Integrating biodiversity assessments into local conservation planning: the importance of assessing suitable data sources

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19 Abstract

20 Strategic Environmental Assessment (SEA) of land-use planning is a fundamental tool to minimize 21 environmental impacts of artificialization. In this context, Systematic Conservation Planning (SCP) tools 22 based on Species Distribution Models (SDM) are frequently used for the elaboration of spatially 23 exhaustive biodiversity diagnostics. Despite the paradigm of "garbage in - garbage out" that emphasises 24 the importance of testing the suitability of data for SDM and priority conservation areas, the assessment 25 of database sources remains relatively rare. In addition, the lack of practical recommendations for the 26 use of open-access databases by SEA stakeholders remains a problem. The aim of this study is to explore 27 the quality of data sources that can be used by stakeholders in SEA to assess priority conservation areas 28 in SEA. The study was done usingused data for nine taxonomic groups (commonly used in inventories for 29 environmental impact assessment) and three databases available to SEA stakeholders. Three local 30 administrative entities in very different socio-ecological contexts were used to examine three main issues 31 : (i) the suitability of local versus national regional or country databases for assessing conservation 32 priorities, (ii) differences among taxonomic groups or territories in terms of the suitability of databases, 33 (iii) the importance of the quality of databases for the application of SDM to assess priority conservation 34 areas. Our study provides several clear messages for potential users of open-access databases. First, the 35 need for prudence in the interpretation of biodiversity maps. Second, for SDM, the collection of individual 36 databases at the national country scale is necessary to complete local data and ensure the suitability of 37 modelsSDM in a local context. Third, a data driven approach can lead to the use of notably different 38 species communities to identify priority conservation areas when compared to the community in the 39 original database. Finally, we propose a workflow to guide SEA stakeholders through the process of data 40 rationalization and use in conservation planning.

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42 *Keywords:* Data-driven approach, Species Distribution Models (SDM), Strategic Environmental Assessment (SEA),
 43 Systematic Conservation Planning (SCP)

1 - Introduction

Land-use change, in particular urban land expansion, leads to artificialization of habitats and soils and is one of the major causes of the loss of biodiversity (Maxwell et al., 2016; IPBES, 2019). The reduction and fragmentation of natural habitats leads to population declines and species extinction (Fahrig, 1997; Horváth et al., 2019; Lino et al., 2019), as well as biotic homogenization that, i.e. mostly the extinction of specialist species and the introduction of exotic species, which involves an increase in genetic, taxonomic and functional similarity (Olden and Rooney, 2006; Zambrano et al., 2019)(Olden and Rooney, 2006; Zambrano et al., 2019).

52 A major tool to limit artificialization is the mitigation hierarchy used in environmental assessments studies. 53 This approach consists of three sequential steps: "avoid" impacts, "reduce or minimize" impacts not avoided 54 and "offset" residual impacts (Bull et al., 2016; Maron et al., 2016). However, in the current application of the 55 mitigation hierarchy several weaknesses prevent it from achieving the goal of "No Net Loss" of biodiversity 56 (Quétier et al., 2014; Bezombes et al., 2019). Avoidance is poorly implemented despite the fact that it is the 57 first and most efficient step of the hierarchy (Bigard et al., 2017; Phalan et al., 2018). What is more, the 58 mitigation hierarchy is mostly applied in a project-by-project approach without scaling up (Pope et al., 2013; 59 Bigard et al., 2017), which limits proper consideration of fragmentation issues (Gontier et al., 2006) and 60 cumulative impacts (Whitehead et al., 2017), including those of multiple small projects (Bigard et al., 2017).

61 To anticipate avoidance measures, Strategic Environmental Assessment (SEA) of land-use planning is a 62 global and fundamental tool to minimize environmental impacts (Baker et al., 2005). SEA provides for the 63 integration of avoidance measures early in the land-use planning process through environmental assessment 64 of policies, plans and programs (Fundingsland Tetlow and Hanusch, 2012; Bigard et al., 2020). However, the 65 implementation of SEA is often based on biodiversity diagnostic maps that are rarely complete and exhaustive. 66 Indeed, biodiversity diagnostics are rarely exclusively based on empirical observations from field surveys 67 (Phalan et al., 2018) and usually use areas and documents already known (e.g., protected areas and green 68 infrastructures).

69 Spatial modelling provides a tool for the elaboration of spatially exhaustive diagnostics of biodiversity maps 70 for land-use and conservation planning (Almenar et al., 2019; Tarabon et al., 2019; Bigard et al., 2020; Tulloch 71 et al., 2019; Baker et al., 2021; Boileau et al., 2022). Among these methods of biodiversity modelling, Species 72 Distribution Models (SDM) are widely used to predict suitable habitat for species based empirical observations 73 (Guisan et al., 2017; Zurell et al., 2020) and are increasingly used in conservation planning (Guisan et al., 2017; 74 Domisch et al., 2019; Baker et al., 2021). Systematic Conservation Planning (SCP) tools are also particular 75 pertinent to identify priority biodiversity stakes and avoid the adoption of an ad hoc approach (Margules and 76 Pressey, 2000; Pressey and Bottrill, 2008) in order to inform SEA (Tulloch et al., 2019).

77 The management of databases and their use for conservation planning is a critical issue for the application 78 of such methods to practical conservation planning. The databases available for SEA stakeholders (i.e. decision 79 makers, environmental consultants and conservation managers) are often limited because of data sensitivity 80 or ownership issues, although more and more programmesprograms contain data that are publicly available 81 and use of them can be made without any particular attention to their quality (Costello and Wieczorek, 2014; 82 Tittensor et al., 2014)- and they are generally unfamiliar to SEA stakeholders. Surprisingly however, despite the 83 prevailing recognition of the "garbage in – garbage out" that emphasises the critical importance of the quality 84 of data (Sanders and Saxe, 2017; Canbek, 2022), an examination of data suitability is relatively rare in local 85 conservation planning (Rondinini et al., 2006; Hermoso et al., 2015a). In this context, some authors argue the 86 necessity of examining the sensitivity of model results to the nature of the datasets that are used (Sanders and 87 Saxe, 2017; Clare et al., 2019; Velazco et al., 2020). SDM studies generally use data that has not been designed 88 specifically for this type of analysis, and is often comprised of presence-only data, hence the need for a rigorous 89 assessment of sampling biases (Beck et al., 2014; Botella et al., 2018; Guisan et al., 2017). Another particularly 90 important point that can influence distribution modelling is the spatial extent of the data, and in particular the 91 question of whether to use only local data or those collected on a larger scale (Baker et al., 2021; Meyer, 2007). 92 The choice and possible combination of data sources is part of this problem due to the fact that they often 93 vary considerably in their design, the gradients covered, and potential sampling biases (Fletcher et al., 2019; 94 Boyd et al., 2023). Basically, the use of available databases requires a rigorous test of their quality and 95 pertinence (Zuckerberg et al., 2011), especially when used for analyses such as SDM (Tulloch et al., 2016; 96 Domisch et al., 2019). Confidence in the models must be assessed through the use of metrics adapted to the 97 data (Guisan et al., 2017; Leroy et al., 2018). As recognised by Clare et al., (2019), the lack of practical

98 recommendations for the use of databases that differ in terms of their quality and pertinence by public 99 authorities or other institutions remains a serious problem.

100 The overall goal of this study is to test the influence of different database sources that can be used by SEA 101 stakeholders to map priority conservation areas in SEAs based on SCP. To do so, we studied three local 102 administrative territories that occur in different socio-ecological contexts in France. The study has three main 103 objectives. First, in terms of the quantity of data available we assess the content of three open-access 104 databases for nine taxonomic groups commonly used in naturalist inventories in environmental impact 105 assessment studies, we assess the variation in the . We evaluate their suitability of the available databases in 106 terms of data quantity for SDM application, at three scales (local, regional and national) for application of SDM. 107 Second, by focusing oncountry). SDM and SCP analyses were performed for two taxonomic groups with 108 different dispersion capacities, home range sizes and environmental (Aves and Papilionidae) to test the 109 hypothesis that sampling bias and differences in ecological response scales, of species may influence the 110 identification priority conservation areas. Second, we explore the influence of databases for on the application 111 of SDM to assess priority conservation areas. Third, we analyse the influence of this data-driven approach on 112 the composition of species communities that are ultimately considered in priority conservation areas relative 113 to the actual communities in the original databases. Third, we analyse the influence of this data-driven 114 approach on the composition of species communities that are ultimately consideredused in the 115 analysisidentification of priority conservation areas relative to the actual communities in the original 116 databases.

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2 - Methods

118 **2.1 - Study sites**

119 To assess the availability and suitability of pertinent data sources, we selected three French local 120 administrative entities in charge of land-use planning: Lodévois-Larzac (T1), Brocéliande (T2), La Rochelle (T3) 121 (Figure 1, Table 1). We selected these territories). We selected these study sites on the basis their contrasting 122 social, ecological and geographical contexts in order to examine patterns of variation of data suitability among 123 sites. (Figure 1, Table 1). For example, each of the three territories is composed of study sites have different 124 ecosystems and bioclimates and the sites vary in terms of urbanization pressures (from 3% to 28% artificial 125 land-use cover for sites T1 and T2 respectively) and . This is due to the presence of protected areas a major 126 city (La Rochelle) in their territories (from 2% to 70% of the protected territory the T2 study site. The major 127 towns in the other two study sites are smaller however there is a major city less than 50 km away for T3both 128 of them (Montpellier and T1Rennes respectively). Only site T1 has an important cover of protected areas 129 (Natura 2000) area.



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Territory	Study site	Lodévois-Larzac (T1)	La Rochelle (T2)	Brocéliande (T3)		
Country / Regio	nAdministrative <u>;ion</u>	France / Occitanie	France / Nouvelle-Aquitaine	France / Bretagne		
Main city c	coordinates	43°43'57"N 3°19'02"E	46°09'35"N 1°09'05"W	48°00'08"N 2°05'48"W		
Ar	ea	554 km ²	327 km ²	298 km ²		
Clin	nate	Mediterranean	Oceanic	Oceanic		
Major habitats / land-use		Mediterranean forest and scrubland	Intensive agricultural field, swamps, few forests	Pastures, temperate forest		
% land-use ⁽¹⁾	Artificial <u>–</u> <u>urban areas</u> Farmland Semi-natural	3% 3% 94%	28% 55% 17%	7% 29% 64%		
Landscape a	nd dynamics	Rural, woodland expansion	Coastal, intensive agriculture, urban expansion	Rural, agricultural intensification		
Urbanizati	on <u>context</u>	Close to a major city , across (<u>Montpellier), linked</u> by a highway	Economic and tourist dynamism <u>, major city</u> <u>presence</u>	Close to a major city , across (<u>Rennes), linked</u> by a highway		
	Regulatory	0%	327 ha – 0.99%	0%		
% protected	Land control	1618 ha – 0.6%	148 ha – 0.45%	80 ha – 0.25%		
area (2)	Contractual	38096 ha – 68.8%	3534 ha – 10.7%	586 ha – 2%		
	Total	38602 ha - 69 7%	3546 ha - 10 8%	592 ha - 2%		

(1) source: OSO Land Cover (Inglada et al., 2019). Artificial is impermeable surfaces; crop field is annual crop and orchards and vineyards; semi-natural is hardwood and soft wood and grasslands and water.

(2) source: https://inpn.mnhn.fr/

138 **2.2 – Workflow of analysis**

A methodological framework was developed to test the influence of different database sources in mapping priority conservation areas in SEAs thanks to a SCP approach, all steps are summarized in Figure 2- and in the following text.

142 **2.3 – DatabaseDatabases** available for SEA stakeholders

143 2.3.1 – Content of available databases

144 We focused on nine taxonomic groups commonly used for naturalist inventories for environmental impact 145 assessment studies: Amphibia, nesting Aves (hereafter name Aves), Chiroptera, Flora, Mammalia aptera (hereafter name Mammalia), Orthoptera, Odonata, Papilionidae and Reptilia (Bigard et al., 2017; Guillet et al., 146 147 2019; Iorio et al., 2022). The three spatial scales used for data collection are depicted in Figure 1: the local scale 148 (i.e. study site with a 10km buffer around), the regional scale (i.e. French administrative regions); and the 149 country scale (i.e. continental France). We selected three open-access databases containing these groups that 150 can be widely used by SEA stakeholders for the assessment and hierarchy of conservation priorities (Figure 2, 151 step 1.1). This study thus directly addresses SEA stakeholders (i.e. decision makers, environmental consultants 152 and conservation managers) using the databases available to them. 153 The first of these databases concerns the French Natural and Landscape Information System (SINP) that is

153 The first of these databases concerns the French Natural and Landscape information system (SNP) that is 154 structured at the regional scale in France.scale of French administrative regions in charge of data extraction 155 requests. Each site has its own database, can be collected only at local scale due to the limited extent of data 156 requests, without the need for a special request (maximum 2000 km², i.e. nearby 10km buffer zone around 157 the study site). This database is composed of opportunist observations and only contains presence data for

- 158 taxa for which identification is confirmed by experts (Jomier et al., 2018). Access to this database requires a
- 159 data extraction request to the regional administrative structure in charge of the database. Generally, a special
- 160 request is necessary above a 10km buffer zone around the study site (maximum 2000 km2), the data remains
- accessible only locally. Each site has its own database (see Appendix (see Appendix A.1).
 The second database is the Global Biodiversity Information Facility (GBIF) an international pla
- 162 The second database is the Global Biodiversity Information Facility (GBIF) an international platform for the 163 provision of biodiversity data that is based on information collected from various databases (Telenius, 2011).
- 164 It is composed of observation data that are not based on protocols and for which presence data and

Table 1 - Description of three study sites in France

identification are not subject to expert confirmation. The data downloading is autonomous from the website(see Appendix A.1).

