**Long after the event, or four things we (should) know about bird invasions**

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**Abstract**

The most significant single event in the study of alien bird invasions occurred in 1981, with the publication of John L. Long’s seminal book “Introduced Birds of the World”. The significance of this book derives not just from its content, but also from its timing, coincident with the 1982 SCOPE programme on the ecology of biological invasions. It was not long before studies started to appear that exploited the data in Long (1981) to address the questions posed by SCOPE regarding alien invasions. As a result, we arguably have a more complete understanding of the invasion process for birds than for any other taxon. Nevertheless, there are still some key issues in the study of bird invasions where understanding is not all it should be. We aim to highlight four of these issues, by arguing that (1) We don’t know half so much about bird invasions as we think, (2) Propagule pressure promotes invasions, (3) Colonization pressure matters, and (4) There is no evidence that escape from parasites promotes alien (bird) invasions. We expect some of the views expressed to be controversial, and others less so, but either way we hope this paper will stimulate others to provide better evidence for or against our propositions.

**Introduction**

The most significant single event in the study of alien bird invasions to date occurred in 1981, with the publication of John Long’s seminal book “Introduced Birds of the World”. The book is subtitled “The worldwide history, distribution and influence of birds introduced to new environments”, but it is the second of these three components that really mark out the book. The bulk of Long (1981) consists of an astonishing catalogue of occasions where bird species have been deliberately or accidentally liberated into areas beyond the limits of their native geographic ranges by human activities. Introductions (*sensu* Blackburn et al. 2011a) of more than 400 bird species are recorded in this catalogue, of which more than 200 species were considered to have established self-sustaining alien populations. Long (1981) mapped the native and alien distributions of these species, and provided a wealth of supplementary information on the circumstances of the introductions. Long (1981) was about as ideal a dataset for the study of human-mediated biological invasions as it would have been possible to conceive of at the time.

The significance of Long (1981) derives not just from its content, but also from its timing. While many people identify the publication of Charles Elton’s seminal 1958 monograph “The Ecology of Invasions by Animals and Plants” as the founding point of the modern study of human-mediated biological invasions, in reality a greater influence was probably a programme on this topic initiated in 1982 by the Scientific Committee on Problems of the Environment (SCOPE, an arm of the International Council of Scientific Unions). The request to SCOPE to fund a programme on the ecology of biological invasions grew out of observations by some of the world’s most influential ecologists on the impacts that alien species could have on natural ecosystems (Simberloff 2011). The programme aimed to answer three main questions (Williamson et al. 1986; Drake et al. 1989): 1) What factors determine whether a species becomes an invader or not? 2) What site properties determine whether an ecological system will resist or be prone to invasions? 3) How should management systems be developed to best advantage given the knowledge gained from studying questions 1 and 2? The SCOPE programme led to a large number of national initiatives, spawned several influential edited volumes, and introduced many senior and young scientists to the field of invasion biology (Simberloff 2011). It therefore prompted scientists to think about questions for which a large dataset on alien species introductions would have been particularly useful, at exactly the moment when just such a dataset (Long 1981) was published.

It was not long before studies started to appear that exploited the data in Long (1981) to answer questions about alien invasions. The first of which we are aware was by Moulton & Pimm (1983), who analysed data from Long (1981) and elsewhere to test for evidence of interspecific competition. They found that per species extinction rates increased as more alien species were introduced to islands in the Hawaiian archipelago, which was consistent with an influence of interspecific competition on the probability of species persistence. Subsequently, Long (1981) has formed the basis for a wide range of studies on facets of human-mediated bird invasions, including on the influence of competition (Moulton 1993), numbers of birds introduced (Cassey et al. 2004), life history traits (Cassey 2002; Sol et al. 2005) and location of introduction (Blackburn & Duncan 2001) on establishment success, on the types of species introduced (Lockwood 1999), and the reasons for introduction (Blackburn et al. 2009a). The data in Long (1981) have been particularly effective in advancing understanding of biological invasions since it was realized that the invasion process was best studied as a sequential series of stages, with success at each stage filtering the identities and characteristics of species entering the subsequent stage (e.g. Williamson 1996; Richardson et al. 2000; Blackburn et al. 2011a).

