Extinctions stemming from environmental change often trigger trophic cascades and coextinctions. Bottom–up cascades occur when changes in the primary producers in a network elicit flow-on effects to higher trophic levels. However, it remains unclear what determines a species’ vulnerability to bottom–up cascades and whether such cascades were a large contributor to the megafauna extinctions that swept across several continents in the Late Pleistocene. The pathways to megafauna extinctions are particularly unclear for Sahul (landmass comprising Australia and New Guinea), where extinctions happened earlier than on other continents. We investigated the potential role of bottom–up trophic cascades in the megafauna extinctions in Late Pleistocene Sahul by first developing synthetic networks that varied in topology to identify how network position (trophic level, diet breadth, basal connections) influences vulnerability to bottom–up cascades. We then constructed pre-extinction (~80 ka) network models of the ecological community of Naracoorte, south-eastern Sahul, to assess whether the observed megafauna extinctions could be explained by bottom–up cascades. Synthetic networks showed that node vulnerability to bottom–up cascades decreased with increasing trophic level, diet breadth and basal connections. Extinct species in the Naracoorte community were more vulnerable overall to these cascades than were species that survived. The position of extinct species in the network – tending to be of low trophic level and therefore having relatively narrow diet breadths and fewer connections to plants – made them vulnerable. However, these species also tended to have few or no predators, a network-position attribute that suggests they might have been particularly vulnerable to new predators. Together, these results suggest that trophic cascades and naivety to predators could have contributed to the megafauna extinction event in Sahul.

Keywords: biotic interactions, coextinction, ecological network, extinction event, food web, Late Pleistocene

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Introduction

Of all the extinctions that have ever occurred on Earth, many – potentially most – have been coextinctions (Dunn et al. 2009). In some cases, these coextinctions involved host-specific parasites that were doomed by the extinction of their host species or flowering plants imperilled by the extinction of their pollinators (Koh et al. 2004). Coextinctions have also been mediated through trophic interactions between herbivores and vegetation and between predators and their prey (Estes et al. 2011, Colwell et al. 2012). Therefore, if we are to understand past extinction events and predict future extinctions, we need to be able to infer coextinction cascades accurately.

Changes in the primary-producer component of a community can trigger bottom-up cascades and profoundly alter ecological communities (Kagata and Ohgushi 2006). However, it is unclear which species are most vulnerable to bottom-up cascades. On the one hand, it has been argued that top predators and species from high trophic levels are particularly sensitive to food-web perturbations and reductions in habitat area/primary productivity (Schoener 1989, Didham et al. 1998, Purvis et al. 2000b, Duffy 2003). However, others have concluded that changes in the diversity or nutrient content of primary producers most strongly affect herbivores, and the cascading effects on higher trophic levels are dampened by trophic distance (Dyer and Letourneau 2002, Gruner 2004, Kagata et al. 2005, Scherber et al. 2010). Similarly, extinction-risk assessments by the International Union for Conservation of Nature (IUCN) and recorded recent extinctions suggest that herbivorous terrestrial vertebrates are particularly vulnerable to extinction (Artwood et al. 2020), a pattern that might partly be explained by the sensitivity of lower trophic levels to bottom-up cascades. The uncertainty regarding how vulnerability to bottom-up cascades varies with species traits (such as trophic level) has limited our ability to assess the importance of bottom-up cascades in past extinction events and to predict how such cascades might unfold in the future.

Identifying the vulnerability of species to bottom-up (or top-down) coextinction cascades relies heavily on understanding species interactions within an ecological community. To this end, ecological network modelling is an invaluable tool for representing ecological communities from the perspectives of species interactions and for studying the consequences of changes in these interactions (Säterberg et al. 2013). In ecological network models, organizational groups (e.g. species, age groups, populations or individuals) are represented by nodes, and interactions – which can be weighted or unweighted – are represented by links (edges). The interaction type most frequently used to build ecological network models is trophic interactions (i.e. food webs). For contemporary communities, there is a growing number of studies that use detailed information on species interactions to build network models and to study trophic cascades (Coll et al. 2008, Boit et al. 2012, Pocock et al. 2012). Unfortunately, similar approaches are challenging to apply to palaeo-communities because of the lack of data on ancient trophic interactions. However, by combining contemporary and palaeo-data to infer trophic interactions, this limitation can be overcome and network models of palaeo-communities can be constructed (Roopnarine 2006, Dunne et al. 2014). For example, Pires et al. (2015) used this approach to model Late Pleistocene mammal communities in the Americas, concluding that 1) pre-existing American mammal networks were not especially unstable (i.e. they were similar to modern networks in Africa in terms of population densities re-establishing after simulated perturbations) and 2) the arrival of humans destabilized the networks because this new predator increased network connectance (i.e. the proportion of potential links that are realized). Investigations of palaeo mass extinctions are fortified through the use of network modelling, because these methods provide insights into the causes and consequences of extinction events. Such events can also be used as a means of validating modelling methods because the outcomes (i.e. extinctions) are known. Despite these opportunities, the application of network modelling to investigate palaeo-extinction events remains underutilised.

