

RESEARCH ARTICLE

The influence of seabirds on their breeding, roosting and nesting grounds: A systematic review and meta-analysis

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Abstract

1. Seabird species world-wide are integral to both marine and terrestrial environments, connecting the two systems by transporting vast quantities of marine-derived nutrients and pollutants to terrestrial breeding, roosting and nesting grounds via the deposition of guano and other allochthonous inputs (e.g. eggs, feathers).
2. We conducted a systematic review and meta-analysis and provide insight into what types of nutrients and pollutants seabirds are transporting, the influence these subsidies are having on recipient environments, with a particular focus on soil, and what may happen if seabird populations decline.
3. The addition of guano to colony soils increased nutrient levels compared to control soils for all seabirds studied, with cascading positive effects observed across a range of habitats. Deposited guano sometimes led to negative impacts, such as guanotrophication, or guano-induced eutrophication, which was often observed where there was an excess of guano or in areas with high seabird densities.
4. While the literature describing nutrients transported by seabirds is extensive, literature regarding pollutant transfer is comparatively limited, with a focus on toxic and bioaccumulative metals. Research on persistent organic pollutants and plastics transported by seabirds is likely to increase in coming years.
5. Studies were limited geographically, with hotspots of research activity in a few locations, but data were lacking from large regions around the world. Studies were also limited to seabird species listed as Least Concern on the IUCN Red List. As seabird populations are impacted by multiple threats and steep declines have been observed for many species world-wide, gaps in the literature are particularly concerning. The loss of seabirds will impact nutrient cycling at localized levels and potentially on a global scale as well, yet it is unknown what may truly happen to areas that rely on seabirds if these populations disappear.

KEYWORDS

enrichment, guano, marine-derived, mobile link, vector

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1 | INTRODUCTION

Abiotic and biotic vectors are responsible for the transport of subsidies across ecosystem boundaries, leading to the replenishment of vital resources in recipient communities (Payne & Moore, 2006) and can be profoundly significant by altering ecosystem-level processes and trophic dynamics (Earl & Zollner, 2017). Abiotic vectors, such as wind (e.g. transport of dust from the Sahara Desert to the Atlantic Ocean; Kaufman et al., 2005), gravity (e.g. fallout of marine snow from surface waters to the deep sea; Kiko et al., 2017) and ocean currents (e.g. dispersal of mangrove seedlings from one island to another; Hodel et al., 2018) move materials from one system to another, usually unidirectional, and are determined by the force of the factor (Payne & Moore, 2006). In comparison, biotic vectors, such as mobile organisms, are often less restricted in their movement and can therefore transport subsidies in a multitude of directions and against those common gradients found with abiotic vectors.

Migratory animals are well-known biotic vectors, connecting ecological communities all across the globe by transporting significant quantities of spatial subsidies, with the potential to influence community structure, diversity and ecosystem function at resident habitats (Bauer & Hoyer, 2014). However, the effects of vectors can be highly variable. Migrants can transport parasites and pathogens (e.g. highly pathogenic avian influenza virus (H5Nx); Wille et al., 2019) as well as propagules (e.g. Old World fruit bats *Cynopterus sphinx* consume fruits and deposit the seeds through their excreta; Shilton et al., 1999), but perhaps the most common transport effects are nutrients, toxicants and energy. Migrants act as transport mechanisms for these subsidies by feeding in one ecosystem, moving to another system, and then offloading the materials via a range of outputs (e.g. defecation; Bauer & Hoyer, 2014; Blais et al., 2005). Vectors can be incredibly beneficial—and the ecological effects can be much more pronounced—when the transport of subsidies is from a nutrient-rich ecosystem to an oligotrophic system (Savage, 2019).

In the marine environment, transfer of spatial subsidies to terrestrial ecosystems has been documented for Pacific salmon (*Oncorhynchus* spp.; Payne & Moore, 2006), sea lions (e.g. *Zalophus wollebaecki*; Fariña et al., 2003) and numerous seabird species (Kolb et al., 2015). Seabirds feed on marine prey (e.g. fish, squid, zooplankton) and transport nitrogen (N) and phosphorous (P) from their marine feeding grounds to their roosting and breeding habitats on land (Daher et al., 2019; Dominguez et al., 2017; Sanchez-Pinero & Polis, 2000). Here, seabirds congregate in dense colonies and deposit these marine-derived nutrients (MDN) in huge quantities primarily through guano, but also through other allochthonous inputs of feathers, eggs, boluses (regurgitated pellets) and carcasses (Adame et al., 2015; Duffy, 1994; Szpak, Longstaffe, et al., 2012). For these reasons, seabirds are one of the most significant vectors for the transportation of these vital nutrients (Marmen et al., 2017).

Humans have been aware of the many benefits of seabird guano and have exploited it for the purposes of fertilizing crops as early as the 16th century (Cushman, 2013). The most well-known areas of guano mining were along the Pacific Margin in Peru and northern Chile

where thick deposits of guano (>50m in extreme cases) were mined extensively from islands (Lucassen et al., 2017; Szpak, Longstaffe, et al., 2012). The trade of guano peaked at 20 million tons/year during the mid-19th century before declining as guano ran low, prices soared and chemical fertilizers became widely available (Mathew, 1970; Szpak, Longstaffe, et al., 2012). When left to accumulate naturally in the wild, the benefits of guano as a fertilizer can include significantly increased soil N and P, and can also impact soil pH, moisture and salinity (Wait et al., 2005). These changes in soil characteristics leads to myriad positive changes for terrestrial ecosystems and have been widely documented (e.g. Magnússon et al., 2014; Mosbech et al., 2018; Sanchez-Pinero & Polis, 2000). The positive impacts provided by seabird guano can include enhanced primary and secondary productivity, increased invertebrate abundance and diversity, and changes to ecological communities (Buelow et al., 2018). Referred to as ornithogenic soils, the effects of nutrient enrichment derived from birds can be exceptionally distinct compared to areas without seabirds and can last for centuries (Mizutani et al., 1991).

On islands, the effect of seabirds on their environment can be more pronounced compared to mainland systems. Islands often receive few external nutrient subsidies besides what the birds transport, thus seabirds can play an integral role in the functioning and resilience of islands (Buelow et al., 2018), and the communities that live within and around colonies (Sanchez-Pinero & Polis, 2000). For example, when a new volcanic island, Surtsey, formed off the coast of Iceland in 1963, the establishment of a gull colony which improved soil nutrient status was the main agent of ecosystem development (Magnússon et al., 2014). Seabirds can therefore facilitate the colonization and regeneration of marginal and nutrient-limited habitats, such as barren, polar, post-glacial or volcanic habitats (Şekericioğlu, 2006).

While the positive benefits of seabirds to a terrestrial system are plentiful, high seabird abundance in an area can also contribute to negative consequences (Wootton, 1991). Guano adds valuable nutrients to a system, but excessive deposition can alter plant and invertebrate communities, kill vegetation or cause dead zones (Gillham, 1961; Martín-Vélez et al., 2019; Saifutdinov & Korobushkin, 2020). This 'guano-trophication', or seabird-induced eutrophication, occurs when the level of nutrients added to a system exceeds what is required (Signa et al., 2015). As apex predators, many seabirds also exhibit high contaminant loads due to biomagnification and bioaccumulation along the marine food chain (Shoji et al., 2019). Chemicals, including persistent organic pollutants (POPs) and trace elements, can pass through individuals via feathers, eggs and guano (Furness & Camphuysen, 1997; Otero et al., 2018). Like the transport of nutrients, many recipient habitats do not already contain these pollutants prior to the arrival of seabirds, and thus the birds introduce pollutants in, at times, substantial quantities (Michelutti et al., 2009; Santamans et al., 2017). In addition to POPs and trace elements, many seabird species consume plastic debris and deposit these pollutants within their colonies via several means. Micro-plastics (<5mm in diameter; Barnes et al., 2009) can be excreted with guano; however, it is unknown whether plastics can be digested or broken up to small enough pieces to be passed via guano, or whether it is only micro- and nano-sized particles (ingested at that size) that

TABLE 1 List of all search terms applied when using the *ISI Web of Science* and *Scopus* databases. Each topic includes a list of all synonyms relevant to that topic and each term (within each topic) was connected by the Boolean operator 'OR'. A total of 265 papers were returned by this search in *Web of Science* and 316 from *Scopus*. An asterisk represents a string of any characters and is used when the word could have different endings, for example, transport* could be transporting or transportation. A question mark represents a single character and is used when a letter within the word could change, for example, fertili?* could be fertilization or fertilization. Please see Supporting Information for the entire search string with all syntax included

Topic 1: Seabird	AND	Topic 2: Vector	AND	Topic 3: Nutrient/pollutant	AND	Topic 4: Guano	NOT	TOPIC 5: Poultry
Seabird		Transport*		Contamin*		Guano		Poultry
'Marine bird'		Marine-derived		Toxin		Faeces		
Avian		Route		Organic		Feces		
		Vector		Inorganic		Dropping*		
		Enrich*		Metal		Excre*		
		Fertili?*		'Trace element'		Manur*		
		Path		'Persistent organic pollutant'*				
		Translocat*		Plastic				
		Biotransport*		Debris				
		Deposit*		Nutrient				
		Engineer		Pollut*				

can be passed (see Bourdages et al., 2021; Gil-Delgado et al., 2017; Provencher et al., 2018; Reynolds & Ryan, 2018). While deposition of plastics via guano is a relatively new topic, it has been known for some time that seabirds can deposit plastics in colonies through other means, such as regurgitation or the death of birds with high loads of ingested plastics (Buxton et al., 2013; Grant et al., 2021; Nel & Nel, 1999).