167 The third database is a French biodiversity monitoring scheme (Vigie Nature) dedicated to assess spatio-168 temporal populations trends. Within this monitoring scheme data collection is based on a standardized 169 biodiversity survey. Despite local spatial distribution heterogeneities, the sampling plan ensures a 170 representation of the current-national distribution of habitats and landscapes across France (Julliard and 171 Jiguet, 2002; Mariton et al., 2022). Homogeneity in identification criteria and compliance with the protocol are 172 ensured by offering training to volunteers. This database is composed of presence/absence and abundance 173 data. Access to these databases requires a data extraction request to the person in charge (see Appendix A.1). 174 These three databases were combined in two ways: "All databases" (i.e. the combination of SINP, GBIF and 175 Vigie Nature) used in section 2.3 (i.e. for assess which taxa are enough documented within each dataset) and

"National databases" (i.e. <u>the combination of GBIF and Vigie Nature</u>, <u>which are available at country scale</u>) used
 specifically in section 2.4 (i.e. <u>forto</u> test the effect of database sources on SDM performance).

178 The databases were collected for continental France except for the SINP that was collected in 10km buffer 179 zone around the study sites due to the access restrictions explained above. The databases were collected over a period of 10 years (i.e. from 01/01/2010 to 31/12/2020) and data with spatial inaccuracy greater than 50 181 meters were not considered. We made a series of operations to standardize, correct and homogenize species 182 names, transform data into occurrences, limit their spatial biases and identify the presence of species into each study site (Appendix A.1).

184 The databases were collected for continental France except for the SINP that was collected in 10km buffer 185 zone around the study sites due to the access restrictions explained above. The databases were collected over 186 a period of 11 years (i.e. from 01/01/2010 to 31/12/2020) and data with spatial inaccuracy greater than 50 187 meters were not considered. We made a series of operations to standardize, correct and homogenize taxa 188 names at the specific taxonomic level using the French taxonomic reference "TAXREF.V14" (Gargominy et al., 189 2021). We transformed data into occurrences and limited their sampling biases by geographical filtering using 190 "spThin" package (Aiello-Lammens et al., 2015). In each study site, we identify the presence of one species at 191 least five observations from "All databases" combined at local scale and defined as present species in France 192 by TAXREF.V14 (Appendix A.1).

193 2.3.2 – Quantitative description of databases for SDM

194 For the nine taxonomic groups, four metrics were selected to quantitatively describe the amount and thus 195 the suitability of each database (i.e. SINP, GBIF, Vigie Nature and <u>"All databases}"</u>) for the realization of SDM 196 in presence-only (Figure 2, step 1.2): (i) the number of species observed in the study site; (ii) the proportion of 197 species with < 15 observations which represents the minimal threshold for the utility of SDM with more 198 accurate predictions than in a random model (Støa et al., 2019); (iii) the proportion of species with between 199 15 and 50 observations, i.e. the minimum number of presences recommended for SDM (Merow et al., 2014 in 200 Guisan et al., 2017); (iv) the proportion of species with > 50 observations, i.e. highly suitable for modelling 201 (Støa et al., 2019). These metrics were calculated at three different scales: (a) for each territorystudy site 202 including a 10km buffer zone (local scale) that is the maximum extent for a SINP data request; (b) on a regional 203 scale that is used for the structure of biodiversity data in France; (c) for continental France. SINP database is 204 analysed individually only at the local scale due to the previously mentioned restriction of access, nevertheless 205 it integrated the three scales of the "All databases.".

206 **2.4 - Systematic conservation planning process**

207 For the three study sites, we identified priority conservation areas with a Systematic Conservation Planning 208 (SCP) tool based on Species Distribution Models (SDM) (Figure 2, steps 2.1 and 2.2). Several variants of SDM 209 were made using different database sources (i.e. GBIF, Vigie Nature and "National databases}") and two 210 methods of generating pseudo-absences (named individual database or mixed databases). Among the nine 211 taxonomic groups studied above, only Aves and Papilionidae taxa were analysed to compare the tests in the 212 SCP process. The data available in France for these groups seems to be sufficient in quantity to realize SDM 213 with each database. The use of these two groups allows for a comparison between one group of highly mobile 214 taxa with a large home range (Aves) and another group with a smaller home range and whose movement 215 closely tracks local environmental variation (Papilionidae). These two taxonomic groups thus have different

biological traits associated with their dispersal and function, hence we predict differences in in terms of the
 spatial resolution of their distribution.

218 2.4.1 - Species distribution modelling (SDM)

We modelled favourable habitats for birds and butterflies in the three study sites using SDM (Figure 2, step 220 2.1). A resolution of 50m was used to meet the needs of the SEA of land-use planning. A buffer zone of 10km 221 around each of the study sites (i.e. local scale) was used for the SDM prediction to limit any border edge effects 222 and to increase the number of species that could be modelled and evaluated. Indeed, species with less than 223 15 data points for the calibration (threshold explain above, Støa et al., 2019) and/or less than 10 data for 224 performance evaluation (threshold defined by expert opinion) were not modelling.

Biodiversity data used for the SDM came from the databases <u>decribeddescribed</u> above at the nationalcountry scale, according to the results of section 2.3 (Table 2). These data were separated into two independent datasets that allow for robust validations with independent data (Matutini et al., 2021). NationalCountry data without local data were used for model calibration and local data were used only for model performance evaluation. Therefore, the SINP database, which is only available at local scale, was not used for model calibration.

231 For model calibration, pseudo-absences were generated with two methods-, separately for each taxonomic 232 group. First, methods to generate pseudo-absences in the individual databases for all their data (i.e. GBIF, Vigie 233 Nature and "National databases}."). For databases with a protocol for sampling (i.e. Vigie Nature) to optimize 234 species detection (day and year periods), the absence points were defined as all the points without the 235 observed species. For databases without such sampling protocols for all their data (i.e. GBIF and "National 236 databases),"), pseudo-absence data were generated with the target-group (TG) approach, which infers the 237 sampling bias from the aggregated occurrences of (TG) species, i.e. the respective taxonomic groups (Ponder 238 et al., 2001; Anderson, 2003; Phillips et al., 2009). Second, a method to mix the presence data in the GBIF and 239 "National databases" with the absence from Vigie Nature (named mixed databases) was applied (Hermoso et 240 al., 2015a).

Three types of environmental variables <u>were</u>used for SDM: <u>biogeographicgeographic</u>, human occupancy and pollution and fragmentation (Appendix <u>A-2</u>A.2).

SDM were <u>calibrate_calibrated</u> by Random Forest down-sample (Valavi et al., 2021a) which according to Valavi et al. (2021b) is among the best performing models for presence-only data. Although the Random Forest is not very sensitive to the non-independence of the variables and over-parametrization (Matsuki et al., 2016; Srisa-An, 2021), in order to be parsimonious, the collinear variables were removed (Pearson >0.7, Appendix A.2A.2, Brun et al., 2020). Thirty bootstraps were performed for each SDM (Guisan et al., 2017) using a calibration for 70% of the data at the <u>nationalcountry</u> scale outside of local scale. The thirty Random Forest bootstraps were combined with mean to provide an ensemble prediction of habitat suitability for all species.

The performance evaluation of the models was done using the Boyce's index (CBI), the most suitable metric for model in presence-only (Boyce et al., 2002; Leroy et al., 2018), with local data (i.e. <u>"All databases"</u> combining SINP, GBIF and Vigie Nature). Dubos et al. (2022) reveal the CBI turns out to be misleading in some cases, thus we used a threshold of 0.3 to define good or poor model quality.

254 2.4.2 – Systematic conservation planning (SCP) tool

255 Priority conservation areas in the three territoriesstudy sites were analysed from SDMs for each database 256 source in using a SCP tool (Figure 2, step 3.1). To meet the needs of SEAs, the study site was restricted to 257 administrative boundaries with a buffer zone of 1km to maintain coherence between administrative entities 258 and a resolution of 50m. The SEAsaim of SEA biodiversity conservation strategies aim-is to establish priorities 259 for the whole territorystudy site as a whole and all the cells have the same cost value of 1. The objective was 260 a maximum coverage objective that seeks to maximize the number of features, i.e. the SDMs (Church et al., 261 1996). The features were only the predictions of SDM with a Boyce's index > 0.3 to limit the influence of 262 poorgood quality models defined previously. The priority conservation areas decisions were between 0 and 1. 263 To obtain a priority gradient, we cumulated Ferrier importance scores (Ferrier et al., 2000) from nine targets 264 of the total amount of each feature (from 0.1 to 0.9 every 0.1). The We used the package "prioritizr" was used 265 (Hanson et al., 2021) with the open-source solver SYMPHONY (Kim et al., 2023).

266 2.5 - Comparative analysis of SDM and priority conservation areas

267 The We analysed the influence of database sources on SDM predictions and priority conservation areas 268 were analysed (Figure 2, step 2.2 and 3.2). SDM performance evaluations were analysed between database 269 sources. The SDM prediction and priority conservation areas maps were compared with the Spearman's rank 270 coefficient (Phillips et al., 2009) and the Schoener's D index as a measure of projection overlap (Schoener, 271 1968) which was calculated with the ENMTool R package (Warren et al., 2008; Warren and Dinnage, 2022).

272 2.6 - Species community analysis

273 We assessed the influence of the complete data driven workflow on the composition of species 274 communities, i.e. differences between the original community (i.e. all species observed in study site) in the 275 database and the final community used to identify priority conservation areas (Figure 2, step 4).

276 To do so, we developed a workflow resistance score for each of the methodological filters for all species. 277 A score of 1 is allocated to species observed in the study area that did not cross any of the stepwise filters. A 278 score of 2 is allocated to species with sufficient data to calibrate SDM, i.e. > 15 national country observations, 279 or evaluate evaluate the performance of SDM, i.e. > 10 local observations. A score of 3 is allocated to species 280 with sufficient data to calibrate and evaluate the performance of the SDM. A score of 4 is for species that were 281 present in the final analysis as a priority species for conservation planning (i.e. with the two previous filters 282 and a Boyce's index > 0.3). In order to assess species composition bias ultimately considered in priority 283 conservation areas, species communities were analysed through traits that can influence species detection 284 (mass, displacement capacity, period activity) and ecological traits (habitats, specialisation) (Appendix 285 A.3). A.3). Missing data were completed with a trait imputation procedure generated using the R package 286 "missForest" (Stekhoven and Bühlmann, 2012) by considering evolutionary relationships in the imputation 287 process (see Carmona et al., 2021) using the R script of Toussaint et al. (2021). Due to the nature of the 288 response variable (i.e. ordinal scoring including four modalities), we used ordinal regression mixed models with 289 cumulative link using the clmm function of "ordinal" R package (Christensen, 2022). We adapted the link 290 function to the data distribution-for Aves and Papilionidae, using thea "cauchit" link for Aves and a "logit" links, 291 respectivelylink for Papilionidae. Species traits were used as fixed effects, while the random effects selected 292 were the study sites for Aves and the combination for study sites and database sources for Papilionidae. Finally, 293 we evaluated the quality of the full model by comparing to the null model with Akaike's information criterion 294 (AIC) (Mac Nally et al., 2018).

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Thus, our models were structured in the following way:

296 Resistance ~ $\beta_0 + \beta_i$.'trait'_i + +(1|StudySites) For Aves 297 For Papilionidae Resistance ~ $\beta_0 + \beta_i$.'traits'_i++(1|StudySites_/Databases) 298 Where β is the parameter estimates, i correspond to the variables of 'traits' using in fix effect, 0 is the shift

299 between ordinal class of resistance (i.e. 1|2, 2|3 and 3|4) and "1|" is the random effect.





Figure 2 - Methodological framework applied to test the influence of different database sources to map conservation planning areas in SEAs based on SCP. The green and blue boxes are the appendices detailing the methodology and the results of the analyses, respectively. The red boxes "R." are the methodological filters used for generate workflow resistance score.

3 - Results

308 3.1 - Quantitative description of databases for SDM

309 The use of individual observation databases provided a limited number of observation data for distribution 310 modelling of many species, regardless of the taxonomic group (Table 2, Appendix 8.1). B.1). None of the GBIF 311 and SINP individual databases have more than 50 observations for all nine taxonomic groups, and there are 312 currently no programs for three of the nine taxonomic groups in the Vigie Nature databases. Local databases 313 and, to a lesser extent, regional databases are not equivalent in terms of the amount of data available for 314 different taxa for the three study sites. The difference is particularly pronounced for the SINP databases, with 315 no data available for five taxonomic groups in the T3 study site, whereas in the T1 study site, six taxonomic 316 groups have sufficient data to model over 50% of the species. At the regional scale, the number of observations 317 per species is highly variable among territoriesstudy sites for the GBIF databases and is more similar for the 318 Vigie Nature databases.

The use of combined databases (i.e. <u>"</u>All databases<u>}"</u>) increased the total number of species that can be studied. At the local scale, the proportion of species in each suitability class showed only small changes, while at the <u>nationalcountry</u> scale of France, it allowed a significant gain in species with suitable data (Table 2). Indeed, on a <u>nationalcountry</u> scale, the GBIF and Vigie Nature databases are complementary with each other. For example, the GBIF has few Chiroptera data, which is complemented by Vigie Nature data, and vice versa for Amphibian data.

