea limbosperma</i>	22377
1028	<i>Oreopteris limbosperma</i>	22909
1029	<i>Thelypteris limbosperma</i>	5910
1030	<i>Orthilia secunda</i>	4054

1031	Orthilia secunda	4054
1032	Pyrola secunda	23440
1033	Oxalis stricta	22973
1034	Oxalis fontana	4065
1035	Oxalis stricta	22973
1036	Oxytropis montana agg.	4080
1037	Oxytropis jacquinii	4084
1038	Oxytropis montana agg.	4080
1039	Papaver dubium	10700
1040	Papaver dubium	10700
1041	Papaver dubium subsp. lecoqii	22999
1042	Pedicularis palustris	4158
1043	Pedicularis palustris	4158
1044	Pedicularis palustris subsp. palustris	13828
1045	Persicaria amphibia	23027
1046	Persicaria amphibia	23027
1047	Polygonum amphibium	4413
1048	Persicaria dubia	11734
1049	Persicaria dubia	11734
1050	Persicaria mitis	23041
1051	Polygonum mite	4429
1052	Persicaria hydropiper	23032
1053	Persicaria hydropiper	23032
1054	Polygonum hydropiper	4422
1055	Persicaria lapathifolia	23034
1056	Persicaria lapathifolia	23034
1057	Persicaria lapathifolia subsp. pallida	13950
1058	Polygonum lapathifolium	4423
1059	Persicaria maculosa	23038
1060	Persicaria maculosa	23038
1061	Polygonum persicaria	4435
1062	Persicaria minor	23039
1063	Persicaria minor	23039
1064	Polygonum minus	4428
1065	Petrorrhagia prolifera agg.	4185
1066	Petrorrhagia prolifera	4187
1067	Tunica prolifera	24402
1068	Petrorrhagia saxifraga	4189
1069	Petrorrhagia saxifraga	4189
1070	Tunica saxifraga	24403
1071	Phegopteris connectilis	23065
1072	Gymnocarpium phegopteris	12299
1073	Phegopteris connectilis	23065
1074	Thelypteris phegopteris	5912
1075	Phleum alpinum agg.	4215
1076	Phleum alpinum	25110
1077	Phleum alpinum agg.	4215
1078	Phleum phleoides	4222
1079	Phleum boehmeri	23073
1080	Phleum phleoides	4222
1081	Phleum pratense agg.	4223
1082	Phleum bertolonii	4224
1083	Phleum pratense	4225
1084	Phleum pratense agg.	4223
1085	Phleum pratense subsp. pratense	27013
1086	Phragmites australis	4229
1087	Phragmites australis	4229
1088	Phragmites communis	23084
1089	Phyteuma orbiculare	26610
1090	Phyteuma orbiculare	26610
1091	Phyteuma orbiculare subsp. orbiculare	24940
1092	Phyteuma spicatum	4264
1093	Phyteuma spicatum	4264

1094	<i>Phyteuma spicatum</i> subsp. <i>occidentale</i>	14758
1095	<i>Phyteuma spicatum</i> subsp. <i>spicatum</i>	4266
1096	<i>Phyteuma spicatum</i> var. <i>coeruleum</i>	23091
1097	<i>Picea abies</i>	4269
1098	<i>Picea abies</i>	4269
1099	<i>Picea excelsa</i>	23106
1100	<i>Picris hieracioides</i>	4274
1101	<i>Picris hieracioides</i>	4274
1102	<i>Picris hieracioides</i> agg.	4272
1103	<i>Pimpinella major</i>	4277
1104	<i>Pimpinella major</i>	4277
1105	<i>Pimpinella major</i> subsp. <i>major</i>	23108
1106	<i>Pimpinella saxifraga</i> agg.	4279
1107	<i>Pimpinella saxifraga</i>	4282
1108	<i>Pimpinella saxifraga</i> agg.	4279
1109	<i>Plantago lanceolata</i>	4320
1110	<i>Plantago lanceolata</i>	4320
1111	<i>Plantago lanceolata</i> subsp. <i>sphaerostachya</i>	23153
1112	<i>Plantago lanceolata</i> var. <i>sphaerostachya</i>	25683
1113	<i>Plantago major</i>	4321
1114	<i>Plantago intermedia</i>	23150
1115	<i>Plantago major</i>	4321
1116	<i>Plantago major</i> subsp. <i>intermedia</i>	4322
1117	<i>Plantago major</i> subsp. <i>major</i>	4323
1118	<i>Plantago maritima</i> agg.	4325
1119	<i>Plantago alpina</i>	4326
1120	<i>Plantago maritima</i>	27766
1121	<i>Plantago strictissima</i>	23161
1122	<i>Plantago media</i> agg.	4332
1123	<i>Plantago media</i>	4333
1124	<i>Plantago media</i> agg.	4332
1125	<i>Poa annua</i> agg.	4343
1126	<i>Poa annua</i>	4344
1127	<i>Poa annua</i> agg.	4343
1128	<i>Poa supina</i>	4345
1129	<i>Poa pratensis</i> agg.	4366
1130	<i>Poa angustifolia</i>	4367
1131	<i>Poa humilis</i>	7372
1132	<i>Poa pratensis</i>	4368
1133	<i>Poa pratensis</i> agg.	4366
1134	<i>Poa pratensis</i> subsp. <i>angustifolia</i>	23193
1135	<i>Poa trivialis</i>	26611
1136	<i>Poa trivialis</i>	26611
1137	<i>Poa trivialis</i> agg.	4372
1138	<i>Poa trivialis</i> subsp. <i>trivialis</i>	7161
1139	<i>Polygala amara</i> agg.	4390
1140	<i>Polygala amara</i>	4391
1141	<i>Polygala amara</i> agg.	4390
1142	<i>Polygala amara</i> subsp. <i>amarella</i>	23205
1143	<i>Polygala amarella</i>	4394
1144	<i>Polygala vulgaris</i>	4405
1145	<i>Polygala oxyptera</i>	23212
1146	<i>Polygala vulgaris</i>	4405
1147	<i>Polygala vulgaris</i> s. str.	10745
1148	<i>Polygala vulgaris</i> subsp. <i>oxyptera</i>	4406
1149	<i>Polygala vulgaris</i> subsp. <i>vulgaris</i>	4407
1150	<i>Polygonatum odoratum</i>	4410
1151	<i>Polygonatum odoratum</i>	4410
1152	<i>Polygonatum officinale</i>	27134
1153	<i>Polygonum aviculare</i> agg.	4415
1154	<i>Polygonum aviculare</i>	4417
1155	<i>Polygonum aviculare</i> agg.	4415
1156	<i>Populus × canadensis</i>	50080