Here we critically review the literature regarding seabirds as vectors for nutrients and pollutants from marine to terrestrial ecosystems. While our review captured literature encompassing multiple pathways of nutrients and pollutants as provided by seabirds (i.e. guano, feathers, eggshells, regurgitations including boluses/pellets, carrion/carcasses), we focused on guano as this is the most significant and common pathway. We highlight global patterns and areas of high research activity and provide recommendations for consistent data collection and reporting in the future studies. We assess the various nutrients and pollutants associated with guano and determine if differences lie between seabird orders. Finally, we conduct a meta-analysis on the impact of guano deposition on soils. We did this using published studies that compare common nutrients in colony soils and control soils. We hypothesize that colony soils will be enriched in nutrients compared to control soils because of the deposition of guano in colonies.

2 | MATERIALS AND METHODS

2.1 | Literature searching

We conducted a systematic review of the available literature on the online platform *Web of Science* and using the Core Collection database, containing the following Citation Indexes: Science

Citation Index Expanded: 1945–2021; Social Sciences Citation Index: 1956–2021; Arts & Humanities Citation Index: 1975–2021; Conference Proceedings Citation Index–Science: 1990–2021; Conference Proceedings Citation Index–Social Science & Humanities: 1990–2021; and Emerging Sources Citation Index: 2015–2021. We also conducted an identical search on the online database *Scopus*. We followed established methods outlined in Haddaway et al. (2018) with the aim to capture all published literature that is listed on *Web of Science* and *Scopus* on the topic of seabirds as vectors for nutrients and pollutants to terrestrial environments. 'Terrestrial', in the case of this review, is defined as any land that seabirds nest, roost, breed or loaf in and therefore is a site where guano is deposited persistently in space and time. This also includes intertidal regions. Papers that studied terrestrial habitats and freshwater habitats (e.g. stream, lake, wetlands) simultaneously were also included. Purely marine-based studies are not included in this review.

The search was conducted on 26 October 2021 and included key terms synonymous with the four broad topics, namely seabird, vector, nutrient/pollutant and guano (Table 1, see Supporting Information for full search string), and captured published literature up until 26 October 2021. We included all relevant peer-reviewed original research papers. Review articles, book chapters and grey literature were excluded.

In addition, we performed backward literature searching on all papers included in the full review to gain more articles relevant to this topic that were not picked up through searching *Web of Science* or *Scopus*. For this, the reference list of each paper was combed through and any paper with a relevant title was extracted and added to the results (Figure 1).

2.2 | Literature screening

The searches returned a combined total of 581 papers, with 265 from *Web of Science* and 316 from *Scopus*. Once duplicate records were removed, there were 396 unique papers from *Web of Science* and *Scopus*. Backwards literature searching produced a further 218 papers (Figure 1). Each paper underwent a preliminary filtering stage where titles and abstracts were read and if the research was not directly related to the role of seabirds as vectors to terrestrial ecosystems, then it was removed. The remaining papers then went through full-text processing to remove any more irrelevant articles (Figure 1; Figure S1). For papers to be included in the final dataset, they had to demonstrate that the focal seabirds influence their terrestrial environment. This meant that papers had to do two things: (a) measure seabird influence (i.e. analyse guano, measure presence/absence) and (b) measure the impact of seabirds on recipient environment (e.g. soil, vegetation, invertebrates). Papers that studied historical trends in pollutants in sediments, seabirds as vectors of viruses, transport of seabird-derived nutrients back to marine environments, atmospheric emissions (above seabird colonies) or laboratory-based fertilization studies were not included (Figure S1).

We used the International Ornithological Congress World Bird List v11.2 (Gill & Donsker, 2021) for classification of species and for defining 'seabirds'. Papers were included only if the focal species were in the following orders: Charadriiformes (gulls, terns, auks, shorebirds, skuas), Phaethontiformes (tropicbirds), Sphenisciformes (penguins), Procellariiformes (petrels, storm-petrels, diving petrels, shearwaters, albatrosses) and Suliformes (boobies, gannets, cormorants, frigatebirds, anhingas).

The total number of peer-reviewed articles after the completion of title and abstract processing and full-text screening was 181, with 101 from database searching (*Web of Science* and *Scopus*) and 80 from backwards literature searching. All studies used in the systematic review and meta-analysis are listed in the Data Sources section. M.L.G. and J.L.L. performed the literature screening.

2.2.1 | Screening for meta-analysis

To demonstrate how powerful guano can be, we ran meta-analyses by comparing nutrient concentrations in colony soils (where guano is deposited) to control soils (where guano is not deposited). For the meta-analysis, we decided to use studies that compared nutrients in soils because this was the most common abiotic factor that was sampled across all studies. Thus, this decision was made post hoc (i.e. after the studies were collected for review). The nutrients included within the meta-analyses were ammonium (NH_4^+ ; $\mu\text{g/g}$), nitrate (NO_3^- ; $\mu\text{g/g}$), total nitrogen (%N) and total phosphorous (%P). All nutrients were measured in dry weight. For papers to be included, they had to:

1. Include a control site(s) along with a site with seabirds (i.e. a seabird colony). The control site(s) must be a strictly defined area without seabirds present. Many papers included a control site that was along a gradient of seabird influence, but as these are not strict control sites, they were excluded from analysis.
2. Measure one or more of the above nutrients in soils in both colony and control soils.
3. Report mean nutrient/pollutant concentrations in colony and control soils, standard deviations (SD) and sample sizes (n), or allow for them to be calculated. For studies that only reported these values within plots, we used GetData Graph Digitizer v.2.26 to extract the relevant data.

After following these conditions, our list of papers that could be included within the meta-analysis was reduced to 32 for nutrients in soils (see Figure 1 for more details, Figure S2). However, not all these papers analysed contain all the chosen nutrients (e.g. a paper may have analysed total nitrogen only). As such, for each of the different nutrients, there were 14 to 22 papers.

2.3 | Data extraction

From the 181 papers, we then extracted data and information on the following: location (island or mainland, coordinates) and habitat of the study (as defined by the IUCN habitat classification scheme; The International Union for Conservation of Nature, 2012), study duration (single or multi-year studies as well as the number of years), study site information (number of sites, whether was there a control/s site, that is, a nearby site without the presence of seabirds to compare against), seabird focal species, what metrics were used to measure seabirds' influence (presence/absence or guano) and if guano was used we included the analysis type (nutrient content, elemental analysis, guano cover, isotope analysis or from literature), what the seabirds were documented to be transporting (nutrients, inorganic pollutants, organic pollutants or physical pollutants) and what abiotic and biotic factors were sampled to determine the effect of seabirds. Due to the diversity of biotic and abiotic factors sampled, most were put into broad categories: terrestrial vegetation (e.g. grass, trees and moss), vertebrates (e.g. fish and reptiles), terrestrial invertebrates (e.g. insects and tardigrades), benthic/aquatic vegetation (e.g. macroalgae, seagrass), intertidal organisms (e.g. molluscs and bivalves) and plankton (e.g. zooplankton and phytoplankton). Soil, water, detritus/leaf litter and sediment were retained in their own categories. The overall effect of the focal species on the given area (as stated in the results or conclusion of the paper, for example, soil properties improved, species abundance increased) was also recorded. As guano was the focus of this study, we extracted information on trace elements analysed within guano and the respective concentrations of those elements, as well as nutrient concentrations in guano.