Megafauna (animals > 44 kg) extinctions swept across several continents during the Late Pleistocene (126 000–12 000 years ago), with the highest proportions of genera lost from Sahul (landmass including Australia and New Guinea) and the Americas (Barnosky 2004, Koch and Barnosky 2006). While the causes of these extinctions are still debated, most evidence points towards the arrival of anatomically modern humans and/or climate change (Lorenzen et al. 2011, Metcalf et al. 2016, Saltré et al. 2019). Irrespective of the root causes, large extinction events such as these always involve both primary and secondary (or co-) extinctions (Colwell et al. 2012). Indeed, it has been argued that the loss of prey species led to large predators going extinct in the Late Pleistocene (Ripple and Van Valkenburgh 2010, Galetti et al. 2018). Although the arrival of modern humans and/or climate change have been identified as the most likely ultimate causes of megafauna extinctions in the Late Pleistocene, vegetation change associated with human arrival and/or climate change has been identified as a pathway through which these ultimate causes could have triggered extinctions (i.e. via bottom-up trophic cascades) (Miller 2005, Faith 2011, Villavicencio et al. 2016, Araújo et al. 2021).

The Late Pleistocene megafauna assemblage of Sahul was distinct from that of other continents in that all the large mammals were marsupials or monotremes (Johnson 2006). Giant reptiles and birds were also a prominent component of the continent’s megafauna (Stuart 2015). While Sahul’s megafauna included many species over the standard body-mass threshold of 44 kg, the term ‘megafauna’ is often extended to include species with a body mass above that of their surviving relatives (Johnson 2006) – a definition we have adopted here. Identifying the pathways by which Sahul’s unique megafauna were lost is challenging because their extinctions happened much earlier in Sahul than elsewhere (Johnson et al. 2016). To characterize such ancient extinction events, a sufficient number of dated fossilized
remains is necessary (Peters et al. 2019). The most detailed and well-studied fossil record spanning the megafauna extinction event in Sahul comes from the Naracoorte region (Meintangk Country) in south-eastern Australia (Fig. 1). This fossil record offers an exceptional picture of the species living in the region over the past 500 000 years, including the ecological community at the time of the main megafauna extinction event that occurred approximately 44 000 years ago in Naracoorte (Saltré et al. 2019). Thus, the Naracoorte fossil record is the best platform available from which to model the ecological and environmental processes potentially involved in megafauna extinctions in Sahul.

We assessed how vulnerability to bottom–up cascades varies with network-position attributes and whether bottom–up coextinction cascades stemming from the loss of basal resources (i.e. primary producers/plants) could have played a role in the megafauna extinctions of Sahul. First, we built small, synthetic networks (2–20 nodes) varying in topology (i.e. structure of connections) and calculated each node’s coextinction vulnerability using two methods: 1) simulation and 2) Bayesian networks. This allowed us to test the general influence of trophic level, diet breadth and number of connected basal resources on vulnerability to bottom–up coextinction cascades.

Next, we used Naracoorte as a model system to assess whether bottom–up coextinction cascades could explain which species went extinct during the Late Pleistocene event. We constructed an entire terrestrial, palaeo-vertebrate assemblage (including all terrestrial vertebrate classes) and combined this assemblage with palaeo and contemporary data to infer trophic interactions and build network models (Fig. 2). These network models consisted of nodes (species) with directed, unweighted links. We then computed each species’ vulnerability to coextinction via bottom–up cascades using the simulation method we validated with the synthetic networks, and we compared the coextinction vulnerabilities (and the traits influencing them) between extinct and extant (surviving into the Holocene) species. In addition to vulnerability to bottom–up coextinction cascades, we also tested for differences in the network positions of extant and extinct species to determine if the relative position could have made extinct species more vulnerable in other ways (e.g. more vulnerable to

Figure 1. Map showing Sahul (top right insert) and the Naracoorte bioregion/Naracoorte Caves in south-eastern Sahul (main figure). These maps show coastline/sea level as they were during the Last Glacial Maximum (LGM; approximately 19 000–26 500 years ago). The grey area and thick outline indicate the land area during the LGM, the thinner lines show present–day coastlines and borders between countries and Australian states and territories, and the green area highlights the Naracoorte bioregion.
a new predator). By incorporating interactions involving all vertebrate species, we adopted a holistic approach to studying megafauna extinctions of the Late Pleistocene.

Methods

We aimed to: 1) identify how vulnerability to bottom–up cascades varies with trophic level, diet breadth and basal connections using synthetic networks and 2) develop and interrogate ecological network models representing the Naracoorte ecosystem prior to the main megafauna-extinction pulse and the arrival of humans in the region (~44 000 years ago) (Saltré et al. 2019). Below, we describe how we generated the synthetic networks, measured node vulnerability to bottom–up cascades, built ecologically realistic models of the Naracoorte network and tested whether bottom–up coextinction cascades could explain which species were lost from the Naracoorte network (see the Supporting information for a flowchart of methods for the construction and analysis of the Naracoorte network).

Synthetic networks

We randomly generated 1000 ‘synthetic’ networks that differed in topology. The networks varied in size (number of nodes: 2–20), number of links (1–92) and connectance (0.06–0.5). Within these networks, nodes varied in trophic level (1–4), number of ‘in’ links (1–8) and the number of basal nodes to which non-basal nodes were directly or indirectly connected through ‘in’ links (1–8). We generated these networks to test how a node’s vulnerability to the removal/extinction of basal nodes (analogous to plants/primary producers) varied depending on the node’s trophic level, diet breadth (number of ‘in’ links) and the number of basal resources to which it was connected. We calculated a consumer-node’s trophic level as the average trophic level of its resource nodes + 1, and we assigned basal/plant nodes trophic level 1.

