

1 **How to best threshold and validate stacked species assemblages? Community**
2 **optimisation might hold the answer**

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19 **ABSTRACT**

- 20 1. The popularity of species distribution models (SDMs) and the associated stacked
21 species distribution models (S-SDMs), as tools for community ecologists, largely
22 increased in recent years. However, while some consensus was reached about the best
23 methods to threshold and evaluate individual SDMs, little agreement exists on how to
24 best assemble individual SDMs into communities, i.e. how to build and assess S-SDM
25 predictions.
- 26 2. Here, we used published data of insects and plants collected within the same study
27 region to test (1) if the most established thresholding methods to optimize single
28 species prediction are also the best choice for predicting species assemblage
29 composition, or if community-based thresholding can be a better alternative, and (2)
30 whether the optimal thresholding method depends on taxa, prevalence distribution
31 and/or species richness. Based on a comparison of different evaluation approaches we
32 provide guidelines for a robust community cross-validation framework, to use if
33 spatial or temporal independent data are unavailable.
- 34 3. Our results showed that the selection of the “optimal” assembly strategy mostly
35 depends on the evaluation approach rather than taxa, prevalence distribution, regional
36 species pool or species richness. If evaluated with independent data or reliable cross-
37 validation, community-based thresholding seems superior compared to single species
38 optimisation. However, many published studies did not evaluate community
39 projections with independent data, often leading to overoptimistic community
40 evaluation metrics based on single species optimisation.
- 41 4. The fact that most of the reviewed S-SDM studies reported over-fitted community
42 evaluation metrics highlights the importance of developing clear evaluation guidelines

43 for community models. Here, we move a first step in this direction, providing a
44 framework for cross-validation at the community level.

45 INTRODUCTION

46 Past and future environmental changes may not only lead to shifts in species distributions
47 (e.g., Parmesan & Yohe 2003; Thuiller *et al.* 2005; Dullinger *et al.* 2012), but also to changes
48 in species assemblages and interactions (e.g., Van der Putten, Macel & Visser 2010; Nogues-
49 Bravo & Rahbek 2011; Blois *et al.* 2013; Alexander *et al.* 2016). Information about
50 communities, here defined as a taxonomic assemblage of distinct populations of species that
51 co-occur in a given space at a given time (Begon, Harper & Townsend 1996), is therefore
52 essential to make informed decisions for conservation prioritisation (D'Amen *et al.* 2011;
53 Guisan *et al.* 2013; Mateo *et al.* 2013) and to create biodiversity indices (e.g., Essential
54 Biodiversity Variables; Pereira *et al.* 2013) for policy decisions (Fleishman, Noss & Noon
55 2006; Granger *et al.* 2015).

56 Different approaches to model communities are available, using either correlative (e.g.,
57 Ferrier & Guisan 2006; Guisan & Rahbek 2011) or mechanistic techniques (e.g., Kearney &
58 Porter 2009; Mokany & Ferrier 2011), with some predicting only macro-ecological properties
59 such as species richness (e.g., Currie *et al.* 2004; Gotelli *et al.* 2009; Dubuis *et al.* 2011) and
60 others also predicting community composition (see D'Amen *et al.* 2017b for a review). In this
61 study, we focused on correlative approaches based on individual species distribution models
62 (SDMs), as they are the most common technique applied to conservation strategies (Guisan *et al.*
63 *al.* 2013), and to predict future patterns of biodiversity in the face of global change (Nogues-
64 Bravo & Rahbek 2011; D'Amen *et al.* 2017b). Niche-based SDMs quantify the relationship
65 between available species occurrences and different environmental factors to analyse and
66 predict distributional patterns (Guisan & Thuiller 2005; Elith & Leathwick 2009; Guisan,
67 Thuiller & Zimmermann 2017). By additionally stacking individual SDMs (S-SDMs), one
68 can produce spatiotemporal projections of species richness and composition (Ferrier & Guisan
69 2006; Guisan & Rahbek 2011).

70 While there is a vast and now long-standing literature on advances and limitations of single
71 species predictions (e.g., Guisan & Thuiller 2005; Guisan *et al.* 2006; Maggini *et al.* 2006;
72 Elith & Leathwick 2009; Meier *et al.* 2010; Zimmermann *et al.* 2010; Merow *et al.* 2014),
73 studies exploring how to improve community predictions based on aggregated information
74 from individual SDMs emerged more recently (e.g., Mateo *et al.* 2012; Benito, Cayuela &
75 Albuquerque 2013; Cord *et al.* 2014; Mod *et al.* 2015; but see Ferrier *et al.* 2002). A
76 fundamental difference among the proposed solutions is whether to maintain the information
77 on species composition in the final predictions. For instance, the simple sum of probabilities
78 of individual SDM predictions usually gives better estimates of species richness, but the
79 information on species identity is lost (Dubuis *et al.* 2011; Calabrese *et al.* 2014). Therefore,
80 predictions of community composition have mainly been achieved so far by thresholding the
81 individual continuous SDM predictions (e.g., probability or suitability index) to obtain binary
82 maps (Liu, White & Newell 2013) and then stacking the latter at the assemblage level (e.g.,
83 Pottier *et al.* 2013; D'Amen *et al.* 2015; D'Amen, Pradervand & Guisan 2015).

84 There are several examples in the literature of optimizing thresholding methods for single
85 species predictions (e.g., Liu *et al.* 2005; Jimenez-Valverde & Lobo 2007; Freeman & Moisen
86 2008; Liu, White & Newell 2013). These led to a mounting consensus about the most
87 appropriate methods, with the majority of SDM studies published nowadays using either an
88 approach maximising the true skills statistics (Max.TSS) or based on the curve in a receiver
89 operating characteristic plot (Opt.ROC, related to AUC) (see Guisan, Thuiller &
90 Zimmermann 2017; Table S1). However, the threshold selection can strongly influence the
91 reliability of the predicted richness and composition of S-SDMs assemblages (Pineda & Lobo
92 2009; Benito, Cayuela & Albuquerque 2013). It is thus relevant to explore which thresholding
93 approach provides the best performance in assemblage estimates, and if alternatives exist that
94 can improve the assemblage prediction from individual SDMs.

95 Studies focussing on S-SDMs tend to over-predict species richness when based on
96 (thresholded) binary predictions (e.g., Pineda & Lobo 2009; Dubuis *et al.* 2011; Mateo *et al.*
97 2012; Pottier *et al.* 2013; Pouteau *et al.* 2015), with some exceptions (e.g., D'Amen,
98 Pradervand & Guisan 2015; Distler *et al.* 2015). Different factors have been proposed to
99 explain this over-prediction: (1) a statistical bias in thresholding site-level occurrence
100 probabilities for each species (Calabrese *et al.* 2014); (2) the implicit assumption of
101 unsaturated communities not assuming an ecological limit for species numbers in assemblages
102 (environmental carrying capacity; Guisan & Rahbek 2011); (3) the lack of considering
103 different constraints on community composition (i.e., ecological, evolutionary, historical, or
104 biological biodiversity drivers; see Mateo, Mokany & Guisan 2017).

105 The commonly used approach to get binary maps from continuous SDM predictions is to use
106 a species-specific threshold, i.e. each species has a single threshold across all sites ("species
107 threshold", Calabrese *et al.* 2014). Recently, another community-based approach, called
108 probability ranking rule (PRR), was proposed to predict assemblage composition from
109 individual SDMs (D'Amen *et al.* 2015). This method does not require a species-specific
110 threshold, therefore preventing over-prediction, but site-by-site ecological constraints (e.g.,
111 macro-ecological models) are applied to assemblages to predict species richness ("site-
112 threshold").

113 Surprisingly, studies aiming to test and improve S-SDM have used very different approaches
114 to evaluate the predicted assemblages (Cord *et al.* 2014; Hespanhol *et al.* 2015; Pouteau *et al.*
115 2015; Thuiller *et al.* 2015; Zurell *et al.* 2016) and this evaluation aspect of the community
116 modelling procedure has not yet received all the attention it deserves. In most studies,
117 assemblage predictions are not adequately evaluated because the data used for the evaluation
118 were already used for individual model fitting, not allowing anymore a correct cross-
119 validation at the community level. Ideally, the best evaluation method should use spatial or

120 temporal independent data (Elith *et al.* 2006; Guisan, Thuiller & Zimmermann 2017), but if
121 not available, an appropriate cross-validation approach should at least be set up.

122 Here, we used published high-resolution data of insects (butterflies and grasshoppers) and
123 plants (forests and grasslands sites), collected within the same study region to (1) test if the
124 most established thresholding methods for optimal single species prediction (i.e., Max.TSS
125 and Opt.ROC) are also the best choice for species assemblages, (2) investigate if the optimal
126 thresholding method depends on taxa, prevalence distribution (Allouche, Tsoar & Kadmon
127 2006), and/or species richness and (3) provide guidelines for a correct community cross-
128 validation framework, to be used if spatially- or temporally- independent data are unavailable.

129 MATERIALS AND METHODS

130 Community data and environmental variables

131 *Study area*

132 The data on all taxa were collected within the same study area located in the western Swiss
133 Alps of the canton Vaud (Fig. 1; 46°10' to 46°30' N; 6°50' to 7°10' E), covering an area of ca.
134 700 km², with elevation ranging from 375 to 3210 m a.s.l. and forested areas up to 1900 m a.s.l.
135 For centuries, agriculture (farming and pasturing) has maintained grasslands among forests and
136 altered the position of the treeline. The highly variable topography and diverse land use of the
137 study area, in combination with our high-resolution environmental data (25 x 25 m cell size),
138 provide a huge range of complex species-environment relationships to test our modelling
139 framework.