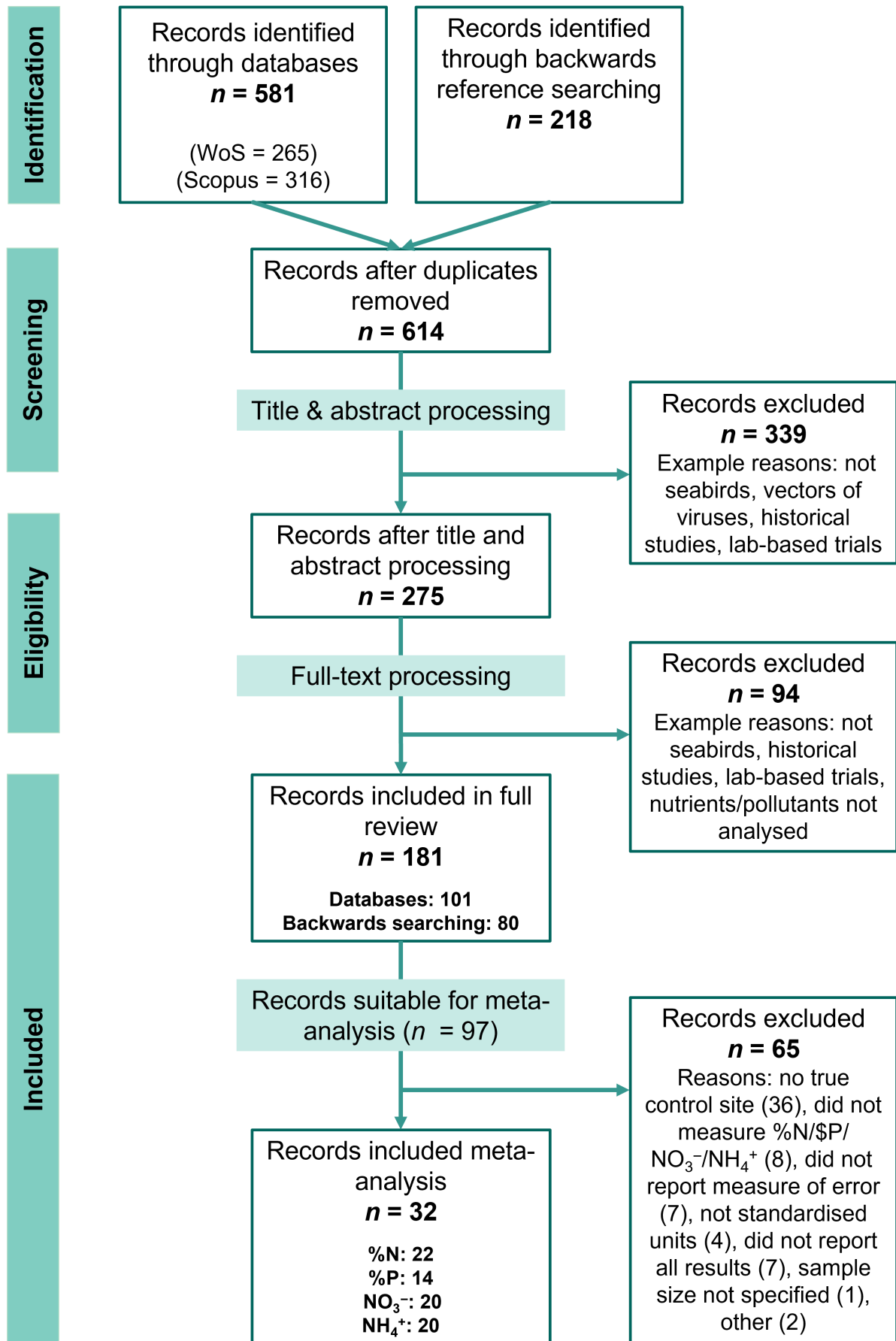


FIGURE 1 Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA; Page et al., 2021) flow diagram for study selection. Diagram depicts the number of studies retained and discarded at each step. The total number of records included in the meta-analysis was 32; however, many publications studied multiple nutrients simultaneously and thus the sum of records for nutrient types does not equal 32

2.3.1 | Data extraction for meta-analysis

For each of the 32 papers suitable to be included in the meta-analysis, we extracted the following data: mean $\pm$ SD concentration of nutrient/s (NH_4^+ , NO_3^- , %N and %P) in seabird soil and in control soil, as well as the number of soil samples for each group (in addition to data extracted for each paper in Section 2.3). For this subset of papers, we also recorded the latitudinal zone the study was undertaken in (polar = $>60^\circ\text{N}$ or S; temperate = $30\text{--}60^\circ\text{N}$ or S; and tropical = $<30^\circ\text{N}$ or S). If a paper measured nutrient concentrations in soils of more than one seabird species, then this was recorded for each species separately.

2.4 | Meta-analysis

2.4.1 | Effect size calculations

We calculated the log response ratio (LRR, also known as Log(Ratio of Means)) as our effect size since nutrient concentrations are continuous positive variables. LRR is a common effect size calculated in ecological meta-analyses because it quantifies the proportionate change between the groups (Hedges et al., 1999). It is calculated by:

$$\text{LRR} = \ln \frac{C_{\text{colony}}}{C_{\text{control}}},$$

where C refers to the concentration of the nutrient in question. When LRR is greater than zero, the concentration of the nutrient is greater in colony soils than in control soils, and when LRR is less than zero, the concentration of the nutrient is higher in control soils than in colony soils. Variance is calculated as:

$$\text{var}(\text{LRR}) = \frac{(SD_{\text{colony}})^2}{N_{\text{colony}}(C_{\text{colony}})^2} + \frac{(SD_{\text{control}})^2}{N_{\text{control}}(C_{\text{control}})^2},$$

where SD and N represent the standard deviation and sample size of each group from each study.

2.4.2 | Statistical analysis

We ran all statistical analyses using the METAFOR package (v 2.4-0; Viechtbauer, 2010) in R version 4.0.2. For each nutrient, we calculated a summary LRR and 95% confidence intervals (CI_{95}) via a random effects model by using the LRR and variances derived from each individual study. We used a random effects model to account for

variation within each study as well as between all the studies, as there was considerable variation in study methodologies. Random effects models are commonly used in ecological meta-analyses because they allow the estimated effect sizes to vary due to differences observed from sampling error and from true ecological differences between the studies (Anderson et al., 2015). For each nutrient, we calculated the Q-score and I^2 index of heterogeneity to determine the amount of unexplained between-study variation and we measured this with a restricted maximum likelihood estimator. When Q-scores and I^2 were significant ($p < 0.05$), we conducted subgroup analyses using meta-regressions with the following moderators: island vs. mainland, geographic location of sites (polar, temperate and tropical) and seabird order, to determine if the variation in effect sizes could be explained. For each moderator group, we hypothesize the following:

1. Island versus mainland: Island ecosystems are isolated from mainland systems, thus are often dependent on subsidies vectored by mobile organisms, such as seabirds (Buelow et al., 2018). As such, we hypothesize that studies on island environments will be more enriched than studies on mainland environments.
2. Geographic location: studies in nutrient-limited environments, such as within polar regions, will be more enriched than studies in regions not limited by nutrients.
3. Seabird order: Enrichment of soils in seabird colonies will vary with seabird order and with nutrients.

Further partitioning (e.g. down to seabird family or genus) was not possible due to our limited sample sizes. We considered the overall effect size and the effect size for each study to be significant if the 95% CI did not overlap zero.

3 | RESULTS

3.1 | Overall trends in publications

A total of 181 papers were critically reviewed and were published over 53 years (1968–2021). Overall, there was an increase in the number of papers published in recent years, with more than half of the reviewed literature published within the last 15 years (64.1%, $n = 116$) with 2015 ($n = 16$) and 2013 ($n = 11$) having the most publications. The proportion of single- and multi-year studies were similar (44.2% and 41.4% respectively; Table 2); however, 26 studies (14.4%) did not report the number of years of sampling. For multi-year studies, most were 2 years in duration (22.1%, $n = 40$), and the two longest studies were 8 years.