Coextinction vulnerability in synthetic networks

We inferred vulnerability of nodes to bottom–up cascades via: 1) simulations and 2) a Bayesian network method (Eklöf et al. 2019). We inferred vulnerability using the following steps:

1. **Simulations:** We performed simulations to estimate the expected loss of nodes in the network. Each simulation involved the removal of a random node from the network, followed by the calculation of the number of nodes that would be lost due to the cascading effect of the removal.
2. **Bayesian network method:** We used a Bayesian network method to infer the most likely sequence of node removals that would lead to the observed extinctions. This method involved constructing a probabilistic model of the network and using this model to infer the most likely sequence of events that led to the observed extinctions.

Figure 2. Example of an inferred ecological network model of the Late Pleistocene Naracoorte assemblage. Small points represent vertebrate species (nodes) and lines represent trophic interactions (links). Point colour shows trophic group (e.g. herbivores, carnivores etc.), and extinct nodes are black. Due to uncertainty regarding trophic interactions, we inferred and analysed 1000 versions/models of the Naracoorte network. Plants, invertebrates and fish are shown as single large points in this figure because the study focuses on terrestrial vertebrates.
We applied two different approaches because there is uncertainty regarding the best methods to infer vulnerability to bottom–up cascades (Colwell et al. 2012), so consensus using both approaches would provide more robust results than relying on one method alone. In the simulation method, primary extinctions occur by randomly removing basal nodes from the network, after which coextinctions are simulated by removing nodes that had lost all their ‘in’ links. Based on 1000 iterations of each network, we calculated the average coextinction vulnerability of each node as the proportion of total basal resources remaining when coextinction occurred.

Bayesian networks are (acyclic) networks where nodes correspond to a set of variables, and links indicate the conditional dependencies among variables. Thus, in a Bayesian representation of a food web, nodes correspond to species (with an associated variable indicating whether the species is extinct or extant), while links correspond to feeding interactions. The value attributed to each node variable (i.e. the extinct/extant status of a given species) is a function of the status of its resources (i.e. the parent nodes). Such a function models how consumers respond to resource loss, with possible responses being linear and non-linear, or topological (consumers’ probability of coextinction is 0 until all resources are lost, in which case, the probability goes to 1). Bayesian networks can be ‘solved’, in that there are various methods available to compute the exact values for each node. This gives an analytical measure of the coextinction vulnerability of all species in a food web depending on the extinction probability of their resources. The approach also assigns a specific baseline extinction probability to each node, representing species’ intrinsic vulnerability to (primary) extinction, i.e. the probability of going extinct regardless of resource availability (Eklöf et al. 2013). We applied this approach to our synthetic networks. Since we were interested in exploring the effects of primary extinctions in the plant community specifically, we assigned a baseline extinction probability of 0.1 to each basal node and of 0 to all other nodes, and we modelled co-extinction using the topological approach.

Analysis of synthetic networks

We fit mixed-effects models to the results from the simulation and Bayesian network approaches separately using the lme4 (Bates et al. 2015) and MuMIn packages (Barton 2020) to assess if the two methods yielded similar results in terms of the effects of trophic level, diet breadth and basal connections on vulnerability to bottom–up coextinction cascades. That is, we did regression analyses to identify the effects of network position on coextinction vulnerability measured by the two methods and tested whether these effects were consistent across methods. Prior to fitting, we scaled the independent variables so that the units of the regression coefficients were the same for all variables. The full/global mixed-effects model had coextinction vulnerability as the dependent variable (response), trophic level, diet breadth, number of connected basal nodes and the interactions between these traits as independent variables (fixed effects), and network identity as a random effect. We compared support for the full models with every combination of nested/reduced model using Akaike’s information criterion weights (\(\Delta AIC\)). If including trophic level, diet breadth and number of basal connections resulted in models with higher \(\Delta AIC\), this suggests these variables affect vulnerability. We calculated model-averaged (via \(\Delta AIC\)) coefficients for the independent variables to identify how each variable influenced vulnerability. We also extracted marginal R\(^2\) from three mixed-effects models that had either trophic level, diet breadth or basal connections as the only independent variable to estimate how much variation in vulnerability each of these variables explained.

Naracoorte study region

The World Heritage-listed (from 1994) Naracoorte Caves in south-eastern South Australia (Meintangk Country; 37°02′24″S, 140°48′00″E; Fig. 1) encompass a series of limestone caves that opened and closed to the surface at different locations, and at different times, over the last 500 000 years (Reed and Bourne 2009). These openings acted as natural pitfall traps, capturing snapshots of Naracoorte’s biodiversity at different periods from at least 500 000 years ago to the present. In addition to the fossils of the many animals that fell into these natural traps, there are remains of species that lived in the caves, such as owls and bats, and their prey (Macken and Reed 2013). Consequently, the Naracoorte Caves provide an ideal platform from which to build palaeo-ecological network models to gain insight into how these long-lost ecosystems functioned and changed over time.