1157	<i>Populus × canadensis</i>	50080
1158	<i>Populus × euamericana</i>	10760
1159	<i>Potentilla argentea</i> agg.	4490
1160	<i>Potentilla argentea</i>	4491
1161	<i>Potentilla argentea</i> agg.	4490
1162	<i>Potentilla erecta</i>	4511
1163	<i>Potentilla erecta</i>	4511
1164	<i>Potentilla tormentilla</i>	23354
1165	<i>Potentilla heptaphylla</i> agg.	4515
1166	<i>Potentilla heptaphylla</i>	4517
1167	<i>Potentilla rubens</i>	23350
1168	<i>Potentilla incana</i>	25757
1169	<i>Potentilla arenaria</i>	4539
1170	<i>Potentilla cinerea</i>	25853
1171	<i>Potentilla cinerea</i> subsp. <i>incana</i>	15377
1172	<i>Potentilla incana</i>	25757
1173	<i>Potentilla × subarenaria</i>	50084
1174	<i>Potentilla palustris</i>	4528
1175	<i>Comarum palustre</i>	21352
1176	<i>Potentilla palustris</i>	4528
1177	<i>Potentilla verna</i> agg.	4538
1178	<i>Potentilla neumanniana</i>	4541
1179	<i>Potentilla pusilla</i>	4542
1180	<i>Potentilla tabernaemontani</i>	20053
1181	<i>Potentilla verna</i>	23355
1182	<i>Potentilla verna</i> agg.	4538
1183	<i>Potentilla verna</i> subsp. <i>vulgaris</i>	28077
1184	<i>Primula veris</i>	4570
1185	<i>Primula officinalis</i>	23378
1186	<i>Primula veris</i>	4570
1187	<i>Primula veris</i> subsp. <i>veris</i>	4575
1188	<i>Primula × digenea</i>	92537
1189	<i>Primula elatior</i> × <i>vulgaris</i>	92534
1190	<i>Primula × media</i>	50085
1191	<i>Primula elatior</i> × <i>veris</i>	92533
1192	<i>Pritzelago alpina</i>	26680
1193	<i>Hutchinsia alpina</i>	3000
1194	<i>Pritzelago alpina</i>	26680
1195	<i>Prunus avium</i>	4582
1196	<i>Cerasus avium</i>	21216
1197	<i>Prunus avium</i>	4582
1198	<i>Prunus avium</i> subsp. <i>avium</i>	24942
1199	<i>Prunus cerasus</i> agg.	4585
1200	<i>Cerasus vulgaris</i>	21220
1201	<i>Prunus cerasus</i>	4586
1202	<i>Prunus mahaleb</i>	4592
1203	<i>Cerasus mahaleb</i>	21219
1204	<i>Prunus mahaleb</i>	4592
1205	<i>Prunus padus</i>	4593
1206	<i>Padus avium</i>	22980
1207	<i>Prunus padus</i>	4593
1208	<i>Prunus spinosa</i> agg.	4597
1209	<i>Prunus × fruticans</i>	4598
1210	<i>Prunus spinosa</i>	4599
1211	<i>Prunus spinosa</i> subsp. <i>fruticans</i>	23396
1212	<i>Pseudolysimachion longifolium</i>	23400
1213	<i>Pseudolysimachion longifolium</i>	23400
1214	<i>Veronica longifolia</i>	6261
1215	<i>Pseudolysimachion spicatum</i>	23404
1216	<i>Pseudolysimachion spicatum</i>	23404
1217	<i>Veronica spicata</i>	6286
1218	<i>Pseudorchis albida</i>	4601
1219	<i>Leucorchis albida</i>	3390