Arguably, we have a more complete understanding of the invasion process for birds than for any other taxon (Blackburn et al. 2009b), and much of the credit for this must go to John Long (Long himself was always surprised by what people made of his book; Mawson 2003). Nevertheless, more than thirty years on from Long’s influential tome, there are still some key issues in the study of bird invasions where understanding is not all it should be. Here, we aim to highlight four of these. Two of the issues we discuss are things we believe that we do know, but which we do not think are widely enough appreciated, or about which debate continues to rumble on in the literature. The other two issues are things that we believe we don’t know as much about as we would like to think. We expect some of the views we express to be controversial, and we hope this will stimulate others to provide better evidence either for or against our positions. We would like to think that some of the uncertainties will be put to rest by our comments below, but we are not holding our collective breath.

**We don’t know half so much about bird invasions as we think**

Long (1981) catalogued 426 bird species recorded as having been introduced somewhere in the world as a result of human activities. Many of these were deliberate introductions carried out in the nineteenth and early twentieth centuries by Acclimatization Societies – organizations specifically formed to promote the establishment of alien species (McDowall 1994) – and concerned species introduced for hunting or ornamental purposes. As a result, there is a preponderance of gamebirds (Order Galliformes: 64 species), wildfowl (Anseriformes: 27 species) and pigeons (Columbiformes: 31 species) on Long’s list, relative to the species richness of these groups (Blackburn & Duncan 2001), and a preponderance of species in these taxa also amongst successful introductions (Lockwood 1999). Long’s list also documents relatively large numbers of parrots (Psittaciformes: 51 species) and sparrows and their allies (Passeridae: 56 species) (Lockwood 1999; Blackburn & Duncan 2001). The species in these last two families were primarily moved beyond their natural range limits by the cage-bird trade – for example, 20% of all traded birds are parrots (FAO 2011) – and the large numbers of them in Long (1981) reflects the fact that cages are permeable barriers to incursions by alien species.

As noted above, Long’s (1981) catalogue has been a hugely influential source of information for studying the invasion process, yet it is now more than 30 years old. In the absence of an updated list, it would be easy to fall into the trap of thinking that bird introductions have become a thing of the past. The realignment of social (or at least scientific and political) attitudes against alien introductions has effectively killed the Acclimatization Societies. The last of them died out in New Zealand in the 1980s. More recently, fears about the cage-bird trade as a route by which human pathogenic strains of influenza might be imported into Europe have led to bans on this trade into European Union (EU) countries (Cooney & Jepson 2006). Both of these developments will undoubtedly have led to fewer species being moved beyond their natural range limits, and the global trade in wild birds declined from an estimated 7.5 million birds a year in 1975 to around 1.5 million in 2007 (Leader-Williams & Tibanyenda, 1996; http://www.cites.org/eng/news/pr/2007/070111\_EU\_bird\_ban.shtml). Unfortunately, new evidence suggests that these developments have not stopped incursions by alien bird species.

Dyer & Blackburn (unpublished) have brought Long’s (1981) list up to date by compiling a spatially and temporally explicit database on the distributions of exotic birds, called GAVIA (Global AVian Invasions Atlas). As of March 2014, the GAVIA database comprised 27,741 distribution records for exotic bird species. It is based on almost 700 published references and substantial unpublished information derived from consultation with more than 600 organisations and experts worldwide. This database is, as far as we are aware, the most comprehensive resource on the global distribution of exotic species in any major taxon, and allows the spatial and temporal dynamics of exotic bird population spread to be examined. GAVIA includes records of introduction for 973 bird species, of which 420 have established apparently viable populations (Dyer & Blackburn unpublished). Thus, GAVIA more than doubles the number of known introduced bird species, relative to the information in Long (1981), and increases the number of established species known by a similar percentage (Long lists just over 200 species as definitely or probably established).

GAVIA records dates of first introduction for species, when available, which allows us to explore temporal trends in introductions. The first known introduction dates for species listed in Long (1981) span the range 500 – 1993 AD (some dates are after the publication of Long, because the first *known* introduction date for a species in Long is for a *population* introduced after Long was published), with a median date of 1898 (N = 395). For species in GAVIA but not in Long, the dates span the range 1750 – 2010, with a median of 1979 (N = 347). A graph of bird introductions versus time for species with known first introduction dates shows that the rate at which species are first introduced is not slowing down (Figure 1), even though the pool of species available for first introduction is decreasing.