Species data

To build a species assemblage list (i.e. to identify the nodes to include in the network models), we used two data sources: FosSahul 2.0 (Rodriguez-Rey et al. 2015, Peters et al. 2019) and the Atlas of Living Australia. FosSahul 2.0 is a database of dated fossil records from Sahul, including an automated quality-rating method for date reliability (Rodriguez-Rey et al. 2015, Peters et al. 2019). We extracted and vetted records from the Naracoorte region (defined as the region between 35°32′48″S and 38°6′50″S, and between 139°10′42″E and 141°0′21″E) from FosSahul 2.0 that had high- or intermediate-quality dates (A*, A or B) (Peters et al. 2019) and whose age was younger than 200 000 years before present. We chose this threshold age to provide a large-enough period to include dated fossils from all/most megafauna species that lived in the region immediately prior to the main extinction event; if we had made the period too narrow, few megafauna species would have been captured despite their likely presence in the region at the time. Fossil records suffer from taphonomic biases (biases in the accumulation and preservation of different organisms), and, consequently, some species that were present in Late Pleistocene Naracoorte are unlikely to be represented in the fossil record. Furthermore, there are biases for studying and dating particular groups of species due to academic and amateur interests (Starrfelt and Liow 2016).
Extinct megafaunal species were not included in any of the aforementioned databases, so for these species, we obtained body mass and diet data from the literature (Supporting information). The 423 vertebrate species in the Naracoorte assemblage included 125 that consumed vertebrates, 249 that consumed plants, 362 that consumed invertebrates and 48 that consumed fish (Supporting information). Of the 423 species, 273 consumed more than one of these resource groups. Mean species body mass ranged from 0.4 g to 2700 kg (Supporting information).

Inferring trophic links

For almost all extant species, information on trophic interactions is incomplete but the lack of knowledge regarding trophic interactions is even greater for long-extinct species. To overcome this limitation and build realistic ecological network models, various methods have been developed for inferring trophic links using species’ traits such as body size (i.e. larger predators tend to eat larger prey, and predators are usually bigger than their prey) (Gravel et al. 2013). Others have built on this approach to improve prediction accuracy (Eklöf et al. 2012, Brousseau et al. 2018, Brose et al. 2019, McLeod et al. 2020) by adding phylogenetic and physiological information, prohibiting impossible or unlikely links, incorporating specific morphological features such as biting force and cuticular thickness and taking into account abundance. However, most of this research focuses on fish or invertebrates, with the effectiveness of these methods rarely applied or validated for terrestrial vertebrates (but see Morales-Castilla et al. 2015, Pires et al. 2015).

We therefore developed and validated a method based on the body-size trophic-niche model (Gravel et al. 2013) to infer trophic links between terrestrial vertebrates. The body-size trophic-niche model that we adapted consists of two quantile regressions: 1) one defining the upper prey-size limit given predator mass and 2) the other defining the lower prey-size limit given predator mass. If a species falls within the upper and lower limits for a particular predator, it is inferred as potential prey for that predator. We used a large predator–prey interaction data set to identify these body-size relationships between terrestrial predators and their prey and tested whether these relationships varied depending on the predator’s taxonomic class (i.e. did including predator class as an independent variable improve the fit of the body size quantile regressions?).

We extracted the interaction dataset from GloBI, an online repository of biotic interactions (Poelen et al. 2014). The data set consisted of 3893 records: 958 records of predation by non-marine mammals, 2711 by birds, 199 by reptiles and 25 by amphibians (Supporting information; Llewelyn et al. 2020). We extracted data on mean body mass for the species from the same databases we used to add this information to the Naracoorte species list. Once we identified the best trophic-niche model (i.e. the combination of quantile regressions that best fit the upper and lower prey-size limits) using the entire GloBI data set and Bayesian information criterion (BIC; Supporting information), we validated this method...
of assigning trophic links by comparing model performance when applied first to the GloBI data divided into training and validation data sets and then to a well-resolved trophic network from the Serengeti (de Visser et al. 2011) (see also S. de Visser unpubl.; Supporting information). We used the true skill statistic to evaluate model performance and found that the top-ranked model (according to BIC) also performed best at assigning links in the validation step (Supporting information; Llewelyn et al. 2020). We used the best trophic-niche model to identify potential prey for each predator of vertebrates (Supporting information, Llewelyn et al. 2020).

While including additional species traits could improve the accuracy of inferred predator–prey interactions, we used only three readily available traits (diet – whether they consumed plants, invertebrates, terrestrial vertebrates and/or fish; body size; and predator taxonomic class – ‘mammals’, ‘birds’, or ‘reptiles and amphibians’). We used only these three traits because: 1) using few traits is compatible with the quantile regression framework; 2) they can easily be extracted for vertebrate species in most assemblages (and therefore the method can be widely applied); and 3) our validation steps demonstrated the resulting performance of the trophic-niche model is sufficient (true skill statistic [TSS] = 0.6 when applied to the Serengeti assemblage; TSS varies from −1 to 1, with a score of 0 indicating no better than random; Supporting information).

**Naracoorte networks**

To build realistic ecological networks for the Naracoorte assemblage, we applied the trophic-niche model to the species list, removed excess links between vertebrates (to account for overestimating the number of predator–prey links) and added links to non-terrestrial vertebrate food resources. However, there is uncertainty regarding which vertebrate predator/prey links to delete as well as how many links to add from non-terrestrial vertebrate food resources to vertebrates. To address this uncertainty, we used a randomization approach in the link-removal and -addition steps described below and generated 1000 versions of the network. That is, we randomly removed (for the vertebrate predator–prey links) or added (for the herbivores, invertivores and piscivores) links in the range indicated as realistic based on contemporary species’ diet breadths. Because we do not know exactly where to add or remove these links, we generated 1000 versions of the network so that our results were not skewed by the particular links selected.

