Network structure of vertebrate scavenger assemblages at the global scale: drivers and ecosystem functioning implications


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The organization of ecological assemblages has important implications for ecosystem functioning, but little is known about how scavenger communities organize at the global scale. Here, we test four hypotheses on the factors affecting the network structure of terrestrial vertebrate scavenger assemblages and its implications on ecosystem functioning. We expect scavenger assemblages to be more nested (i.e. structured): 1) in species-rich and productive regions, and 2) regions with low human impact, because the most efficient carrion consumers that promote nestedness are large vertebrate scavengers, which are especially sensitive to human persecution. 3) We also expect climatic conditions to affect assemblage structure, because some scavenger assemblages have been shown to be more nested in colder months. Finally, 4) we expect more organized assemblages to be more efficient in the consumption of the resource. We first analyzed the relationship between the nestedness of the scavenger assemblages and climatic variables (i.e. temperature, precipitation, temperature variability and precipitation variability), ecosystem productivity and biomass (i.e. NDVI) and degree of human impact (i.e. human footprint) using 53 study sites in 22 countries across five continents. Then, we related structure (i.e. nestedness) with its function (i.e. carrion consumption rate). We found a more nested structure for scavenger assemblages in regions with higher NDVI values and lower human footprint. Moreover, more organized assemblages were more efficient in the consumption of carrion. However, our results did not support the prediction that the structure of the scavenger assemblages is directly related to climate. Our findings suggest that the nested structure of vertebrate scavenger assemblages affects its functionality and is driven by anthropogenic disturbance and ecosystem productivity worldwide. Disarray of scavenger assemblage structure by anthropogenic disturbance may lead to decreases in functionality of the terrestrial ecosystems via loss of key species and trophic facilitation processes.

Keywords: carrion, consumption rate, ecological networks, global change, macroecology, NDVI

Introduction

Community structure is a key determinant of ecosystem functioning (Hooper et al. 2005, Bannar-Martin et al. 2017), biodiversity conservation (Tylanakis et al. 2010) and community stability (Landi et al. 2018). In turn, the non-random structure of communities (Guillemaut et al. 2011, de Miguel et al. 2016) is driven by numerous biological, ecological, anthropogenic and evolutionary forces (Dalsgaard et al. 2013, Martin-González et al. 2015, Ley et al. 2017). Thus, disentangling the factors affecting the organization of ecological communities is essential for understanding ecosystem functioning and prioritizing conservation efforts.

The structure of animal and plant communities can be characterized in different ways (Verhoef and Morin 2010). Traditionally, studies have focused on describing species composition and richness, but in the last few decades, analytical approaches that provide additional information have gained the attention of ecologists. Network analyses summarize communities and assemblages numerically, making community structure statistically comparable (Bascompte and Jordano 2013). Networks have thus been widely used to compare the structure of ecological communities at the global scale (Dalsgaard et al. 2013, Schleuning et al. 2014, Song et al. 2017), and several studies have revealed that the structure of ecological networks may be affected by global-scale factors such as current and past climate (Martín-González et al. 2015, Sebastián-González et al. 2015, Song et al. 2017, Albouy et al. 2019, Mendoza and Araújo 2019), or human impact (Sebastián-González et al. 2015, Mendoza and Araújo 2019). However, the effect of each factor is contingent upon the type of ecological assemblage under study (e.g. it is different for seed-dispersal and for pollination assemblages, Sebastián-González et al. 2015, Song et al. 2017). Thus, it is important to identify the processes driving these large-scale patterns for different types of assemblages.