At the <u>nationalcountry</u> scale, aggregation of the databases seem to provide the most suitable setup (databases and scale) for SDM analysis. Using these compiled, <u>nationalcountry</u> databases provides a large amount of data for a large number of species present in the three <u>territoriesstudy sites</u> (Table 2 and Appendix B.1).

329 3.2 - SDM and priority conservation areas analysis

330 The evaluation of SDM revealed differences among the database sources; none of which produced more 331 than 87% of satisfactory models for the two studied taxonomic groups and some had less than 20% of 332 satisfactory models (Table 3, Appendix B.2). Use of the GBIF data led to a higher proportion of well-evaluated 333 SDM, ranging between 48 and 79% of satisfactory models for the species in the two taxonomic groups. GBIF 334 data are also more suitable than Vigie Nature data, they produced between 11% and 37% more satisfactory 335 models than the latter database (Table 3). Nevertheless, between 4% and 9% of species provide well-evaluated 336 models from Vigie Nature and poorly evaluated by GBIF database. The combination of "National databases" 337 (i.e. GBIF and Vigie Nature) decreased the performance of SDMs with GBIF data, but still yield better results 338 than SDM based on the Vigie Nature database. The substitution in individual databases of pseudo-absences 339 for the absences from Vigie Nature (i.e. mixed databases) reduced the performance of models based on GBIF, 340 but increased the performance of the combined "National databases" (Figure 3, Appendix B-2). B.2). Regardless 341 of the database used, our analyses revealed significant differences between territoriesstudy sites (Table 3). No 342 SDMs for butterflies in T2 could be evaluated due to insufficient local data. For the T3 study site, over 50% of 343 the Papilionidae and Aves models perform poorly, whereas for the T1 study site poor models occur in less than 344 50% of the evaluations.

Although important differences in model performance between the database sources used for SDM showed a high degree of overlap, as indicated by Shoener's D index with values above 0.8, the ranking of habitat suitability was highly variable. This was in particularly the case for the GBIF and Vigie Nature databases that had median spearman's rank coefficient values between 0.2 and 0.5 and a very wide distribution (Appendix B.3). The substitution of pseudo-absence data in GBIF and <u>"National databases</u>" by absence data from Vigie Nature, showed a similar situation (Appendix B.4).

For priority conservation areas, whatever the individual databases used, the overlaps with Schoener's D index were above 0.72 and similar in each <u>territorystudy site</u>. Nevertheless, Sperman's rank coefficients showed a greater difference in prioritization ranks in particular between GBIF and Vigie Nature and for the T3 study site (Table 4). Between maps of priority conservation areas, we observed similarities in overlap, despite a significant difference in the hierarchy of areas to be prioritized (Figure 4, Appendix B.5). <u>The list of species is</u> presented in Appendix B.7, where it can be seen that there are no difficulties with respect to invasive species which are very few in the data sets.

358 **3.3 - Species community analysis**

359 The distribution of workflow resistance scores showed that only 30% and 42% respectively of Aves and 360 Papilionidae species were integrated in priority conservation areas maps. Among species not integrated, the 361 workflow steps filtering the most species concern the amount of suitable data for model evaluation followed 362 by the quality of the models (Figure 5). The analysis of the species community composition observed in each 363 of the three study sites in comparison with the species community integrated in priority conservation 364 areasarea identification revealed significant differences for all three study sites. For Aves communities, the 365 differences concern an under-representation of nocturnal species, large species with high dispersal capacity, 366 and species of swamp habitats and deciduous forests in relation to the observed species community in the 367 databases for the three study sites. Conversely, species that favour urban habitats, shrubland, grassland and 368 coniferous forests are over-represented in the species community of the final maps, as are species with 369 specialized diets and foraging strata (Table 5). For Papilionidae communities, common species with long flight 370 periods are over-represented in the final community used for analysis. Species related to anthropogenic and 371 thermo/meso Mediterranean habitats, and species that use a wide range of hostplants are over-represented 372 in relation to the original species community, while supra-Mediterranean species and those of montane 373 environments are under-represented (Table 5).

Table 2 - Quantitative description of the observation databases (Local SINP, GBIF, Vigie Nature) for nine taxonomic groups in the three study sites ("T1": Lodévois-Larzac, "T2": La Rochelle, "T3": Brocéliande) for "local" (study site with buffer of 10km), "Regional" (administrative region) and "FranceCountry" (continental France). "nSp" is the number of species observed at least 3 times in a given territorystudy site within a 10 km buffer zone, ">15obs" is the percentage of species with more than 15 observation and ">50obs" the percentage with more than 50 observations (red < 25%, orange 25-75%, green >75%, grey - no data).

						GBIF						SINP		Vigie Nature						All databases combined											
1	axa		Local		I	Regiona	al	Fran	ice Cou	ntry		Local			Local]	Regiona	1	Fra	nceCou	<u>ntry</u>		Local]	Regiona	ıl	Fra	nceCou	ntry
		T1	T2	T3	T1	T2	T3	T1	T2	T3	T1	T2	T3	T1	T2	Т3	T1	T2	T3	T1	T2	T3	T1	T2	T3	T1	T2	T3	T1	T2	Т3
	nSp	11	1	11	13	2	11	14	2	11	13	2	0		-	-	_	-	-	-			14	2	11	14	2	11	14	2	11
Amp	>15obs																Ν	o progra	ım												
-	>50obs																														
	nSp	100	73	77	198	159	120	204	170	131	197	94	82	114	155	101	182	143	120	201	170	132	205	171	132	205	166	132	205	171	132
Ave	>15obs																														
	>50obs																														
	nSp	4	0	5	13	13	5	19	19	21	21	6	18	25	22	24	25	22	24	25	22	25	26	22	26	26	22	26	26	22	26
Chi	>15obs																														
	>50obs																														
	nSp	786	476	639	1428	1093	825	1573	1261	995	1703	1335	843	286	177	136	825	657	413	1336	1105	848	1844	1460	1093	1844	1158	1093	1844	1460	1093
Flo	>15obs																														
	>50obs	1.0								• •							L			l											
	nSp	10	1	14	23	6	21	23	6	28	23	7	30										23	1	30	23	6	30	23	1	30
Mam	>15obs																N	o progra	ım												
	>50obs	20					20	67		20					1					T					20			20			20
Ode	nSp	38	5	23	56	/	28	57	/	30	57	1	0	0	1	24	56	/	27	57	/	30	57	/	30	57	/	30	57	/	30
Odo	>1300s																														
		53	3	24	93	25	26	95	25	34	91	8	0	25	19	19	25	19	19	25	19	19	98	25	34	98	25	34	98	25	34
Ort	>15obs	55	5	24	75	25	20	15	25	54	71	0	0	25	17	17	25	17	17	23	17	17	70	25	54	70	23	54	70	25	54
on	>50obs																														
		115	15	33	144	18	43	147	18	52	154	11	0	0	0	50	132	18	51	149	18	52	154	18	52	154	18	52	154	18	52
Pap	>15obs	115	15	55	111	10	-15	117	10	52	151		Ū	Ŭ	0	50	152	10	51	112	10	52	154	10	52	154	10	52	154	10	52
1	>50obs																														
	nSp	18	2	7	20	4	7	21	4	7	21	4	0										22	4	7	22	4	7	22	4	7
Rep	>15obs																N	o progra	ım												
-	>50obs																														

Table 3 - Proportion of well-evaluated SDM corresponding to Boyce's index greater than 0.3. "National" combines GBIF and Vigie Nature databases.

Tava	Study	Number of]	Individual datab	ase	SDM with Vigie Nature absence				
Idxd	site	species	GBIF	Vigie Nature	<u>"</u> National <u>"</u>	GBIF	<u>"National"</u>			
	T1	114	79 %	61 %	70 %	87 %	82 %			
Aves	T2	87	68 %	31 %	47 %	57 %	46 %			
	T3	56	48 %	38 %	29 %	52 %	43 %			
Danilianidaa	T1	127	75 %	52 %	64 %	75 %	73 %			
Papiliolliuae	T3	25	52 %	16 %	20 %	12 %	20 %			



Figure 3 - Proportion of SDM with Boyce's index (CBI) greater than 0.3 by database source ("Individual" and "Mixed" with Vigie Nature absence), by combining territoriesstudy sites (T1, T2, T3) and taxonomic groups (Aves, Papilionidae). "National" combines GBIF and Vigie Nature databases.

Table 4 - Overlap of priority conservation areas between database sources using two metrics: Schoener's D index (D) and Spearman's rank coefficient (S cor). VN is Vigie Nature database and "National" combines GBIF and Vigie Nature databases.

Study sites	<u>"</u> Nationa	Ind 1 <u>"</u> – GBIF	lividual da <u>"</u> Nation	atabase al <u>"</u> – VN	GBIF	- VN	Pseud datal GE	lo-absenc base - abs 3IF	e of indi ence fro <u>"</u> Nati	vidual m VN onal <u>"</u>
	D	S cor	D	S cor	D	S cor	D	S cor	D	S cor
T1	0.82	0.56	0.81	0.53	0.78	0.36	0.75	0.88	0.79	0.38
T2	0.82	0.74	0.76	0.49	0.72	0.42	0.76	0.64	0.79	0.64
T3	0.77	0.34	0.81	0.51	0.78	0.29	0.73	0.19	0.79	0.39
T1	0.85	0.64	0.84	0.65	0.83	0.59	0.72	0.66	0.82	0.66
Т3	0.80	0.49	0.77	0.31	0.78	0.22	0.76	0.17	0.72	0.06
	Study sites T1 T2 T3 T1 T3 T3	Study sites "National D T1 0.82 T2 0.82 T3 0.77 T1 0.85 T3 0.80	Study sites Ind "National" - GBIF D S cor T1 0.82 0.56 T2 0.82 0.74 T3 0.77 0.34 T1 0.85 0.64 T3 0.80 0.49	Study "Mational" - GBIF Mational" Sites D S cor D T1 0.82 0.56 0.81 T2 0.82 0.74 0.76 T3 0.77 0.34 0.81 T1 0.85 0.64 0.84 T3 0.80 0.49 0.77	Study sites Individual database "National" - GBIF "National" - VN D S cor D T1 0.82 0.56 0.81 0.53 T2 0.82 0.74 0.76 0.49 T3 0.77 0.34 0.81 0.51 T1 0.85 0.64 0.84 0.65 T3 0.80 0.49 0.77 0.31	Study sites Individual database <u>"National" - GBIF</u> <u>"National" - VN</u> GBIF D S cor D S cor T1 0.82 0.56 0.81 0.53 0.78 T2 0.82 0.74 0.76 0.49 0.72 T3 0.77 0.34 0.81 0.51 0.78 T1 0.85 0.64 0.84 0.65 0.83 T3 0.80 0.49 0.77 0.31 0.78	Study sites ''National'' - GBIF ''National'' - VN GBIF - VN D S cor D S cor T1 0.82 0.56 0.81 0.53 0.78 0.36 T2 0.82 0.74 0.76 0.49 0.72 0.42 T3 0.77 0.34 0.81 0.51 0.78 0.29 T1 0.85 0.64 0.84 0.65 0.83 0.59 T3 0.80 0.49 0.77 0.31 0.78 0.22	Study sites Mational" - GBIF Mational" - VN GBIF · VN GBI GBIF D S cor D S cor D T1 0.82 0.56 0.81 0.53 0.78 0.36 0.75 T2 0.82 0.74 0.76 0.49 0.72 0.42 0.76 T3 0.77 0.34 0.81 0.51 0.78 0.29 0.73 T1 0.85 0.64 0.84 0.65 0.83 0.59 0.72 T3 0.80 0.49 0.77 0.31 0.78 0.22 0.76	Study "National" - GBIF "National" - VN GBIF - VN GBIF - VN GBIF D S cor D S cor D S cor S cor <td>$\begin{array}{c c c c c c c c c c c c c c c c c c c$</td>	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $



Figure 4 - Maps of priority conservation areas for Aves and Papilionidae of tree study sites (T1, T2, T3) from different individual database source (All, GBIF, Vigie Nature).



Figure 5 - Workflow resistance scores for Papilionidae and Aves. 1 - species observed in the study area that did not cross any of the stepwise filters. 2 - species with sufficient data to calibrate SDM, i.e. > 15 nationalcountry observations, or evalueevaluate the performance of SDM, i.e. > 10 local observations. 3 - species with sufficient data to calibrate and evaluate the performance of the SDM. 4 - species present in the final analysis, i.e. Boyce's index > 0.3.

Table 5 - Parameter estimates (β), standard error (se) and P-values for the full model of Aves and Papilionidae species resistance to the workflow. Appendix B.6, the evaluation of the quality of the model.