TABLE 2 Proportion of publications ($n = 181$) included in this review with the variables and reported metrics extracted. Habitat types are based on the IUCN Habitat Classification Scheme which includes 16 broad habitat types at level 1, and a further 119 types listed at level 2 (The International Union for Conservation of Nature, 2012)

Variable	Reported metrics	% of publications
Location	Island	76.8
	Mainland	23.2
Ecosystem	Terrestrial	91.2
	Intertidal	3.9
	Both	5.0
Habitat ^a	Grassland	
	Tundra	18.2
	Subantarctic	5.0
	Temperate	3.9
	Subtropical/Tropical Dry	0.6
	Forest	
	Temperate	13.8
	Boreal	5.5
	Subtropical/Tropical Dry	3.9
	Subtropical/Tropical Mangrove	1.7
	Subtropical/Tropical Moist Lowland	0.6
	Subantarctic	0.6
	Desert	
	Cold	17.1
	Hot	3.3
	Shrubland	
	Mediterranean	7.7
	Subantarctic	0.6
	Boreal	0.6
	Subtropical/Tropical Dry	0.6
	Temperate	0.6
	Marine Intertidal	
	Tidepools	3.3
	Rocky Shoreline	1.1
	Mangrove (Submerged Roots)	0.6
	Wetlands	
	Permanent Freshwater Lakes (> 8 ha)	1.1
	Permanent Freshwater Pools (< 8 ha)	1.1
	Tundra Pools	0.6
	Bogs, Marshes, Swamps, Fens, Peatlands	0.6
	Marine Coastal/Supratidal	
	Sea Cliffs/Rocky Offshore Islands	1.1
	Coastal Brackish/Saline Lakes/Pools	0.6
	Artificial Aquatic	
	Ponds	0.6
	Irrigated Land	0.6
	Savanna (Dry)	0.6
	Multiple	4.4
Control site	Yes	75.1
	No	22.1
	Not mentioned	2.8

TABLE 2 (Continued)

Variable	Reported metrics	% of publications
Number of sampling sites	1	7.7
	2–4	38.1
	5–7	17.7
	8–10	11.0
	11–15	6.1
	16–20	8.8
	>21 (maximum 64)	7.7
	Not mentioned	2.8
Multi-year study	Yes	41.4
	No	44.2
	Not mentioned	14.4
Study duration (years)	<1	44.2
	2	22.1
	3	12.7
	>4 (maximum 8)	6.6
	Not mentioned	14.4
Number of focal species	1	56.9
	2	16.0
	3	8.3
	>4 (maximum 29)	15.5
	Not mentioned	3.3
Was guano measured?	Yes	47.0
	No	53.0
Measure of guano? (<i>n</i> = 85 studies) ^b	Nutrient content	32.4
	Elemental analysis	21.6
	Stable isotope analysis	18.0
	Defecation rate	12.6
	Guano cover (%)	5.4
	Pre-determined from literature	5.4
	Plastic load	2.7
	Presence of guano	1.8
Other measures of seabird influence	Presence /Absence of seabirds	81.8
	Feathers	7.7
	Eggs/shells	4.4
	Carrion/carcasses	3.3
	Boluses/pellets	2.2
Seabirds are vectors of ^c	Nutrients	84.5
	Inorganic pollutants	24.3
	Organic pollutants	5.5
	Physical pollutants	1.7
Overall effect of seabirds	Positive	49.2
	Negative	19.9
	Mixed	22.7
	Neutral	8.3

(Continues)

TABLE 2 (Continued)

Variable	Reported metrics	% of publications
Factors sampled ^c	Soil	64.1
	Terrestrial vegetation	44.2
	Water	19.3
	Invertebrates	17.1
	Sediment	13.3
	Detritus/leaf litter	9.9
	Vertebrates	9.9
	Benthic/aquatic vegetation	9.9
	Intertidal organisms	5.5
	Plankton	4.4
	Other	14.9

^aFor definitions of habitat classes, readers are directed to <https://www.iucnredlist.org/resources/habitat-classification-scheme>

^b20 studies measured guano using two different analyses, and three studies used three different analyses

^cThe variables 'Seabirds are vectors of' and 'Factors sampled' do not sum to 100% as many publications studied nutrients and pollutants simultaneously or sampled multiple factors

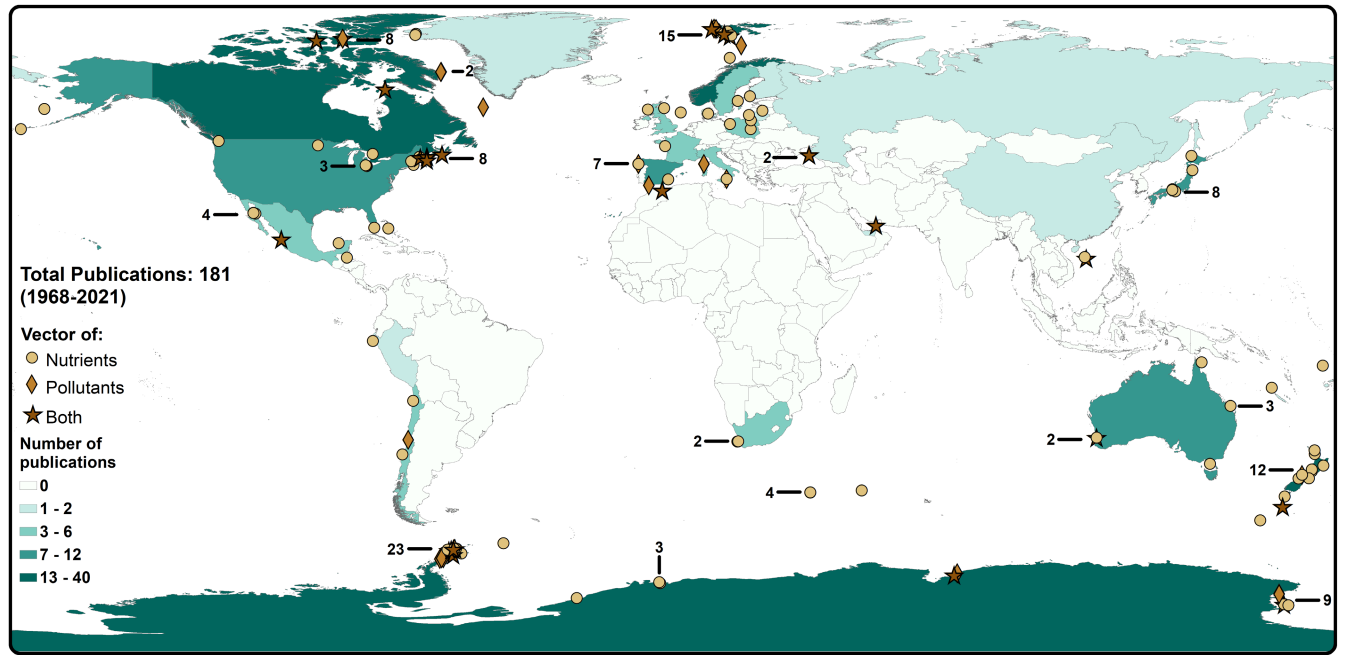


FIGURE 2 Most publications on seabirds as vectors of nutrients and pollutants were completed in the Northern Hemisphere, with a particular focus on regions in the Arctic. However, the country with the most studies was Antarctica ($n = 40$). Unique markers are used for publications examining nutrients (yellow circle), pollutants (tan diamond) or a combination of both (brown star; one marker per publication). Countries are shaded teal depending on the number of studies published. Several locations were studied frequently (e.g. King George Island, Antarctica) and for these areas a number is included to indicate how many studies took place in that region

3.2 | Distribution of publications

From the 181 publications, a total of 1,418 sites were examined from 30 different countries (Figure 2). Antarctica was the most represented (22.1%, $n = 40$ publications), followed by Canada (10.5%, $n = 19$) and Norway (9.9%, $n = 18$). Six publications spanned multiple countries while all others included only one country. Research was more heavily focused in the Northern

Hemisphere, with 104 studies (57.5%). Research spanned both terrestrial (91.2%, $n = 165$ publications) and intertidal (3.9%) environments (purely oceanic studies were not included in this review), and a small handful of studies incorporated both (5.0%). A diversity of habitats was covered, including grassland (27.6%, $n = 50$ publications; Table 2), forest (26.0%, $n = 47$) and desert (cold: 17.1%, $n = 31$; hot: 3.3%, $n = 6$), and several papers studied multiple environments (4.4%, $n = 8$). Islands (76.8%, $n = 139$

publications) were studied more frequently than mainland habitats (23.2%, $n = 42$).