140 *Plant data*

141 The forest data were part of a forest inventory of the canton Vaud conducted between 1988 and
142 2002 (mostly 1990 to 1994) and consisted of 3076 sites. The forest sites were distributed on a
143 400 m grid all across the forested area of the canton and had a circular area of 314 m² (Fig. 1;
144 for details see Hartmann, Fouvy & Horisberger 2009). In total, 703 plant species were recorded,
145 but only 312 (44%) had enough occurrence data (> 20 occurrences) across the dataset for
146 modelling purposes (see Table 1 for more detailed statistics on the datasets).

147 The grassland dataset was collected between 2002 and 2009 following an equal random-
148 stratified sampling of non-forested areas in the study area. In total, 911 vegetation sites of 4 m²
149 were sampled (Fig. 1; for more information see Dubuis *et al.* 2011). A total of 905 plant species
150 were recorded but only the 212 most frequent (>20 occurrences) were selected for modelling
151 (Table 1).

152 To predict the distribution of the plant species we used five environmental variables: growing

153 degree-day (above 0 °C), moisture index over the growing season (difference between
154 precipitation and potential evapotranspiration), the sum of potential solar radiation over the
155 year, slope (in degrees), and topographic position (unit-less, indicating the ridges and valleys).
156 All these variables were at a 25 m resolution and have been shown to be useful predictors for
157 plant species in mountain environments (see Dubuis *et al.* 2011; D'Amen *et al.* 2015; Scherrer
158 *et al.* 2017 for details on predictors).

159 *Insect data*

160 Data on butterflies and grasshoppers were respectively collected in 192 and 202 squares of 50
161 m x 50 m across all the elevational range of the study area (Fig. 1; see Pellissier *et al.* 2012;
162 Pradervand *et al.* 2013, for more information). In total, 131 butterfly and 41 grasshopper
163 species were observed, but due to model limitations only the most common 67 butterfly and
164 20 grasshopper species (≥ 20 occurrences) were considered for modelling (Table 1).

165 For our SDMs we used the same predictors as D'Amen, Pradervand and Guisan (2015): four
166 bioclimatic variables (solar radiation, summer temperature, annual degree-days and annual
167 average number of frost days during the growing season), an index of vegetation productivity,
168 i.e. normalized difference vegetation index (as proxies for trophic resources), and the distance
169 to forest. These variables were selected as they are not highly correlated (< 0.7 ; Dormann *et al.*
170 2013) and considered ecologically important for insects (e.g., Turner, Gatehouse & Corey
171 1987; Hawkins & Porter 2003).

172 **The modelling framework**

173 Our modelling framework used three different S-SDM based community modelling pathways
174 (“single species cross-validation”, “independent data” and “community cross-validation”)
175 representing the most commonly reported practices in the literature (see Fig. 2 and
176 “Evaluating community predictions” section).

177 *Single species modelling, thresholding and evaluation*

178 Individual species models were run by generalised linear models (GLM; McCullagh & Nelder
179 1989), generalised additive models (GAM; Hastie & Tibshirani 1990), random forest (RF;
180 Breiman 2001) and boosted regression trees (BRT; Elith, Leathwick & Hastie 2008). Models
181 for species with more than 50 occurrences were fitted by simple SDMs using all five selected
182 predictors, followed by a weighted (AUC) ensemble forecast (Marmion *et al.* 2009). Species
183 having only between 20 and 50 occurrence records were fitted by an ensemble bivariate
184 approach optimised for rare or under-sampled species (Lomba *et al.* 2010; Breiner *et al.*
185 2015): individual models were calibrated on bivariate combinations of the selected predictors
186 with all four modelling techniques, followed by a consensus forecast from all the resulting
187 “small models” weighted by their AUC scores. We used a repeated split-sample procedure
188 (N=25) for model evaluation, followed by a weighted (AUC) ensemble forecast (across
189 techniques and split-sample runs).

190 The projected probability outputs of the ensemble models were binarised using two
191 thresholding schemes: (1) *species-specific-thresholds* (a single threshold calculated for each
192 species) and (2) *site-specific-thresholds* (differing for each site on the basis of additional
193 community information, i.e. species richness predictions). We selected seven different
194 species-specific-thresholding techniques, which can be classified in four major groups: single-
195 index based, sensitivity and specificity combined, model-building data-only-based, and
196 predicted probability-based (see Table S1; Liu *et al.* 2005; Nenzen & Araujo 2011 for details
197 on classification). As the thresholding techniques showed minimal within-group variance (see
198 Figure S1 and S2), we decided to only present the results for one thresholding technique per
199 group in the main manuscript. The chosen techniques were: Cohen’s Kappa maximization
200 approach (*Max.Kappa*; single-index based), TSS maximization approach (*Max.TSS*,
201 sensitivity and specificity combined), observed prevalence (*Obs.Preval*; model-building data-

202 only-based approach), and average probability approach (*AvgProb*; predicted probability-
203 based approach; for details on techniques see Table S1). In addition, we applied two site-
204 thresholds (community-based approaches) using species richness (SR) predictions in
205 combination with a probability ranking rule (PRR). These methods selected a number of
206 species equal to the predicted SR on the basis of decreasing probabilities of presence
207 calculated by the SDMs (D'Amen *et al.* 2015; D'Amen, Pradervand & Guisan 2015).
208 Therefore, the species with the highest probabilities in a site are selected (considered present)
209 in decreasing order until the SR predicted for the site is reached. The SR predictions were
210 derived by either summing the per site probabilities of individual SDMs, obtaining a
211 prediction of richness for each site (pS-SDM; Dubuis *et al.* 2011) or by a macro-ecological
212 model (MEM; see D'Amen, Pradervand & Guisan 2015 for details), directly modelling the
213 richness of the sites. As results from the two site-thresholds were concordant, we only show
214 here the former (*pS-SDM+PRR*).

215 To evaluate the threshold independent performance of our individual species models, the area
216 under the curve of a Receiver-Operating Characteristic (ROC) plot (AUC; Fielding & Bell
217 1997) was calculated based on a repeated split sampling cross-validation (Thuiller, Georges &
218 Engler 2013). Additionally, based on our independent/cross-validation data we calculated five
219 threshold dependent metrics for each thresholding technique: the overall accuracy (PCC; i.e.
220 proportion of correctly classified presence and absences; Fielding & Bell 1997), sensitivity
221 (proportion of correctly predicted presences), specificity (proportion of correctly predicted
222 absences), the true skill statistic (i.e. [(sensitivity + specificity) - 1]; TSS; Allouche, Tsoar &
223 Kadmon 2006) and Cohen's Kappa (Kappa; i.e., overall accuracy but corrected for chance
224 performance; Cohen 1968).

225 *Evaluating community predictions*

226 All the community predictions were built by stacking binary SDMs of individual species (S-
227 SDMs; Dubuis *et al.* 2011; Guisan & Rahbek 2011). The three modelling pathways (Fig. 2)
228 were identical regarding the modelling procedure for single species, thresholding and
229 community assemblage and only varied in the selection of the data for community calibration
230 and evaluation.

231 - The “single species cross-validation” (SSCV) approach (Fig. 2) has not fully
232 “unused/independent” data for community evaluation (i.e. sites not used for the
233 calibration of any single species). Here, in the process of the cross-validation of all
234 individual SDMs (i.e. across all species), different sites are selected at each resampling
235 iteration and for each species, so that all sites are most likely used in at least one split-
236 sampling run and their information incorporated in the final ensemble model. This
237 approach cannot thus be considered based on fully independent data. The SSCV
238 approach has been to date the most common way to model and evaluate communities
239 predictions based on S-SDMs (Fig. 2; e.g., Dubuis *et al.* 2011; Calabrese *et al.* 2014;
240 D'Amen, Pradervand & Guisan 2015; Distler *et al.* 2015). As no independent data is
241 set aside for community evaluation, this approach usually gets evaluated with all the
242 sites used for calibration. However, to avoid bias in the results due to different
243 numbers of evaluation sites, we evaluated the SSCV approach only on 30% of the
244 available sites (identical to the ID and CCV approach below).

245 - The (spatial or temporal) “independent data” (ID) approach (Fig. 2) starts with two
246 completely independent datasets. One is used for the calibration of the SDMs (i.e.
247 70% of the sites) and the other set is used (only) to evaluate the performance of the
248 community predictions (i.e. 30% of the sites; Fig 2; e.g., Benito, Cayuela &
249 Albuquerque 2013; Pottier *et al.* 2013; Cord *et al.* 2014; D'Amen *et al.* 2015; Zurell *et*
250 *al.* 2016).

251 - The “community cross-validation” (CCV) approach (Fig. 2) uses a repeated split
252 sampling of sites (100 repetitions) dividing the available sites into calibration (70%)
253 and evaluation sets (30%) to perform all the modelling procedure from the single
254 species prediction to the community assembly (Fig. 2). In contrast to the previous ID
255 pathway (above), which only uses one (spatial or temporal) fixed independent
256 evaluation dataset, in the CCV approach all SDMs are fitted at each split-sample
257 iteration using the same training and test sets for all species, thus minimizing the risk
258 of bias in the evaluation data (i.e. if the training and test sets differ across species, as
259 in the ID approach). This repeated cross-validation also allows the
260 estimation/simulation of confidence intervals for community predictions instead of
261 just a single value per community. To our knowledge, no study used this community
262 cross-validation method so far.

263 To compare the community model performance among thresholding techniques and
264 modelling pathways, we calculated eight different community agreement metrics: 1) the
265 deviation of the predicted from the observed species richness (SR.deviation), 2) the
266 proportion of species correctly predicted as present (community sensitivity), 3) the proportion
267 of species correctly predicted as absent (community specificity), 4) community accuracy
268 (PCC; i.e. the percent correctly classified species, present or absent), 5) the community TSS
269 (here measured for a site across all species, rather than for a species across all sites as in
270 single SDM evaluation; Pottier *et al.* 2013) , 6) the community kappa (same as for TSS, for a
271 site across species; Pottier *et al.* 2013), and 7) the Sørensen similarity (Sørensen 1948).