		Aves			Papilioniadae							
	Variables	β	se	P-value	Variables	β	se	P-value				
	β₀ 1 2	-7.84	1.12	/	β ₀ 1 2	-2.54	1.28	/				
	β ₀ 2 3	0.14	0.33	/	β ₀ 2 3	1.39	1.26	/				
	β ₀ 3 4	1.4	0.34	/	β ₀ 3 4	2.97	1.27	/				
	Mass	-0.21	0.13		WingspanM	0.01	0.09					
	Avian hand-wing index	-0.31	0.07	***	FMoMean	0.38	0.11	***				
	Nocturn	-2.28	0.59	***	Hostplant N	0.23	0.11	*				
	Deciduous	-0.27	0.15		Hostplant Spe	0.16	0.11					
	Coniferous	0.38	0.15	*	HPG Bi	0.28	0.50					
	Woodland	0.17	0.14		HPG Th	-0.21	0.22					
del	Shrub	0.55	0.13	***	HPG Sb	-0.32	0.23					
noo	Grassland	0.47	0.13	**	HPG Tr	0.22	0.45					
ll r	Mountain meadows	0.23	0.24		HPG Li	0.02	0.36					
Fu	Reed	0.08	0.33		SSI	0	0					
	Swamps	-0.84	0.28	**	AltVeg 'A'	0.06	1.12					
	Rocks	-0.14	0.21		AltVeg 'Mo'	-0.93	0.31	**				
	Urban	0.83	0.14	***	AltVeg 'SupMed'	-0.79	0.46					
	Spe. Diet	0.92	0.35	**	AltVeg 'ThMeMed'	1.37	0.37	***				
	Spe. Foraging behav.	-0.08	0.4		Rarity '2'	0.86	0.29	**				
	Spe. Diet strat	1.49	0.4	***	Rarity '3'	1.42	0.34	***				
	Spe. Habitat	-0.14	0.38									
	Spe. Nest	-0.28	0.72									
	Spe. Mean	-2.26	1.63									

P-value: *** P < 0.001, ** P < 0.01, * P < 0.05, P < 0.1, / P-value not applicable

Trait description: Spe. is the specialization, FMoMean is duration of yearly flight period, Hostplant N is hostplant specificity, HostplantSpe is hostplant specificity index, HPG is hostplang growth form (Bi: short herb, Th: tall herb, Sb: shrub, Tr: tree, Li: Liana), SSi is Species Specialization Index, AltVeg is altitudinal vegetation (A=Anthropogenic, Co: Foothill, TheMeMed: Thermo/Meso-Mediterranean, Med: Mediterranean, SupMed: Supra-Mediterranean , Mo: Montane, Asa: Alpine and Subalpine). For more details on traits see Appendix A.3

4 - Discussion

408 The absence of recommendations for the use of available databases that differ in terms of their quality 409 and pertinence by public authorities or other institutions remains a serious problem for local conservation 410 planning (Clare et al. 2019). The goal of this study was to test the suitability of different database sources 411 that can be used by public stakeholders to map priorities for biodiversity stakes in SEAs and SCP. We found 412 that the compilation of databases at the national country scale is the most suitable procedure to apply SDM 413 to a large number of species. For Aves and Papilionidae, the GBIF database provided the highest proportion 414 of well-assessed SDM. We detected a significant overlap in species distributions in different database 415 sources despite significant variability in the order of habitat suitability and similar spatial predictions for 416 priority conservation areas. Finally, we've showed that the composition of the species community used 417 for priority conservation areas in all three study sites were clearly not representative of the observed 418 species communities in the original database (in terms of ecological traits). species and ecological traits). 419 Finally, despite important differences among the study sites in terms of the proportion of artificial land 420 cover and protected areas we found no particular differences between the three study sites. Clearly, the 421 data sources are the most important factor influencing the results.

422 Open-access biodiversity data provide a valuable source of information for decision makers, 423 environmental consultants and conservation and land-use planningmanagers (i.e. SEA stakeholders;); they 424 contain vital information on species locations compared to expert knowledge and unshared datasets that 425 are inaccessible to most users (Sousa-Baena et al., 2014; Meyer et al., 2015). Through their use and careful 426 application of SDM, they contribute to the estimation of relative habitat suitability in a given study site 427 (Baker et al., 2021). Our study showed however that SDM for a large majority of species observed locally 428 requires their compilation on a national country scale. Local and regional data are not suitable for model 429 calibration but remain important for assessing the suitability of models in a local context. Indeed, this 430 performance evaluation step is one of the most restrictive filters in the workflow we proposed, as 431 evidenced by SDM's for butterflies in the T2 study site, where no species could be evaluated. The 432 spatial extent of data collection can influence distribution modelling (Meyer, 2007), and our study 433 emphasises this importance for SDM and the use of data available on national country scale in France.

434 Different types of databases are constructed in different way - with opportunistic data collection or, in 435 some cases, as part of a scientific monitoring scheme – allowing the use of as complementary data sources 436 (Beck et al., 2013; Shirey et al., 2021). The three individual databases examined in this study are indeed 437 complementary in that when they are combined they provide a suitable source for modelling the 438 distribution of many species. Nevertheless, some groups commonly have a low numberamount of data in 439 such bases, e.g. Insecta (Troudet et al., 2017). The study of data with a fine spatial grain, as required for 440 SEA and SCP (Guisan et al., 2013), reveals information gaps for more taxonomic groups at the global scale, 441 e.g. Amphibians and Mammals (Witté and Touroult, 2017). The construction of an overall database at 442 national country scale is therefore the most appropriate way to have suitable data for SDM of different 443 taxonomic groups. Furthermore, there is a dilemma between protocolized and opportunistic data. 444 Although protocolized data are recommended for SDM (Guisan et al., 2017; Guillera-Arroita et al., 2015), 445 very often the amount of such data is low, which can be detrimental at the local scale, particularly for 446 model evaluation with data having the same sampling bias. For opportunistic data, their large number is of 447 course a positive point, however the estimating their sampling bias can be a real challenge (Botella et al., 448 2018; Fithian et al., 2015; Matutini et al., 2021) to ensure the reliability of the results.

449 SDM of Aves and Papilionidae species clearly revealed differences between the databases used for 450 modelling, with differential impacts on the identification of conservation priorities. Indeed, the high 451 overlap in species distribution overlap is high between databasedata sources, as indicated by the 452 Schoener's Schoener's D index, i.e. indicates that, regardless of the data source, species, are predicted in 453 similar environments (Warren et al., 2008). However, Spearman's ranking of habitat suitability between 454 data sources was highly variable, indicative that species' responses to environments are highly variable, as 455 are the location of favourable habitats (Warren et al., 2008). Although the use of presence-absence data is 456 advocated for SDM (Guillera-Arroita et al., 2015; Valavi et al., 2021b; Dubos et al., 2022)(Guillera-Arroita 457 et al., 2015; Valavi et al., 2021b; Dubos et al., 2022), we showed that opportunist data from GBIF provided 458 a greater number of well-assessed models at the local scale. Models using opportunist data with a target-459 group approach to generate pseudo-absences provides a sufficient quality of information on species

460 distribution (Phillips et al., 2009; Barber et al., 2022) and can be correctly used in SCP (Sofaer et al., 2019; 461 Baker et al., 2021). The lack of data at the regionallocal scale, whatever the database, does not allow us to 462 explain a better fit of models using GBIF data. Evaluating the models with a large proportion of opportunist 463 data could however bias the evaluation, but only independent data were used, which provides robust 464 validation of SDM (Matutini et al., 2021). Moreover, in contrast to Hermoso et al. (2015a), we found that 465 mixing presence-only data with absence data increased the number of misjudged models. In addition to 466 the use of the ROC curve (AUC) as a presence-only model evaluation metric by Hermoso et al. (2015a), the different results can be explained by different sampling biases between the two data types (Baker et al., 467 468 2022; Barber et al., 2022). Finally, the GBIF data seem to be more adapted to model the distribution of a 469 large number of species.

470 The notion of "garbage in – garbage out" emphasises the critical importance of the quality of data 471 (Sanders and Saxe, 2017), nevertheless, the examination of data suitability for conservation planning 472 remains rare. In addition to the above issues our study revealed the importance of attention that should 473 be paid to the representativeness of the species communities used in the models compared with the actual 474 species communities observed in the study sites. This is particularly important in the light of the finding 475 that there are marked differences between conservation priorities when different database sources are 476 employed. Indeed, the number and composition of species in the community used can influence 477 conservation priorities. Elsewhere it has been shown the difference will decline as the number of species 478 increases (Kujala et al., 2018). The methodology tested in our study is based on a data-driven approach 479 that attempts to use all available biodiversity data. This approach is data intensive, but is necessary to 480 ensure the best representation of the observed local biodiversity. We revealed that such an approach can 481 nevertheless induce a significant bias in the species community that is ultimately studied. Indeed, the 482 prevalence of data affects the composition of the modelled species as well as the accuracy of the models 483 and the evaluation of the species response (Fukuda and De Baets, 2016). Particular attention should thus 484 be paid to the representativeness of the species communities used in the models in relation to the actual 485 species communities observed in the territory under study- site.

486 Our study presents a workflow (Figure 2) for identifying biodiversity stakes using a data-driven 487 approach from open-access database sources. Land-use planningSEA stakeholders can use this workflow 488 as a step towards the rationalization of data in order to reduce the biases mentioned above. The 489 confrontation of the limits of such a workflow with the needs of SEA stakeholders could illustrate how to 490 precisely target new sources of database that should be collected according to the suitability of current 491 databases for priority groups. This workflow could be compared with the data context of another country 492 to compare our findings. Hermoso et al. (2015b) revealed that evaluation models using a new collection of 493 field data does not necessarily reduce the problems of model uncertainty. However, other databases can 494 be examined by SEA stakeholders as well as other monitoring schemes (e.g. "PopAmphibien" for Reptilian 495 and Amphibian populations in France http://lashf.org/popamphibien-2/) or negotiate the use of databases 496 that are not yet shared. An important issue is thus the integration of To overcome this data sharing 497 problem, the structuring of networks of different contributors of data and users of the databases and 498 ambitious regional policies is necessary. As evidenced by our three study sites, the quantity of local data 499 available is correlated with the number of years the SINP has been implemented. An important issue is 500 thus the integration of SEA stakeholders in the workflow we propose, and their appropriation of the 501 procedure. This could be done by a form of participatory modelling (Lagabrielle et al., 2010; Lees et al., 502 2021), where stakeholders are consulted for issues and choices such as the species to be examined. In such 503 participatory modelling it is important to avoid arbitrary choices that are neither reproducible nor 504 representative of local diversity, but rather the result of administrative or political interest. Finally, it is 505 currently recommended to use these tools to elaborate a more holistic approach to SCP (Cadotte and 506 Tucker, 2018).

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5 – Perspectives: operational implementation by SEA stakeholders

509 Spatially exhaustive and ecologically representative priority conservation areas are crucial for the 510 elaboration of SEAs that aim to limit artificialization impacts as early as possible in the planning process. 511 Empirical observations are major sources of information on biodiversity that are still rarely used by SEAs. 512 The collection of open-access databases for SEA territories provides important but incomplete knowledge 513 on species occurrence. Furthermore, their use is particularly interesting to help strategically direct 514 inventory campaigns (especially for under sampled taxa and areas), that go beyond the emphasis on rare, 515 threatened and emblematic species. These additional data would clearly improve the assessment of the 516 SDMs suitability in local territories administrative entities as our study sites. What is also interesting here 517 for SEA stakeholders is that the process of filtering species and attributing them a score allows us to 518 identify for the identification of different groups of species in terms of their needs for additional data in 519 order to undertake SDM. 520 The influence of database sources on the identification of priority conservation areas reveals the 521 importance of examining their suitability. Thus, it is necessary to be prudent in the interpretation of 522 biodiversity maps. The integration of local experts may help limit any misjudgements in the workflow

523 procedure.

524 The influence of database sources on the identification of priority conservation areas reveals the 525 importance of examining their suitability. In our study this is true for three highly contrasting study areas 526 that differ markedly in terms of the cover of protected areas and artificialisation. The problem of data 527 sources is thus typical of many areas. Thus, it is necessary to be prudent in the interpretation of biodiversity 528 maps. The integration of local experts may help limit any misjudgements in the workflow procedure. 529 Indeed, the integration of "expert" knowledge and local studies is valuable information, which is important 530 to share, and which it is important to consider in order to complete our proposal. In future studies, species 531 conservation issues for spatial prioritization could be considered by focusing on (for example) the issues 532 associated with threatened and/or invasive species. The multiple dimensions of biodiversity could be 533 analyzed within a context of limited data access and the complementarity of different facets (functional 534 and phylogenetic) in addition to a classical species-based approach (Brumm et al., 2021; Cadotte and 535 Tucker, 2018).

A data-driven approach that considers as many species as possible requires a large amount of data, biases the species communities considered and does not highlight species of particular interest as their threats and regulatory protections. It is therefore necessary to rationalize this approach, by integrating the needs and issues of local SEA stakeholders.

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Data, scripts, code, and supplementary information availability

Observation data from SINP and Vigie nature databases are not available for confidentiality reasons,
 but their link and the request process are detailed in Appendix A1. The data of GBIF are available online:
 https://doi.org/10.15468/dl.ry6uw7.

558 The data, R scripts, outputs of steps 1.2 to 4 of figure 2 are available online: 559 https://doi.org/10.5281/zenodo.7883973.