Terrestrial vertebrate scavengers (i.e. carrion consumers) are key providers of ecosystem functions as they regulate disease and pest expansion, accelerate the cycling of nutrients, and stabilize food webs (Beasley et al. 2019). Moreover, supplementing these ecosystem services provided by scavengers requires costly carcass collection and transportation (Morales-Reyes et al. 2015). However, the efficiency of the scavenging assemblage at consuming carrion varies widely depending on factors such as scavenger species richness, composition and abundance (Mateo-Tomás et al. 2017), vegetation type (Turner et al. 2017), temperature (DeVault et al. 2004), human impact (Huijbers et al. 2015), or carcass characteristics (i.e. size; Selva et al. 2005, Moleón et al. 2015). In addition, most studies that attempt to disentangle the
factors driving scavenger assemblages have focused on small spatial scales with only one or a few study sites (Selva and Fortuna 2007, Turner et al. 2017). Recently, in a global study, Sebastián-González et al. (2019) found the number of scavenger species in an assemblage was related to the degree of human impact, with fewer scavenger species found in areas with higher human impacts. However, our understanding of the factors driving the structure of these assemblages at the global scale is still very limited.

Here, we use network analyses and the largest compilation of data to date on vertebrate scavenger assemblages at the global scale (53 studies across five continents, Fig. 1, Supplementary material Appendix 1) to identify the effects of macroecological factors in structuring vertebrate scavenger assemblages. We focus on nestedness, a feature of community organization that has already been used to describe scavenger assemblages (Selva and Fortuna 2007). In a nested assemblage, the species feeding on carcasses visited by few consumers are subsets of those species feeding on carcasses visited by more consumers. In scavenging assemblages, this pattern can emerge through facilitation, when the use of carrion by some species facilitates its use by others, such as hyaenas tearing through tough skin and thereby opening a large carcass for smaller scavengers (Álvarez et al. 1976, Kane and Kendall 2017). Previous studies evaluating the network structure of vertebrate scavenger assemblages have found a more nested organization of those assemblages featuring greater competition for resources. This can occur for example during the cold season when food resources are more valuable (Selva and Fortuna 2007), in assemblages with specialized (e.g. vultures,
different individuals of each species detected scavenging at tive abundance as the maximum number of unequivocally tive information was available, we computed scavenger rela in this study). For 47 of the 53 datasets for which quantita that were confirmed to consume carrion were included vertebrate scavengers fed on a monitored carcass (only spe gathered between 1991 and 2019.

The field or the carcass was removed by a scavenger. Data were uously monitored until only bones and/or skin remained in were prey remains after predator kills. They were continu remained six studies were based on either direct sighting which are functionally dominant through their efficiency at consuming carrion, but are especially sensitive to anthropo

Methods

Dataset

We compiled information from 53 studies in 22 countries across five continents (Fig. 1, Supplementary material Appendix 1) to describe the structure of terrestrial vertebrate scavenger assemblages using 2629 carcasses (mean number of carcasses per study ± SD: 49.6 ± 53.6; range: 6–267). We monitored the vertebrate species consuming herbivore carcasses of a wide size range (from rodents and birds weighing < 0.01 kg to ungulates weighing > 100 kg), mainly using automatic camera traps (47 out of the 53 studies) with stand-

Carcass monitoring was summarized at each study site by constructing a matrix where carcass i was represented by rows and scavenging species j by columns. We first analyzed qualitative information (i.e. species occurrence at a carcass) where each matrix element a_{ij} indicated whether scavenger species j had been detected consuming carcass i or not (scored 1 or 0). Also, for the weighted matrices, each matrix element a_{ij} was filled with the number of different individuals of each species detected at a single carcass (i.e. scavenger relative abundance).

For each carcass, we also calculated carcass depletion time as the total time (in hours) required to completely consume or remove each carcass since it was located or placed in the field. We considered it fully consumed when only parts of the skeleton were left or if taken away from the camera focus by a large scavenger (Moleón et al. 2015). In the last case, we only considered carcasses in which the part taken was small enough to be consumed completely by the scavenger. Then, we calculated the average depletion time at each site, and estimated the carcass consumption rate as the kilogram of carcass consumed per hour by dividing the average carcass weight by the average depletion time. We only calculated the carcass consumption rate for those sites that were monitored with camera traps to avoid biases.