272 *Correlation of single species and community evaluation metrics*

273 For each combination of dataset, modelling pathway and thresholding method ($4 \times 3 \times 9 =$
274 108) we calculated the average evaluation metric for all five single species metrics and all
275 seven community metrics. We then calculated the Spearman correlation of all possible

276 combinations of our five single species and seven community evaluation metrics. The
277 resulting correlation matrix tells us if methods (modelling pathways or thresholding methods)
278 that yield the highest scores in a certain single species metric also yield the highest score in
279 the corresponding community evaluation metric.

280 **RESULTS**

281 **Performance of individual SDMs**

282 As expected the evaluation scores of the individual SDMs were similar to earlier studies
283 published with the same data (D'Amen *et al.* 2015; D'Amen, Pradervand & Guisan 2015;
284 Scherrer *et al.* 2017) and their performance was not affected by the chosen community
285 evaluation approach (Table 1, Table S3). Despite their differences in site SR, prevalence
286 distribution and species pool the average performance of individual SDMs was similar across
287 all taxa (Table 1, Table S3). Additionally, the often reported effect of species prevalence on
288 model performance was only marginal in our study, with rare and common species having
289 similar average model performance within a given taxonomic group (Fig. S3).

290 **Correlation of single species and community evaluation metrics**

291 The correlation between the single species and corresponding community metrics was highest
292 ($\text{cor} > 0.93$; Table 2) for some combinations of metrics based on partial information from the
293 contingency table comparing predictions to observations (i.e. PCC, specificity and sensitivity)
294 and considerably lower for the metrics accounting for all dimensions of the contingency table,
295 such as TSS and Cohen's Kappa ($\text{cor} = 0.73$; Table 2). Correlations between non-
296 corresponding single species and community metrics (i.e. Sørensen and SR deviation) tended
297 to be even lower, with the exception of Kappa versus Sørensen (Table 2).

298 **Species richness and compositional similarity**

299 The deviation in species richness between observed and predicted communities was strongly
300 dependent on the chosen thresholding method (Fig. 3). The thresholding approach that uses
301 the average predicted probability (*AvgProb*) showed the highest amount of over-prediction
302 followed by the combined sensitivity and specificity approach (*Max.TSS*). The other three
303 thresholding methods (*Obs.Preval*, *Max.Kappa* and *pS-SDM+PRR*) performed very similar
304 and showed overall no tendency to over-predict species richness. There were no significant
305 differences between the three modelling pathways for any of the studied taxa (Fig. 3). The
306 absolute number of over-predicted species was strongly related to the average number of
307 species per plot (SR) and therefore differed among the taxa (Fig. 3). However, when corrected
308 for the differences in SR the over-prediction did not significantly vary anymore across taxa.

309 The compositional similarity (Sørensen similarity index) varied significantly both among
310 thresholding techniques and modelling pathways (Fig. 4). The compositional similarity was
311 expectedly always much higher with the “single species cross-validation” (SSCV) pathway
312 compared to the “independent data” (ID) or the “community cross-validation” (CCV)
313 pathways, which both performed similarly. There was also a strong interaction between
314 modelling pathway and thresholding technique. Using the SSCV pathway, thresholding by
315 *Obs.Preval* and by *Max.Kappa* performed better (Fig. 4). However, if independent sites were
316 available for the community evaluations (ID and CCV pathways), the community based
317 approaches (*pS-SDM+PRR*) performed better than the *Obs.Preval* and *Max.Kappa* thresholds
318 (Fig. 4). The similarity between predicted and observed communities was higher in the two
319 insect datasets than in the two plant datasets (Fig. 4), which is likely due to the lower number
320 of insect species compared to plant species modelled. Surprisingly, the most established
321 thresholding methods for single species SDMs based on sensitivity and specificity (i.e.
322 *Max.TSS*, *Opt.ROC* and *SenSpec*; Fig. 4 and Fig. S1 and S2) never ranked highest, as one or

- 323 more of the other thresholding method always ranked above them, both for community
324 composition and for species richness.

325 **DISCUSSION**

326 **Do the most established thresholds for single species work as well for community** 327 **predictions?**

328 In this paper, we asked if the most established methods for single species thresholding are
329 also the optimal choice for making predictions at the community level and if there is a direct
330 link between the individual species predictions and the corresponding community metrics.
331 Our results confirm the existence of such a link for single-index based metrics such as
332 sensitivity, specificity and accuracy. However, these results should be interpreted with caution
333 as maximising sensitivity or specificity can simply be achieved by predicting the species as
334 present or absent (respectively) everywhere. In our study system, most of the modelled
335 species have a low prevalence (i.e. are absent at most sites), thus accuracy (PCC) can often be
336 improved by predicting the species as “absent” nearly everywhere.

337 The two most commonly used community evaluation metrics, Sørensen similarity index and
338 deviation in species richness, were only weakly correlated with most evaluation metrics used
339 for individual species. The most established thresholding methods for individual species
340 predictions (i.e., *Max.TSS*, *Opt.ROC*, *SenSpec*) did show lower performance when applied to
341 community-level predictions. This is likely due to the fact that both TSS and ROC try to find
342 the best trade-off between sensitivity and specificity (Guisan, Thuiller & Zimmermann 2017).
343 As most of the species have a prevalence far below 50% (i.e., are absent in many more sites
344 than present), adding a few more presences might have a big effect on the sensitivity (by
345 increasing the chance of finding the few real presences) but only marginally affects the
346 specificity. By definition, increasing sensitivity also increases TSS, but with the drawback of
347 a slight over-prediction. While this might not matter much on a single species basis, for
348 community-level predictions the over-prediction will accumulate when summing binarised
349 maps across all species, leading to the often observed over-estimation of species richness in S-

350 SDMs (e.g., Pineda & Lobo 2009; Dubuis *et al.* 2011; Mateo *et al.* 2012; Pottier *et al.* 2013;
351 Pouteau *et al.* 2015; Zurell *et al.* 2016). It is important to remark, that in the rare case of an
352 ecosystem mostly comprising of widespread species (i.e., prevalence >50 %) this will turn
353 into the opposite as TSS and ROC will optimise absences leading to an underestimation of
354 species richness. The strength of the over/under prediction bias is therefore linked to the
355 prevalence distribution of the modelled species assemblages. However, in the vast majority of
356 natural systems, both the site SR and the regional species pool are driven by a large number of
357 rare (low prevalence species) compared to a few widespread species (Preston 1948; Magurran
358 & Henderson 2003).

359 The community-based thresholding methods based on the selection of the most probable
360 species (through a probability ranking) up to the predicted site richness (*MEM+PRR*, *pS-*
361 *SDM+PRR*) can overcome this problem, because they are able to constrain species predictions
362 based on a different value of species richness in each site (i.e. making them site-specific
363 thresholding methods). Therefore, these methods prevent over-prediction while still allowing
364 the analyses of species composition. Our results thus support the conclusion that, when the
365 final goal is to optimize community composition, community-thresholding methods are the
366 best option. Yet, as discussed in the next section, two single-species thresholding methods –
367 *maximized Kappa* and *observed prevalence* – also showed good results for predicting
368 communities (close to the community-based approaches). However, as community-based
369 thresholds combine the optimisation of species richness prediction and a probability ranking
370 rule (PRR), they would always select the species with the highest predicted probabilities in
371 each site (D'Amen, Pradervand & Guisan 2015). This could seem logic and straightforward,
372 but there might be a bias when the species in the community have varying prevalence
373 (D'Amen *et al.* 2017a). In fact, the maximum predicted probability is depending on the
374 prevalence of the species, thus the common species will tend to always have greater

375 maximum predicted probabilities than rare species and, as a result, will be considered present
376 an over-proportionate number of time in the final community compositions. This bias will
377 produce high similarity scores (Sørensen index) in the prediction evaluation, as the most
378 common species are correctly predicted in most sites. However, the drawback is that the rarest
379 species will be often omitted in the community predictions, which can be for instance
380 problematic if the final goal of the modelling exercise has conservation implications.

381 **Is there a “best” threshold for community S-SDMs?**

382 We also tested if different methods for binarising community S-SDMs could be superior
383 depending on the taxonomic group, prevalence distribution or species richness. While we
384 observed significant differences between the different groups (i.e. taxa), there is no simple
385 statistical way to assess if these differences are attributable to the biology of the taxa
386 themselves or simply to the differences in site species richness and prevalence distributions.
387 Nevertheless, when we standardized the deviation in species richness by the total number of
388 modelled species (regional species pool), no significant difference was any more visible
389 among the different taxonomic groups. The differences in species richness deviation seem
390 therefore a direct cause of the regional species pool. The same also seems correct for the
391 Sørensen similarity index, as datasets with higher species richness and species pool have
392 lower similarity scores. This likely results from the fact that the more species need to be
393 predicted correctly, the more difficult it becomes to predict the whole communities.

394 A similar ranking of thresholding methods was overall observed across taxonomic group
395 within a given modelling pathway, while among the pathways there were clear shifts in the
396 ranking of thresholding methods: with no independent community evaluation data (SSCV),
397 the *Obs.Preval* and *Max.Kappa* threshold showed superior results, while the pathways using
398 independent community evaluation data (ID and CCV) indicated the community-based
399 thresholding to be superior (*pS-SDM+PRR*). This observation is in line with published

400 literature, where studies not using independent community data usually report a good
401 performance of single species optimisations methods (e.g. D'Amen, Pradervand & Guisan
402 2015; Distler *et al.* 2015; Thuiller *et al.* 2015), while studies using independent data usually
403 have better results using community constraints (e.g. D'Amen *et al.* 2015). Yet, it is
404 remarkable to notice that, although previously much criticized in the literature (e.g.,
405 McPherson, Jetz & Rogers 2004; Allouche, Tsoar & Kadmon 2006), maximized Kappa
406 (together here with the observed prevalence) did indeed perform well as a thresholding
407 method for predicting both single species and communities, being nearly always superior to
408 the sensitivity-specificity thresholding methods supporting earlier findings of Manel,
409 Williams and Ormerod (2001).