It should not really come as a surprise that bird introductions are an ongoing, and indeed increasing, phenomenon. First of all, there is abundant evidence from a variety of other taxa and spatial scales that the rate of invasions is not slowing down. For example, marine species invasions into San Francisco Bay classically show an accelerating rate of increase in the period 1850 – 1990 (Cohen & Carlton 1998). The number of new mammal species introduced to Europe shows exponential growth since 1500AD, as does the number of new species per year (Genovesi et al. 2009). The rate at which alien terrestrial invertebrate species establish in Europe also continues to grow, and in the period 2000 – 2007 was double that of 1950 – 1974 (Roques et al. 2009). In fact, the number of new bird species introduced into Europe has grown relatively modestly in this context, (Figure 2, with an exponential model fitting no better than a linear one). Worldwide, however, the temporal change in bird species introductions is well modeled by an exponential increase (R2 = 0.98, F3,6 = 190.3, P < 0.001; Figure 1). However, it is better modeled by a piecewise regression with a break point at 1860 (R2 = 0.996, F4,5 = 560, P < 0.001): the rate at which bird species introductions occur takes off rapidly at almost exactly the same time that the first Acclimatization Societies were founded (McDowall 1994), and it never looks back.

Second, the data catalogued by Long (1981) present a very Eurocentric view of the process of bird introductions. It is undoubtedly true that much of the history of bird introductions, in terms of quantity if not length, has been a tale of European endeavours deliberately to naturalize species (Crosby 1993), but the focus has shifted dramatically in recent years. Nowadays, most bird introductions are accidental, primarily the result of escaping pets (Lever 2005). The data in Long (1981) already identify this shift in composition (Blackburn et al. 2009a). Further, the most important correlate of variation in the number of invasive alien species across countries is the volume of merchandise imports (Westphal et al. 2008), suggesting that international trade is now the primary driver of species introductions. This is likely to be a result of higher colonization pressure (Lockwood et al. 2009; and see below). To understand modern patterns of bird introductions, we should be looking to regions with cultures of bird-keeping and high levels of international (and within biogeographically diverse countries, national) trade. In these regards, East and South-East Asia stand out.

Bird-keeping is deeply rooted in East Asian culture, and includes such activities as bird-walking and bird competitions (e.g. comparing birds on condition or song quality; Jepson & Ladle 2005). Eastern religions also practice prayer animal release (e.g. Environment and Animal Society of Taiwan 2009), and adherents buy and liberate enormous numbers of animals, including birds, for the purpose of accruing karma. These activities support an extensive bird trade, moving thousands of birds of hundreds of species beyond their natural range limits. A good example in these respects is provided by Taiwan. In a recent survey of a sample of Taiwanese bird shops, Su et al. (2014, unpub.) recorded more than 26,000 individual birds of more than 240 species for sale (see also Shieh et al. 2006). The total market value of these birds was over US$0.75 million. Over two-thirds of the species for sale were alien, and there is clear potential for the deliberate and accidental release of such birds to contribute to avian invasions. Indeed, at least 90 alien bird species are known from the wild in Taiwan, and around 35 alien species have breeding records there (Shieh et al. 2006; Su et al. unpub.). Nine of these species with breeding records are not listed in Long (1981) at all. Alien bird species have already proven to be a conservation threat in Taiwan through actual and potential hybridization with native bird species (Severinghaus 2007, Yao 2007; Li et al. 2010), and so the ongoing trade in cage birds is likely to be of significant conservation concern. This will be true not only in Taiwan, but also in other countries with active and developing markets in wild birds, such as China (Li & Jiang 2014) and Brazil (Regueira & Bernard 2012).

Overall, it is worrying that while scientists and policy makers know well the dangers posed by alien species (see Roy et al. 2014), the range of bird species (and species in most other taxa) with recorded introductions continues to grow, with no sign of the rate slowing down (Figure 1). It is also worrying that most introductions nowadays stem from the pet bird trade: this trade is likely to grow further worldwide because keeping birds is related to prosperity in developing regions (e.g. Jepson & Ladle 2005). It would be easy to imagine that bird introductions are no longer a problem because the cultures responsible for the “golden age” of acclimatization have abandoned this practice. Unfortunately, nothing could be further from the truth. The fact that these ongoing invasions provide new opportunities and questions for scientists studying the invasion process might be viewed as scant consolation for the problems that they are likely to cause.