We used the trophic-niche model to assign potential predator–prey links in the Naracoorte species assemblage. Although trophic-niche models are good at identifying potential links, they almost always overestimate the number of realized links (Marple et al. 2018). This is because predators are unlikely to consume all prey within their size range – some species are not palatable, are dangerous, too rare, difficult to capture, use different microhabitats or have other ecological characteristics that make them unsuitable for regular consumption (Marple et al. 2018). To build a network with a more realistic structure, we assigned a probability to each interaction based on the prey’s position in the predator’s prey-size range and a Gaussian distribution centred on this range (with a standard deviation equal to one-quarter of the predator’s prey-size range); the highest probability was for prey close to the centre of the prey-size range (i.e. centre of the distribution) and decreased the closer the prey was to the predator’s limits. For each predator, we randomly sampled from a density kernel fit to published carnivore diet breadths (n = 12; sampled between 1 and twice the maximum diet breadth in the Supporting information) to select the number of potential prey that were ‘realised’. In assigning the sampled diet breadths to individual predators, predators with more potential prey (indicated by the trophic-niche model) were assigned larger diet breadths than were those with fewer potential prey. To account for different degrees of dietary specialization, we Poisson-resampled the number of potential prey for each predator before ranking predators according to their number of potential prey, slightly shuffling relative diet breadths between network models. Then, using the assigned diet breadths, we randomly selected from the potential prey, taking into account the probability of the predator–prey interaction. This method resulted in a vertebrate network with realistic connectance (proportion of potential links that are realized) and with most of each predator’s prey closer to the centre, rather than the limits, of their prey-size range.

Terrestrial vertebrates not only consume other terrestrial vertebrates but they also consume invertebrates, plants, fungi and fish. In addition to inferring trophic links among terrestrial vertebrates, we therefore needed to add links to vertebrates from these other food resources. However, we did not include detail on individual species within these resource groups because: 1) our study focusses on terrestrial vertebrate species; 2) invertivore, plant and fungal diversities are not well resolved for most ecosystems (including for Late Pleistocene Naracoorte); and 3) fish only constitute a small part of the Naracoorte community in terms of biodiversity and biomass. We therefore generated a pool of n species for each of these groups (n = 1300 for plants, 6000 for invertivores and 23 for freshwater fish), with the number of invertivore and plant species calculated based on the described diversity in these groups relative to terrestrial vertebrate diversity in Australia (Chapman 2009), and the number of fish determined by the diversity of freshwater fish recorded in the Naracoorte Coastal Plains bioregion in Atlas of Living Australia (<ala.org.au>; accessed 6 April 2021). To assign links to vertebrates from species in these groups, we used published records of vertebrate diet breadth for 20 herbivores, 6 invertivores and 9 piscivores (Supporting information). We fit kernel densities to the invertivore, herbivore and piscivore diet breadth data and randomly sampled (within a diet breadth range of 1 to twice the maximum diet breadth recorded for that trophic guild in the Supporting information) from these distributions to assign the number of links between each resource group and vertebrate consumer in each of the 1000 network models. However, because the published piscivore diet breadths (Supporting information) came
from terrestrial Australian predators that are not exclusively piscivorous, we multiplied the number of fish in the diets of these predators by the number of resource groups from which the predator fed before fitting and sampling from the kernel densities. We made this modification to offset the adjustment accounting for inflated diet breadths in omnivores.

For vertebrates that fed from more than one group (i.e. omnivores, which represent over half the vertebrates in this network), we proportionally adjusted the number of ‘in’ links (food resources) depending on how many food groups they fed. For example, if a species consumed from two groups (e.g. vertebrates and invertebrates), we randomly deleted half of the links from each group; if they fed from three groups, we randomly removed two-thirds of the links from each food group (and so on). We made these deletions to prevent omnivores from having inflated diet breadths.

In some cases involving water birds, we used empirical evidence to avoid assigning unrealistic trophic links. For example, pelicans are large and, consequently, the trophic-niche model predicts that pelicans take large vertebrate prey. However, we know that pelicans are not birds of prey and do not consume large terrestrial animals. Thus, we restricted the allocation of trophic links for such birds to prevent them from feeding on unrealistically large terrestrial vertebrates (they were prevented from consuming prey that weighed over a third of their body mass).

We completed the networks by generating links from plants to invertebrates and from plants to fish. To determine diet breadth for these herbivores, we sampled from a Pareto distribution ($\alpha = 1.02$, truncated at 52) following Forister et al. (2015). The $\alpha$ and truncation values were based on a temperate woodland system (i.e. similar to Naracoorte) (Fraser and Wells 2006, Reed 2012, Forister et al. 2015).

**Analysis of Naracoorte models**

We estimated the coextinction vulnerability of Naracoorte vertebrates using the same simulation approach as we applied to the synthetic networks, i.e. we iteratively (and randomly) removed plant nodes to simulate primary extinctions, and coextinctions were triggered when animals lost all their food resources. We measured each species’ coextinction vulnerability as the proportion of plant nodes remaining when coextinction occurred. We repeated the simulations 1000 times for each of the 1000 network models to test whether: 1) the effects of trophic level, diet breadth and basal connections (direct and indirect) on coextinction vulnerability were consistent with results obtained from the synthetic models and 2) vulnerability to bottom–up cascades differed between extinct and extant species (i.e. comparing the relative vulnerability of species).