410 It is important to notice that the shift in ranking between modelling pathways was likely due
411 to a lower degree of overfitting and therefore a lower decrease in performance when
412 predicting to independent data.

413 **Summing up: How to evaluate community predictions correctly?**

414 Our results show that the “single species cross-validation” approach (SSCV), the most
415 commonly used in the literature to evaluate community predictions (e.g., Dubuis *et al.* 2011;
416 Calabrese *et al.* 2014; Distler *et al.* 2015), yields overoptimistic and thus not fully realistic
417 measures of predictive power. While this approach is usually able to provide satisfying
418 evaluation for single species, as revealed by the cross-validation of individual species runs, it
419 shows a clear degradation of predictions when measured at the level of communities. This
420 occurs likely because “all” sites are used at least once at some stage across all modelling runs
421 of the split-sampling procedure, and thus no observation (or very few in the best cases)
422 remains fully independent (i.e. unused) for the final evaluation at the community level.
423 Additionally, the sets of training sites used at each run differ among the species, making the
424 results not entirely comparable across species.

425 The second approach found in the literature builds on the first one (SSCV; thus including an
426 internal cross-validation evaluation), but uses spatially or temporally independent data (ID)
427 for the assessment (thus an external evaluation), thus (unlike SSCV) using the same set of
428 evaluation sites for all species (e.g., Benito, Cayuela & Albuquerque 2013; Pottier *et al.* 2013;
429 Cord *et al.* 2014). When such independent data are available, this method provides the best
430 possible evaluation, provided that the evaluation data are representative of the area where the
431 models apply. This approach – with both internal and external evaluation - is also the one
432 considered as optimal in James *et al.* (2013), and recently promoted in the field of SDMs by
433 Guisan, Thuiller and Zimmermann (2017).

434 The third approach (CVV), newly presented here, repeats the ID approach a large number of
435 times within a cross-validation procedure at the community-level (no example of this
436 approach known in the literature). By doing this, the risk of bias in the evaluation data,
437 inherent to the selection of a single evaluation data set, is minimized compared to the simple
438 ID approach. Additionally, the repeated cross-validation allows the assessments of uncertainty
439 and confidence intervals around the community predictions' performance metrics. However,
440 as this approach selects the same sites for all species, its application is only possible under
441 specific circumstances. First, all the species data need to be collected in the same sites (i.e.
442 true 'community data'). Second, as this approach leads to an unequal number of
443 presences/absences between different cross-validation runs for the same species, it can lead to
444 models failing for very rare (low sample size) species in some of the cross-validation runs if
445 not enough presence sites are selected in the training set.

446 According to our results and despite the potential limitations we advise the use of the
447 proposed community cross-validation approach (CCV) to evaluate community models in
448 future studies. In fact, we clearly showed that the common practice of evaluating the
449 community predictions on the same dataset used for calibration process (SSCV) leads to

450 overoptimistic estimations of model performance. In the commonest case of unavailability of
451 truly spatial (i.e., different region) or temporal (i.e., different sampling period) independent
452 data, often independent datasets are “created” by randomly splitting the initial dataset in two
453 parts. However, we advocate against this practise and instead promote the community cross-
454 validation approach, which minimizes the artefacts of randomly splitting the initial data and
455 allows the estimation of uncertainty associated with the community evaluation metrics.

456

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464 the SIB Swiss Institute of Bioinformatics.

465 **Authors' contributions**

466 DS and AG conceived the ideas; RF and MD analysed the plant and insect data; DS and RGM
467 developed the modelling framework; DS led the writing and all authors contributed critically
468 to the drafts and gave final approval for publication.

469 **Data Accessibility**

470 A generalised version of the community cross-validation algorithm is available in the ecospat
471 R package (Cola *et al.* 2016) on GitHub (ecospat.CCV;
472 <https://doi.org/10.5281/zenodo.1287805>). All species and environmental data are available on
473 Dryad: <https://doi.org/10.5061/dryad.28d4k> (Grassland species and environmental predictors
474 for plants; Guisan, Dubuis & Vittoz 2011) and <https://doi.org/10.5061/dryad.nf925ps> (forest,
475 insect species and environmental predictors for insects; Guisan *et al.* 2018).

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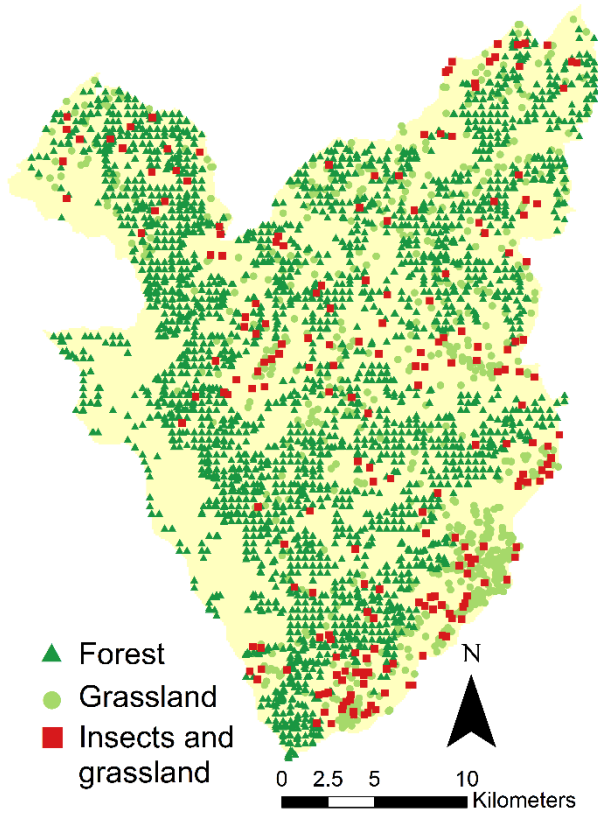
677 **Figure legends**

678 **Figure 1:** Map of the study area with the forested sites (dark green triangles, N=3076), the
679 grassland sites (light green circles and red squares, N=903) and the insect sites (red squares,
680 butterflies N=192, grasshoppers N=202).

681 **Figure 2:** The modelling framework illustrating the three different community modelling
682 approaches: “single species cross-validation” (SSCV), “independent data” (ID) and
683 “community cross-validation” (CCV).

684 **Figure 3:** Deviation in site specific species richness between observations and predictions for
685 the four different datasets (top to bottom) and the three different modelling pathways (left to
686 right). The boxplots are sorted by the median and the colours indicate the different
687 thresholding techniques used to binarise predictions. The line in the box indicates the median,
688 boxes range from the 25th to the 75th percentile and the whiskers indicate ± 2 standard
689 deviations. Letters above the boxplots indicate significant differences (Wilcoxon rank sum
690 test, $p < 0.05$).

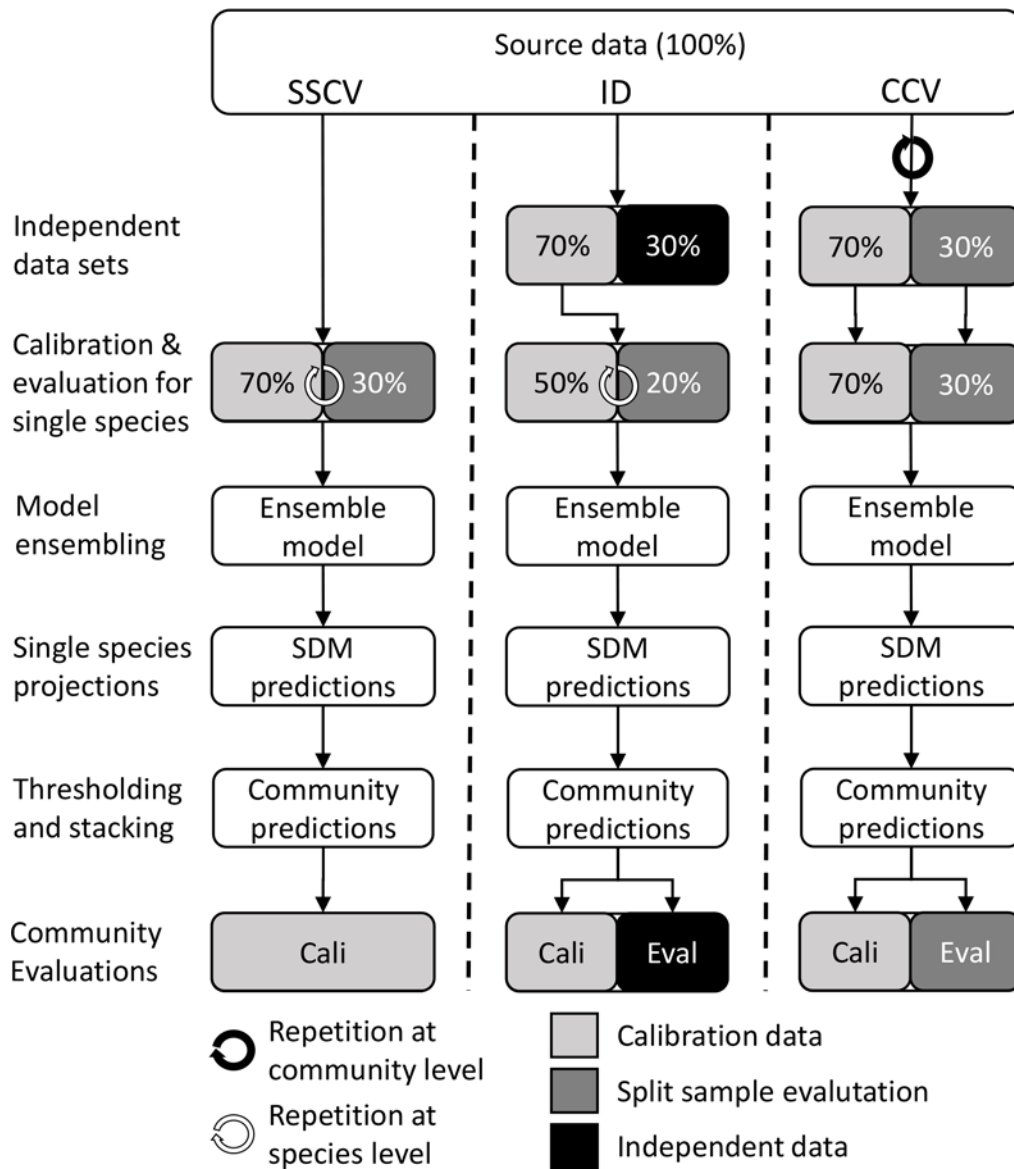
691 **Figure 4:** Sørensen similarity between observations and predictions for the four different
692 datasets (top to bottom) and the three different modelling pathways (left to right). The
693 boxplots are sorted by the median and the colours indicate the different thresholding
694 techniques. The line in the box indicates the median, boxes range from the 25th to the 75th
695 percentile and the whiskers indicate ± 2 standard deviations. Letters above the boxplots
696 indicate significant differences (Wilcoxon rank sum test, $p < 0.05$).