**Propagule pressure promotes invasions**

The data in Long (1981) and GAVIA (Dyer & Blackburn unpub.) identify a large number of established alien bird populations, but also show that most introductions do not result in establishment. Understanding why some introductions succeed while others fail was a primary motivation for the first two questions in the SCOPE programme on the ecology of biological invasions. However, it turns out that a key element of the answer relates neither to characteristics of the species (question 1), nor to characteristics of the location (question 2).

A basic fact of population biology is that populations comprising fewer individuals are more likely to go extinct in any given period of time than populations with more individuals; in other words, extinction probability is negatively related to population size. The reasons for this are well known: small populations are more likely to suffer from effects of demographic and genetic stochasticity, more likely to be extirpated by environmental stochasticity, and more likely to suffer from Allee effects. Small population size is a major concern for conservation biologists, and the lower the number of individuals of a species, the higher the category of threat on the IUCN Red List to which it is assigned (IUCN 2014). It makes sense to assume that alien species released in lower numbers would also be more likely to go extinct, and hence be less likely to establish a persistent population at the location of introduction. Numerous empirical analyses, including many on data for introductions of alien birds compiled in Long (1981) and elsewhere (e.g. Thomson 1922), are consistent with this expectation (e.g. Dawson 1984; Cassey et al. 2004; Lockwood et al. 2005; Sol et al. 2005, 2012; Colautti et al. 2006; Hayes & Barry 2008; Blackburn et al. 2009a, 2011b, 2013; Simberloff 2009; Duncan et al. 2014).

Given that a positive relationship between establishment success and number of animals released (we term this the “propagule pressure effect”) is expected on theoretical grounds, is observed in data, and has been verified in experimental releases (albeit in taxa other than birds), one would not imagine that it could be controversial. Indeed, arguably it is not! Yet, there has recently been an attempt to make it so by a small group of collaborators working on historical bird introductions, who argue that propagule pressure does not drive establishment success (Moulton et al. 2011, 2012a,b, 2013).

Moulton and colleagues have used several lines of argument to promote this view, mostly based around the data for intentional bird introductions to New Zealand. First, they attempted to undermine the propagule pressure effect (Moulton et al. 2010, 2012a), arguing that historical data have a number of uncertainties and errors that render the conclusions of previous analyses of them insecure – “all that can be garnered from the historical record is a list of the species that were introduced, and some indication of when the individuals were released” (Moulton & Cropper 2014). This is incorrect, as recent detailed examination of the historical data demonstrate (Pipek et al. 2015), but even were it not, it would be interesting that the uncertainties and errors all apparently favour the propagule pressure effect. It is also interesting that Moulton et al. are happy enough to use these data, showing that the smallest releases for species successfully introduced to New Zealand were smaller than the largest releases for failed introductions, which they argued shows that release size does not determine success. However, this assumes that success is a deterministic function of release size (it clearly is not because some failed introductions are larger than those for some successes in many, if not most, species), and Moulton et al.’s analysis cannot recover a propagule pressure effect in simulation models where the effect is explicitly specified (Blackburn et al. 2011b). A robust re-analysis of Moulton et al.’s data on release sizes showed that the propagule pressure effect is indeed supported (Blackburn et al. 2011b).

Second, Moulton et al. argued that the propagule pressure effect arises because establishment success encourages people to release more animals, rather than because releasing more animals increases the likelihood of establishment. Blackburn et al. (2013) countered this by identifying six predictions of the hypothesis that establishment success drives propagule pressure, and demonstrating that all six predictions could be falsified. For example, if success is unrelated to the number of birds released, and successful populations had larger releases because their initial success led to more releases, then there should be no relationship between establishment success and propagule pressure, controlling for the number of releases. Yet, there is. Moulton & Cropper’s (2014) response, that this could be because both numbers introduced and number of introductions could be higher if undertaken by organizations with more resources, is no more than special pleading.