Measuring network structure

We measured network structure using nestedness, which can be calculated both with occurrence and abundance (called weighted nestedness) data. We first calculated the nestedness of the assemblage at each study site based on species occurrences using the metric NODF (nestedness overlap and decreasing fills; Almeida-Neto et al. 2008), then weighted nestedness based on relative abundances of individuals of each species using the metric weighted NODF (WNODF; Almeida-Neto and Ulrich 2011). These metrics estimate the degree of nestedness of the matrix, ranging from 0 to 100 (i.e. perfectly nested).

All network metrics must be normalized for comparison. To do so, we compared the observed value of each matrix with the values of matrices constructed following a null model. The null model for NODF controls for the effects of species richness, number of sampled carcasses, and keeps the heterogeneity in the number of interactions across species and carcasses. In the null model for WNOF; the probability that a species consumes a carcass or that a carcass is consumed are proportional to the number of carcasses where the species was detected and the number of individuals of each species detected consuming each carcass, respectively. We used 1000 simulations for both models. We normalized the nestedness values following Montesinos-Navarro et al. (2017) as:
**rMETRIC** = \(\frac{\text{value} - \text{mean}(\text{null model})}{\text{mean}(\text{null model})}\)

where mean(null model) is the mean of all the metric values calculated using each null model. We refer to these metrics as relative nestedness (rNODF) and relative weighted nestedness (rWNODF).

We included matrix size (i.e. the total number of columns and rows in each study) and connectance (i.e. the proportion of realized interactions in relation to the total number of possible interactions) as covariates in our models (see Statistical analyses) to control for the effects of these network properties on nestedness values (Fortuna et al. 2010, Song et al. 2017). We calculated nestedness

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Table 1. Summary of the stated biogeographical hypotheses, expected pattern, the underlying processes behind it and the main supporting references. NDVI: normalized difference vegetation index, HF: human footprint.

<table>
<thead>
<tr>
<th>Hypothesis</th>
<th>Expected pattern</th>
<th>Underlying processes</th>
<th>Main references</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Network structure will be related to climatic conditions (i.e. seasonality, temperature and precipitation)</td>
<td>There is a more nested organization in cold seasonal regions where there is greater competition for resources due to limited food availability. When temperature and humidity are high, decomposers can monopolize some carcasses, affecting vertebrate scavenger's structure.</td>
<td>Selva and Fortuna 2007 DeVault et al. 2011 Allen et al. 2014 Ray et al. 2014 Sebastián-González et al. 2016</td>
</tr>
<tr>
<td>II</td>
<td>Increase in nestedness with an increase in productivity and biomass (i.e. NDVI)</td>
<td>There is a more nested organization in productive regions where scavenger rich assemblages promote higher inter specific competition for resources and facilitation processes.</td>
<td>Bailey et al. 2004 Selva and Fortuna 2007 Cusens et al. 2012 Allen et al. 2014 Moleón et al. 2014 Sebastián-González et al. 2016 Mateo-Tomás et al. 2017</td>
</tr>
<tr>
<td>III</td>
<td>Decrease in nestedness with an increase in human impact (i.e. HF)</td>
<td>The species lost more rapidly from the scavenger assemblage due to anthropogenic threats are those that consume carrion faster and promote nestedness (e.g. vultures, large carnivores).</td>
<td>Mateo-Tomás et al. 2017</td>
</tr>
<tr>
<td>IV</td>
<td>Increase in carcass consumption rate in more nested assemblages</td>
<td>The facilitation processes promoting a nested assemblage increase the efficiency in carrion consumption.</td>
<td>Sebastián-González et al. 2016</td>
</tr>
</tbody>
</table>

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Figure 2. Conceptual representation on how the loss of key species due to human impact may affect the structure of the network. Red circles represent scavenger species and grey circles individual carcasses. Solid connecting lines indicate that a scavenger species was detected consuming a specific carcass. Dashed grey lines represent secondary lost interactions caused by the loss of other species. The left graph represents a hypothetical perfectly nested assemblage and the blue lines show all the carcasses consumed by a key scavenger species (here, a vulture). On the right graph the vulture is lost due to human impacts, having several consequences on the network: First, the upper and middle carcasses are no longer available to some of the other consumers (e.g. because their skin is very thick and some species are unable to open and consume it). Second, the carcass at the bottom of the network is no longer consumed by vertebrates (and will then be consumed by invertebrates and decomposers, which are slower). Consequently, the structure of the assemblage is no longer nested. Silhouettes from The Noun Project <https://thenounproject.com>, authors: S. Laing, A. Bearne, M. Turan Ercan, H. Richir, Bluertip Design and P. Lehmann.
values with the ‘bipartite’ package of R (Dormann et al. 2009).