698

699 **Figure 1:** Map of the study area with the forested sites (dark green triangles, N=3076), the
700 grassland sites (light green circles and red squares, N=903) and the insect sites (red squares,
701 butterflies N=192, grasshoppers N=202).

702 **Figure 2**



703

704 **Figure 2:** The modelling framework illustrating the three different community modelling

705 approaches: “single species cross-validation” (SSCV), “independent data” (ID) and

706 “community cross-validation” (CCV).

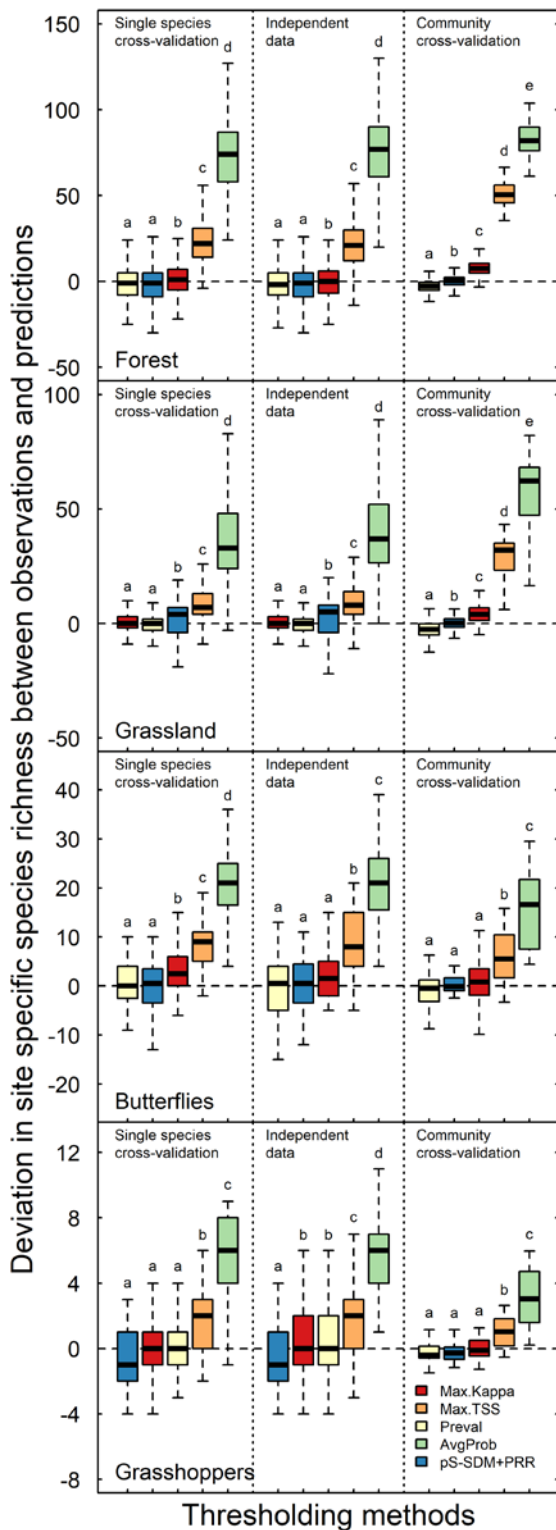


Figure 3: Deviation in site specific species richness between observations and predictions for the four different datasets (top to bottom) and the three different modelling pathways (left to right). The boxplots are sorted by the median and the colours indicate the different thresholding techniques used to binarise predictions. The line in the box indicates the median, boxes range from the 25th to the 75th percentile and the whiskers indicate ± 2 standard deviations. Letters above the boxplots indicate significant differences (Wilcoxon rank sum test, $p < 0.05$).

Figure 4

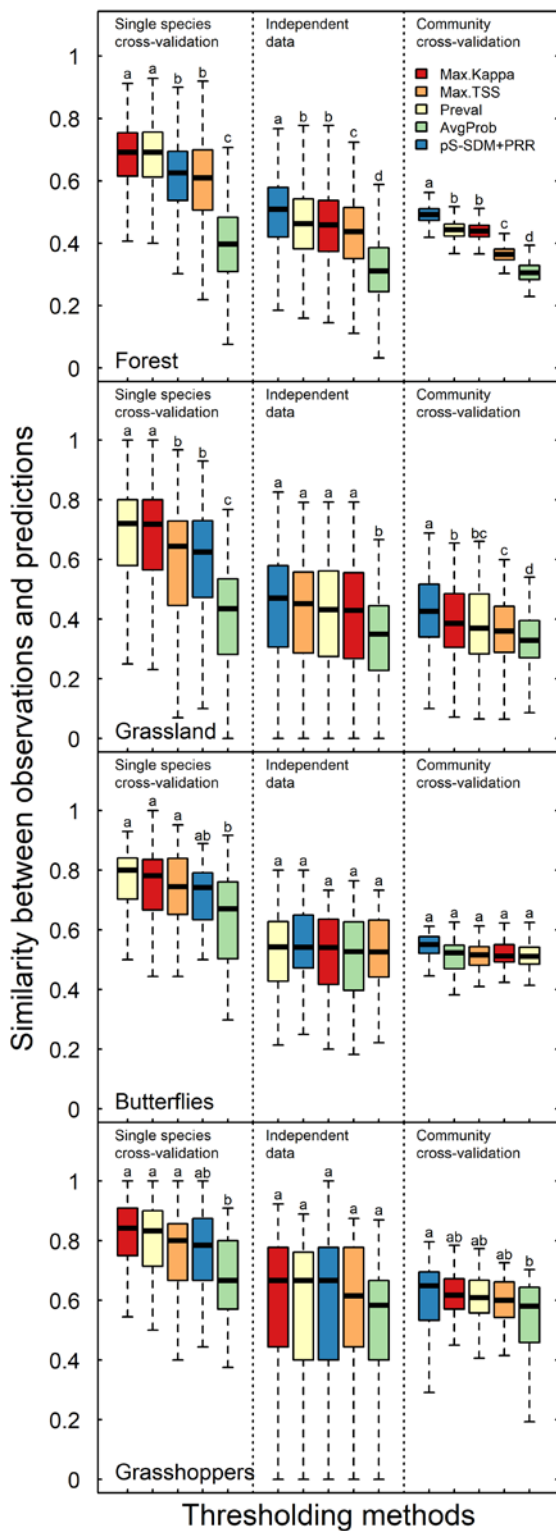


Figure 4: Sørensen similarity between observations and predictions for the four different datasets (top to bottom) and the three different modelling pathways (left to right). The boxplots are sorted by the median and the colours indicate the different thresholding techniques. The line in the box indicates the median, boxes range from the 25th to the 75th percentile and the whiskers indicate ± 2 standard deviations. Letters above the boxplots indicate significant differences (Wilcoxon rank sum test, $p < 0.05$).

Table 1: Basic statistics of the data sets used for the case study and the evaluation metrics (AUC) for the individual species distribution models using the three different community evaluation approaches. SSCV = Single species cross-validation, ID = Independent data, CCV = Community cross-validation

Data set	Number of species modelled (recorded)	Prevalence (mean \pm sd)	Species richness (mean \pm sd)	AUC SSCV (mean \pm sd)	AUC ID (mean \pm sd)	AUC CCV (mean \pm sd)
Forest	312 (703)	0.044 \pm 0.090	29.5 \pm 11.8	0.80 \pm 0.09	0.80 \pm 0.08	0.79 \pm 0.09
Grassland	212 (905)	0.098 \pm 0.089	23.5 \pm 13.8	0.82 \pm 0.07	0.83 \pm 0.06	0.81 \pm 0.06
Butterflies	77 (131)	0.235 \pm 0.137	18.1 \pm 9.2	0.76 \pm 0.10	0.75 \pm 0.12	0.76 \pm 0.10
Grasshoppers	20 (41)	0.256 \pm 0.193	5.1 \pm 3.3	0.84 \pm 0.07	0.86 \pm 0.08	0.84 \pm 0.06

Table 2: Pearson Correlation of single species and community evaluation statistics. The asterisks indicate the significance level. Correlations of the single species evaluation metrics and their corresponding community evaluation metric are highlighted in bold.

Single species	Community metrics						
	Accuracy	Sensitivity	Specificity	KAPPA	TSS	Sørensen similarity	SR deviation
Accuracy	1.00 ***	-0.37 *	0.95 ***	0.70 ***	0.37 *	0.37 *	-0.58 ***
Sensitivity	-0.36 **	0.93 ***	-0.54 ***	0.01 n.s.	0.56 ***	0.18 n.s.	-0.44 ***
Specificity	0.97 ***	-0.53 ***	0.99 ***	0.64 ***	0.20 n.s.	0.31 *	-0.63 ***
KAPPA	0.41 **	0.50 *	0.27 *	0.79 ***	0.72 ***	0.82 ***	-0.3 *
TSS	0.06 n.s.	0.85 ***	-0.14 n.s.	0.35 n.s.	0.79 ***	0.38 **	-0.20 n.s.