Moulton and colleagues subsequently shifted their position on the propagule pressure effect, arguing that “there is a somewhat increased probability of failure…for very small numbers introduced”, but that in many cases, more individual birds were introduced than needed to guarantee establishment (Moulton & Cropper 2014). That claim may be true, but is impossible to prove, and beside the point. For a population to have established, at least one individual within that population must have descendants surviving at a location after some period of time (Caswell 2001; Fox 2005). The probability that at least one individual leaves a surviving lineage depends on the birth and death rates of individuals in the population, and hence on features of the species and environment concerned, but inevitably increases with the number of individuals initially present (Leung et al. 2004; Jerde & Lewis 2007; Duncan et al. 2014). It is hard to conceive of realistic situations where introducing more birds would lead to lower per capita establishment probabilities, as would be necessary for success not to be a positive function of propagule pressure, at least for the release sizes we tend to see in birds (typically in the range 10 – 500, median = 50 for deliberate releases; Blackburn et al. 2009a).

Ultimately, therefore, logic dictates that propagule pressure must enhance the likelihood of establishment success, and success is indeed consistently higher for bird populations where larger numbers have been introduced, within species, within countries, across regions and globally (Blackburn et al. 2009a, 2011b). Moulton & Cropper (2014) argue that “That tendency explains little of the historical pattern of establishment outcomes for passeriform birds”. Propagule pressure does indeed explain only a fraction of the variance in establishment success, and characteristics of the species and environment clearly matter (Duncan et al. 2014), but propagule pressure is the nevertheless the most robust and consistent predictor of establishment success that we know (Cassey et al. 2005).

**Colonization pressure matters**

While propagule pressure is the most consistent predictor of alien bird establishment success, other factors must also play a role. Duncan et al. (2003) distinguished between three broad categories of determinants of success: location-, species-, and event-level factors (the last category being effects that can differ between two introductions of a given species to a given location, such as propagule pressure). It makes sense that whether a population succeeds or fails will depend on characteristics of the species concerned, and of the location to which it is introduced. Indeed, these effects were the primary scientific focus of the SCOPE programme (questions 1 & 2), and resulted in substantial literatures on these issues (Simberloff 2011).

Focusing on location-level effects, Shea & Chesson (2002) distinguished three broad features of the area of introduction that are likely to influence establishment, hypothesizing that fewer enemies, more resources and benign environmental conditions should all promote success. These predictions make sense, and are easy to state, but in practice there are so many axes along which their effects might act that studies rarely address them directly. Instead, most studies of location-level effects consider surrogate variables, such as the latitude at which a location lies, whether it is on an island or continental mainland, or whether it represents “pristine” or disturbed environments (Blackburn et al. 2009a). Spatial structure in the distribution of the bird species listed in Long (1981) identifies rich assemblages of aliens at mid-latitudes, and on archipelagoes such as the Hawaiian Islands, New Zealand and the Greater Antilles (Figure 3). These patterns, together with formal analyses of alien species richness, have been used to conclude that areas with simpler biotic communities (i.e. islands, high latitudes) are easier to invade than species rich areas (i.e. the continental tropics), as long as those simpler communities are not in environmentally harsh (i.e. polar or desert) regions.

Unfortunately, however, there is an analytical problem that has often been overlooked by studies of location-level effects on establishment success, and which may invalidate their conclusions. The number of alien species established at a location, *E*, is necessarily some fraction of the total number of species introduced there (or close to there), *I*, such that

*E = IS* Eqn. 1

where *S* is the establishment probability (Lonsdale 1999). Lockwood et al. (2009) term *I* “colonization pressure”. Thus, the hotspots of alien species richness in Figure 3 may be hotspots (high *E*) because they are easy to invade (high *S*), or because they are no easier (or even harder) to invade, but had higher colonization pressure (high *I*). For example, the Hawaiian Island of Maui has *E* = 27 alien bird species established, whereas Mauritius has *E* = 19. It would be easy to conclude that Maui is easier for birds to invade than Mauritius. However, the colonization pressure for Maui was *I* = 47 species, while for Mauritius *I* = 33 species. Thus, *S* is more or less identical for these two islands (0.57 vs. 0.58, respectively); Maui only appears easier to invade because of its higher colonization pressure (Lockwood et al. 2009).