**Macroeological variables**

For each site, we calculated six macroecological variables (see Table 2 for details) related to our hypotheses. Some of these variables were selected because 1) they characterize the climatic conditions of a region: mean annual temperature (°C), temperature seasonality, mean annual precipitation (mm) and precipitation seasonality; 2) they are related to resource availability: normalized difference vegetation index (NDVI), a proxy of the primary productivity and biomass; or 3) they represent the degree of human impact: human footprint (HF), an indicator of human population pressure, human land use and infrastructure, and human access. HF has been calculated for two years, so we assigned to each study site the HF value closest to the date when the study was performed: 1993 (n = 3) and 2009 (n = 50).

We calculated these variables within a 20 km buffer radius around the coordinates of each study site to represent site conditions where carcasses were located. This 125 664 ha sampling area also represents habitat characteristics of the study areas where sites were located, at a biogeographical scale that minimizes dilution from nearby areas with different land uses (mean size ± SD of study sites: 106 324 ± 297 867 ha; see Supplementary material Appendix 1 for the sizes of all study sites). Most of these variables derived from a model and have a large spatial resolution, so we did not expect them to show substantial variability among scales. However, we also calculated all variables using 10 and 30 km buffers around the center of each study site to determine the spatial consistency of our results at smaller and larger resolutions, respectively. We then correlated each of the variables at the three spatial scales (e.g. temperature at 10 versus 20 km, 10 versus 30 km, 20 versus 30 km) to assess their potential variability using Pearson correlations.

**Statistical analyses**

Previous studies indicate that some variables, such as carcass weight, presence of obligate scavengers and spatial autocorrelation, may influence scavenger assemblage structure. Large carcasses are consumed by more nested scavenger assemblages (Moleón et al. 2015), thus we included the average weight of monitored carcasses (in kg) at each study site as a covariate in the model. Assemblages of obligate scavengers, such as vultures, are also known to be more nested (Sebastián-González et al. 2016), so we also included the presence of obligate scavengers as a covariate. Finally, to account for spatial autocorrelation in the structure of the scavenger assemblages, we added a spatial autocovariate term. The autocovariate term was computed from the weighted average distance of all samples, indicating the degree of spatial clustering among dependent variables. We used the ‘autocov_dist’ function from the ‘spdep’ library (Bivand 2015).

We first evaluated if the covariates affected the structure of the scavenger assemblage. We tested the effect of five covariates: two variables associated with the experimental design and study site: (a) carcass weight and (b) presence of obligate scavengers; two variables related to the network metrics: (c) matrix size and (d) matrix connectance, and one variable to account for the spatial autocorrelation of the data: (e) spatial autocovariance. To do so, we fitted one-predictor generalized linear models (GLMs) relating the covariates with the network metrics (see results of this analysis in Supplementary material Appendix 2 Table A2.2). Variables that were significantly related (p < 0.05) to scavenger assemblage structure were included as covariates in the models performed with the macroecological variables (see below).

Relationships between network structure and predictor variables may be non-linear and thus we compared linear and quadratic one-predictor models for each macroecological variable and our dependent variables using an AIC-based model selection approach. We determined whether each macroecological predictor variable should be included as quadratic in the models for assemblage structure (i.e. temperature seasonality for rNODF and both temperature seasonality and temperature for rWNODF).