The asterisks indicate the significance level (n.s.= not significant, * p<0.05, ** p<0.01, *** p<0.001)

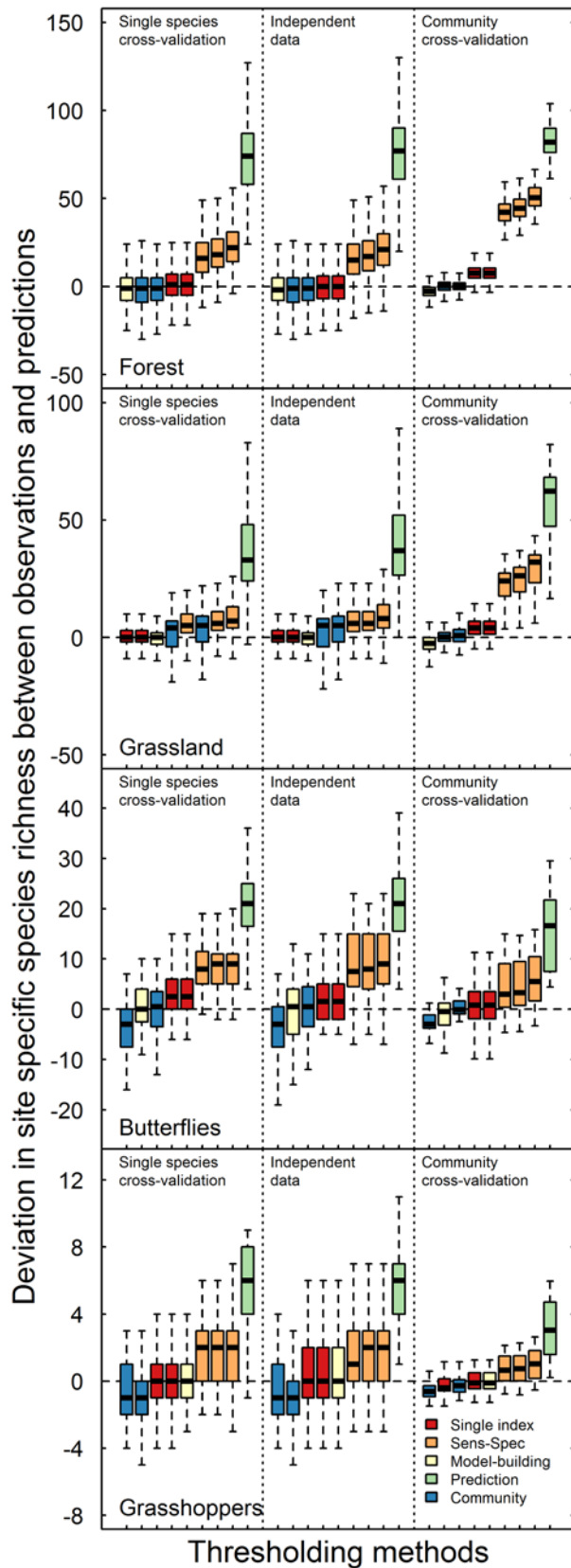


Figure S1: Deviation in site specific species richness between observations and predictions for the four different datasets (top to bottom) and the three different modelling pathways (left to right). The boxplots are sorted by the median and the colours indicate the different thresholding techniques. The line in the box indicates the median, boxes range from the 25th to the 75th percentile and the whiskers indicate ± 2 standard deviations. For details on the method used within each threshold group see Table S1.

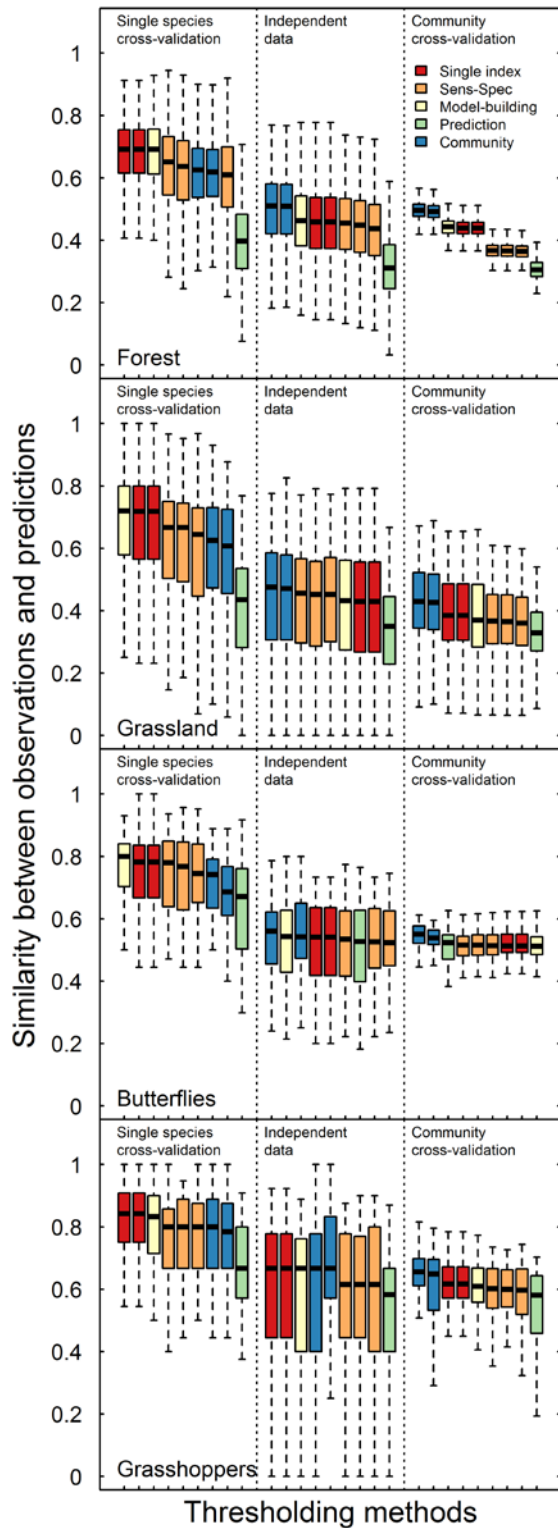


Figure S2: Sørensen similarity between observations and predictions for the four different datasets (top to bottom) and the three different modelling pathways (left to right). The boxplots are sorted by the median and the colours indicate the different thresholding techniques. The line in the box indicates the median, boxes range from the 25th to the 75th percentile and the whiskers indicate ± 2 standard deviations. For details on the method used within each threshold group see Table S1.

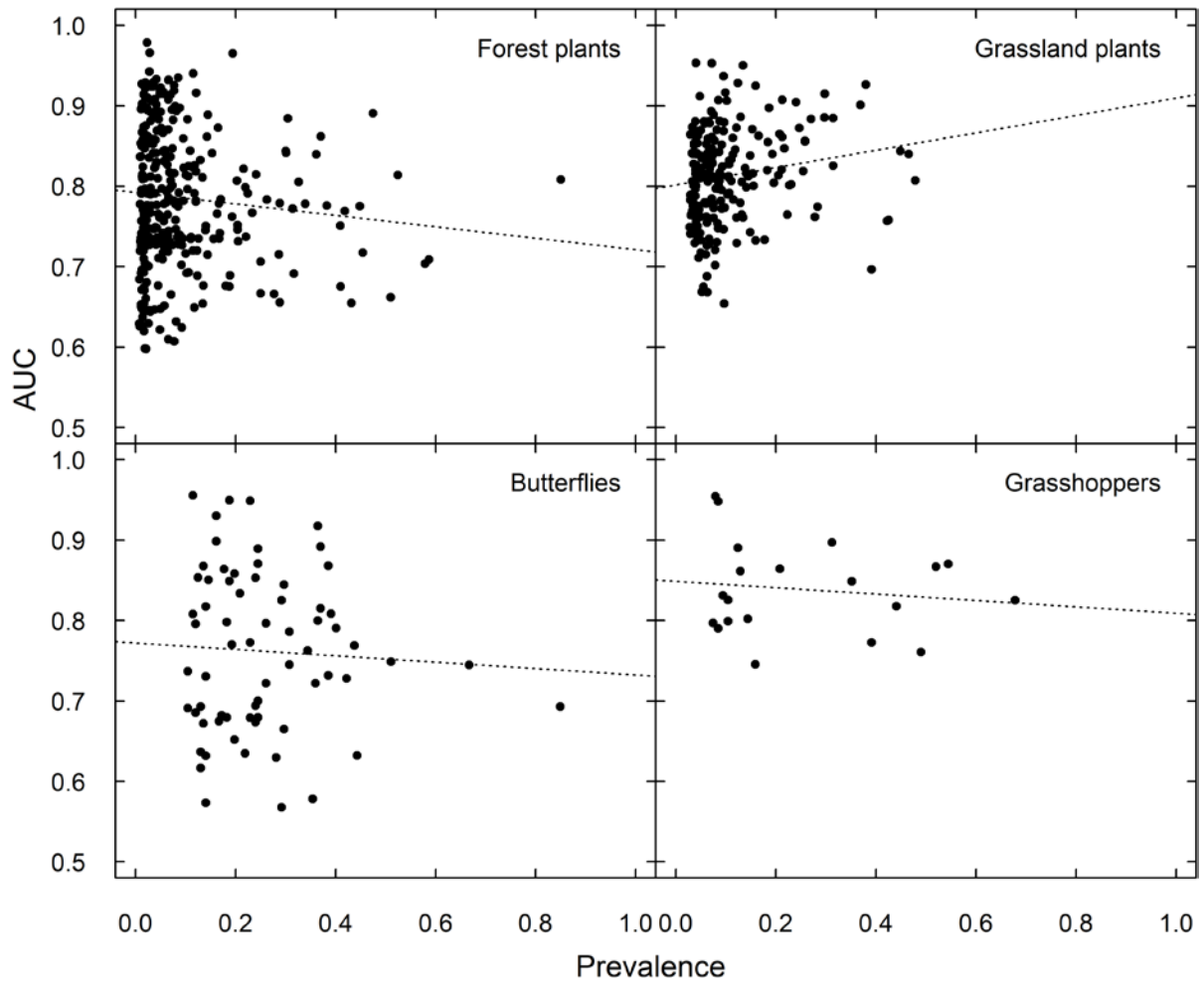


Figure S3: The relationship of the prevalence of a species (i.e., percentage of sites inhabited) to the performance of the SDMs (i.e., as measured by AUC) for the four studied data sets (taxa).