Colonization pressure makes life difficult for anyone wishing to understand causes of spatial or temporal variation in establishment success, or in determinants of alien species richness. *E* and *I* in Equation 1 are not independent, because *E* is a subset of *I*, and hence is always constrained to be <*I*. The null hypothesis for the relationship between *E* and *I* is therefore not zero, but a positive relationship (see Brett 2004). This means that one cannot analyze causes of variation in *E* directly, but must instead model the number of established species out of the total number of introduced species as a binomial response variable (Lonsdale 1999; Lockwood et al. 2009). This is straightforward enough for some questions, such as causes of variation in establishment success in different acclimatization regions (Blackburn et al. 2011b; see above), or in temporal variation in establishment success at a location (Lockwood et al. 2009). However, it creates substantial problems for attempts to answer other questions.

For example, consider tests of hypotheses for the spatial variation in alien species richness in Figure 3. Any such test would need to account for colonization pressure. However, most locations where invasive species occur are not locations where the species were introduced, because the species have subsequently spread. Richness variation across regions may be attributable to colonization pressure (i.e. Maui vs Mauritius), whereas richness variation within regions may or may not (e.g. *E* can and does vary across Maui, but *I* = 47 for the whole island), but models are likely to need to address causes of richness at both levels.

Failing to account for colonization pressure has led to some erroneous conclusions in the invasion literature. For example, Brown (1989) showed a negative relationship between the number of native and number of alien bird species in four regions (Hawaii, New Zealand, Australia and North America) as evidence that historically isolated areas with low native species diversity are relatively easily invaded. However, this is not a valid conclusion in the absence of data on colonization pressure. Subsequently, Sol (2000) showed that the percentage success rates for bird introductions did not differ between Australia and New Zealand, although they did between Hawaii and North America. Sol (2000) nevertheless further showed that the latter difference could be attributed to the characteristics of the different species introduced at the different locations: there was no location-level difference in the success rate for those species introduced to both Hawaii and North America. The idea that simpler communities, and especially oceanic islands, are easier to invade is a seductive one, but robust evidence in support of such statements that also accounts for colonization pressure is surprisingly elusive.

**There is no evidence that escape from parasites promotes alien (bird) invasions**

While colonization pressure undoubtedly needs to be factored in to any consideration of location-level effects on establishment success, there is still evidence that success varies across locations (Duncan et al. 2014). One or all of Shea & Chesson’s (2002) broad categories of determinants of success (fewer enemies, more resources or benign environmental conditions) is therefore likely to be the cause.

There is abundant evidence that biotic interactions regulate natural populations of animals and plants, and so it comes as little surprise that these processes are frequently invoked to explain the success or otherwise of human-mediated biological invasions (e.g. Elton 1958). One mechanism by which this success might be mediated is now generally termed the Enemy Release Hypothesis (ERH) (Keane & Crawley 2002). This hypothesis proposes that alien species escape the negative impacts on reproduction and survival they experience from their natural enemies, including predators/herbivores and parasites/pathogens, left behind in their native geographic ranges. In consequence, they experience lower levels of enemy impact than native species in their new ranges. If the reduction in enemy regulation translates into increased population growth, then alien species may be able to increase rapidly in numbers, escaping the stochastic effects that afflict small populations (Allendorf et al. 2012) and hence increasing the probability of establishment and invasive spread.

The last few years have seen an upsurge of interest in the role of parasites (including pathogens) in the context of enemy release, following an influential paper by Torchin et al. (2003; cited more than 760 times by December 2014 according to Google Scholar). Torchin et al. showed that parasite species richness and prevalence was generally lower in the alien than the native range for a variety of introduced species, including molluscs, crustaceans, fishes, amphibians, reptiles, mammals and birds. For example, 44 parasite species have been recorded on European starlings *Sturnus vulgaris* in their native range, but only 9 in their North American alien range. Torchin et al. (2003) do not explicitly conclude that the observed escape from parasites causes invasion success, but they do note the implications of their findings for the potential for biocontrol of invasive species.

Subsequently, a range of studies has assessed evidence of release from parasites in biological invasions (reviewed by Colautti et al. 2004; Mitchell et al. 2006), with outcomes that depend on the type of study undertaken (Colautti et al. 2004). Biogeographical studies compare rates of parasitism (richness and/or prevalence) in native and introduced populations of the same species (as Torchin et al. 2003), and tend to support the ERH. Thus, introduced plant populations are exposed to fewer species of pathogens, are less impacted by pathogens and herbivores, and are less negatively impacted by soil organisms than are native conspecific populations (Callaway et al. 2004; Mitchell et al. 2006; Inderjit & van der Putten 2010). In contrast, community studies compare rates of parasitism for native and introduced populations of different species within the same community, and are much more equivocal in their support for the ERH (Colautti et al. 2004).