We then evaluated the relationships between the structure of the scavenger assemblage given by each of our two network metrics (i.e. rNODF and rWNODF) and the six macroecological variables measuring climatic conditions, productivity and human impact (Table 2) by means of GLMs. To do so, we first estimated correlations among the predictor variables to identify highly correlated ones. Since none were highly correlated (r < 0.7 in all cases), we included all variables in the analyses. Then, we calculated variance inflation factors (VIF) for the macroecological predictor variables using the ‘car’ package (Fox and Weisberg 2011) to further assess collinearity. The only variable exhibiting VIF values exceeding three (Zuur et al. 2010) was eliminated from the model (i.e. temperature seasonality for the two metrics). In each model, predictor variables were standardized to the same scale (i.e. they were transformed to have a mean of 0 and standard deviation of 1) and matrix size was log-transformed to meet normality. We also looked for influential outliers using the ‘car’ package. We detected one outlier in the model for rNODF (a dataset with small-sized birds from USA, ID = 13, in Supplementary material Appendix 1) and we thus ran the analysis with and without it for comparison.

We fitted all possible combinations and subsets of the predictor variables and the significant covariates for each of the two network metrics separately. We selected the model with the lowest AICc, but when there was more than one model with a ΔAICc < 2 relative to the best model, we implemented a model-averaging function in the ‘MuMIn’ package (Bartoń 2013). This function averages parameter estimates across all considered models for each dependent variable where the respective parameter appeared, weighted by the relative importance of each model. We also calculated the percentage of explained deviance (i.e. the amount of variability
Table 2. Summary of the variables used in this study, their description, spatial resolution, data years, source data, related hypothesis and main reference. NDVI: normalized difference vegetation index, HF: human footprint. * In latitudes higher than 50°, CHRIPS data were not available, so the variables were calculated using this second source. ** HF was calculated at two years, so we assigned to each study site the HF value closest to the date when the study was performed: 1993 (n = 3) and 2009 (n = 50).

Table 3. Coefficient of determination (R²) for all the models. a: Supplementary material Appendix 1. b: Supplementary material Appendix 10. All analyses were performed in R 3.5.2 (R Development Core Team).

Results

Across all studies, nestedness (rNODF) values were generally positive, with only three negative values (mean rNODF ± SD: 1.08 ± 0.71; range: −0.71 to 2.90, Supplementary material Appendix 1), suggesting scavenger assemblages are generally organized in a nested pattern. However, there was not a clear structural trend for weighted nestedness (mean rWNODF ± SD: −0.10 ± 0.46, range: −1.00, 1.06), as this index presented a similar proportion of both positive and negative values. Also, rNODF and rWNODF were positively correlated (GLM, coefficient: 1.085, p < 0.0001; Supplementary material Appendix 2 Fig. A2.1).

All the macroecological variables were highly correlated at the three spatial scales studied (all r > 0.87, all p < 0.0001; Supplementary material Appendix 2 Table A2.1) indicating little effect of the spatial scale. Thus, we used the 20 km scale for all our analyses.

From the covariates assessed, both rWNODF and rNODF were related to the weight of the monitored carcasses, but none of the metrics were related to the spatial autocovariance or the presence of obligate scavengers (Fig. 3, Supplementary material Appendix 2 Table A2.2). Carcass weight affected the structure of scavenger assemblages by increasing nestedness. Also, when we evaluated the effect of the covariates associated to network properties, both nestedness metrics were related to network connectance and matrix size. Whereas network connectance decreased nestedness, larger matrices were more nested (Fig. 3, Supplementary material Appendix 2 Table A2.2).

Contrary to our Hypothesis I, most of the variables describing climatic conditions (i.e. mean temperature, temperature seasonality and precipitation seasonality) did not affect the structure of scavenger assemblages. Only precipitation was negatively related with nestedness, but this relationship was not significant after one influential outlier with a very low nestedness was removed (ID = 13 in Supplementary material Appendix 1), or when we excluded studies based on observations (Supplementary material Appendix 2 Table A2.3–A2.5). However, our Hypothesis II was supported by the data, since assemblages in areas with higher NDVI (i.e. considered a proxy of productivity) values were more explained) of each model. We estimated the relative importance of each predictor variable (w) by summing the AIC weights across all models in the set where a given variable occurred. Finally, we re-ran all the models including only the data taken using camera traps to account for differences in sampling methodology in the results.