Table S1: Description of the ten thresholding methods based on Liu *et al.* (2005) and Nenzen and Araujo (2011).

Approach	Acronym	Definition	Reference
Single index-based approaches			
1. Kappa maximization approach	Max.Kappa	Kappa statistic is maximized	(Huntley <i>et al.</i> 1995; Guisan, Theurillat & Kienast 1998)
2. Maximum commission error	MCE05	Allowed a maximum commission error of 5%	(Mateo <i>et al.</i> 2012)
Sensitivity and specificity-combined approaches			
3. TSS maximization approach	Max.TSS	TSS statistic is maximized	(Allouche, Tsoar & Kadmon 2006)
4. Sensitivity-specificity equality approach	SensSpec	Difference of sens-spec is minimized	(Cantor <i>et al.</i> 1999)
5. ROC plot-based approach	Opt.ROC	ROC statistic is maximized	(Cantor <i>et al.</i> 1999)
Model-building data-only-based approach			
6. Prevalence approach	Preval	Prevalence of the calibration data	(Cramer 2003)
Predicted probability-based approaches			
7. Average probability approach	AvgProb	Taking the average predicted probability of the model-building data as threshold	(Cramer 2003)
Community based approaches			
8. pS-SDM+PRR	pS-SDM+PRR	Probability stacked SDM	(Dubuis <i>et al.</i> 2013)
9. MEM+PRR	MEM+PRR	Macroecological model for SR	(Guisan & Rahbek 2011)

Table S2 : Community evaluation metrics used in this study.

Metric	Definition
Species richness	
Deviation in species richness	$Dev.SPR = n_{pred} - n_{obs}$
Prediction success	
Sensitivity	$Sens = \frac{TP}{TP + FA}$
Specificity	$Spec = \frac{TA}{TA + FP}$
Community accuracy	$Acc = \frac{TP + TA}{N}$
Community TSS	$TSS = Sens + Spec - 1$
Community Kappa	$K = \frac{Acc - p_e}{1 - p_e}$
Community composition	
Sørensen	$S = \frac{2 * TP}{2 * TP + FP + FA}$

n_{pred} = Number of species predicted

n_{obs} = Number of species observed

N = Number of events

TP = Correctly predicted present species

TA = Correctly predicted absent species

FP = Falsely predicted present species

FA = Falsely predicted absent species

$p_e = \frac{(TP+FA)(TP+FP)+(TA+FP)(TA+FA)}{N^2}$

2 **Table S3** : Evaluation scores of individual SDMs by TSS (A), Kappa (B), PCC (C), Sensitivity (D) and Specificity (E) for the three community
3 evaluation approaches and four datasets. SSCV = Single species cross-validation, ID = Independent data, CCV = Community cross-validation,
4 FO = Forest plants, GL = Grassland plants, BF = Butterflies, GH = Grasshoppers.

5 (A) TSS

Thresholding Approach	SSCV				ID				CCV			
	FO	GL	BF	GH	FO	GL	BF	GH	FO	GL	BF	GH
Max.Kappa	0.2 ± 0.14	0.27 ± 0.2	0.31 ± 0.23	0.42 ± 0.17	0.21 ± 0.17	0.25 ± 0.21	0.3 ± 0.22	0.43 ± 0.26	0.23 ± 0.13	0.28 ± 0.14	0.31 ± 0.18	0.37 ± 0.15
MCE05	0.3 ± 0.17	0.32 ± 0.17	0.27 ± 0.18	0.43 ± 0.16	0.28 ± 0.17	0.34 ± 0.17	0.27 ± 0.23	0.45 ± 0.21	0.25 ± 0.17	0.34 ± 0.12	0.31 ± 0.17	0.42 ± 0.12
Max.TSS	0.35 ± 0.2	0.38 ± 0.14	0.34 ± 0.24	0.47 ± 0.12	0.33 ± 0.22	0.38 ± 0.21	0.34 ± 0.23	0.5 ± 0.26	0.35 ± 0.14	0.39 ± 0.11	0.35 ± 0.18	0.44 ± 0.12
SensSpec	0.32 ± 0.16	0.36 ± 0.14	0.33 ± 0.18	0.51 ± 0.18	0.31 ± 0.2	0.37 ± 0.21	0.34 ± 0.24	0.51 ± 0.26	0.35 ± 0.13	0.38 ± 0.11	0.35 ± 0.18	0.45 ± 0.11
Opt.ROC	0.34 ± 0.21	0.36 ± 0.19	0.33 ± 0.17	0.44 ± 0.26	0.32 ± 0.21	0.37 ± 0.21	0.34 ± 0.23	0.49 ± 0.25	0.35 ± 0.13	0.38 ± 0.11	0.35 ± 0.18	0.44 ± 0.12
Preval	0.18 ± 0.15	0.27 ± 0.15	0.3 ± 0.19	0.41 ± 0.23	0.2 ± 0.16	0.26 ± 0.2	0.31 ± 0.22	0.4 ± 0.23	0.18 ± 0.14	0.23 ± 0.15	0.3 ± 0.17	0.37 ± 0.17
AvgProb	0.43 ± 0.16	0.5 ± 0.12	0.41 ± 0.21	0.55 ± 0.14	0.47 ± 0.16	0.53 ± 0.13	0.38 ± 0.23	0.56 ± 0.15	0.44 ± 0.16	0.49 ± 0.11	0.4 ± 0.18	0.54 ± 0.14
pS-SDM+PRR	0.12 ± 0.19	0.17 ± 0.24	0.28 ± 0.24	0.28 ± 0.22	0.14 ± 0.18	0.2 ± 0.24	0.24 ± 0.24	0.29 ± 0.28	0.14 ± 0.17	0.19 ± 0.21	0.27 ± 0.23	0.27 ± 0.22
MEM+PRR	0.16 ± 0.17	0.2 ± 0.23	0.25 ± 0.24	0.3 ± 0.24	0.14 ± 0.18	0.2 ± 0.24	0.22 ± 0.24	0.32 ± 0.28	0.14 ± 0.17	0.2 ± 0.22	0.25 ± 0.22	0.3 ± 0.22

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7 (B) KAPPA

Thresholding Approach	SSCV				ID				CCV			
	FO	GL	BF	GH	FO	GL	BF	GH	FO	GL	BF	GH
Max.Kappa	0.2 ± 0.14	0.24 ± 0.18	0.28 ± 0.22	0.35 ± 0.19	0.21 ± 0.15	0.24 ± 0.19	0.29 ± 0.22	0.42 ± 0.25	0.2 ± 0.14	0.24 ± 0.15	0.29 ± 0.18	0.36 ± 0.15
MCE05	0.21 ± 0.13	0.23 ± 0.15	0.24 ± 0.22	0.31 ± 0.16	0.21 ± 0.13	0.28 ± 0.15	0.27 ± 0.22	0.41 ± 0.18	0.11 ± 0.12	0.16 ± 0.14	0.24 ± 0.17	0.32 ± 0.16
Max.TSS	0.19 ± 0.12	0.27 ± 0.17	0.3 ± 0.19	0.4 ± 0.14	0.21 ± 0.14	0.27 ± 0.16	0.3 ± 0.21	0.41 ± 0.23	0.17 ± 0.13	0.22 ± 0.15	0.3 ± 0.17	0.36 ± 0.15
SensSpec	0.21 ± 0.14	0.21 ± 0.17	0.29 ± 0.19	0.41 ± 0.17	0.22 ± 0.14	0.28 ± 0.17	0.3 ± 0.21	0.42 ± 0.23	0.17 ± 0.13	0.23 ± 0.15	0.3 ± 0.17	0.37 ± 0.14
Opt.ROC	0.15 ± 0.15	0.22 ± 0.14	0.32 ± 0.18	0.43 ± 0.17	0.22 ± 0.14	0.27 ± 0.16	0.3 ± 0.21	0.42 ± 0.23	0.17 ± 0.13	0.23 ± 0.15	0.3 ± 0.18	0.37 ± 0.14
Preval	0.2 ± 0.13	0.22 ± 0.18	0.3 ± 0.2	0.34 ± 0.17	0.21 ± 0.15	0.25 ± 0.18	0.3 ± 0.22	0.39 ± 0.22	0.19 ± 0.14	0.23 ± 0.15	0.3 ± 0.17	0.36 ± 0.16
AvgProb	0.17 ± 0.12	0.21 ± 0.14	0.26 ± 0.17	0.38 ± 0.15	0.17 ± 0.13	0.22 ± 0.15	0.26 ± 0.19	0.37 ± 0.18	0.16 ± 0.13	0.22 ± 0.15	0.29 ± 0.16	0.37 ± 0.16
pS-SDM+PRR	0.15 ± 0.16	0.17 ± 0.18	0.26 ± 0.23	0.28 ± 0.25	0.14 ± 0.17	0.18 ± 0.2	0.22 ± 0.23	0.29 ± 0.29	0.14 ± 0.16	0.17 ± 0.19	0.26 ± 0.21	0.28 ± 0.22
MEM+PRR	0.14 ± 0.16	0.17 ± 0.21	0.24 ± 0.22	0.32 ± 0.26	0.14 ± 0.17	0.18 ± 0.2	0.22 ± 0.23	0.34 ± 0.29	0.15 ± 0.16	0.19 ± 0.19	0.25 ± 0.21	0.32 ± 0.22