3.3 | Focal seabird species

A total of 86 seabird species from 12 families and four orders were covered in the literature, with Charadriiformes accounting for 41.9% (families: Alcidae, Chionidae, Haematopodidae, Laridae and Stercorariidae), followed by Sphenisciformes (21.7%; Spheniscidae), Procellariiformes (18.8%; Diomedidae, Hydrobatidae and Procellariidae) and Suliformes (17.6%; Fregatidae, Phalacrocoracidae and Sulidae). Adelie penguins *Pygoscelis adeliae* was the species studied the most (7.1%, $n = 23$ publications), followed by great cormorants *Phalacrocorax carbo* (6.8%, $n = 22$) and gentoo penguins *Pygoscelis papua* (6.5%, $n = 21$). Of the 181 papers reviewed, 56.9% ($n = 103$; Table 2) had a single focal species, while 39.8% included multiple species ($n = 72$, range = 2–29). A handful of publications did not report the seabird species concerned (8.8%, $n = 16$) or only reported a portion of the seabird species included within the study (3.9%, $n = 7$). Research was predominately done on species classified as Least Concern by the International Union for Conservation of Nature's Red List of Threatened Species (IUCN Red List; $n = 267$ occurrences/65 species), while research on species listed as Endangered on the IUCN Red List was minimal in comparison ($n = 15$ occurrences/6 species—*Procellaria westlandica*, *Megadyptes antipodes*, *Phalacrocorax capensis*, *Phoebastria fusca*, *Spheniscus demersus* and *Morus capensis*) and there were no studies on species listed as Critically Endangered.

3.4 | Metrics used to measure seabird influence

Publications were highly variable in terms of what tools and metrics were used to measure the influence of seabirds on a given system. Guano was examined in 85 publications (47.0%), of which 35 analysed samples for nutrient content (N, P or both), 23 quantified non-essential trace elements and 20 included stable isotope analysis (Table 2). When guano was not collected and analysed, proxy measures were used instead. The presence and absence of seabirds was commonly used in publications as a simple measure of guano (i.e. the presence of seabirds indicate that guano is being deposited, while the absence of seabirds infers that there is no guano deposition; 81.8%, $n = 148$), and this was often reported in conjunction with the analysis of guano (30.4%, $n = 55$). A small portion of publications measured other allochthonous inputs by seabirds, such as feathers (7.7%, $n = 14$) and eggshells (4.4%, $n = 8$). Forty publications (22.1%) did not include a control site (i.e. a location without seabirds to compare against).

A diversity of biotic and abiotic samples was measured to determine the effect of seabirds and the flow of subsidies in recipient environments. The most common was soil (64.1%, $n = 116$ publications), followed by vegetation (44.2%, $n = 80$) and followed by water (19.3%, $n = 35$). Most publications measured multiple factors (57.5%, $n = 104$

publications, range = 2–7 factors), while 33.7% ($n = 61$) sampled only one factor and 8.8% ($n = 16$) did not sample any (i.e. only measured guano).

3.5 | Seabirds as vectors

Nutrients were the only subsidies recorded in the early literature (1968–1990; Figure 3) and accounted for 84.5% ($n = 153$) of all studies considered in this review. In contrast, the first publication to consider seabirds as vectors of pollutants was in 1991 (inorganic pollutants; Godzik, 1991). Since that time, there has been an increase in the number of publications investigating the role of seabirds as vectors of pollutants; however, there has been a similar increase in the number of publications focusing on nutrients (43.8% of nutrient papers were published in the past decade; Figure 3) and scientific publishing in general. Overall, inorganic pollutants transported by seabirds were documented more in the literature (24.3%, $n = 44$) than organic pollutants (5.5%, $n = 10$). Several papers measured nutrients and pollutants simultaneously (nutrients and inorganic pollutants: $n = 27$; nutrients and organic pollutants: $n = 5$). Three publications measured physical pollutants (i.e. plastic particles) in guano.

A total of 89 publications (49.2%; Table 2) concluded that seabirds had a positive influence on the terrestrial environment, while 36 (19.9%) returned a detrimental result (see Section 4.4 below). Some publications returned mixed results (22.7%, $n = 41$), showing both negative and positive effects, while a small portion did not detect a significant change or reported neutral results (8.3%, $n = 15$).

3.5.1 | Contents of guano

There were 35 papers that analysed the nutrient content of guano. All but two of these papers analysed the total percentage of N or P (or both) in guano. Seabirds from the order Suliformes had higher levels of P than all other orders, with a $M \pm SD$ of $15.08 \pm 1.07\%$ P (range = 14.32–15.83%P; Table 3), whereas Procellariiformes typically exhibited guano with a higher N content (19.13 ± 3.46 , range = 15.19–21.70%N).

Across the 23 papers that analysed guano for trace elements, there were a total of 30 different elements analysed. The most common elements analysed were zinc and cadmium, examined in 17 papers, followed by copper ($n = 16$) and lead ($n = 15$). The concentrations of these elements, plus the next three most analysed (arsenic, manganese and nickel; $n = 13$, 10 and 10 respectively), were extracted from the relevant papers. Guano was high in zinc across all seabird orders, with a $M \pm SD$ concentration of $261.03 \pm 235.25 \mu\text{g/g}$ (range = 4.07–1,200.00 $\mu\text{g/g}$; Table 4). Copper concentrations were high as well (84.57 ± 100.13 , 1.24–350.00 $\mu\text{g/g}$), followed by manganese (47.25 ± 75.49 , 0.82–300.00 $\mu\text{g/g}$). Across seabird orders, there were some differences; for example, guano from Charadriiformes was high in arsenic, cadmium and lead compared to all other orders (6.48 ± 4.35 , 11.40 ± 25.44 and 15.19 ± 14.21 respectively), whereas Sphenisciformes exhibited concentrations of copper and manganese

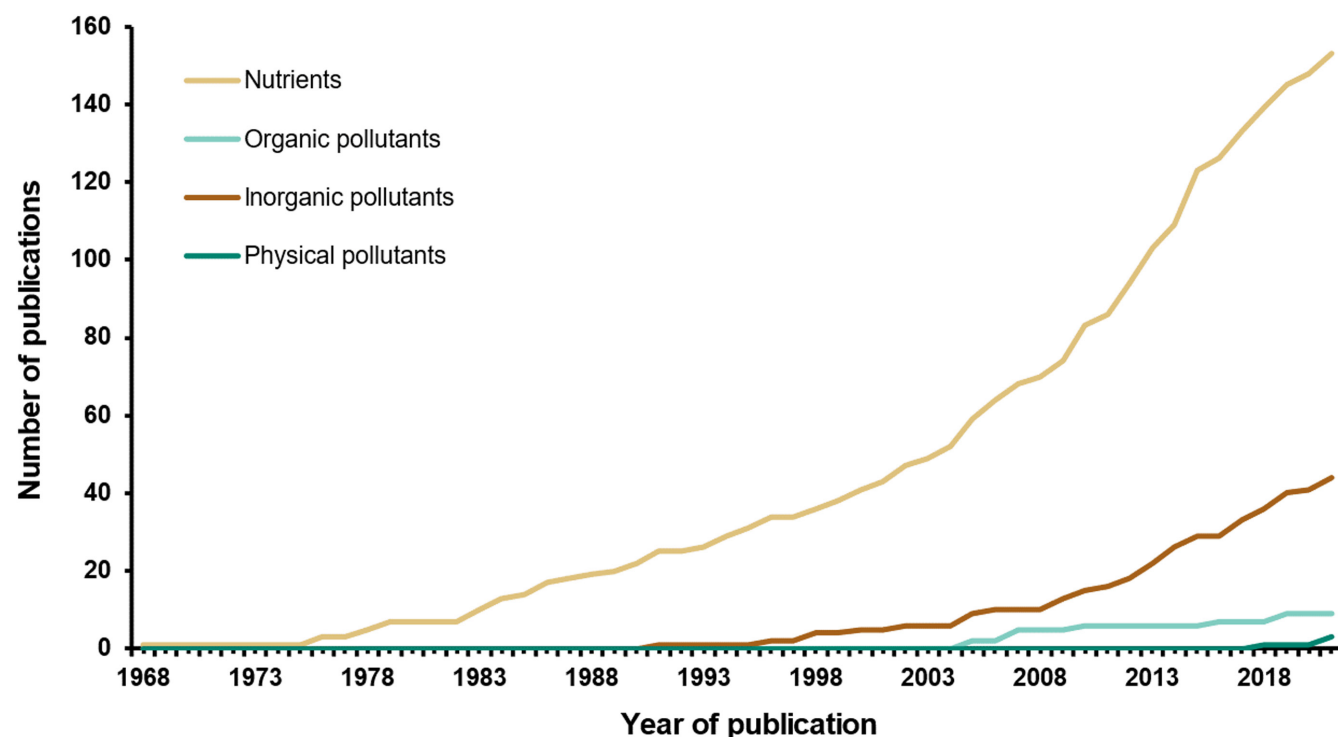


FIGURE 3 Growth in publications reporting on the different subsidies (nutrients, inorganic, organic or physical pollutants) transported by seabirds to land. Of the 181 publications reviewed, the majority explored nutrients. Inorganic pollutants were first reported in 1998, while organic pollutants were first reported in 2005. Physical pollutants (i.e. plastics) were not reported until 2018. A number of papers reported multiple subsidies (e.g. nutrients and inorganic pollutants), thus lines do not sum to 181