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899	Appendix
900	Appendix A - Additional material and methods
901	A.1 - Collect and series of operation on species occurrences
902 903 904 905 906 907 908 909 910 911 912 913 914 915	 Databases collection: Access to source databases: French Natural and Landscape Information System (SINP, https://inpn.mnhn.fr), each of the three study sites has its own database: https://sinp-occitanie.fr/atlas/ for Lodévois-Larzac, (T1), https://obv-na.fr/ and https://observatoire-fauna.fr/ for Brocéliande,La Rochelle (T2), https://data.biodiversite-bretagne.fr for La RochelleBrocéliande (T3). GBIF: GBIF.org (04 October 2022) GBIF Occurrence Download https://doi.org/10.15468/dl.ry6uw7 French biodiversity monitoring scheme (Vigie Nature): https://www.vigienature.fr/, the database used are the STOC for Aves, the Vigie-Chiro for Chiroptera and Orthoptera, the Vigie-Flore for Flora, the STELI for Odonata, the STERF for Papilionidae Period of 1011 years, from 01/01/2010 to 31/12/2020 Extent: continental France (without island, Corsica and overseas regions) Spatial accuracy of 0 to 50m
916 917 918 919	 Standardization of species names : French taxonomic reference TAXREF.V14 (Gargominy et al., 2021)(Gargominy et al., 2021). Keep only the species and subspecies Groups data at the species level
920 921 922 923 924	 Taxonomic filtering: from TAXREF.V14 Keep only species in nine taxonomic groups selected: Amphibia, Aves, Chiroptera, Flora, Mammalia aptera (Mammalia), Orthoptera, Odonata, Papilionidae and Reptilia Delete species not present in France so keep: "Present", "Endemic", "Subendemic", "Cryptogenic", "Introduced", "Invasive introduced"
925 926 927 928	 Data transformation and filtering: Transformation data into occurrence, i.e. transform data into presence only Geographical filtering for reduce sampling biases and delete duplicate data: spatial thinning at a distance of 50m with "spThin" package (Aiello Lammens et al., 2015)(Aiello-Lammens et al., 2015)
929 930	Identify the presence of species into each study site: at least five observations from all databases in the study site to certify the presence of the species
931 932	Standardization of the observation period between databases: for Aves filtering observations between 1 March and 15 July corresponding to the STOC period.

A.2 - Environmental variables selected for Species Distribution Models (SDM) of Aves and Papilionidae species. Variables used for SDM are the variables with collinearity removed (Pearson >0.7).

Type of varia	bles	Variables	Database	Year	Raw precision	Source	Variables used for SDM
	Climatic	19 bioclimatic variables	Chelsa	1981- 2010	1 km	Karger et al., 2021, 2017	Bio 2 (temp. range), Bio 5 (temp. warmest month), Bio 6 (min. temp. coldest month), Bio 9 (temp. driest quarter), Bio 13 (prec. wettest month), Bio 14 (prec. driest month), Bio 15(prec. seasonality)
		Elevation, mean in a buffer of fifty meters					Elevation
BiogeographyGeography	Elevation	Slope, mean in a buffer of 50 meters	BD ALTI, IGN	2014	25 m	IGN, 2022	Slope
		(TPI), mean in a buffer of 50 meters					TPI
	Wetland	Area of wetland potential: very high in a buffer of 50, 500, 2000 meters	INRAE Wetland potential	2014	50 m	Berthier et al., 2014	Wetland in buffer of 50 and 500 m
	Waterway	Linear meters of waterway in a buffer of 50, 500, 2000 meters	BD TOPO, IGN	2022	Vector	IGN, 2022	Waterway in buffer of 50 and 500m
Human occupancy	Land-use	Area of artificial, conifer forest, crop, deciduous forest, land, lawn, mineral, prairie, water in buffer of 50, 500 and 2000 meters	OSO Land Cover	2018	25 m	Inglada et al., 2019	Artificial, conifer, crops, deciduous, fruit culture, land, mineral, prairie, water in a buffer of 50 and 500 m Lawn in a buffer of 500 m, Sand in a buffer of 50, 500, 2000 m
Tunian occupancy	Hedge line	Linear meters of hedge in a buffer of 50, 500, 2000 meters	BD TOPO, IGN	2022	Vector	IGN, 2022	Hedge line in buffer of 50, 500 m
	NDVI mean	NDVI mean in buffer of 50, 500, 2000 meters	DHI NDVI	2022	10 m	CESBIO, 2021, 2022	DHI NDVI in buffer of 50 and 500m
Pollution /	Light pollution	Mean value in buffer of 500 meters of night time light average masked of 2015	VIIRS night time lights	2015	500 m	Elvidge et al., 2021	Night time light
Fragmentation	Road fragmentation	Linear meters of fragmenting of main road in a buffer of 50, 500, 2000 meters	BD TOPO, IGN	2022	Vector	IGN, 2022	Main road in a buffer of 50, 500, 2000 m

936 A.3 - Species traits of Aves and Papilionidae. Missing traits were imputed by considering evolutionary 937 938 relationships following the process of Carmona et al. (2021) with the help the R scripts of Toussaint et al. (2021).

938

	Traits	Description	Reference
	Mass	Body mass using data from males, females, and/or unspecified adults in g	Storchová and Hořák, 2018
	HWI	Avian hand-wing index (HWI), an estimate of wing shape as a proxy for dispersal ability in birds	Sheard et al., 2020
	Nocturn	Nocturnal activity	Wilman et al., 2014
Aves	Habitats (Deciduous Coniferous, Woodland, Shrub, Grassland, Mountain meadows, Reed, Swamps,	Species occupies habitat in breeding area, 1: yes; 0: no	Storchová and Hořák, 2018
	Rocks, Urban) Spe. Diet, Spe. Foraging behav., Spe. Diet strat, Spe. Hab. Spo. Nost. Spo. Maan	Specialization respectively of diet, foraging behaviour, foraging substrate, habitat, nesting site and mean of five	Morelli et al., 2020
	Wingsnan	Average length of male wingspan in mm	
	FMoMean	Average duration of vearly flight period	
	HostplantN	Hostplant specificity, number of hostplants	Middleton-Welling et al.,
	HostplantSpe	Hostplant specificity index	2020
dae	HPG	Hostplant growth form: Short herb/grass (<1 m) (Bi), tall herb/grass (>1 m) (Th), shrub (Sb), tree (Tr), liana (Li)	
oni	SSI	Species Specialization Index	Essens et al., 2017
Papili	AltVeg	Altitudinal vegetation: A=Anthropogenic, Co=Foothill, TheMeMed=Thermo/Meso-Mediterranean, Med=Mediterranean, SupMed=Supra-Mediterranean, Mo=Montane, ASa=Alpine and Subalpine	Dupont et al., 2013
	Rarity	Rarity in France between 1 very rare to 3 common species	In Moussus et al., 2019 from oreina.org and faune.france.org

The phylogenetic trees used to consider evolutionary relationships were from Jetz et al. (2012) and Wiemers et al. (2019, 2020) for the Aves and Papilionidae, respectively.

939

941 Appendix B - Supplementary results

B.1 - Description of observation databases (Local SINP, GBIF, Vigie Nature) available nine taxonomic groups
in the three study sites (T1: Lodévois-Larzac, T2: Brocéliande, T3: La Rochelle) for "local" (Local" (study
site with a buffer zone 10 km), "Regional" (administrative region) and "FranceCountry" (continental
France) scales. Statistics: "nSp" is the number of species with at least threefive observations on the
study site within a 10 km buffer; "pSp<15.00bs" and "pSp>500bs" are the percentage of
species with <15, 15 to 50 and >50 observations respectively and "-" represents no data available.

		,,		,			Loc	al					
Tovo	Statista		GBIF			SINP			Vigie Nature	nbined			
Taxa	Statiste	T1	T2	Т3	T1	T2	Т3	T1	T2	Т3	T1	T2	Т3
	nSp	11	1	11	13	2	0			-	14	2	11
A	pSp<15obs	82%	100%	100%	23%	50%	-		N		29%	50%	100%
Amphibia	pSp15.50obs	18%	0%	0%	15%	50%	-		No program		7%	50%	0%
	pSp.>50obs	0%	0%	0%	62%	0%	-				64%	0%	0%
	nSp	100	73	77	197	94	82	114	155	101	205	171	132
Avoc	pSp<15obs	87%	93%	95%	37%	96%	85%	56%	50%	54%	37%	47%	59%
Aves	pSp15.50obs	10%	6%	4%	23%	3%	15%	32%	17%	20%	22%	22%	13%
	pSp.>50obs	3%	1%	1%	40%	1%	0%	12%	33%	27%	41%	31%	28%
	nSp	4	0	5	21	6	18	25	22	24	26	22	26
Chinemtone	pSp<15obs	100%	-	100%	10%	83%	50%	12%	41%	21%	12%	41%	27%
Chiroptera	pSp15.50obs	0%	-	0%	38%	0%	44%	16%	36%	46%	19%	36%	34%
	pSp.>50obs	0%	-	0%	52%	17%	6%	72%	23%	33%	69%	23%	39%
	nSp	786	476	639	1703	1335	843	286	177	136	1844	1460	1093
Flore	pSp<15obs	81%	85%	81%	52%	48%	74%	100%	100%	100%	52%	50%	72%
FIOTA	pSp15.50obs	16%	14%	17%	30%	25%	14%	0%	0%	0%	28%	24%	17%
	pSp.>50obs	3%	1%	3%	18%	27%	12%	0%	0%	0%	20%	25%	11%
	nSp	10	1	14	23	7	30				23	7	30
Mammalia	pSp<15obs	100%	100%	100%	78%	86%	60%	No program			74%	86%	57%
iviaiiiiiaiia	pSp15.50obs	0%	0%	0%	17%	14%	20%				22%	14%	20%
	pSp.>50obs	0%	0%	0%	4%	0%	20%				4%	0%	23%
	nSp	38	5	23	57	1	0	0	1	24	57	7	30
Odonata	pSp<15obs	74%	100%	100%	28%	100%	-	-	100%	100%	28%	100%	97%
Ouoliata	pSp15.50obs	26%	0%	0%	37%	0%	-	-	0%	0%	30%	0%	3%
	pSp.>50obs	0%	0%	0%	35%	0%	-	-	0%	0%	42%	0%	0%
	nSp	53	3	24	91	8	0	25	19	19	98	25	34
Orthoptore	pSp<15obs	92%	100%	100%	71%	100%	-	24%	58%	16%	56%	68%	53%
Ormopiera	pSp15.50obs	8%	0%	0%	23%	0%	-	24%	32%	52%	18%	24%	29%
	pSp.>50obs	0%	0%	0%	6%	0%	-	52%	10%	32%	26%	8%	18%
	nSp	115	15	33	154	11	0	0	0	50	154	18	52
Papilionidaa	pSp<15obs	60%	100%	94%	21%	100%	-	-	-	88	21%	100%	73%
rapinolituae	pSp15.50obs	30%	0%	6%	19%	0%	-	-	-	12	18%	0%	27%
	pSp.>50obs	10%	0%	0%	60%	0%	-	-	-	0	60%	0%	0%
	nSp	18	2	7	21	4	0				22	4	7
Pontilio	pSp<15obs	67%	100%	100%	24%	100%	-		No program		27%	75%	100%
керина	pSp15.50obs	28%	0%	0%	33%	0%	-	l	no program		32%	25%	0%
	pSp.>50obs	6%	0%	0%	43%	0%	-				41%	0%	0%

948 B.1.A - Local scale (10km buffer zone).

949 B.1.A - Regional scale.

Shar Scare	•	I				Regional				
			GBIF		I	Vigie Natur	e	All da	tabases con	nbined
Taxa	Statistic	T1	T2	Т3	T1	T2	T3	T1	T2	Т3
	nSp	13	2	11				14	2	11
Amphihio	pSp<15obs	8%	0%	73%		No massage		7%	0%	73%
Ampinoia	pSp15.50obs	23%	50%	18%		No program		14%	50%	18%
	pSp.>50obs	69%	50%	9%				79%	50%	9%
	nSp	198	159	120	182	143	120	205	166	132
A	pSp<15obs	21%	21%	40%	31%	29%	32%	12%	16%	19%
Aves	pSp15.50obs	12%	12%	22%	21%	15%	27%	11%	10%	28%
	pSp.>50obs	67%	67%	38%	48%	56%	41%	77%	74%	53%
	nSp	13	13	5	25	22	24	26	22	26
Chiroptera	pSp<15obs	100%	100%	100%	0%	0%	8%	0%	0%	15%
	pSp15.50obs	0%	0%	0%	4%	0%	17%	8%	0%	15%
	pSp.>50obs	0%	0%	0%	96%	100%	75%	92%	100%	69%
	nSp	1428	1093	825	825	657	413	1844	1158	1093
	pSp<15obs	43%	38%	47%	95%	94%	94%	32%	40%	47%
Flora	pSp15.50obs	19%	19%	20%	5%	6%	6%	26%	19%	23%
	pSp.>50obs	38%	49%	33%	0%	0%	0%	42%	41%	31%
	nSp	23	6	21				23	6	30
	pSp<15obs	39%	33%	62%				30%	33%	50%
Mammalia	pSp15.50obs	26%	0%	33%		No program		22%	0%	23%
	pSp.>50obs	35%	67%	5%				48%	67%	27%
	nSp	56	7	28	56	7	27	57	7	30
	pSp<15obs	18%	14%	64%	77%	71%	100%	9%	14%	53%
Odonata	pSp15.50obs	30%	14%	25%	23%	29%	0%	21%	14%	37%
	pSp.>50obs	52%	71%	11%	0%	0%	0%	70%	72%	10%
	nSp	93	25	26	25	19	19	98	25	34
0.1	pSp<15obs	42%	24%	58%	4%	0%	11%	30%	4%	27%
Orthoptera	pSp15.50obs	39%	40%	42%	4%	0%	16%	22%	0%	29%
	pSp.>50obs	19%	36%	0%	92%	100%	74%	48%	96%	44%
	nSp	144	18	43	132	18	51	154	18	52
	pSp<15obs	12%	0%	37%	42%	11%	55%	7%	0%	34%
Papilionidae	pSp15.50obs	19%	11%	42%	30%	22%	41%	13%	6%	31%
	pSp.>50obs	69%	89%	21%	28%	67%	4%	81%	94%	35%
	nSp	20	4	7				22	4	7
D	pSp<15obs	20%	0%	14%				18%	0%	14%
Reptilia	pSp15.50obs	30%	25%	57%		No program		28%	25%	57%
	pSp.>50obs	50%	75%	29%				59%	75%	29%
Reptilia	pSp15.50obs pSp.>50obs	30% 50%	25% 75%	57% 29%		No program		28% 59%	25% 75%	

B.1.C - France<u>Country</u> scale.