We tested if assemblage structure was related to its scavenging function by fitting GLMs with rNODF or rWNODF values as response variables and carcass consumption rate as a predictor variable. We also included the significant covariates in the model. Consumption rate was log-transformed. We used a Gaussian distribution of errors for all GLM analyses. All analyses were performed in R 3.5.2 (R Development Core Team).
nested (Table 3, Fig. 4). Also, assemblages showed a less nested structure in regions with greater human impact (i.e. areas with higher human footprint scores; Table 3, Fig. 4). Nevertheless, the relationship between network structure and some macroecological factors, especially precipitation, was weak overall and the data showed a large dispersion (see raw data in Fig. 4). None of our hypotheses were corroborated with the weighted nestedness metrics (Table 4). Finally, carcass consumption rate was positively related to both rNODF and marginally to rWNODF, suggesting that more organized assemblages are also functionally more efficient at consuming carcasses (Table 5).
Table 3. Model (GLM) relating nestedness (rNODF) with macroecological variables. We present the model-averaged coefficients, standard error (SE) and the relative importance of each variable (w) for models with ΔAICc < 2. Significant p-values (< 0.05) are highlighted in bold. NDVI: normalized difference vegetation index. This model does not include the influential outlier found in the analysis (see Supplementary material Appendix 2 Table A2.3 for results including the outlier). Percentage of explained deviance of the model (i.e. proportion of the variability explained by the model): 46.33. Number of studies included in the model: 53.

<table>
<thead>
<tr>
<th>Hypothesis</th>
<th>Variable</th>
<th>Coefficient</th>
<th>SE</th>
<th>w</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Covariate</td>
<td>Connectance</td>
<td>−0.395</td>
<td>0.066</td>
<td>1.00</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Covariate</td>
<td>Carcass weight</td>
<td>0.065</td>
<td>0.070</td>
<td>0.23</td>
<td>0.372</td>
</tr>
<tr>
<td>I</td>
<td>Precipitation</td>
<td>−0.186</td>
<td>0.091</td>
<td>1.00</td>
<td>0.045</td>
</tr>
<tr>
<td>I</td>
<td>Precipitation seasonality</td>
<td>0.056</td>
<td>0.067</td>
<td>0.21</td>
<td>0.418</td>
</tr>
<tr>
<td>II</td>
<td>NDVI</td>
<td>0.305</td>
<td>0.091</td>
<td>1.00</td>
<td>0.001</td>
</tr>
<tr>
<td>III</td>
<td>Human footprint</td>
<td>−0.153</td>
<td>0.069</td>
<td>1.00</td>
<td>0.031</td>
</tr>
</tbody>
</table>

Discussion

We found that the network structure of scavenger assemblages was affected by global-scale anthropic and environmental factors. As predicted by our Hypotheses II and III, NDVI (as a proxy of biomass and productivity) and human footprint (as a proxy of degree of human impact) affected assemblage nestedness the most, while the white effect of climatic variables on assemblage structure (Hypothesis I) had low statistical support. This suggests that both anthropogenic disturbance and environmental factors are important drivers of the network structure of vertebrate scavengers. There is growing evidence that human impact affects the structure of trophic networks, as already detected for several taxonomic groups, including terrestrial mammals (Mendoza and Araújo 2019), diatoms in lakes (Wang et al. 2019) and reef fish (Ruppert et al. 2018). Our study broadens this result to the functional group of scavengers, revealing detrimental effects for its functioning and subsequent ability to provision ecosystem services. In particular, we found that nested structure was associated with higher consumption efficiency, relating assemblage structure with its function in the ecosystem (e.g. see Bannar-Martin et al. 2017 for other networks).