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9 C) Percentage correct classified (PCC)

Thresholding Approach	SSCV				ID				CCV			
	FO	GL	BF	GH	FO	GL	BF	GH	FO	GL	BF	GH
Max.Kappa	0.91 ± 0.09	0.9 ± 0.07	0.78 ± 0.1	0.83 ± 0.08	0.9 ± 0.09	0.89 ± 0.07	0.77 ± 0.09	0.83 ± 0.1	0.88 ± 0.09	0.87 ± 0.06	0.76 ± 0.08	0.82 ± 0.07
MCE05	0.85 ± 0.15	0.79 ± 0.07	0.66 ± 0.11	0.82 ± 0.08	0.88 ± 0.07	0.87 ± 0.05	0.77 ± 0.09	0.82 ± 0.08	0.59 ± 0.14	0.68 ± 0.09	0.64 ± 0.1	0.73 ± 0.08
Max.TSS	0.85 ± 0.08	0.84 ± 0.07	0.73 ± 0.11	0.83 ± 0.09	0.86 ± 0.1	0.85 ± 0.08	0.73 ± 0.11	0.81 ± 0.09	0.77 ± 0.07	0.79 ± 0.05	0.72 ± 0.08	0.79 ± 0.06
SensSpec	0.79 ± 0.1	0.84 ± 0.04	0.73 ± 0.09	0.81 ± 0.08	0.87 ± 0.09	0.86 ± 0.07	0.73 ± 0.1	0.81 ± 0.08	0.79 ± 0.07	0.81 ± 0.05	0.73 ± 0.07	0.8 ± 0.06
Opt.ROC	0.86 ± 0.08	0.86 ± 0.06	0.72 ± 0.11	0.82 ± 0.05	0.87 ± 0.1	0.86 ± 0.07	0.74 ± 0.1	0.81 ± 0.09	0.79 ± 0.07	0.81 ± 0.05	0.73 ± 0.08	0.8 ± 0.06
Preval	0.92 ± 0.08	0.91 ± 0.05	0.79 ± 0.08	0.83 ± 0.06	0.9 ± 0.08	0.89 ± 0.06	0.77 ± 0.1	0.83 ± 0.09	0.9 ± 0.08	0.89 ± 0.06	0.77 ± 0.08	0.82 ± 0.07
AvgProb	0.71 ± 0.07	0.67 ± 0.08	0.64 ± 0.11	0.74 ± 0.08	0.69 ± 0.08	0.69 ± 0.07	0.64 ± 0.11	0.73 ± 0.08	0.69 ± 0.07	0.69 ± 0.07	0.66 ± 0.09	0.73 ± 0.07
pS-SDM+PRR	0.93 ± 0.09	0.88 ± 0.07	0.76 ± 0.08	0.86 ± 0.09	0.91 ± 0.1	0.89 ± 0.08	0.77 ± 0.11	0.83 ± 0.1	0.91 ± 0.1	0.89 ± 0.08	0.78 ± 0.09	0.84 ± 0.1
MEM+PRR	0.92 ± 0.1	0.9 ± 0.09	0.8 ± 0.09	0.84 ± 0.09	0.91 ± 0.1	0.89 ± 0.09	0.79 ± 0.1	0.86 ± 0.08	0.91 ± 0.1	0.89 ± 0.08	0.79 ± 0.08	0.86 ± 0.09

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12 D) Sensitivity

Thresholding Approach	SSCV				ID				CCV			
	FO	GL	BF	GH	FO	GL	BF	GH	FO	GL	BF	GH
Max.Kappa	0.31 ± 0.18	0.36 ± 0.24	0.47 ± 0.21	0.51 ± 0.26	0.27 ± 0.21	0.32 ± 0.25	0.46 ± 0.24	0.56 ± 0.26	0.32 ± 0.18	0.37 ± 0.18	0.5 ± 0.2	0.52 ± 0.22
MCE05	0.45 ± 0.13	0.61 ± 0.11	0.66 ± 0.18	0.73 ± 0.12	0.35 ± 0.17	0.42 ± 0.17	0.42 ± 0.23	0.57 ± 0.19	0.68 ± 0.05	0.67 ± 0.07	0.7 ± 0.09	0.71 ± 0.11
Max.TSS	0.44 ± 0.28	0.55 ± 0.15	0.62 ± 0.14	0.67 ± 0.15	0.46 ± 0.28	0.52 ± 0.27	0.59 ± 0.22	0.66 ± 0.25	0.57 ± 0.11	0.59 ± 0.11	0.61 ± 0.14	0.63 ± 0.15
SensSpec	0.55 ± 0.25	0.49 ± 0.15	0.58 ± 0.18	0.65 ± 0.13	0.42 ± 0.26	0.49 ± 0.25	0.58 ± 0.22	0.67 ± 0.25	0.54 ± 0.1	0.55 ± 0.11	0.58 ± 0.14	0.63 ± 0.12
Opt.ROC	0.44 ± 0.2	0.54 ± 0.14	0.57 ± 0.21	0.67 ± 0.14	0.44 ± 0.26	0.49 ± 0.26	0.58 ± 0.22	0.65 ± 0.24	0.55 ± 0.1	0.56 ± 0.11	0.59 ± 0.14	0.62 ± 0.14
Preval	0.28 ± 0.17	0.28 ± 0.21	0.46 ± 0.21	0.55 ± 0.25	0.26 ± 0.2	0.32 ± 0.24	0.49 ± 0.22	0.53 ± 0.25	0.24 ± 0.18	0.29 ± 0.18	0.47 ± 0.19	0.5 ± 0.23
AvgProb	0.76 ± 0.11	0.82 ± 0.1	0.77 ± 0.12	0.83 ± 0.1	0.79 ± 0.11	0.85 ± 0.1	0.78 ± 0.17	0.85 ± 0.13	0.76 ± 0.1	0.81 ± 0.06	0.78 ± 0.1	0.84 ± 0.07
pS-SDM+PRR	0.21 ± 0.28	0.26 ± 0.32	0.43 ± 0.36	0.42 ± 0.39	0.21 ± 0.28	0.28 ± 0.33	0.44 ± 0.36	0.45 ± 0.4	0.21 ± 0.27	0.26 ± 0.3	0.45 ± 0.33	0.41 ± 0.36
MEM+PRR	0.21 ± 0.28	0.3 ± 0.31	0.38 ± 0.36	0.41 ± 0.38	0.21 ± 0.29	0.28 ± 0.33	0.38 ± 0.35	0.43 ± 0.38	0.21 ± 0.27	0.28 ± 0.3	0.39 ± 0.32	0.42 ± 0.34

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16 E) Specificity

Thresholding Approach	SSCV				ID				CCV			
	FO	GL	BF	GH	FO	GL	BF	GH	FO	GL	BF	GH
Max.Kappa	0.95 ± 0.09	0.91 ± 0.07	0.82 ± 0.12	0.87 ± 0.1	0.94 ± 0.07	0.93 ± 0.07	0.84 ± 0.12	0.88 ± 0.1	0.91 ± 0.09	0.93 ± 0.07	0.81 ± 0.12	0.85 ± 0.1
MCE05	0.87 ± 0.15	0.81 ± 0.04	0.78 ± 0.11	0.79 ± 0.09	0.93 ± 0.03	0.91 ± 0.03	0.86 ± 0.1	0.88 ± 0.08	0.57 ± 0.15	0.91 ± 0.03	0.61 ± 0.13	0.7 ± 0.1
Max.TSS	0.88 ± 0.09	0.79 ± 0.07	0.73 ± 0.1	0.83 ± 0.09	0.87 ± 0.1	0.86 ± 0.09	0.75 ± 0.13	0.84 ± 0.09	0.78 ± 0.08	0.86 ± 0.09	0.74 ± 0.1	0.8 ± 0.08
SensSpec	0.89 ± 0.07	0.82 ± 0.06	0.77 ± 0.1	0.81 ± 0.09	0.89 ± 0.09	0.88 ± 0.07	0.75 ± 0.12	0.84 ± 0.09	0.81 ± 0.07	0.88 ± 0.07	0.76 ± 0.09	0.82 ± 0.07
Opt.ROC	0.84 ± 0.08	0.87 ± 0.08	0.74 ± 0.11	0.82 ± 0.09	0.88 ± 0.09	0.88 ± 0.08	0.76 ± 0.13	0.84 ± 0.09	0.8 ± 0.07	0.88 ± 0.08	0.76 ± 0.09	0.82 ± 0.07
Preval	0.92 ± 0.07	0.95 ± 0.05	0.85 ± 0.13	0.85 ± 0.09	0.94 ± 0.08	0.94 ± 0.06	0.82 ± 0.13	0.87 ± 0.11	0.94 ± 0.08	0.94 ± 0.06	0.84 ± 0.11	0.87 ± 0.09
AvgProb	0.69 ± 0.08	0.68 ± 0.06	0.63 ± 0.12	0.72 ± 0.09	0.68 ± 0.08	0.68 ± 0.07	0.6 ± 0.12	0.71 ± 0.09	0.68 ± 0.08	0.68 ± 0.07	0.62 ± 0.09	0.69 ± 0.08
pS-SDM+PRR	0.92 ± 0.15	0.91 ± 0.12	0.81 ± 0.19	0.86 ± 0.19	0.93 ± 0.15	0.92 ± 0.12	0.8 ± 0.23	0.84 ± 0.21	0.93 ± 0.14	0.92 ± 0.12	0.82 ± 0.19	0.86 ± 0.18
MEM+PRR	0.93 ± 0.14	0.93 ± 0.1	0.85 ± 0.18	0.91 ± 0.15	0.93 ± 0.15	0.92 ± 0.13	0.85 ± 0.2	0.89 ± 0.16	0.93 ± 0.14	0.92 ± 0.13	0.86 ± 0.17	0.88 ± 0.15

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