					Fra	ance <u>Countr</u>	v			
Tono	Statistic		GBIF			Vigie Natur	re	All da	tabases cor	nbined
Taxa	Statistic	T1	T2	T3	T1	T2	T3	T1	T2	T3
	nSp	14	2	11		-	-	14	2	11
A	pSp<15obs	7%	0%	0%	N.			0%	0%	0%
Amphibia	pSp15.50obs	0%	0%	9%		No program	n	7%	0%	9%
	pSp.>50obs	93%	100%	91%				93%	100%	91%
	nSp	204	170	131	201	170	132	205	171	132
	pSp<15obs	2%	1%	1%	8%	9%	4%	1%	0%	0%
Aves	pSp15.50obs	3%	1%	1%	11%	9%	7%	1%	1%	0%
	pSp.>50obs	95%	98%	98%	81%	82%	89%	98%	99%	100%
	nSp	19	19	21	25	22	25	26	22	26
CI 1	pSp<15obs	90%	89%	90%	0%	0%	0%	0%	0%	4%
Chiroptera	pSp15.50obs	10%	11%	10%	0%	0%	0%	4%	0%	0%
	pSp.>50obs	0%	0%	0%	100%	100%	100%	96%	100%	96%
	nSp	1573	1261	995	1336	1105	848	1844	1460	1093
	pSp<15obs	25%	20%	17%	68%	61%	53%	19%	14%	16%
Flora	pSp15.50obs	18%	18%	16%	19%	23%	27%	21%	17%	16%
	pSp.>50obs	57%	62%	67%	13%	16%	20%	60%	69%	68%
	nSp	23	6	28				23	7	30
	pSp<15obs	13%	16%	18%	No program			9%	14%	23%
Mammalia	pSp15.50obs	17%	17%	28%				13%	29%	17%
	pSp.>50obs	70%	67%	54%				78%	57%	60%
	nSp	57	7	30	57	7	30	57	7	30
	pSp<15obs	2%	0%	0%	11%	0%	0%	0%	0%	0%
Odonata	pSp15.50obs	10%	14%	0%	26%	0%	13%	3%	0%	0%
	pSp.>50obs	88%	86%	100%	63%	100%	87%	97%	100%	100%
	nSp	95	25	34	25	19	19	98	25	34
	pSp<15obs	22%	12%	6%	4%	0%	0%	15%	0%	0%
Orthoptera	pSp15.50obs	28%	20%	21%	0%	0%	0%	25%	4%	9%
	pSp.>50obs	50%	68%	73%	96%	100%	100%	60%	96%	91%
	nSp	147	18	52	149	18	52	154	18	52
	pSp<15obs	4%	0%	0%	13%	5%	0%	1%	0%	0%
Papilionidae	pSp15.50obs	8%	0%	0%	18%	6%	0%	7%	0%	0%
	pSp.>50obs	88%	100%	100%	69%	89%	100%	92%	100%	100%
	nSp	21	4	7				22	4	7
D	pSp<15obs	0%	0%	0%				4%	0%	0%
Reptilia	pSp15.50obs	14%	0%	0%	l	No program	n	5%	0%	0%
	pSp.>50obs	86%	100%	100%				91%	100%	100%

953 B.2 - SDM evaluation of Aves and Papilionidae taxonomic groups for (A) three individual database sources 954 and (B) between SDM using different pseudo-absence sources. Each graphic is a pair of databases-955 (National combined GBIF and Vigie Nature). Each point is an individual species that is listed in both databases, some species are assessed only in one database (missing). A Boyce's index threshold of 0.3 956 957 is used to assess the proportion of satisfactory and unsatisfactory models conjointly for each of the two 958 database sources. The blue quadrant contains species whose SDMs are evaluated in a satisfactory 959 manner in both data bases (Boyce index > 0.3 for both databases) ; the red quadrant contains the 960 species for which SDMs are evaluated in an unsatisfactory manner in both databases (Boyce index < 0.3 for both databases). In the top-left quadrant and the bottom-right quadrants, SDM model 961 962 evaluation was satisfactory for only one database (on the in y and x axes respectively).





B.3 - Overlap between SDM from individual database sources. VN is Vigie Nature database <u>and National</u>
 <u>combined GBIF and Vigie Nature</u>.



968 B.4 - Overlap between SDM from individual database sources and SDM from mixed databases (i.e. using
 969 absence from Vigie Nature). <u>National combine of GBIF and Vigie Nature.</u>



971 B.5 - Maps of conservation planning area for Aves and Papilionidae of tree study sites (T1, T2, T3) from 972 973 different individual database source (<u>GBIF, Vigie Nature and "</u>National databases, GBIF, Vigie Nature" combining the two databases).



974 975

976 *B.6* - Evaluation of the quality of the full model (Table 5) compared to null model using AIC.

Model	Aves	Papilionidae
Full	3057	1322
Null	3357	1574

978B.7 - Species list observed in each study site (T1, T2, T3) and their French status from TAXREF.V14979(Presence, Introduced, Invasive introduced) and IUCN regional red list from BD.STATUT.V16 (LC, NT, VU,
EN, CR, DD, NA) for A) Aves species and B) Papilionidae Species.

981 <u>B.7.A – Aves species</u>

Species	Observed Erench status		IUCN regional red list				
species	<u>T1</u>	<u>T2</u>	<u>T3</u>	French status	<u>T1</u>	<u>T2</u>	<u>T3</u>
Accipiter gentilis	Yes	Yes	Yes	Present	LC	VU	<u>EN</u>
<u>Accipiter nisus</u>	Yes	Yes	Yes	Present	LC	LC	LC
Acrocephalus arundinaceus	Yes	Yes	E .	Present	VU	CR	±.
Acrocephalus schoenobaenus		Yes	Yes	Present		VU	LC
Acrocephalus scirpaceus	Yes	Yes	E .	Present	NT	VU	2
Actitis hypoleucos	Yes	Yes	Yes	Present	EN	CR	NA
<u>Aegithalos caudatus</u>	Yes	Yes	Yes	Present	LC	LC	LC
<u>Aegypius monachus</u>	Yes	E .	E .	Present	CR	2	Ξ.
<u>Aix galericulata</u>	Yes	E .	E I	Introduced	NA	Ξ.	Ξ.
<u>Alauda arvensis</u>	Yes	Yes	Yes	Present	<u>LC</u>	<u>VU</u>	LC
Alcedo atthis	Yes	Yes	Yes	Present	NT	<u>NT</u>	LC
<u>Alectoris rufa</u>	Yes	Yes	Yes	Present	DD	DD	DD
<u>Anas acuta</u>		<u> </u>	Yes	Present		2	VU
<u>Anas crecca</u>	E .	E .	Yes	<u>Present</u>	E .	z –	<u>LC</u>
<u>Anas platyrhynchos</u>	Yes	Yes	Yes	Present	DD	LC	<u>LC</u>
<u>Anser anser</u>	E L	Yes	Yes	Present	E .	VU	NA
<u>Anthus campestris</u>	Yes	Yes	Ξ.	Present	<u>VU</u>	EN	Ξ.
<u>Anthus pratensis</u>	Yes	Yes	Yes	Present	<u>VU</u>	EN	VU
<u>Anthus spinoletta</u>	Ξ	Yes	Yes	Present	Ξ	<u>LC</u>	LC
<u>Anthus trivialis</u>	Yes	Yes	Yes	Present	LC	LC	<u>LC</u>
<u>Apus apus</u>	Yes	Yes	Yes	Present	<u>LC</u>	<u>NT</u>	LC
<u>Aquila chrysaetos</u>	Yes	Ξ	Ξ	Present	<u>VU</u>	Ξ.	Ξ
<u>Aquila fasciata</u>	Yes	Ξ.	Ξ.	Present	CR	Ξ.	Ξ.
<u>Ardea alba</u>	Yes	Yes	Yes	Present	VU	NT	EN
<u>Ardea cinerea</u>	Yes	Yes	<u>Yes</u>	Present	<u>LC</u>	<u>LC</u>	<u>LC</u>
<u>Ardea purpurea</u>	Yes	<u>Yes</u>	Ξ.	Present	<u>EN</u>	<u>VU</u>	Ξ.
<u>Ardeola ralloides</u>	Yes	Ξ	Ξ	Present	<u>vu</u>	Ξ	Ξ.
<u>Arenaria interpres</u>	=	Yes	Ξ.	Present	Ξ.	NA	Ξ.
<u>Asio otus</u>	Yes	Yes		Present	<u>LC</u>	<u>LC</u>	Ξ.
<u>Athene noctua</u>	Yes	Yes	Yes	Present	<u>NT</u>	<u>NT</u>	<u>VU</u>
<u>Aythya ferina</u>	Ξ	Ξ	Yes	Present	Ξ	Ξ	<u>EN</u>
<u>Aythya fuliqula</u>		Ξ.	<u>Yes</u>	Present	=	Ξ.	<u>LC</u>
<u>Botaurus stellaris</u>	Yes	=		Present	EN	<u>-</u>	-
<u>Branta bernicia</u>		<u>Yes</u>	<u>Yes</u>	Present	-	<u>NA</u>	<u>LC</u>
<u>Bubo bubo</u>	Yes	<u>-</u>	<u>-</u>	Present		-	
Bubulcus ibis	Yes	Yes	Yes	Present		LC	<u>N I</u>
Burninus oedichemus	<u>Yes</u>	<u>Yes</u>	<u>-</u>	Present			-
<u>Buteo buteo</u>	<u>Yes</u>	res	res	Present		LC	LC
<u>Calidric alba</u>	res	<u>-</u>	Ξ.	Present	EIN	<u>-</u>	=
<u>Calidria albia</u>	=	Yes	Ξ.	Present	=		=
	<u>-</u>	res	<u>-</u>	Present	-	INA	-
Cardualis cardualis	Voc	<u>-</u> Voc	Voc	Present		<u>-</u>	
<u>Caranis daurisa</u>	Voc	165	Tes	Present		<u>IN I</u>	<u>LC</u>
<u>Certopis dudricu</u> Certhia brachydactyla	Vos			Present		<u>-</u>	<u>-</u>
Corthia familiaris	Vos	105	165	Procont			
Cettia cetti	Vos	<u> </u>	<u>-</u> Voc	Procont		<u>-</u>	<u>-</u>
<u>Cettiu cetti</u> Charadrius alexandrinus	105	Vos	105	Procont	<u>LC</u>	EN	
Charadrius dubius		Vos	-	Procont			=
<u>Charadrius hiaticula</u>	Voc	Voc		Present		VU	
Chlidonias niger	Ves	Ves		Present	NA	CR	-
Chloris chloris	Ves	Ves	Ves	Present	NT	NT	<u>-</u>
Chroicocenhalus ridihundus	Ves	Ves	Ves	Present		VII	10
Ciconia ciconia	Yes	Yes	-	Present	NT	NT	-
Ciconia niara	Yes	-	1	Present	EN	-	Ξ.
Cinclus cinclus	Yes	-		Present		1	1
Circaetus gallicus	Yes	Yes		Present		EN.	1
Circus geruainosus	Yes	Yes	1	Present	VU	VU	1
Circus cvaneus	Yes	Yes	Yes	Present	EN	NT	EN
Circus pyaraus	Yes	Yes	-	Present	EN	NT	
Cisticola iuncidis	Yes	Yes	Yes	Present	LC	NT	LC
Clamator alandarius	Yes	-	-	Present	NT	-	-
ciamator giunuunus	103	-	-	resent	1.0.1	-	-