TABLE 3 Mean $\pm$ SD (range; n) of total nitrogen (%N) and total phosphorous (%P) analysed in guano, separated by seabird order

Order	Nutrient	
	Nitrogen	Phosphorous
Charadriiformes	7.07 $\pm$ 4.47 (1.39–15.20; 9)	1.54 $\pm$ 0.80 (0.47–2.76; 9)
Procellariiformes	19.13 $\pm$ 3.46 (15.19–21.70; 3)	1.16 $\pm$ 0.31 (0.90–1.50; 3)
Sphenisciformes	13.04 $\pm$ 6.03 (0.35–20.85; 10)	3.13 $\pm$ 2.80 (0.16–10.28; 10)
Suliformes	9.89 $\pm$ 9.35 (3.28–16.50; 2)	15.08 $\pm$ 1.07 (14.32–15.83; 2)
Overall	11.30 $\pm$ 6.52 (0.35–21.70; 24)	3.36 $\pm$ 4.22 (0.16–15.83; 24)

that were much higher than all other orders (177.50 ± 106.63 and 87.47 ± 142.21 respectively; Table 4).

3.5.2 | Meta-analysis

The concentrations of all nutrients were significantly higher in seabird colony soils than in control soils (%N: LRR = 1.15, $p < 0.0001$; %P: LRR = 2.11, $p < 0.0001$; NO_3^- : LRR = 2.18, $p < 0.0001$; and NH_4^+ : LRR = 2.81, $p < 0.0001$; Figure 4, Figures S3–S6). Ammonium levels in colony soils showed the biggest difference and were on average

41.1 times higher than control soils, followed by nitrate concentrations (5.3 $\times$ higher), total phosphorous (4.3 $\times$) and total nitrogen (2.1 $\times$). Q-scores and I^2 were significant for all models (all $p < 0.001$; Table S1), suggesting that the heterogeneity in effect sizes between studies was higher than what would be expected by sampling error alone. Subgroup analyses indicated that the impact of guano deposition on island systems was significantly greater for all nutrients in comparison to deposition on mainland systems, while differences in geographic zones or seabird orders was less pronounced and more variable across nutrient types (Figure 4). Studies conducted in polar regions demonstrated significantly higher soil nutrient values across nutrient types, whereas studies in temperate zones showed significant results for all but N. Studies undertaken in tropical zones were minimal thus firm conclusions cannot be drawn. Differences in soil nutrient concentrations due to deposition by different seabird orders were highly variable, with studies examining seabirds from the order Sphenisciformes displaying significantly high levels of ammonium in colony soils (LRR = 4.27, $p < 0.0001$; Figure 4). Overall, the addition of seabird guano to colony soils increases nutrient concentrations compared to control soils.

4 | DISCUSSION

Seabirds have a vast array of ecological functions both on land and at sea. They have been relied upon as ocean sentinels for decades, providing information on fish stocks (Montevecchi, 1993; Piatt et al., 2007) and trends in chemical and physical pollutants at sea

TABLE 4 Mean $\pm$ SD (range; n) of the seven most analysed metals in guano, separated by seabird order. Concentrations are reported in $\mu\text{g/g}$ dry weight. For those where only the mean is reported, $n = 1$

Order	Metal						
	Arsenic	Cadmium	Copper	Manganese	Nickel	Lead	Zinc
Charadriiformes	6.84 $\pm$ 4.35 (1.72–13.07; 8)	11.40 $\pm$ 25.44 (0.33–74.16; 9)	31.88 $\pm$ 19.68 (6.25–60.00; 8)	52.09 $\pm$ 48.42 (19.00–147.81; 7)	9.04 $\pm$ 6.07 (0.50–18.45; 6)	15.10 $\pm$ 14.21 (2.04–40.00; 9)	173.78 $\pm$ 114.43 (64.83–412.47; 10)
Procellariiformes	3.22 $\pm$ 1.50 (0.97–4.05; 4)	7.61 $\pm$ 1.20 (5.86–8.40; 4)	11.83 $\pm$ 7.07 (7.75–20.00; 3)	14.57 $\pm$ 10.40 (4.30–25.51; 5)	0.53 $\pm$ 0.04 (0.50–0.55; 2)	0.09 $\pm$ 0.01 (0.09–0.09; 2)	373.70 $\pm$ 463.01 (140.00–1,200.00; 5)
Sphenisciformes	1.69 $\pm$ 1.58 (0.40–3.81; 6)	7.88 $\pm$ 13.06 (1.67–37.14; 7)	177.50 $\pm$ 106.63 (1.24–350.00; 7)	87.47 $\pm$ 142.21 (0.82–300.00; 4)	10.23	1.76 $\pm$ 1.87 (0.41–5.39; 6)	240.36 $\pm$ 140.74 (4.07–510.00; 8)
Suliformes	6.45	12.14 $\pm$ 8.2 (6.34–17.93; 2)	21.10	20.79	20.00	1.60	390.11 $\pm$ 144.77 (232.94–518.00; 3)
Overall	4.29 $\pm$ 3.67 (0.40–13.07; 19)	9.58 $\pm$ 16.88 (0.33–74.16; 22)	84.57 $\pm$ 100.13 (1.24–350.00; 19)	47.25 $\pm$ 75.49 (0.82–300.00; 17)	8.50 $\pm$ 7.17 (0.50–20.00; 18)	7.83 $\pm$ 11.82 (0.09–40.00; 18)	261.03 $\pm$ 235.25 (4.07–1,200.00; 26)

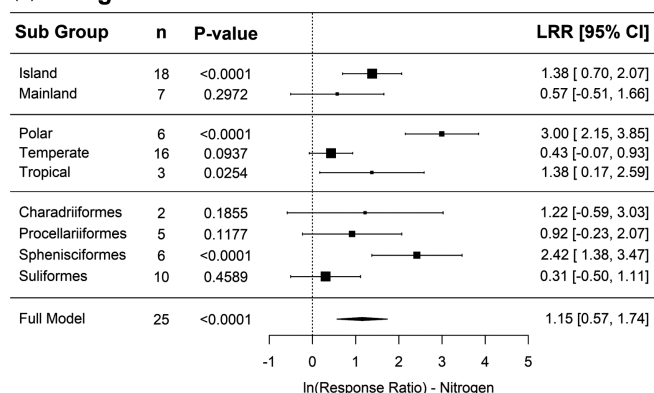
(Burger & Gochfeld, 2004; O'Hanlon et al., 2019). On land, we acknowledge seabirds as ecosystem engineers (Bancroft et al., 2005), capable of changing the physical and chemical conditions in their breeding and roosting grounds. While seabirds as physical engineers, such as burrowing, has been well described (Bancroft et al., 2004; McKechnie, 2006), data on the mechanisms surrounding seabirds as chemical engineers are comparatively limited. For example, only in the past 53 years have data on seabirds as vectors of nutrients or pollutants become available (Figure 3), with nearly 50% of the literature published in the last decade. This influx can be attributed to our growing knowledge of the important ecological services provided by seabirds (Mosbech et al., 2018; Şekercioğlu, 2006). The growth of literature on seabirds as vectors for nutrients has demonstrated their capacity to transport subsidies and alter environments in beneficial ways, but with the increasing awareness of ocean pollution, the birds' abilities to act as vectors for harmful subsidies has also been featured (Bourdages et al., 2021; Grant et al., 2021). This trend is mirrored in other fields globally where there has been a steady increase in pollutant research interest over time in both terrestrial and marine environments.