NDVI was the main variable affecting assemblage structure. Previous studies showed that a nested structure minimizes competition and allows more species to coexist on the carrion resource (Selva and Fortuna 2007, Sebastián-González et al. 2016), and that assemblages in highly productive regions tend to have a larger number of species (Bailey et al. 2004, Cusens et al. 2012). Thus, the high competition for resources in species-rich assemblages may be compensated by a nested organization. Among scavengers, nestedness may arise from inter-specific facilitation processes, which increase the efficiency of carrion consumption. For example, some species are better able than others to locate carcasses and can be used as resource indicators (e.g. mammalian scavengers watching alighting vultures; Kane and Kendall 2017), whereas others are able to open thick skins and provide access to the nutrition-rich interior of carcasses (e.g. large scavengers opening carcasses for small scavengers, Álvarez et al. 1976). These facilitation processes occur more often in species-rich productive environments (Hooper et al. 2005, Loreau and de Mazancourt 2013) and likely promote species persistence (Sebastián-González et al. 2016) and efficient nutrient cycling (Mateo-Tomás et al. 2017).

Another factor that affected the structure of vertebrate scavenger assemblages was human impact. Recently, the number of vertebrate scavenger species observed at carcasses was shown to decrease with higher anthropogenic disturbance (Sebastián-González et al. 2019). Our results suggest that anthropogenic disturbance also affects the way species are organized in the scavenger assemblage. The most nested scavenger assemblages were found in relatively pristine regions, such as Bialowieza Forest (Poland), whereas the most random assemblages were present in regions most impacted by human activities, such as in areas close to large cities (Fig. 3). Species-rich assemblages retaining the most efficient carrion consumers (e.g. vultures and large mammals; Dirzo et al. 2014, Mateo-Tomás et al. 2017) are strongly nested (Selva and Fortuna 2007), whereas the extirpation of these species breaks down this structure. It seems that defaunated assemblages lack the facilitation processes provided by key scavengers, reducing assemblage robustness to high levels of interspecific competition for carrion (Sebastián-González et al. 2016). The fact that vulture presence did not generally influence the degree of nestedness indicates that vultures are not the only key scavengers affecting scavenger assemblages in terrestrial ecosystems. Other species such as large carnivores also seem to be efficient scavengers (Moleón et al. 2015, Mateo-Tomás et al. 2017). In addition, some ‘facilitatory’ species, such as large predators (Moleón et al. 2014) could be especially vulnerable to human disturbance. Thus, further studies should try to identify the traits that characterize these key species, to uncover the most appropriate management strategies aiming to preserve the ecosystem functions provided by scavengers (Buechley and Şekercioğlu 2016).

Besides the effect of species loss, human-impacted regions may provide persisting scavengers with an increased availability of alternative foods through anthropic subsidies (Moreno-Opo and Margalida 2019), reducing competition for carrion (Oro et al. 2013). In these humanized regions, facilitation processes lose their relevance and scavenger assemblages may be more randomly organized. Moreover, the effects of human-related disturbances are expected to increase where human populations continue to grow (United Nations 2019), threatening the maintenance of ecosystem functions and services provided by scavengers (DeVault et al. 2016). The metric
used to measure human impact (HF) is based on a combination of information on population density, infrastructure and human access. That provides a good general descriptor of anthropogenic disturbance but does not permit the identification of specific factors affecting scavenger assemblages. Such factors could include poisoning of carcasses for retaliatory killing of predators (Ogada et al. 2019) or the use of veterinary drugs such as diclofenac (Green et al. 2016). Thus, we recommend further studies to determine the relative importance of specific anthropic drivers affecting the structure of scavenger assemblages (Buechley and Şekercioğlu 2016).

The nested organization of the assemblage was also related to faster carrion consumption rate. This relationship has already been detected for five Spanish sites (Sebastián-González et al. 2016), but this is the first time that it has been recorded at a global scale. Our results support the view that nestedness is driven by facilitation processes among the species that consequently increase carrion consumption efficiency (Selva and Fortuna 2007, Sebastián-González et al. 2016). Moreover, the negative effect of human activities on assemblage structure will also reduce the efficiency in the provisioning of scavenging functions by scavenger species, with important implications for the functioning of the ecosystem and the services it provides. However, further research is needed to explore the comparative influence of assemblage attributes, such as composition and structure, and other external factors on scavenging efficiency of different carcass types and sizes. These studies should consider finer estimates of consumed carrion biomass and additional efficiency-related metrics, such as carcass detection time.