To test whether the coextinction vulnerability results were consistent with those from the synthetic models, we fit linear-regression models to the data with vulnerability to coextinction cascades as the response, and trophic level, diet breadth, basal connections and the interactions between these variables as independent variables (fixed effects). Rather than using the raw data from the 1000 network models, we used each species mean value for each trait and vulnerability for these analyses. We then compared relative model probabilities ($\Delta AIC$) for the full model to all possible reduced models and examined the coefficients of the main effects to determine if they were similar to those obtained from the synthetic networks.

We compared the susceptibility of Naracoorte’s extinct versus extant species to bottom–up coextinction cascades in three steps. First, we compared the groups in terms of their trophic level, diet breadth and number of basal connections to determine if differences in these variables suggest one group would be more vulnerable than the other. Second, to test for an association between coextinction vulnerability and extinction status, we compared $\Delta AIC$, support for two models: a null model with vulnerability to coextinction as the dependent variable and no fixed effects versus a model that was identical to the first, except with extinction status as an independent variable. Third, we ran a randomization test to assess the probability that extinct species were more vulnerable to coextinction than were extant species. Here, we sampled the coextinction vulnerabilities of 10 extinct and 10 extant species (with replacement) from each of the 1000 models 20 times (i.e. using the raw data rather than species means) and each time calculated the mean difference in coextinction vulnerability between the two groups. The purpose of the regression-model comparison and randomization test was to test specifically for an association between extinction status and vulnerability to bottom–up cascades that could explain (or have contributed to) the extinctions that occurred at Naracoorte.

We also assessed the position of extinct versus extant species in the network using 12 different network metrics: trophic level, pageRank, betweenness centrality, eigenvector centrality, closeness centrality (in), coreness (in), degree (in), eccentricity (in), closeness centrality (out), coreness (out), degree (out) and eccentricity (out) (see Supporting information for metric descriptions). We calculated the metrics followed by an ‘in’ or ‘out’ for each node using links pointing towards (in) or away from (out) the focal node. We chose these metrics because they are commonly used, node-level metrics describing the position in the network. For each metric, we calculated the species’ mean value across the 1000 network models. After checking for highly correlated metrics and removing those identified as redundant (i.e. metrics that had a Spearman’s $\rho > 0.8$), we ran a principal component analysis and visually inspected for grouping of extinct and extant species. The reduced list of metrics included closeness centrality (out), eccentricity (out), degree (in), coreness (in), betweenness and PageRank.

**Results**

**Synthetic networks**

The two methods calculating bottom–up coextinction vulnerability (simulation and Bayesian network) yielded similar
results in terms of the effects of trophic level, diet breadth and basal connections on node vulnerability to bottom–up coextinction cascades. \( wAIC \), indicated that the full models (i.e., those that had all three independent variables and their interactions) were more strongly supported than reduced models using both approaches, with the full models having \( wAIC \approx 1 \) and marginal \( R^2 \geq 0.53 \) (Supporting information). The weighted model-averaged coefficients describing the relationships between the three independent variables and coextinction vulnerability were negative, irrespective of which of the two methods we used to calculate vulnerability (Fig. 3a, Supporting information). These negative correlations indicate that vulnerability to bottom–up cascades decreased with increasing number of basal connections, diet breadth and trophic level. Marginal \( R^2 \) of the three reduced models, each of which had one of the three variables as a main effect, suggests that the number of basal connections explained most of the variation in coextinction vulnerability in the synthetic networks (Fig. 3b). We restricted our analyses of the Naracoorte network models to the simulation method because both approaches yielded similar results and the Bayesian network method was prohibitively time-consuming for networks of the size of the Naracoorte models.

Naracoorte network

In terms of the effects of trophic level, diet breadth and basal connections on vulnerability, the patterns in the Naracoorte network were similar to those from the synthetic networks (Supporting information). \( wAIC \), strongly supported the full model over reduced models (\( wAIC \approx 1 \) and adjusted \( R^2 = 0.99 \) for the full model; Supporting information), and the three main effects were negatively correlated with coextinction vulnerability (Fig. 3c, Supporting information). Reduced models that had either trophic level, diet breadth or basal connections as the only independent variable had \( R^2 > 0.23 \), indicating that each of these variables was associated with a substantial proportion of variation in vulnerability (Fig. 3d).

Extinct species had fewer basal connections, narrower diet breadths and came from lower trophic levels, on average, than those species that survived into the Holocene (Fig. 4a–c; mean basal connections: 298 versus 515, diet breadth: 34 versus 79 and trophic level: 2.3 versus 2.9 for extinct versus extant species). Extinct species also had higher coextinction vulnerability than did surviving species (Fig. 4d; mean coextinction vulnerability $\pm 95\%$ confidence interval: 0.045 $\pm$ 0.015 versus 0.008 $\pm$ 0.002 for extinct and extant species, respectively), a result consistent with coextinction vulnerability being higher for species with fewer basal connections, narrower diet breadth and from lower trophic levels.