Coccothraustes coccothraustes	Yes	Yes	Yes	Present	LC	NT	VU
<u>Columba livia</u>	Yes	Yes	Yes	Present	DD	DD	DD
Columba oenas	Yes	Yes	Yes	Present	VU	EN	LC
Columba palumbus	Yes	Yes	Yes	Present	LC	LC	LC
Coracias garrulus	Yes	-	-	Present	NT	-	
Corvus corax	Yes			Present	LC	- I.	- E -
Corvus corone	Yes	Yes	Yes	Present	LC	LC	LC
Corvus frugilegus	-	Yes	Yes	Present	-	LC	LC
Corvus monedula	Yes	Yes	Yes	Present	LC	NT	LC
Coturnix coturnix	Yes	Yes	Yes	Present	NT	VU	LC
Cuculus canorus	Yes	Yes	Yes	Present	LC	LC	LC
Cvanistes caeruleus	Yes	Yes	Yes	Present	LC	LC	LC
Cvanus olor	Yes	Yes	-	Present	NA	LC	-
Delichon urbicum	Yes	Yes	Yes	Present	IC	NT	10
Dendrocopos major	Yes	Yes	Yes	Present	10	10	10
Dendrocopos minor	Yes	Yes	Yes	Present	10	NT	10
Dryoconus martius	Yes	-	Yes	Present		-	10
Earetta aarzetta	Yes	Yes	Yes	Present		10	NT
Elanus caeruleus	-	Ves	-	Present	-	VII	-
Emberiza calandra	Voc	Voc	Voc	Present	īc	VII	EN
Emberiza cirlus	Voc	Voc	Voc	Prosont		10	
<u>Emberiza citrinella</u>	Vos	Vos	Vos	Present	NT	NT	NT
Emberiza bortulana	Voc	165	165	Present		INT	<u>IN I</u>
Emberiza sebeenielus	Vec	<u>-</u>	<u>-</u>	Present		<u>-</u>	<u>-</u>
Emberiza schoenicius	Yes	Yee	Yes	Present			<u>vo</u>
Entracus rubecula	<u>Yes</u>	<u>Yes</u>	<u>Yes</u>	Present	<u>LC</u>	LC	<u>LC</u>
Faico columbarius	<u> </u>	Yes	=	Present		<u>INA</u>	Ξ.
<u>Faico naumanni</u>	<u>Yes</u>	<u>-</u>	Ξ.	Present	<u>vu</u>	-	Ξ.
Faico peregrinus	<u>Yes</u>	<u>Yes</u>	<u>-</u>	Present		<u>CR</u>	 N.T.
<u>Falco subbuteo</u>	<u>Yes</u>	<u>Yes</u>	<u>Yes</u>	Present	<u>NI</u>	NI	<u>NI</u>
Falco tinnunculus	<u>Yes</u>	<u>Yes</u>	Yes	Present	LC	<u>NI</u>	<u>LC</u>
Falco vespertinus	<u>Yes</u>	<u>Yes</u>	Ξ.	Present	NA	<u>NA</u>	Ξ.
<u>Ficedula albicollis</u>	<u>Yes</u>	Ξ.	Ξ.	Present	<u>NT</u>	Ξ.	Ξ.
<u>Ficedula hypoleuca</u>	Yes	=	=	Present	<u>EN</u>	Ξ.	
Fringilla coelebs	<u>Yes</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>	<u>LC</u>
<u>Fringilla montifringilla</u>	Ξ	Ξ	Yes	Present	Ξ	Ξ	DD
<u>Fulica atra</u>	<u>Yes</u>	<u>Yes</u>	<u>Yes</u>	<u>Present</u>	<u>LC</u>	<u>LC</u>	<u>LC</u>
<u>Galerida cristata</u>	Yes	Yes	Ξ.	Present	LC	<u>LC</u>	Ξ.
<u>Gallinaqo qallinaqo</u>	<u>Yes</u>	Yes	<u>Yes</u>	Present	<u>CR</u>	<u>CR</u>	NA
<u>Gallinula chloropus</u>	Yes	Yes	Yes	Present	LC	NT	LC
<u>Garrulus glandarius</u>	Yes	Yes	Yes	Present	LC	LC	LC
<u>Grus grus</u>	Yes	Yes	<u> </u>	Present	CR	<u>CR</u>	
<u>Gyps fulvus</u>	Yes	<u> </u>	=	Present	VU	Ξ.	<u> </u>
<u>Hieraaetus pennatus</u>	Yes	E .	E .	Present	VU	E I	Ξ.
Himantopus himantopus	Yes	Yes	2	Present	LC	NT	<u> </u>
<u>Hippolais polyglotta</u>	Yes	Yes	Yes	Present	LC	<u>LC</u>	LC
<u>Hirundo rustica</u>	Yes	Yes	Yes	Present	NT	<u>NT</u>	LC
<u>Ichthyaetus melanocephalus</u>	-	Yes	=	Present	-	<u>CR</u>	<u> </u>
<u>Jynx torquilla</u>	Yes	Yes	E I	Present	NT	VU	5
Lanius collurio	Yes	Yes	Yes	Present	NT	NT	EN
Lanius excubitor	-	-	Yes	Present	-	-	NA
Lanius meridionalis	Yes	-	-	Present	EN		-
Lanius senator	Yes			Present	NT	- I.	- E -
Larus argentatus	-	Yes	Yes	Present	-	vŪ	VU
Larus canus	_	Yes	-	Present	_	EN	-
Larus fuscus	Yes	Yes	Yes	Present	NA	LC	LC
Larus marinus	-	Yes	-	Present	-	EN	-
Larus michahellis	Yes	Yes	1	Present	LC	VU	- T
Limosa lapponica	-	Yes	-	Present	-	NA	
Limosa limosa		Yes	1	Present	1	CR	- E -
Linaria cannahina	Yes	Yes	Yes	Present	NT	NT	10
Locustella luscinioides	105	Voc	-	Present		EN	-
Locustella naevia	Voc	Voc	Voc	Present	-	VII	10
Lophophapes cristatus	Voc	Voc	Voc	Procont		VU	
	Voc	105	105	Dresont		<u>v0</u>	<u></u>
	Voc	<u> </u>	<u> </u>	Present			<u>-</u>
	Vec	Voc	Voc	Procent			<u>LC</u> \/LL
	Voc	Vac	162	Present			VU
Lusciniu Svecica	res	165	- Voc	Present			-
	=	Ξ.	<u>res</u>	Present	=	Ξ	
Morgus morganos	=	Ξ.	Vec	Present	=	Ξ	LC NIA
<u>iviergus merganser</u>	-	 	res	Present	 N 177	-	<u>INA</u>
<u>ivierops aplaster</u>	res	res	Ξ.	Present	<u>IN 1</u>	VU	Ξ.

Milvus miarans	Yes	Yes	_	Present	IC	1C	_
Milvus milvus	Yes	-	-	Present	EN EN	-	
Monticola savatilis	Voc			Present	VII		
Monticola solitarius	Vos	-	-	Procont		-	-
	<u>res</u>	-	-	Present	<u>vo</u>	-	-
<u>IVIOTACIIIA AIba</u>	<u>Yes</u>	<u>Yes</u>	<u>Yes</u>	Present	<u>LC</u>	<u>LC</u>	<u>LC</u>
<u>Motacilla cinerea</u>	<u>Yes</u>		<u>Yes</u>	Present	LC		<u>LC</u>
<u>Motacilla flava</u>	Yes	Yes	2	Present	<u>NT</u>	LC	Ξ.
<u>Muscicapa striata</u>	Yes	Yes	Yes	<u>Present</u>	LC	NT	LC
<u>Neophron percnopterus</u>	Ξ.	Yes	Ξ.	<u>Present</u>	Ξ	EN	Ξ.
<u>Numenius arquata</u>	Yes	Yes	<u> </u>	Present	CR	EN	±.
<u>Numenius phaeopus</u>	-	Yes		Present	=	NA	=
<u>Nycticorax nycticorax</u>	Yes	Yes	E.	Present	NT	VU	±.
Oenanthe oenanthe	Yes	Yes	Yes	Present	NT	EN	EN
Oriolus oriolus	Yes	Yes	-	Present	LC	LC	-
Otus scops	Yes	-		Present	NT	-	
Pandion haliaetus	Ves			Present	VII		
Parus major	Voc	Voc	Voc	Present		īc	īc
Passar domosticus	Voc	Voc	Voc	Procent		NT	
Passer admesticus	res	res	<u>res</u>	Present	LC		
Passer montanus	<u>Yes</u>	<u>Yes</u>	<u>Yes</u>	Present	<u>IN I</u>	EN	EN
<u>Perdix perdix</u>	Yes	Yes	Yes	Present	<u>NT</u>	DD	DD
<u>Periparus ater</u>	Yes		Yes	<u>Present</u>	LC	Ξ.	NT
<u>Pernis apivorus</u>	Yes	Yes	<u> </u>	Present	<u>LC</u>	VU	<u> </u>
<u>Petronia petronia</u>	Yes	<u> </u>	2	Present	LC	2	2
<u>Phalacrocorax carbo</u>	Yes	Yes	Yes	Present	NA	VU	LC
Phasianus colchicus	Yes	Yes	Yes	Introduced	NA	DD	DD
Phoenicurus ochruros	Yes	Yes	Yes	Present	IC	IC	IC
Phoenicurus phoenicurus	Ves	Ves	Ves	Present	10	10	VII
Phylloscopus hopelli	Vos	Voc	103	Procont		NT	<u>vo</u>
Phylloscopus bollelli	Vee	Vee	<u>-</u>	Present			-
Phylloscopus collybita	<u>Yes</u>	Yes	<u>Yes</u>	Present	<u>LC</u>	<u>LC</u>	LC
Phylloscopus sibilatrix	<u>Yes</u>	=	Yes	Present	EN	Ξ.	<u>NI</u>
<u>Phylloscopus trochilus</u>	<u>Yes</u>	Yes	Yes	<u>Present</u>	<u>NA</u>	<u>CR</u>	EN
<u>Pica pica</u>	<u>Yes</u>	Yes	<u>Yes</u>	<u>Present</u>	<u>LC</u>	<u>LC</u>	<u>LC</u>
<u>Picus canus</u>	Yes	Yes	E I	<u>Present</u>	EN	<u>CR</u>	E I
<u>Picus viridis</u>	Yes	Yes	Yes	Present	LC	<u>LC</u>	<u>LC</u>
<u>Platalea leucorodia</u>	-	Yes	<u> </u>	Present	-	EN	=
Pluvialis apricaria		-	Yes	Present		-	LC
Pluvialis sauatarola		Yes	-	Present		NA	-
Podicens cristatus	Voc	-	Voc	Present	īc	-	īc
Poocilo palustris	Voc	<u> </u>	Voc	Procent		<u>-</u>	NT
	Vee	Tes	165	Present		<u>vu</u>	INT
Prunella collaris	<u>Yes</u>			Present	EN	<u> </u>	<u> </u>
Prunella modularis	Yes	Yes	Yes	Present	<u>LC</u>	LC	LC
<u>Ptyonoprogne rupestris</u>	<u>Yes</u>	Ξ.	Ξ.	<u>Present</u>	<u>LC</u>	Ξ.	Ξ.
<u>Pyrrhocorax pyrrhocorax</u>	Yes	<u> </u>	<u> </u>	<u>Present</u>	VU	<u> </u>	<u> </u>
<u>Pyrrhula pyrrhula</u>	Yes		Yes	Present	VU	2	VU
<u>Rallus aquaticus</u>	Yes	Yes	<u> </u>	Present	LC	VU	=
Recurvirostra avosetta	-	Yes		Present	-	VU	
Reaulus ianicapilla	Yes	Yes	Yes	Present	LC	LC	LC
Regulus regulus	Ves	-	Ves	Present	10		10
Piparia riparia	Vos	Voc	Voc	Procont	EN	NT	
<u>Ripula rubatra</u>	Vec	Vec	165	Present			
<u>Suxicola rubetra</u>	res	res	-	Present			-
	<u>Yes</u>	Yes	Yes	Present	<u>vu</u>	<u>IN I</u>	<u>LC</u>
<u>Scolopax rusticola</u>	<u>Yes</u>	=	=	Present		Ξ.	Ξ.
<u>Serinus serinus</u>	<u>Yes</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>NT</u>	<u>LC</u>
<u>Sitta europaea</u>	Yes	Yes	Yes	<u>Present</u>	LC	LC	LC
<u>Spatula clypeata</u>	Yes	Yes	Yes	<u>Present</u>	DD	VU	<u>LC</u>
<u>Spatula querquedula</u>	<u> </u>	Yes	<u> </u>	Present	<u> </u>	<u>CR</u>	<u> </u>
<u>Spinus spinus</u>	Yes			Present	VU		=
Sterna hirundo	Yes	Yes	-	Present	LC	VU	-
Streptopelia decaocto	Yes	Yes	Yes	Present	LC	LC	LC
Strentonelia turtur	Yes	Yes	Yes	Present	IC	VII	IC
Striv aluco	Vos	Voc	Voc	Procont		10	
Sturnus vulgaria	Voc	Vac	Vac	Brocont			10
<u>Sturnus vulgaris</u>	<u>res</u>	<u>res</u>	<u>res</u>	Present			
<u>Sylvia atricapilla</u>	Yes	<u>Yes</u>	<u>Yes</u>	Present	LC	LC	LC
<u>Sylvia borin</u>	Yes	Yes	Yes	Present	<u>LC</u>	<u>NT</u>	<u>LC</u>
<u>Sylvia cantillans</u>	Yes	Yes	E L	Present	LC	LC	E .
<u>Sylvia communis</u>	Yes	Yes	Yes	Present	<u>LC</u>	<u>NT</u>	<u>LC</u>
<u>Sylvia conspicillata</u>	=	Yes	E .	Present	=	EN	E .
<u>Sylvia hortensis</u>	Yes	±.	±.	Present	LC	E.	Ξ
<u>Sylvia me</u> lanocephala	Yes	-	-	Present	LC	-	-
Sylvia undata	Yes		Yes	Present	VU		LC
Tachybaptus ruficollis	Yes	Yes	Yes	Present	LC	LC	LC