4.1 | Geographic distribution of studies

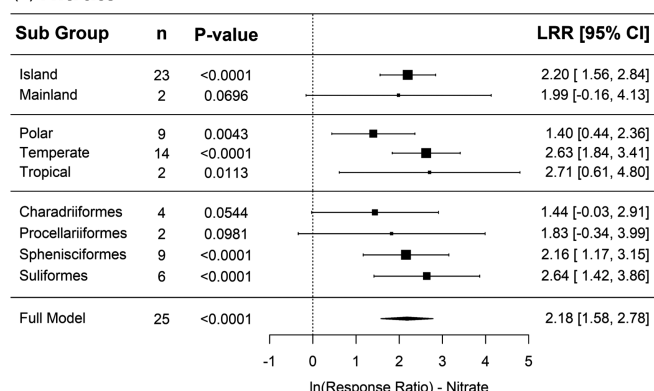
Many studies were undertaken in nutrient-limited environments, such as the Arctic and Antarctic, where external nutrient subsidies are considered integral for maintaining local production as resident sources of subsidies are often very minimal (Adame et al., 2015). For example, a large proportion of studies were undertaken in high latitude regions (>60 N/S; 40.9%, $n = 74$ publications), where terrestrial areas are characterized by cold temperatures and low nutrients (Maron et al., 2006); however, in places where seabirds congregate, there are hotspots of biological productivity and diversity (Brimble et al., 2009; Griffiths et al., 2010; Pereira et al., 2013), and this pattern was observed through our meta-regressions with colony soils in polar regions being significantly more enriched in nutrients compared to control soils (Figure 4). Similar observations were recorded in the Gulf of Mexico, which is known for its relatively productive waters juxtaposed against hyper-arid deserts, where the presence of seabirds increased production in comparison to areas without seabirds (Polis & Hurd, 1996). In general, the dynamics of many components (e.g. vegetation, consumers, soil properties) in terrestrial regions across the globe are highly dependent on nutrient subsidies transported by seabirds (Sanchez-Pinero & Polis, 2000).

Similar to the presence of colonial seabirds in nutrient-limited environments, seabirds are integral to many islands, as these ecosystems are isolated from external nutrient inputs and are often reliant on seabirds and other mobile vectors for the continued replenishment of vital subsidies (Buelow et al., 2018). Overall, island environments were studied more intensively than mainland environments, with 76.8% of publications based on islands. Most studies reported that islands with seabirds typically exhibited increased nutrient concentration in vegetation (Adame et al., 2015; Richardson et al., 2019) and a higher abundance of invertebrates (Markwell & Daugherty, 2002; Sanchez-Pinero & Polis, 2000; Zawierucha et al., 2016). In comparison, islands

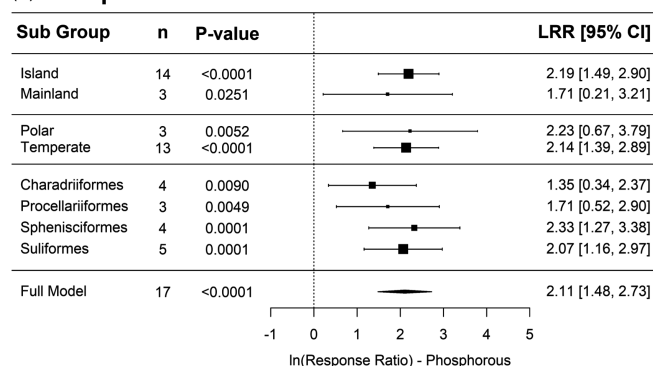
(a) Nitrogen



(c) Nitrate



(b) Phosphorous



(d) Ammonium

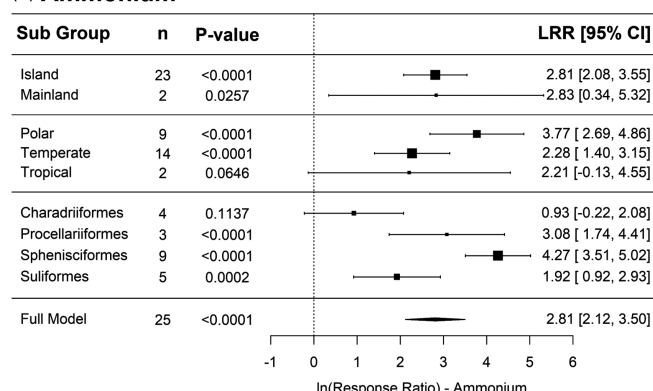


FIGURE 4 Forest plots showing the effect of guano deposition on nutrient concentrations ((a) nitrogen, (b) phosphorous, (c) nitrate and (d) ammonium) in soils. Effect sizes >0 show that nutrient concentrations were greater in seabird soils compared to control soils not impacted by guano deposition. The mean effect size (LRR, log risk ratio) and 95% confidence intervals are shown for the overall result (full model) and for each subgroup analysis: island vs mainland study sites, location of study sites (polar = >60°N or S; temperate = 30–60°N or S; and tropical = <30°N or S) and seabird order (Charadriiformes, Procellariiformes, Sphenisciformes and Suliformes). Confidence intervals that overlap zero are not significant. N and P were measured as percentage dry weight, while NO_3^- and NH_4^+ were measured in $\mu\text{g/g}$ dry weight. The subgroup 'Tropical' for phosphorous is not included as $n = 1$. There were three publications that studied multiple seabird species, thus the n for the full model for each nutrient type does not equal the corresponding n in Figure 1. Four studies included in all plots did not clarify the seabird species, thus the sum of n for the subgroup seabird order does not equal the corresponding n for the full model

without seabirds (e.g. control sites) often reported lower soil N and P (Rajakaruna et al., 2009; De la Peña-Lastra et al., 2020; Figure 4), and hence did not see those same benefits as islands with seabirds present. Across island ecosystems, as well as nutrient-limited environments, the effects of MDN from seabirds are substantial and promote the maintenance of these ecological communities (Polis & Hurd, 1996).

While there were hotspots of research activity in a handful of locations around the world, such as Svalbard, the Gulf of Mexico and King George Island (Antarctica), there are some obvious gaps. There were few studies in Africa (all were based in South Africa), with few data available from south-east and west Asia, as well as the east coast of South America (Figure 2). This is worrying considering that seabird populations around the world are in decline (Croxall et al., 2012; Dias et al., 2019) or have disappeared in some cases (Feare, 1978), which will have implications for recipient habitats as the transfer of MDN will be reduced. It is important to have a wide representation from a range of ecosystems and habitat types, and not just studies from nutrient-limited environments.

4.2 | Focal seabird species

Most seabirds were listed as Least Concern (77.2%) on the IUCN Red List, with very few data available for globally threatened species (Vulnerable, Endangered or Critically Endangered). While it is important to study all species, regardless of population status, it is concerning that very few studies have examined globally threatened species, given the significant influence seabirds have on their terrestrial environments (Rodrigues et al., 2021). Interestingly, seabirds that were historically important guano-producing birds (whose guano was mined for fertilizer) in South America were only studied once (Peruvian Booby, *Sula variegata*; Lucassen et al., 2017), despite two of the species (Guanay cormorant *Phalacrocorax bougainvillii* and the Peruvian brown pelican *Pelecanus thagus*) being listed as Near Threatened on the Red List. The guano produced by these species is evidently laden with nutrients (Szpak, Millaire, et al., 2012), thus a decline in their populations would lead to changes to their habitats, yet as these species have not been studied, we do not know what these changes would be.

Substantial, cascading and often detrimental changes may occur in areas where there once were thriving seabird colonies due to the reduction in MDN transported by the birds (Duda et al., 2020; Maron et al., 2006). The loss or major reduction in seabird colonies could result in the alteration in important ecosystem functions (Şekercioğlu et al., 2004). We could expect to see changes to invertebrate communities and vegetation assemblages, as observed on islands where predators have decimated seabird populations. For example, in the Aleutian Archipelago introduced Arctic foxes *Vulpes lagopus* preyed on seabirds resulting in a reduction in MDN and declining soil fertility, which transformed the habitats on multiple islands from grassland to a dwarf shrub-dominated landscape (Croll et al., 2005). Furthermore, the loss of seabird populations may not just impact nutrient cycling at localized levels, but potentially on a global scale as well (Riddick et al., 2018) as this could lead to a reduction in global N emissions.