To test for an association between species’ extinction status and vulnerability to bottom–up cascades, we compared support for a model with coextinction vulnerability as the dependent variable and extinction status as the only fixed effect versus a null model (a random-intercept model with no fixed effects) and we also did a randomization test comparing the coextinction vulnerability of extinct versus extant species. Both approaches indicated that extinction status was associated with vulnerability to bottom–up cascades, with extinct species more vulnerable than those that survived into the Holocene. The \( wAIC \), for the model with extinction status as a fixed effect was $\approx 1$, indicating that this model was strongly supported over the null model. The randomization test showed that extinct species had a probability of 0.91 of being more vulnerable to bottom–up coextinctions than extant species (Fig. 5).

Extinct species differed from extant species in terms of their position in the network. Principal component analysis of six network-position metrics showed that extinct and extant species fell into two distinct groups according to the second principal component (dimension 2 in Fig. 6a). The main contributors to this principal component are metrics describing a node’s connection to the network through its ‘out’ links, including the closeness centrality (out) and eccentricity (out) metrics (Fig. 6a, Supporting information). Closer examination of the out links (i.e, number of predators) showed that, on average, extinct species had <1 predator, whereas extant species had >3 (Fig. 6b; mean number of predators: 0.2 versus 3.3 for extinct versus extant species). Indeed, the average number of predators was lower for extinct than extant species in all 1000 models of the Naracoorte network.

Discussion

Our analyses demonstrate that a species’ vulnerability to bottom–up coextinction cascades varies depending on its trophic level, diet breadth and number of basal connections. We also found that the position of extinct species in the Naracoorte network – being primarily herbivorous (low trophic level) and therefore having relatively narrow diet breadths and few pathways to basal resources – might have made them more vulnerable to bottom–up coextinction cascades than were co-occurring species that survived into the Holocene. The Naracoorte results suggest that trophic cascades potentially contributed to the megafauna extinction event in southeastern Sahul. However, the extinct species from Naracoorte also had fewer predators than did extant species, a network position attribute that would likely have made them more vulnerable to the arrival of the new ‘super predator’ – humans (Darimont et al. 2015).

The Naracoorte and synthetic network models revealed that vulnerability to bottom–up coextinction cascades precipitated by plant extinctions decreased with increasing trophic level, diet breadth and number of basal connections (Fig. 3, Supporting information). Our results therefore support previous findings that species with narrower diet breadths/fewer pathways to basal resources are more vulnerable to bottom–up coextinction cascades (Purvis et al. 2000a, Kanerdy et al. 2012, Chichorro et al. 2019). However, previous research does not provide a clear expectation regarding the relationship between trophic level and extinction vulnerability. It is often assumed that higher trophic levels are more vulnerable to extinction than are lower levels due to the
cumulative effects of disturbances on lower trophic levels (on which higher trophic levels depend) and direct persecution by humans (Purvis et al. 2000b, Duffy 2003). However, our analyses that specifically tested for the sensitivity of species to primary extinctions in the plant community imply that vulnerability to these bottom–up cascades in fact decreases with trophic level. Consistent with these results, several manipulative experiments of ecological communities have revealed that changes in the plant component of the community most strongly affect herbivores and impacts on higher trophic levels diminish with trophic distance (Kagata et al. 2005, Schädler et al. 2010, Scherber et al. 2010). However,
these manipulative experiments involved plants and invertebrates, so confirmation is still required that the relationship between trophic level and vulnerability to bottom-up cascades observed in these systems also occurs in vertebrate communities.

Theoretical studies have also concluded that the effects of bottom-up cascades dampen with trophic distance. Applying Rosenzweig–MacArthur models and synthetic (but ecologically feasible) networks, the loss of primary producers triggered extinctions in herbivores before doing so in carnivores, and herbivores were more vulnerable to these cascades than were carnivores (Kaneryd et al. 2012). However, our vulnerability scores were based on coextinction being triggered when a consumer lost all food resources. Coextinctions could be triggered at lower thresholds and/or vary between species. While the congruence between our results and those from previous studies support the methods and threshold we used, further investigation into how coextinction threshold influences the effect of network position on node vulnerability, as well as how coextinction thresholds covary with species/community traits, is needed to refine methods for predicting the probability and magnitude of bottom-up cascades. For

Figure 4. Comparison of species that survived into the Holocene (extant shown in green) to those that went extinct in the Late Pleistocene (extinct shown in yellow) from the Naracoorte network. (a) shows the number of basal nodes (plants) connected directly and indirectly to each node via ‘in’ links; (b) shows diet breadth of each node; (c) indicates trophic level and (d) shows the calculated coextinction vulnerability. The panels, which all include density violin plots and scatterplots, are based on each species’ mean score across the 1000 Naracoorte network models.
example, it is plausible that resource-loss thresholds decrease with body size. If such a relationship exists, it would exacerbate the vulnerability of larger-bodied species to habitat disturbance (Bennett and Owens 1997, Cardillo and Bromham 2001). Such a relationship could have made the extinct megafauna of Naracoorte particularly vulnerable to bottom–up cascades (i.e. in addition to the effects trophic level, diet breadth and links to basal resources) and might explain why three predatory species – which were not predicted to be vulnerable to bottom–up cascades based on their trophic level – also went extinct (Fig. 4c).