Our prediction that the structure of scavenger assemblages would be directly related to climatic conditions was not supported by the analyses. Although mean annual rainfall was negatively correlated with nestedness, the effect was weak and the variability large (Fig. 4). Climatic conditions can affect food availability and thus competition for carrion resources (Selva and Fortuna 2007, Sebastián-González et al. 2016), but other non-climatic factors such as productivity and human impact seem to be more important drivers of assemblage structure. Also, climate may indirectly affect the structure of scavenger assemblages through its effects on NDVI values (Pei et al. 2019). Moreover, we focused on vertebrate scavengers because they rapidly consume large amounts of carrion (e.g. > 90% carrion available, DeVault et al. 2011), particularly during colder months. However, at high temperatures decomposers become strong competitors and can monopolize some carcasses (DeVault et al. 2011, Ray et al. 2014), thereby influencing the organization of the vertebrate scavenger assemblage.

Interestingly, the weighted metric of nestedness showed weaker trends than the one based on occurrences. Even if both metrics were significantly related, weighted nestedness was not related to any macroecological variable and did not support any of our hypotheses. This may be partially related to the methodology used to quantify the number of individuals consuming a carcass, which underestimates the abundance of those species that do not have distinct marks, color patterns, age/sexual dimorphism, or are solitary foragers and thus rarely gather at carcasses. Thus, there may be geographical differences in the relative abundances depending on
what species are present in the different parts of the world and if individuals of these species can be easily identified. Moreover, weighted nestedness is less prevalent in ecological assemblages than nestedness (Sebastián-González et al. 2015) because it requires that both the assemblage and its weights (here, relative abundances) are nested (Almeida-Neto and Ulrich 2011). Indeed, in this study, the mean rWNODF value was close to zero, suggesting that vertebrate scavenger assemblages do not follow a weighted nested pattern and thus a lack of macroecological trends is not surprising.

Finally, it is important to underline that the effects of some of the variables were weak and the data were highly variable. This reflects the complexity of scavenger systems across the globe and the challenges of making predictions about the effects of human development and climate change on their structure. That said, our dataset is the largest ever compiled and analyzed for vertebrate scavengers, and included a wide variety of arid and mesic terrestrial ecosystems distributed across temperate and tropical regions, making our results representative of a large diversity of scavenger assemblages around the world. We show for the first time that the network structure of scavenger assemblages is affected by global-scale factors such as the degree of human impact (i.e. HF), a combination of ecosystem productivity and biomass (i.e. NDVI) and nestedness increases scavenger efficiency, an important ecological function and ecosystem service. These findings can help to inform global conservation planning aimed at maintaining the essential ecosystem functions provided by vertebrate scavengers in terrestrial ecosystems.

Data availability statement

All data necessary to reproduce the analyses presented in this study are included in the Supplementary material Appendix 1 and available from Figshare Digital Repository: <https://doi.org/10.6084/m9.figshare.12179880.v1> (Sebastián-González et al. 2020).

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<table>
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<th>Hypothesis</th>
<th>Variable</th>
<th>Coefficient</th>
<th>SE</th>
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<th>p-value</th>
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<td>Covariate</td>
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Table 5. Models (GLM) relating nestedness (rNODF) or weighted nestedness (rWNODF) with carcass consumption rate (log-transformed). We present the model coefficients, standard error (SE) and the p-values. Significant p-values (< 0.05) are highlighted in bold.
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Author contributions – ESG and JASZ designed the research; each author/s collected the data from their site (see Supplementary material Appendix 1); ESG performed the analyses and led the writing; all authors read and revised the manuscript.

References

Appendix 1–2. Supplementary material (available online as Appendix ecog-05083 at www.ecography.org/appendix/ecog-05083>). Appendix 1–2.