1220	<i>Pseudorchis albida</i>	4601
1221	<i>Pulmonaria officinalis</i> agg.	4636
1222	<i>Pulmonaria obscura</i>	4637
1223	<i>Pulmonaria officinalis</i>	4638
1224	<i>Pulmonaria officinalis</i> agg.	4636
1225	<i>Pulsatilla alpina</i>	15385
1226	<i>Pulsatilla alpina</i> agg.	4642
1227	<i>Pulsatilla vulgaris</i>	26617
1228	<i>Anemone pulsatilla</i>	20428
1229	<i>Anemone pulsatilla</i> subsp. <i>grandis</i>	20429
1230	<i>Pulsatilla vulgaris</i>	26617
1231	<i>Pulsatilla vulgaris</i> agg.	4656
1232	<i>Pulsatilla vulgaris</i> subsp. <i>grandis</i>	23437
1233	<i>Pyrus communis</i> agg.	4669
1234	<i>Pyrus communis</i>	6936
1235	<i>Pyrus communis</i> agg.	4669
1236	<i>Pyrus communis</i> subsp. <i>pyraster</i>	23449
1237	<i>Pyrus pyraster</i>	4671
1238	<i>Quercus petraea</i> agg.	4678
1239	<i>Quercus petraea</i>	4680
1240	<i>Quercus sessiliflora</i>	23468
1241	<i>Quercus robur</i>	4685
1242	<i>Quercus pedunculata</i>	23466
1243	<i>Quercus robur</i>	4685
1244	<i>Ranunculus acris</i> agg.	4689
1245	<i>Ranunculus acris</i>	4690
1246	<i>Ranunculus acris</i> agg.	4689
1247	<i>Ranunculus aquatilis</i> agg.	4697
1248	<i>Ranunculus aquatilis</i>	4698
1249	<i>Ranunculus aquatilis</i> agg.	4697
1250	<i>Ranunculus peltatus</i>	12030
1251	<i>Ranunculus auricomus</i> agg.	4709
1252	<i>Ranunculus auricomus</i>	11978
1253	<i>Ranunculus auricomus</i> agg.	4709
1254	<i>Ranunculus auricomus</i> s. l.	4710
1255	<i>Ranunculus biformis</i>	27241
1256	<i>Ranunculus kochii</i>	10821
1257	<i>Ranunculus rectus</i>	6956
1258	<i>Ranunculus bulbosus</i>	4717
1259	<i>Ranunculus bulbosus</i>	4717
1260	<i>Ranunculus bulbosus</i> subsp. <i>bulbosus</i>	12015
1261	<i>Ranunculus ficaria</i>	4721
1262	<i>Ficaria verna</i>	21893
1263	<i>Ranunculus ficaria</i>	4721
1264	<i>Ranunculus ficaria</i> subsp. <i>bulbilifer</i>	4722
1265	<i>Ranunculus flammula</i> agg.	4726
1266	<i>Ranunculus flammula</i>	4727
1267	<i>Ranunculus flammula</i> agg.	4726
1268	<i>Ranunculus reptans</i>	4728
1269	<i>Ranunculus montanus</i> agg.	4743
1270	<i>Ranunculus breyninus</i>	23477
1271	<i>Ranunculus montanus</i>	4747
1272	<i>Ranunculus montanus</i> agg.	4743
1273	<i>Ranunculus polyanthemos</i> agg.	4758
1274	<i>Ranunculus nemorosus</i>	4759
1275	<i>Ranunculus nemorosus</i> subsp. <i>polyanthemophyllus</i>	23501
1276	<i>Ranunculus polyanthemoides</i>	4760
1277	<i>Ranunculus polyanthemophyllus</i>	4761
1278	<i>Ranunculus polyanthemos</i>	4762
1279	<i>Ranunculus polyanthemos</i> agg.	4758
1280	<i>Raphanus raphanistrum</i> agg.	4778
1281	<i>Raphanus raphanistrum</i>	4780
1282	<i>Raphanus sativus</i>	4781

1283	<i>Rhinanthus angustifolius</i>	23544
1284	<i>Rhinanthus angustifolius</i>	23544
1285	<i>Rhinanthus angustifolius</i> subsp. <i>angustifolius</i>	23545
1286	<i>Rhinanthus angustifolius</i> subsp. <i>grandiflorus</i>	6569
1287	<i>Rhinanthus serotinus</i>	4832
1288	<i>Rhinanthus aristatus</i> agg.	4817
1289	<i>Rhinanthus aristatus</i>	23551
1290	<i>Rhinanthus glacialis</i>	4819
1291	<i>Rhinanthus minor</i>	4828
1292	<i>Alectorolophus minor</i>	20304
1293	<i>Rhinanthus minor</i>	4828
1294	<i>Ribes rubrum</i> agg.	4850
1295	<i>Ribes rubrum</i>	4851
1296	<i>Ribes rubrum</i> agg.	4850
1297	<i>Rosa canina</i> agg. s. l.	94740
1298	<i>Rosa canina</i>	26665
1299	<i>Rosa canina</i> agg.	4872
1300	<i>Rosa canina</i> agg. s. l.	94740
1301	<i>Rosa corymbifera</i>	26666
1302	<i>Rosa obtusifolia</i>	4897
1303	<i>Rosa rubiginosa</i> agg. s. l.	92752
1304	<i>Rosa elliptica</i>	4887
1305	<i>Rosa micrantha</i>	4893
1306	<i>Rosa rubiginosa</i>	4902
1307	<i>Rosa rubiginosa</i> agg. s. l.	92752
1308	<i>Rosa spinosissima</i>	23649
1309	<i>Rosa pimpinellifolia</i>	4899
1310	<i>Rosa tomentosa</i> agg. s. l.	92754
1311	<i>Rosa tomentosa</i>	4912
1312	<i>Rosa tomentosa</i> agg.	4907
1313	<i>Rosa villosa</i> agg.	4908
1314	<i>Rubus canescens</i>	4952
1315	<i>Rubus canescens</i>	4952
1316	<i>Rubus tomentosus</i>	25709
1317	<i>Rubus grabowskii</i>	7078
1318	<i>Rubus thyrsanthus</i>	5056
1319	<i>Rubus latiarcuatus</i>	24749
1320	<i>Rubus vulgaris</i> var. <i>mollis</i>	11374
1321	<i>Rubus montanus</i>	24778
1322	<i>Rubus candicans</i>	4951
1323	<i>Rubus montanus</i>	24778
1324	<i>Rubus plicatus</i>	5024
1325	<i>Rubus affinis</i>	4931
1326	<i>Rubus plicatus</i>	5024
1327	<i>Rubus</i> sect. <i>Rubus</i>	11351
1328	<i>Rubus fruticosus</i> s. l.	11886
1329	<i>Rubus</i> sect. <i>Rubus</i>	11351
1330	<i>Rumex acetosa</i> agg.	92799
1331	<i>Rumex acetosa</i>	5073
1332	<i>Rumex thyrsiflorus</i>	5108
1333	<i>Rumex acetosella</i>	26619
1334	<i>Rumex acetosella</i>	26619
1335	<i>Rumex acetosella</i> agg.	5074
1336	<i>Rumex acetosella</i> subsp. <i>acetosella</i>	26618
1337	<i>Rumex acetosella</i> subsp. <i>tenuifolius</i>	27201
1338	<i>Rumex tenuifolius</i>	5077
1339	<i>Rumex arifolius</i>	20080
1340	<i>Rumex alpestris</i>	5078
1341	<i>Rumex arifolius</i>	20080
1342	<i>Rumex pseudoalpinus</i>	23661
1343	<i>Rumex alpinus</i>	5079
1344	<i>Rumex × pratensis</i>	50156
1345	<i>Rumex × pratensis</i>	50156