The extinction pattern observed in the Naracoorte assemblage could have been the result of bottom–up cascades triggered by changes in the plant community, as demonstrated by our vulnerability modelling. This leads to the question: Did vegetation change at the same time as the megafauna disappeared? Studies in other regions of Sahul have detected shifts in vegetation roughly coinciding with, or immediately

![Figure 5](image5.png)

**Figure 5.** Density plot showing results from a randomization test (20 000 replicates) comparing bottom–up coextinction vulnerability of extinct versus extant species from the Naracoorte network models. We calculated each species’ coextinction vulnerability using the simulation method (removing plant nodes and then removing animal nodes that no longer had connections to plant nodes). From these results, we sampled (with replacement) the coextinction vulnerabilities of 10 extinct and 10 extant species and calculated the mean differences in coextinction between the groups. We repeated this process 20 times for the results from each of the 1000 network models to build the density plot. The yellow area of the density plot indicates higher vulnerability for extinct species, and the green area indicates higher vulnerability for extant species.

![Figure 6](image6.png)

**Figure 6.** Comparison of extinct versus extant species in terms of their position in the Naracoorte trophic network. (a) is a biplot of the first two principal components from a principal component analysis of six variables describing species’ positions in the network. (b) is a violin and scatter plot showing the number of species that preyed on each extant versus extinct species. We inferred 1000 models of the Naracoorte network, and from these models, we calculated each species’ mean value of each metric for use in the principal-components biplot and predator-diversity plot.
preceding, megafauna extinction. Hypothesised drivers of these vegetation shifts include land-use changes associated with human arrival (i.e. fire-stick farming) (Turney et al. 2001, Miller 2005) and/or climate change (DeSantis et al. 2017). However, there are no detailed studies on the vegetation of Naracoorte spanning the Late Pleistocene extinction event (but see Bampton 2021 for a review of broad proxies of vegetative change over this period). The megafauna’s disappearance from Naracoorte did, nonetheless, coincide with an unusually cool period (Supporting information) and the arrival of humans (~44 000 years ago) (Saltré et al. 2019), offering the intriguing possibility that changes in climate and/or land use triggered shifts in vegetation that had consequences for higher trophic levels in the network. To evaluate this possibility, more studies are required to model vegetation changes in south-eastern Sahul (including the Naracoorte region), and these must be validated using the pollen record and/or other fossil data.

By considering the network position of all vertebrate species in the assemblage, a clear difference between extinct and extant species emerged – extinct species had fewer predators than did species that survived (mean number of predators: 0.2 versus 3.3 for extinct versus extant species, respectively; Fig. 6b). This predator naivety, coupled with the species’ slow life histories, likely made megafauna especially vulnerable to new predators (Flannery 1990, Johnson 2002, Johnson et al. 2016, Bradshaw et al. 2021) and suggests that hunting by humans could have adversely affected megafauna. Thus, a network modelling approach to assessing extinction vulnerability suggests that bottom–up and/or top–down processes could have selectively removed the now-extinct species from the Naracoorte community. However, there remains scope to address uncertainties regarding the structure of the Naracoorte network and the methods for estimating vulnerability to ecological cascades. As palaeo-vegetation, invertebrate, trophic (including detailed information on the diets of extinct species) and climate data improve, network models can incorporate this information to build more refined networks, and include more detailed top–down and bottom–up forcings to assess the plausibility of the different potential causes and pathways to extinction – including what (if any) role humans played in the megafauna’s demise.

In summary, our network modelling of Late Pleistocene Naracoorte suggests bottom–up coextinction cascades and/or predator naivety and the arrival of humans could have contributed to the megafauna extinction event in Sahul. Indeed, that our network models showed that extinct species were vulnerable to bottom–up cascades and new predation pressures lend support to recent research suggesting that climate change (that can shift vegetation and lead to bottom–up cascades) and human arrival together drove the megafauna extinction trajectories in much of south-eastern Sahul (Saltré et al. 2019). Our research, along with other recent studies (Roopnarine et al. 2007, Dunne et al. 2014, Pires et al. 2015), demonstrates that network modelling is a powerful tool for investigating and understanding ancient extinction events. By developing methods to model whole-community responses to change and validating these methods using ancient extinction events, we can also provide better estimates of the fates of contemporary communities as the sixth mass-extinction event unfolds (Ceballos et al. 2017).

Acknowledgements – We thank E. Reed, C. Carbone, M. Tucker, C. Dickman and V. K. Llewelyn for their constructive input. We acknowledge the Meintangk and Kaurna people, the traditional owners of the Naracoorte Caves and the Adelaide region, respectively. Funding – This work was supported by the Australian Research Council Centre of Excellence for Australian Biodiversity and Heritage (grant number CE170100015).

Author contributions

Project conceptualization by JL, GS, CJAB and FS. Data curation by JL, GS, KJP, CNJ, MCM and SNd-V. Investigation and formal analysis by JL, GS and CJAB. Methodology by JL, GS, CJAB and DBS. Original draft preparation by JL, GS, CJAB and FS. All authors contributed to reviewing and editing the draft.

Transparent Peer Review

The peer review history for this article is available at <https://publons.com/publon/10.1111/ecog.06089>.

Data availability statement

All relevant data and code are available at Zenodo: <https://doi.org/10.5281/zenodo.3756223>.

Supporting information

The supporting information associated with this article is available from the online version.

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