4.3 | Metrics used to measure seabird influence

A range of methods were used to measure the influence of seabirds, ranging from simple methods such as the presence/absence of seabirds or percentage guano cover, to more quantitative, analytical methods (e.g. nutrient concentrations in guano). The presence/absence of seabirds is an easy, low-cost way to test the impact of seabirds in each area, with sites where seabirds are absent functioning as controls. Many studies benefitted from access to islands with and without seabirds (e.g. Markwell & Daugherty, 2002; Powell et al., 1991) while others used a gradient of seabird abundance (e.g. sites were chosen with increasing distance from a colony; Signa et al., 2013; Zmudczynska-Skarbek et al., 2013; Ziolk & Melke, 2014). While there is still merit in these qualitative studies, it can potentially be difficult to identify the exact cause of the observed effects, and definitively say that it was the presence of seabirds that resulted in the observed changes (Kolb et al., 2015), as the effects of seabirds can last long after they have left the area (Mizutani et al., 1991). This is particularly important as the nutrient concentration in guano varies considerably among seabird orders (Table 3) and by trophic level and foraging behaviours, as has been observed between coastal, oceanic and predatory seabirds (Wing et al., 2014) as well as between planktivorous and piscivorous seabirds (Zwolicki et al., 2013, 2016). While our review was limited by the number of studies examining the nutrient content of guano, and thus we only separated guano by order (Table 3), there is some discussion that guano nutrient content is different within seabird species, particularly those foraging in different locations and feeding on a variety of prey types. Several studies (5.4%) based their analyses on nutrient or pollutant levels in guano from previous literature. While there is some benefit in this (e.g. low cost), the results of this study demonstrate the highly variable nature of nutrients and pollutants in guano (Tables 3 and 4), thus authors should attempt to use values from the same species (or family) to ensure accuracy.

Stable isotope analyses were a popular method for tracing MDN in recipient terrestrial environments in many of the studies

(Table 2). Stable isotopes of a variety of elements (C, N, O, H and S) have been used extensively in seabird ecology for inferring foraging locations and providing insights into diet (Bond & Jones, 2009; Callaham et al., 2012; Hoenig et al., 2022) and their application as tracers of nutrient flows is well established (Hebert et al., 2006; Michener & Schell, 1994). They can easily trace seabird-derived nutrients in coastal and island food webs because of the distinction in marine $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ values compared to terrestrial N and C isotopes (Harding et al., 2004). The $\delta^{15}\text{N}$ values of consumers are enriched by 3–5‰ with each trophic transfer along the marine food web (Post, 2002), thus seabirds—being mostly tertiary consumers—generally have high $\delta^{15}\text{N}$ values (Lucassen et al., 2017). As guano contains considerable amounts of marine-derived N, stable isotope analyses can be applied to materials (e.g. soil, vegetation, invertebrates; Hawke & Clark, 2010) from within and outside seabird colonies to trace the flow of nutrients from the guano, and have been used successfully to determine the presence of seabirds in areas where colonies are no longer present (Kameda et al., 2006).

4.4 | Detrimental impacts of seabirds

While most literature focused on the transport of MDN and the substantial positive benefits derived from the addition of seabirds in terrestrial environments, there were some negative impacts associated with seabirds as well. The deposition of excessive amounts of guano can cause guanotrophication, or seabird-induced eutrophication, as has been observed with a range of seabirds, including cormorants and gulls (Kolb et al., 2012; Otero et al., 2015). An excess in nutrients can indirectly lead to decreases in abundance and diversity of faunal groups (Signa et al., 2015), or directly lead to the destruction of vegetation through poisoning (Molina-Montenegro et al., 2013), but the effects of guanotrophication are most often detected in waterbodies adjacent to seabird colonies (e.g. coastal lakes, rockpools; McColl & Burger, 1976; Portnoy, 1990; Martín-Vélez et al., 2019).

Guanotrophication is not the only negative consequence derived from guano deposition. Guano can contain high concentrations of inorganic pollutants (i.e. heavy metals) and POPs which can bioaccumulate in flora and fauna in recipient environments (Godzik, 1991; Otero et al., 2018). The literature regarding seabirds as vectors for such pollutants commonly focused on metals with very few studies exploring POPs (Figure 3). Several metals, particularly cadmium, copper and lead, were consistently analysed in guano samples (e.g. Celis et al., 2014; Espejo et al., 2014; Otero, 1998). These metals are often the focus of other environmental pollutant studies as they are well known for their toxicity, ability to bioaccumulate in organisms and their persistence and longevity in the environment (Šerić Jelaska et al., 2014). Many studies demonstrated the impact of pollutant laden guano on recipient environments by measuring metal concentrations in soils, vegetation and invertebrates, with results showing higher contaminant levels in samples from colonies compared to samples from control sites (Headley, 1996; Santamans et al., 2017; Shoji et al., 2019). Furthermore, of the few papers that

did explore POP transport by seabirds, the majority of papers established a similar pattern, with higher concentrations of POPs in soils and organisms within colonies in comparison to areas without seabirds (Choy et al., 2010; Evenset et al., 2004; Foster et al., 2011; Roosens et al., 2007). The transport of POPs by seabirds is 30 times more efficient than atmospheric transport, with potential risks to human health (Evenset et al., 2007).

While this review only returned three papers that examined microplastics in guano (<5mm diameter; Provencher et al., 2018; Bourdages et al., 2021; Hamilton et al., 2021), a small number of papers (not included in this review) have also demonstrated that it is not just microplastics that can be transported to seabird colonies via guano, with meso- (5–20mm) and macro-plastics (>20mm; Barnes et al., 2009) being deposited through boluses and carcasses (Buxton et al., 2013; Grant et al., 2021), or collected from sea or shorelines and used as nesting materials (Grant et al., 2018; Van De Crommenacker et al., 2021). These papers indicate that the incidence of plastics in seabird colonies is likely widespread and increasing. The literature surrounding the movement of plastics by seabirds to their colonies has increased in recent years, like the increase in pollutant papers detected in this review (Figure 3). It is unknown whether these plastics negatively impact the terrestrial systems they are deposited in; however, research in other fields (e.g. agriculture) suggest that plastics in soils can significantly affect temperature and N cycling (Seeley et al., 2020; Lavers et al., 2021) and microbial activity (de Souza Machado et al., 2018). As the deposition of guano increases the N content of soils (Figure 4a,c,d), the potential for concurrent positive and negative effects, if guano contains microplastics, is plausible, or may lead to a reduction in the incorporation of guano-derived N. Research in this field is much needed, as plastic pollution is predicted to increase in coming years and, subsequently, the incidence of plastics in colonies is also likely to increase.

4.5 | Future directions

Many papers do not report the most basic information, including the year(s) sampled, exact location studied or the seabird species involved (Table 2). It is critical that all published studies contain sufficient information so that the work may be repeated, included in meta-analyses, and results compared (Cassey & Blackburn, 2006). Furthermore, papers often did not report variance, presenting only mean values, or did not include detailed information on samples sizes. This made it challenging to conduct the meta-analysis, as papers that did not include these measures had to be excluded, drastically reducing the number of papers that could be synthesized (Figure 1). Soil types vary considerably (WRB, 2015) and play a significant role in the flux of nutrients and pollutants (Tian et al., 2017), yet this information was rarely presented across papers, particularly in those exploring the effects of guano addition in colony soils. The integration of soil science with pollution research in the context of ornithogenic transport would be a useful avenue to explore in the future studies. In addition to these recommendations of reporting basic but necessary information, we

recommend focusing research efforts on declining seabird species (e.g. those listed as Critically Endangered or Endangered on the IUCN Red List), the transport of pollutants by seabirds, particularly POPs, as this has been minimally studied over the years (Figure 3), as well as longer studies (i.e. >2 years) to measure annual trends.

5 | CONCLUSIONS

While we have highlighted the role of seabirds as vectors of both beneficial nutrients and potentially detrimental pollutants, seabirds remain integral to the continued success of many environments all around the world. In addition to providing nutrients, seabirds can regulate habitats through seed dispersal, physical ecosystem engineering, and assist with carcass and waste disposal. More broadly, birds (both terrestrial and marine) are well known to provide a highly diverse range of ecological functions and services, through regulation and supportive measures (Şekercioğlu, 2006). The potential decline and loss of these keystone species and the critical services they provide could be devastating for habitats globally (Bauer & Hoyer, 2014; Ellis, 2005; Şekercioğlu et al., 2004). Considering that many bird groups are currently in decline, with the biggest declines seen in seabirds, it is of utmost importance to continue to study avian groups, so that we can anticipate and mitigate potential consequences.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

AUTHORS' CONTRIBUTIONS

All authors conceived the ideas and designed the methodology; M.L.G. and J.L.L. collected the data; M.L.G. analysed the data and led the writing of the manuscript. All authors contributed critically to the drafts and gave final approval for publication.

DATA AVAILABILITY STATEMENT

The datasets supporting this article and relevant R code are freely available from the Institute for Marine and Antarctic Studies (IMAS) Data Portal: <https://doi.org/10.25959/ENWE-GB98> (Grant et al., 2022).

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