1346	Rumex crispus × obtusifolius	92814
1347	Salix fragilis agg.	5167
1348	Salix × rubens	5169
1349	Salix fragilis	5168
1350	Salix myrsinifolia	5185
1351	Salix myrsinifolia	5185
1352	Salix nigricans	23730
1353	Salix repens agg.	27193
1354	Salix repens	27194
1355	Salix repens agg.	27193
1356	Salix repens s. l.	5191
1357	Salix repens subsp. repens	5193
1358	Salix retusa agg.	5196
1359	Salix retusa	5198
1360	Salix retusa agg.	5196
1361	Salix serpillifolia	5199
1362	Salix × multinervis	27195
1363	Salix × multinervis	27195
1364	Salix aurita × cinerea	92857
1365	Salix × smithiana	27196
1366	Salix × smithiana	27196
1367	Salix caprea × viminalis	92873
1368	Salix × wimmeriana	50108
1369	Salix caprea × purpurea	92872
1370	Salsola kali	5208
1371	Salsola kali subsp. tragus	5211
1372	Salsola tragus	23749
1373	Sanguisorba minor	5231
1374	Sanguisorba minor	5231
1375	Sanguisorba minor subsp. minor	5232
1376	Scabiosa columbaria agg.	5337
1377	Scabiosa columbaria	5338
1378	Scabiosa columbaria agg.	5337
1379	Scabiosa lucida	5342
1380	Scleranthus annuus agg.	5376
1381	Scleranthus annuus	5377
1382	Scleranthus annuus agg.	5376
1383	Scleranthus polycarpos	5378
1384	Scleranthus × intermedius	92980
1385	Scleranthus annuus × perennis	92978
1386	Securigera varia	21382
1387	Coronilla varia	1662
1388	Securigera varia	21382
1389	Sedum rupestre agg.	5429
1390	Sedum reflexum	5433
1391	Sedum rupestre	23907
1392	Sedum rupestre subsp. reflexum	23910
1393	Sedum sexangulare	5437
1394	Sedum boloniense	23894
1395	Sedum mite	23901
1396	Sedum sexangulare	5437
1397	Sedum telephium agg.	5440
1398	Hylotelephium maximum	7126
1399	Sedum maximum	5441
1400	Sedum telephium	27746
1401	Sedum telephium agg.	5440
1402	Sedum telephium subsp. maximum	23912
1403	Senecio alpinus	5467
1404	Senecio alpinus	5467
1405	Senecio cordatus	23945
1406	Senecio aquaticus agg.	5468
1407	Senecio aquaticus	5469
1408	Senecio aquaticus agg.	5468