Tachymarptis melba	Yes	E.	E .	Present	VU	±.	±.
<u>Tadorna ferruginea</u>	<u> </u>	Yes	E I	Present		NA	<u> </u>
<u>Tadorna tadorna</u>	Yes	Yes	Ξ	Present	LC	LC	<u>-</u>
<u>Tetrax tetrax</u>	Yes	±.	Ξ	Present	NT	<u> </u>	<u>-</u>
Thalasseus sandvicensis	Yes	Yes	E I	Present	VU	NT	<u> </u>
<u>Tichodroma muraria</u>	Yes	E I	E I	Present	CR	± 1	<u> </u>
<u>Tringa erythropus</u>	Ξ.	Yes	Ξ	Present	E .	NA	2
<u>Tringa nebularia</u>	E .	Yes	E .	Present	E .	NA	<u> </u>
<u>Tringa ochropus</u>	Yes	Yes	Yes	Present	NA	NA	DD
<u>Tringa totanus</u>	Yes	Yes	E .	Present	EN	VU	<u> </u>
Troglodytes troglodytes	Yes	Yes	Yes	Present	LC	LC	LC
<u>Turdus iliacus</u>	Yes	±.	Yes	Present	NA	E S	DD
<u>Turdus merula</u>	Yes	Yes	Yes	Present	LC	LC	LC
Turdus philomelos	Yes	Yes	Yes	Present	LC	LC	LC
<u>Turdus pilaris</u>	Yes	±.	Yes	Present	VU	E S	DD
<u>Turdus torquatus</u>	E .	Yes	E .	Present	E .	LC	<u> </u>
Turdus viscivorus	Yes	Yes	Yes	Present	LC	NT	LC
<u>Tyto alba</u>	E .	Yes	Yes	Present	E .	VU	DD
<u>Upupa epops</u>	Yes	Yes	Yes	Present	LC	<u>LC</u>	<u>LC</u>
<u>Vanellus vanellus</u>	<u> </u>	Yes	Yes	Present	=	VU	VU

983 <u>B.7.A – Papilionidae species</u>

Species	Obse	erved	Franch status	IUCN regio	onal red list
<u>species</u>	<u>T1</u>	<u>T3</u>	French status	<u>T1</u>	<u>T3</u>
<u>Aqlais io</u>	Yes	Yes	Present Present	LC	<u>LC</u>
Anthocharis cardamines	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
Anthocharis euphenoides	Yes	E S	Present	LC	<u>-</u>
<u>Apatura ilia</u>	Yes	E S	Present	LC	<u>-</u>
<u>Apatura iris</u>	Yes	Yes	Present	<u>NT</u>	<u>NT</u>
Aphantopus hyperantus	Yes	Yes	Present	LC	NT
<u>Aporia crataegi</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
<u>Araschnia levana</u>	Yes	Yes	Present	LC	LC
<u>Arethusana arethusa</u>	Yes	E S	Present	<u>LC</u>	<u> </u>
<u>Argynnis pandora</u>	Yes	E S	Present	<u>LC</u>	<u> </u>
<u>Argynnis paphia</u>	Yes	Yes	Present	LC	LC
<u>Aricia agestis</u>	Yes	Yes	Present	LC	LC
<u>Aricia montensis</u>	Yes	E S	Present	DD	<u>-</u>
<u>Boloria dia</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
<u>Boloria euphrosyne</u>	Yes	± 1	Present	<u>LC</u>	<u> </u>
<u>Boloria selene</u>	Yes	E S	Present	<u>NT</u>	<u> </u>
<u>Brenthis daphne</u>	Yes	E S	Present	<u>LC</u>	<u> </u>
Brenthis hecate	Yes	± 1	Present	VU	<u> </u>
<u>Brintesia circe</u>	Yes	E S	<u>Present</u>	<u>LC</u>	<u> </u>
<u>Cacyreus marshalli</u>	Yes	E S	Invasive introduced	NA	<u> </u>
<u>Callophrys avis</u>	Yes	E S	<u>Present</u>	<u>LC</u>	<u> </u>
<u>Callophrys rubi</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
<u>Carcharodus alceae</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
<u>Celastrina argiolus</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
<u>Charaxes jasius</u>	Yes	Ξ.	<u>Present</u>	<u>LC</u>	±
<u>Chazara briseis</u>	Yes		<u>Present</u>	<u>VU</u>	E .
<u>Coenonympha arcania</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
<u>Coenonympha dorus</u>	Yes		<u>Present</u>	<u>LC</u>	E .
<u>Coenonympha pamphilus</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
<u>Colias alfacariensis</u>	Yes	Yes	Present	<u>LC</u>	LC
<u>Colias crocea</u>	Yes	Yes	Present	<u>LC</u>	LC
<u>Cupido alcetas</u>	Yes		<u>Present</u>	<u>LC</u>	E .
<u>Cupido argiades</u>	Ξ.	Yes	<u>Present</u>	<u> </u>	<u>NT</u>
<u>Cupido minimus</u>	Yes	Ξ	Present	<u>LC</u>	
<u>Cupido osiris</u>	Yes	Ξ	<u>Present</u>	<u>NT</u>	<u> </u>
<u>Cyaniris semiarqus</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>NT</u>
<u>Erebia aethiops</u>	Yes	Ξ	<u>Present</u>	<u>NT</u>	<u> </u>
<u>Erebia epistyqne</u>	Yes	Ξ	<u>Present</u>	EN	<u> </u>
<u>Erebia meolans</u>	Yes	E .	Present	<u>LC</u>	2
<u>Erebia neoridas</u>	Yes	Ξ	Present	<u>NT</u>	_
<u>Erynnis tages</u>	Yes	Yes	Present	<u>LC</u>	LC
<u>Euchloe crameri</u>	Yes	E .	Present	<u>LC</u>	2
<u>Euphydryas aurinia</u>	Yes	E .	Present	<u>NT</u>	2
Fabriciana adippe	Yes	Ξ	Present	<u>NT</u>	

<u>Fabriciana niobe</u>	Yes	5	Present	<u>NT</u>	±.
Glaucopsyche alexis	Yes	z –	Present	<u>LC</u>	± 1
Glaucopsyche melanops	Yes	2	Present	LC	2
Gonepteryx cleopatra	Yes	2	Present	LC	E .
<u>Gonepteryx rhamni</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
<u>Hamearis lucina</u>	Yes	Ξ.	Present	<u>LC</u>	<u> </u>
<u>Hesperia comma</u>	Yes	E S	Present	<u>LC</u>	
<u>Hipparchia alcyone</u>	Yes	Ξ.	<u>Present</u>	DD	<u> </u>
<u>Hipparchia faqi</u>	Yes	Ξ.	<u>Present</u>	<u>LC</u>	<u> </u>
<u>Hipparchia fidia</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	LC
<u>Hipparchia genava</u>	Yes	<u> </u>	<u>Present</u>	<u>NT</u>	<u> </u>
<u>Hipparchia semele</u>	Yes	Yes	Present	<u>LC</u>	EN
<u>Hipparchia statilinus</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	EN
<u>Hyponephele lupina</u>	Yes	Ξ.	<u>Present</u>	EN	Ξ.
<u>Hyponephele lycaon</u>	Yes	Ξ.	<u>Present</u>	EN	Ξ.
Iberochloe tagis	<u>Yes</u>	Ξ	<u>Present</u>	EN	Ξ.
Iphiclides podalirius	<u>Yes</u>	Ξ	<u>Present</u>	<u>LC</u>	Ξ.
<u>Issoria lathonia</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
Laeosopis roboris	Yes		Present	<u>LC</u>	Ξ.
<u>Lampides boeticus</u>	Yes	<u> </u>	<u>Present</u>	<u>LC</u>	Ξ.
Lasiommata maera	Yes	Yes	Present	LC	CR
<u>Lasiommata megera</u>	Yes	Yes	Present	LC	LC
Leptidea sinapis	Yes	Yes	Present	LC	LC
Leptotes pirithous	<u>Yes</u>	Ξ.	Present	<u>LC</u>	Ξ.
<u>Libythea celtis</u>	<u>Yes</u>		Introduced	<u>LC</u>	
<u>Limenitis camilla</u>	<u>Yes</u>	Yes	Present	LC	<u>LC</u>
<u>Limenitis reducta</u>	Yes	Ξ.	Present	LC	Ξ.
Lycaena alciphron	Yes	Ξ.	Present	LC	<u> </u>
Lycaena dispar	Yes	<u>-</u>	Present	<u>NI</u>	-
Lycaena phiaeas	Yes	<u>Yes</u>	Present		
Lycaena tityrus	<u>Yes</u>	Yes	Present Present		<u>LC</u>
Lysanara bellargus	<u>Yes</u>	Ξ.	Present Present		Ξ.
Lysandra bispapa	<u>Yes</u>	=	Present		=
Lysunara hispana	Vec	<u>-</u>	Present		-
Malanaraja galathaa	Voc	Voc	Present		
Melanaraja lachesis	Voc	165	Present		
Melanaraja occitanica	Voc		Present		
Melanaraja russiae	Ves	-	Present	VII	
Melitaea athalia	Ves	-	Present		
Melitaea celadussa	Yes		Present		
Melitaea cinxia	Yes	Yes	Present		10
Melitaea deione	Yes	-	Present		-
Melitaea diamina	Yes	1	Present	NT	
Melitaea didyma	Yes	1	Present	LC	
Melitaea parthenoides	Yes		Present	LC	
Melitaea phoebe	Yes	Yes	Present	LC	LC
Minois dryas	Yes	-	Present	LC	-
Muschampia floccifera	Yes	-	Present	NT	
Muschampia lavatherae	Yes	- 2 -	Present	NT	
Muschampia proto	Yes		Present	NT	
Nymphalis antiopa	Yes	=	Present	<u>NT</u>	5
Nymphalis polychloros	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
Ochlodes sylvanus	Yes	Yes	Present	LC	<u>LC</u>
<u>Papilio machaon</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	LC
<u>Pararge aegeria</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
<u>Parnassius apollo</u>	Yes	Ξ.	Present	<u>VU</u>	<u> </u>
<u>Phengaris alcon</u>	Yes	2	Present	VU	2
Phengaris arion	Yes	2	Present	<u>NT</u>	2
<u>Pieris brassicae</u>	Yes	Yes	Present	<u>LC</u>	LC
<u>Pieris mannii</u>	Yes	E .	Present	DD	Ξ.
<u>Pieris napi</u>	Yes	Yes	Present	LC	<u>LC</u>
<u>Pieris rapae</u>	Yes	Yes	Present	LC	LC
<u>Plebejus arqus</u>	Yes	Yes	Present	LC	<u>NT</u>
<u>Piebejus idas</u>	Yes	Yes	Present	<u>NT</u>	CR
Polygonia c-album	Yes	Yes	Present	LC	<u>LC</u>
Polyommatus amandus	Yes	Ξ.	Present	VU	Ξ.
Polyommatus daphnis	Yes	Ξ.	Present		Ξ.
Polyommatus donulas	<u>Yes</u>	Ξ.	Present		Ξ.
Polyonimatus adrylas	<u>res</u>	Ξ.	Present		Ξ.
<u>POIVOITITIULUS ESCHERI</u>	res	2	Present		<u> </u>

Polyommatus icarus	Yes	Yes	Present	LC	<u>LC</u>
Polyommatus thersites	Yes	<u> </u>	Present Present	DD	<u>-</u>
Pontia daplidice	Yes	<u> </u>	Present	LC	<u>_</u>
Pseudophilotes baton	Yes	<u> </u>	Present	LC	<u>_</u>
<u>Pyrqus alveus</u>	Yes	<u> </u>	Present	<u>NT</u>	± 1
<u>Pyrqus armoricanus</u>	Yes	<u> </u>	Present Present	LC	<u>-</u>
<u>Pyrgus carthami</u>	Yes	2	Present	<u>NT</u>	<u>-</u>
<u>Pyrqus cirsii</u>	Yes	<u> </u>	Present Present	<u>VU</u>	<u>-</u>
<u>Pyrqus foulquieri</u>	Yes	<u> </u>	Present Present	EN	<u>-</u>
<u>Pyrgus malvae</u>	Yes	<u> </u>	<u>Present</u>	DD	<u> </u>
Pyrgus malvoides	Yes	E.	Present	LC	<u> </u>
<u>Pyrqus onopordi</u>	Yes	<u> </u>	<u>Present</u>	DD	<u> </u>
<u>Pyronia bathseba</u>	Yes	<u> </u>	<u>Present</u>	<u>LC</u>	<u> </u>
<u>Pyronia cecilia</u>	Yes	<u> </u>	<u>Present</u>	<u>LC</u>	<u> </u>
<u>Pyronia tithonus</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
<u>Quercusia quercus</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
<u>Satyrium acaciae</u>	Yes	Ξ.	<u>Present</u>	<u>LC</u>	±
<u>Satyrium esculi</u>	Yes	Ξ.	<u>Present</u>	<u>LC</u>	±
<u>Satyrium ilicis</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
<u>Satyrium pruni</u>	Yes	Ξ.	<u>Present</u>	DD	±
<u>Satyrium spini</u>	Yes	<u> </u>	<u>Present</u>	<u>LC</u>	<u> </u>
<u>Satyrium w-album</u>	Yes	2	Present	LC	± 1
<u>Satyrus actaea</u>	Yes	2	Present	VU	± 1
<u>Satyrus ferula</u>	Yes	Ξ.	<u>Present</u>	<u>VU</u>	±
<u>Speyeria aqlaja</u>	Yes	Ξ.	<u>Present</u>	<u>LC</u>	±
<u>Spialia sertorius</u>	Yes	Ξ.	<u>Present</u>	<u>LC</u>	±
<u>Thecla betulae</u>	Yes	Ξ.	<u>Present</u>	<u>LC</u>	±
Thymelicus acteon	Yes	<u> </u>	<u>Present</u>	<u>LC</u>	<u> </u>
<u>Thymelicus lineola</u>	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
Thymelicus sylvestris	Yes	Yes	<u>Present</u>	<u>LC</u>	<u>LC</u>
<u>Vanessa atalanta</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
<u>Vanessa cardui</u>	Yes	Yes	Present	<u>LC</u>	<u>LC</u>
Zerynthia polyxena	Yes	Ξ	Present	<u>LC</u>	± 1
<u>Zerynthia rumina</u>	Yes	<u> </u>	<u>Present</u>	<u>LC</u>	Ξ.