1409	<i>Senecio erraticus</i>	5470
1410	<i>Senecio incanus</i>	5485
1411	<i>Senecio carniolicus</i>	23942
1412	<i>Senecio jacobaea</i>	5494
1413	<i>Jacobaea vulgaris</i>	25789
1414	<i>Senecio jacobaea</i>	5494
1415	<i>Senecio nemorensis</i> agg.	5496
1416	<i>Senecio fuchsii</i>	5498
1417	<i>Senecio hercynicus</i>	5499
1418	<i>Senecio nemorensis</i>	94860
1419	<i>Senecio nemorensis</i> agg.	5496
1420	<i>Senecio ovatus</i>	23947
1421	<i>Senecio ovatus</i> subsp. <i>alpestris</i>	7006
1422	<i>Serratula tinctoria</i>	27047
1423	<i>Serratula tinctoria</i>	27047
1424	<i>Serratula tinctoria</i> agg.	5524
1425	<i>Seseli libanotis</i>	5540
1426	<i>Libanotis montana</i>	22460
1427	<i>Libanotis sibirica</i>	22463
1428	<i>Seseli libanotis</i>	5540
1429	<i>Sesleria varia</i> agg.	5558
1430	<i>Sesleria albicans</i>	23996
1431	<i>Sesleria albicans</i> subsp. <i>albicans</i>	27051
1432	<i>Sesleria albicans</i> var. <i>albicans</i>	15414
1433	<i>Sesleria caerulea</i>	25028
1434	<i>Sesleria caerulea</i> subsp. <i>calcaria</i>	13441
1435	<i>Sesleria varia</i>	24991
1436	<i>Setaria pumila</i>	24008
1437	<i>Setaria glauca</i>	5563
1438	<i>Setaria pumila</i>	24008
1439	<i>Silene</i>	60691
1440	<i>Lychnis</i>	60681
1441	<i>Silene</i>	60691
1442	<i>Silene acaulis</i> agg.	5578
1443	<i>Silene acaulis</i>	5579
1444	<i>Silene acaulis</i> agg.	5578
1445	<i>Silene coronaria</i>	11625
1446	<i>Lychnis coronaria</i>	3551
1447	<i>Silene flos-cuculi</i>	24031
1448	<i>Lychnis flos-cuculi</i>	3552
1449	<i>Silene flos-cuculi</i>	24031
1450	<i>Silene latifolia</i>	7012
1451	<i>Melandrium album</i>	22627
1452	<i>Silene alba</i>	5581
1453	<i>Silene latifolia</i>	7012
1454	<i>Silene latifolia</i> subsp. <i>alba</i>	22546
1455	<i>Silene pratensis</i>	20058
1456	<i>Silene noctiflora</i>	5603
1457	<i>Melandrium noctiflorum</i>	22630
1458	<i>Silene noctiflora</i>	5603
1459	<i>Silene nutans</i> agg.	5604
1460	<i>Silene nutans</i>	5606
1461	<i>Silene nutans</i> agg.	5604
1462	<i>Silene viscaria</i>	11627
1463	<i>Lychnis viscaria</i>	3554
1464	<i>Silene viscaria</i>	11627
1465	<i>Viscaria vulgaris</i>	24569
1466	<i>Silene vulgaris</i>	5629
1467	<i>Silene cucubalus</i>	24029
1468	<i>Silene inflata</i>	25277
1469	<i>Silene vulgaris</i>	5629
1470	<i>Silene vulgaris</i> agg.	20067
1471	<i>Silene vulgaris</i> subsp. <i>vulgaris</i>	5634

1472	<i>Silene vulgaris</i> var. <i>humilis</i>	24046
1473	<i>Soldanella alpicola</i>	24100
1474	<i>Soldanella pusilla</i> s. l.	5678
1475	<i>Solidago virgaurea</i>	5682
1476	<i>Solidago virgaurea</i>	5682
1477	<i>Solidago virgaurea</i> subsp. <i>virgaurea</i>	5684
1478	<i>Sonchus asper</i>	5690
1479	<i>Sonchus asper</i>	5690
1480	<i>Sonchus asper</i> subsp. <i>asper</i>	5691
1481	<i>Sorbus aria</i> agg.	5696
1482	<i>Sorbus aria</i>	5697
1483	<i>Sorbus aria</i> agg.	5696
1484	<i>Sorbus aucuparia</i>	5700
1485	<i>Sorbus aucuparia</i>	5700
1486	<i>Sorbus aucuparia</i> subsp. <i>aucuparia</i>	5701
1487	<i>Spergula pentandra</i> agg.	5730
1488	<i>Spergula morisonii</i>	5731
1489	<i>Spergula vernalis</i>	24154
1490	<i>Spergularia media</i>	5734
1491	<i>Spergularia maritima</i>	24145
1492	<i>Spergularia media</i>	5734
1493	<i>Stellaria alsine</i>	5769
1494	<i>Stellaria alsine</i>	5769
1495	<i>Stellaria uliginosa</i>	5782
1496	<i>Stellaria aquatica</i>	24917
1497	<i>Myosoton aquaticum</i>	3805
1498	<i>Stellaria aquatica</i>	24917
1499	<i>Stellaria media</i> agg.	5774
1500	<i>Stellaria media</i>	5775
1501	<i>Stellaria media</i> agg.	5774
1502	<i>Stellaria pallida</i>	5777
1503	<i>Stellaria nemorum</i>	5778
1504	<i>Stellaria nemorum</i>	5778
1505	<i>Stellaria nemorum</i> subsp. <i>nemorum</i>	5780
1506	<i>Stellaria palustris</i>	11573
1507	<i>Stellaria glauca</i>	24184
1508	<i>Stellaria palustris</i>	11573
1509	<i>Stipa calamagrostis</i>	24197
1510	<i>Achnatherum calamagrostis</i>	47
1511	<i>Stipa pennata</i> agg.	5787
1512	<i>Stipa joannis</i>	5797
1513	<i>Stipa pennata</i>	24203
1514	<i>Stipa pennata</i> var. <i>mediterranea</i>	13411
1515	<i>Stipa pulcherrima</i>	13746
1516	<i>Symphytum officinale</i>	26629
1517	<i>Sympphytum officinale</i>	26629
1518	<i>Sympphytum officinale</i> agg.	5823
1519	<i>Tanacetum corymbosum</i> agg.	5841
1520	<i>Chrysanthemum corymbosum</i>	21294
1521	<i>Tanacetum corymbosum</i>	5843
1522	<i>Tanacetum parthenium</i>	5845
1523	<i>Chrysanthemum parthenium</i>	21315
1524	<i>Tanacetum parthenium</i>	5845
1525	<i>Tanacetum vulgare</i>	5846
1526	<i>Chrysanthemum vulgare</i>	21314
1527	<i>Tanacetum vulgare</i>	5846
1528	<i>Taraxacum</i> sect. <i>Alpina et Hamata et Ruderalia</i>	93219
1529	<i>Taraxacum</i> sect. <i>Alpina</i>	7424
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1532	<i>Taraxacum</i> sect. <i>Ruderalia</i>	7434
1533	<i>Taraxacum officinale</i>	14513
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1543	Senecio helenitis	5482
1544	Tephroseris helenitis	23949
1545	Tephroseris integrifolia	23953
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1547	Tephroseris integrifolia	23953
1548	Teucrium scorodonia	5889
1549	Teucrium scorodonia	5889
1550	Teucrium scorodonia subsp. scorodonia	5891
1551	Thalictrum minus agg.	5896
1552	Thalictrum minus	5898
1553	Thalictrum minus agg.	5896
1554	Thalictrum minus subsp. saxatile	5902
1555	Thlaspi caerulescens	5928
1556	Noccaea caerulescens	22785
1557	Thlaspi cepaeifolium	25000
1558	Thlaspi rotundifolium	5940
1559	Thlaspi perfoliatum	5938
1560	Microthlaspi perfoliatum	22667
1561	Thlaspi perfoliatum	5938
1562	Thymus praecox agg.	5955
1563	Thymus praecox	5957
1564	Thymus praecox agg.	5955
1565	Thymus praecox subsp. polytrichus	5959
1566	Thymus praecox subsp. praecox	5960
1567	Thymus serpyllum subsp. praecox	13239
1568	Thymus pulegioides agg.	5964
1569	Thymus pulegioides	5965
1570	Thymus pulegioides agg.	5964
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1572	Thymus serpyllum subsp. chamaedrys	13255
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1574	Thymus angustifolius	24265
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1577	Tragopogon pratensis	7141
1578	Tragopogon minor	5999
1579	Tragopogon orientalis	6000
1580	Tragopogon pratensis	7141
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1582	Tragopogon pratensis subsp. orientalis	24325
1583	Tragopogon pratensis subsp. pratensis	7100
1584	Trichophorum cespitosum	27072
1585	Trichophorum cespitosum	27072
1586	Trichophorum cespitosum subsp. germanicum	24330
1587	Trifolium dubium agg.	6028
1588	Trifolium dubium	6029
1589	Trifolium dubium agg.	6028
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1591	Trifolium pratense	6057
1592	Trifolium pratense	6057
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1595	Matricaria inodora	22000
1596	Matricaria maritima	22580
1597	Matricaria perforata	22003

1598	<i>Tripleurospermum inodorum</i>	20061
1599	<i>Tripleurospermum maritimum</i>	6098
1600	<i>Tripleurospermum perforatum</i>	6097
1601	<i>Ulmus glabra</i>	6137
1602	<i>Ulmus glabra</i>	6137
1603	<i>Ulmus montana</i>	24424
1604	<i>Ulmus scabra</i>	24425
1605	<i>Ulmus minor agg.</i>	6139
1606	<i>Ulmus campestris</i>	24420
1607	<i>Ulmus minor</i>	6140
1608	<i>Utricularia minor agg.</i>	6152
1609	<i>Utricularia minor</i>	6154
1610	<i>Utricularia minor agg.</i>	6152
1611	<i>Vaccinium oxycoccos agg.</i>	6161
1612	<i>Oxycoccus palustris</i>	4070
1613	<i>Vaccinium oxycoccos</i>	6163
1614	<i>Vaccinium oxycoccos agg.</i>	6161
1615	<i>Vaccinium uliginosum</i>	6166
1616	<i>Vaccinium uliginosum</i>	6166
1617	<i>Vaccinium uliginosum agg.</i>	6164
1618	<i>Valeriana officinalis agg.</i>	6177
1619	<i>Valeriana officinalis</i>	6178
1620	<i>Valeriana officinalis agg.</i>	6177
1621	<i>Valeriana pratensis</i>	6179
1622	<i>Valeriana procurrens</i>	6180
1623	<i>Valeriana wallrothii</i>	6183
1624	<i>Veronica agrestis agg.</i>	93432
1625	<i>Veronica agrestis</i>	6226
1626	<i>Veronica persica</i>	6271
1627	<i>Veronica polita</i>	6272
1628	<i>Veronica anagallis-aquatica agg.</i>	6231
1629	<i>Veronica anagallis-aquatica</i>	6232
1630	<i>Veronica anagalloides</i>	6233
1631	<i>Veronica catenata</i>	6234
1632	<i>Veronica austriaca agg.</i>	6237
1633	<i>Veronica austriaca</i>	13751
1634	<i>Veronica austriaca subsp. dentata</i>	24471
1635	<i>Veronica austriaca subsp. teucrium</i>	25019
1636	<i>Veronica prostrata</i>	13869
1637	<i>Veronica teucrium</i>	6243
1638	<i>Veronica chamaedrys</i>	13752
1639	<i>Veronica chamaedrys</i>	13752
1640	<i>Veronica chamaedrys agg.</i>	6248
1641	<i>Veronica hederifolia</i>	26766
1642	<i>Veronica hederifolia</i>	26766
1643	<i>Veronica hederifolia agg.</i>	6257
1644	<i>Veronica hederifolia subsp. hederifolia</i>	7101
1645	<i>Veronica sublobata</i>	6259
1646	<i>Veronica verna agg.</i>	6292
1647	<i>Veronica dillenii</i>	6293
1648	<i>Veronica verna</i>	6294
1649	<i>Veronica verna agg.</i>	6292
1650	<i>Vicia cracca agg.</i>	6301
1651	<i>Vicia cracca</i>	6302
1652	<i>Vicia cracca agg.</i>	6301
1653	<i>Vicia tenuifolia</i>	6306
1654	<i>Vicia sativa agg.</i>	6328
1655	<i>Vicia angustifolia</i>	6329
1656	<i>Vicia sativa</i>	6334
1657	<i>Vicia sativa agg.</i>	6328
1658	<i>Vicia sativa subsp. nigra</i>	24521
1659	<i>Vicia sativa var. nigra</i>	24518
1660	<i>Vicia tetrasperma agg.</i>	6338

1661	<i>Vicia tenuissima</i>	6340
1662	<i>Vicia tetrasperma</i>	6341
1663	<i>Vicia tetrasperma</i> agg.	6338
1664	<i>Vincetoxicum hirundinaria</i> agg.	6349
1665	<i>Cynanchum vincetoxicum</i>	21465
1666	<i>Vincetoxicum hirundinaria</i>	6351
1667	<i>Vincetoxicum officinale</i>	24536
1668	<i>Viola canina</i> agg.	24994
1669	<i>Viola canina</i>	6367
1670	<i>Viola canina</i> subsp. <i>canina</i>	6368
1671	<i>Viola persicifolia</i>	6386
1672	<i>Viola persicifolia</i>	6386
1673	<i>Viola stagnina</i>	6394
1674	<i>Viola sylvatica</i> agg.	94746
1675	<i>Viola × bavarica</i>	7420
1676	<i>Viola reichenbachiana</i>	6390
1677	<i>Viola reichenbachiana</i> × <i>riviniana</i>	93487
1678	<i>Viola riviniana</i>	6391
1679	<i>Viola sylvatica</i>	24559
1680	<i>Viola tricolor</i> agg.	6397
1681	<i>Viola arvensis</i>	6398
1682	<i>Viola arvensis</i> × <i>tricolor</i>	93457
1683	<i>Viola arvensis</i> subsp. <i>arvensis</i>	7208
1684	<i>Viola tricolor</i>	6402
1685	<i>Viola tricolor</i> subsp. <i>tricolor</i>	6405
1686	<i>Viola tricolor</i> var. <i>arvensis</i>	12796
1687	x <i>Festulolium krasanii</i>	93548
1688	<i>Festuca arundinacea</i> × <i>Lolium multiflorum</i>	91392
1689		

1690 Extended Data Table 3 | List of all taxon names that were adapted within projects, in
 1691 addition to the harmonisation across all projects, as shown in Extended Data Table 2.
 1692 PROJECT_ID and Project_Name refer to the project in Extended Data Table 1, RS_PLOT is the
 1693 plot resurvey ID, which identifies the groups of plots compared in time, RELEVNE_NR is the
 1694 plot observation ID in the Turboveg 2 database. Taxon_name_old is the name given by the
 1695 original author, while Taxon_name_new_1 and Taxon_name_new_2 refer to newly assigned
 1696 taxon names. In case of two new names the cover values of the old taxon were equally
 1697 divided among the two new taxa.

PROJECT_ID	Project_Name	RS_Plot	RELEVNE_NR	Taxon_name_id	Taxon_name_ne_w_1	Taxon_name_ne_w_2
	Bode				Cuscuta	
6	(2005)	HEIDE	331	Cuscuta	epithymum	
	Bode				Fagus	
6	(2005)	HEIDE	270	Fagus	sylvatica	
	Bode				Fagus	
6	(2005)	HEIDE	271	Fagus	sylvatica	
	Jandt & Leonhardt				Tragopogon	
23	(unpubl.)	32	138	Tragopogon	pratensis	
	Jandt & Leonhardt				Pulsatilla	
23	(unpubl.)	63	190	Pulsatilla	pratensis	
	Jandt & Leonhardt				Anthericum	
23	(unpubl.)	56	182	Anthericum	ramosum	
	Jandt & Leonhardt				Epipactis	
23	(unpubl.)	80	214	Epipactis	atrorubens	
	Jandt & Leonhardt				Trifolium	
23	(unpubl.)	123	6	Trifolium	repens	
	Jandt & Leonhardt				Epipactis	
23	(unpubl.)	138	36	Epipactis	atrorubens	
	Jandt & Leonhardt				Achillea	
23	(unpubl.)	146	38	Achillea	millefolium	
	Jandt & Leonhardt				agg.	
23	(unpubl.)	146	38	Tragopogon	pratensis	
	Jandt & Leonhardt				Tragopogon	
23	(unpubl.)	205	118	Tragopogon	pratensis	
	Jandt & Leonhardt				Epipactis	
23	(unpubl.)	246	120	Epipactis	atrorubens	
	Jandt & Leonhardt				Epipactis	
23	(unpubl.)	251	128	Epipactis	atrorubens	
	Jandt & Leonhardt				Tragopogon	
23	(unpubl.)	367	166	Tragopogon	pratensis	
	Jandt & Leonhardt				Tragopogon	
23	(unpubl.)	373	176	Tragopogon	pratensis	
	Hagen				Festuca	
16	(1996)	G04	180	Festuca	rupicola	
	Hagen				Pulsatilla	
16	(1996)	G09	171	Anemone	vulgaris	
	Hagen				Pulsatilla	
16	(1996)	G15	183	Anemone	vulgaris	
	Hagen				Epipactis	
16	(1996)	G23	290	Epipactis	atrorubens	

	Hagen				
16	(1996)	Z20	377	Epipactis	<i>Epipactis atrorubens</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	A_CF5	857	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_16A	2014	Orchis	<i>Orchis x hybrida</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_14C	2004	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_16B	2015	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_18A	2026	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_18D	2029	Orchis	<i>Orchis purpurea</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_18E	2030	Orchis	<i>Orchis purpurea</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_19A	2032	Orchis	<i>Orchis purpurea</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_19B	2033	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_19D	2035	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_20A	2038	Platanthera	<i>Platanthera chlorantha</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_20C	2040	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_20D	2041	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_21B	2045	Orchis	<i>Orchis x hybrida</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_21C	2046	Orchis	<i>Orchis purpurea</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_22A	2050	Orchis	<i>Orchis purpurea</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_22B	2051	Platanthera	<i>Platanthera chlorantha</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_22E	2054	Orchis	<i>Orchis militaris</i>
	Heinrich,				
	Marstaller				
	& Voigt				
43	(2012)	CAT_23A	2056	Orchis	<i>Orchis purpurea</i>

	Heinrich, Marstaller & Voigt					
43	(2012)	CAT_23B	2057	Platanthera	Platanthera chlorantha	
	Heinrich, Marstaller & Voigt					
43	(2012)	CAT_28A	2086	Orchis	Orchis militaris	
	Heinrich, Marstaller & Voigt					
43	(2012)	M_CF3	471	Ulmus	Ulmus glabra	
	Heinrich, Marstaller & Voigt					
43	(2012)	M_CF3	495	Ulmus	Ulmus glabra	
	Heinrich, Marstaller & Voigt					
43	(2012)	M_CF3	519	Ulmus	Ulmus glabra	
	Heinrich, Marstaller & Voigt					
43	(2012)	M_CF4	568	Orchis	Orchis x hybrida	
	Heinrich, Marstaller & Voigt					
43	(2012)	M_CF6	570	Orchis	Orchis militaris	
	Horchler (unpubl.)	67	93	Callitriches	Callitriches palustris agg.	
	Horchler (unpubl.)	68	94	Callitriches	Callitriches palustris agg.	
	Hüllbusch et al.					
19	(2016)	MW9	28	Tragopogon	Tragopogon pratensis	
	Kuhn et al. (2011)	4350724.00_5487257 .00	1608	Campanula	Campanula rapunculoides	
	Kuhn et al. (2011)	4365134.20_5531963 .60	1300	Primula	Primula veris	
	Kuhn et al. (2011)	4365183.00_5532063 .00	1301	Campanula	Campanula rapunculoides	
	Kuhn et al. (2011)	4365308.00_5532308 .00	1302	Campanula	Campanula rapunculoides	
	Kuhn et al. (2011)	4377860.10_5547643 .00	1357	Primula	Primula veris	
	Kuhn et al. (2011)	4395362.00_5271657 .00	2024	Primula	Primula elatior	
	Kuhn et al. (2011)	4415784.00_5532303 .00	832	Primula	Primula veris	
	Kuhn et al. (2011)	4420009.00_5433917 .00	3580	Agrimonia	Agrimonia eupatoria	
	Kuhn et al. (2011)	4421750.00_5573096 .00	34	Ajuga	Ajuga reptans	
	Kuhn et al. (2011)	4450612.00_5509207 Meineke & Menge	769	Ajuga	Ajuga reptans	
55	(2010)	DBF 4	16	Alchemilla	Alchemilla vulgaris agg.	
	Meineke & Menge					
55	(2010)	DBF 4	17	Alchemilla	Alchemilla vulgaris agg.	

	Meineke & Menge				
55	(2010)	DBF 4	18	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 4	19	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 8	36	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 8	37	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 8	38	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 8	40	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 10	46	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 10	47	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 10	48	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 10	50	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 11	51	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 11	52	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 11	53	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 11	55	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 13	61	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 13	62	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 13	63	Alchemilla	Alchemilla vulgaris agg.
	Meineke & Menge				
55	(2010)	DBF 13	65	Alchemilla	Alchemilla vulgaris agg.
	Müller				
29	(2002)	K/SH1	119	Thymus	Thymus praecox agg.
	Peppler- Lisbach &				
	Könitz				
32	(2017)	N 86-192	56	Euphorbia stricta	Euphrasia stricta
	Peppler- Lisbach &				
	Könitz				
32	(2017)	N 86-193	57	Euphorbia stricta	Euphrasia stricta
	Raehse				
36	(2001)	M546/50	880	Fragaria	Fragaria vesca

	Schmidt et al.				
33	(unpubl.)	Hünstollen_KF1_37	1825	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_40	1828	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF1_41	1829	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF1_42	1830	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_43	1831	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_44	1832	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_45	1833	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_46	1834	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_47	1835	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF1_48	1836	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_49	1837	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_50	1838	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF1_51	1839	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF2_30	1869	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF2_32	1871	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF2_34	1873	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF2_35	1874	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF2_37	1876	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone ranunculoides
33	(unpubl.)	Hünstollen_KF2_41	1880	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF2_42	1881	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF2_43	1882	Anemone	Anemone nemorosa
	Schmidt et al.				Anemone nemorosa
33	(unpubl.)	Hünstollen_KF2_44	1883	Anemone	Anemone nemorosa
	Strubelt & Zacharias				Campanula trachelium
47	(2015)	8	41	Campanula	

1 **Supplementary Methods**

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