Differences in the general outcomes of biogeographical versus community studies has resulted in the validity of these tests being questioned in the literature. On the one hand, Colautti et al. (2004) argued that biogeographical tests are flawed because they compare the parasite richness of introduced populations with that of entire native species, rather than with the specific source population(s) for the introduction. However, this bias should not affect some measures of prevalence, or experimental tests of pathogen impact (Inderjit & van der Putten 2010). In contrast, Mitchell et al. (2006) argued that many within-community comparisons of plants, at least, are poor tests of the ERH because they rely on estimates of visible damage from natural enemies, rather than on measures of performance or demographics. They also tend to compare congeneric species of native and aliens, and will thus be conservative tests of ERH if invaders with close relatives in the native community are more likely to acquire new enemies from them. Whether biogeographical or community studies best represent the true situation with respect to escape from parasites and the ERH therefore remains an open question.

There is, however, a much more fundamental problem with evidence from both biogeographical and community studies for the ERH in terms of escape from parasitism: none of it actually informs about the causes of invasion success. A species must pass through a succession of stages – transport, introduction, establishment and spread – in order to negotiate the pathway from native to alien invasive, and it may fail to become an invader if it falls at barriers imposed at any of these stages (Blackburn et al. 2011a). The ERH is likely primarily to operate at the establishment and spread stages. In terms of establishment, increased population growth arising from a reduction in enemy regulation may increase the likelihood that an introduction can escape the perils of small population size (and so establish). In terms of spread, release from enemies may increase the likelihood that an established population can produce enough colonists to enable dispersal and establishment at further new locations For the ERH to influence establishment, however, it is necessary for populations that successfully establish to have benefitted more from release from the impacts of parasites than those that fail to establish. As most tests are framed in terms of parasite species richness, this would equate to successful species having fewer parasites (or to have escaped more) than those that fail to establish. Similarly, if the ERH influences the ability of an established population to spread, we would expect a positive relationship between extent of spread and degree of release from parasitism. However, neither biogeographical nor community studies inform about these comparisons: they say nothing about the level of parasitism in those populations that failed to establish relative to those that succeeded, nor do they tell us whether level of parasitism influences tendency to, or extent of, spread.

To be informative about the influence of the ERH on invasion success, therefore, studies need to compare the extent of escape from parasitism in populations that are introduced and become established versus those that are introduced but do not, or in populations that establish and spread to varying extents. [In terms of the framework for comparisons in invasion studies proposed by van Kleunen et al. (2010), this is comparisons of type B, rather than A (community studies) or F1 (biogeographic studies).] We are aware of only two studies that have explored the ERH in this fashion for parasites (Mitchell & Power 2003; van Kleunen & Fischer 2009), and in alien plant species rather than birds. Mitchell & Power (2003) showed that, among plant species listed as natural area invaders, species that experienced more complete pathogen release were more widely invasive. Here, however, invasive means the number of states and non-governmental organizations that list the species as an invasive problem in natural areas, and so is not a direct measure of extent of spread. Moreover, the relationship was only found when non-noxious and non-invasive alien species were excluded from the data set, and even then the analysis is not robust to the inclusion of an outlying data point. Van Kleunen & Fischer (2009) showed that the geographic spread of alien plants introduced from North America to Europe was negatively associated with their release from fungal pathogens, in direct opposition to the ERH. Additionally, Klironomos (2002) showed that interactions between plants and soil microbes were more likely to be positive for invasive alien than rare native plant species, and that the strength and sign of such interactions was positively related to abundance for plants in an old field meadow community in Canada. Callaway et al. (2004) showed that plant-soil interactions were more likely to be positive in the alien than the native range of the plant *Centaurea maculosa*. Taken together, these studies suggest that the extent of plant invasions may be related to the degree of escape from negative interactions with soil biota, but they stop short of demonstrating such a relationship. To date, therefore, there is arguably no study that can be considered to support the ERH in terms of the influence of escape from parasitism on invasion success, not only for birds, but for any taxon.

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Figure 1. A graph of bird introductions over time for species with known first introduction dates. Dates are grouped into 50 year intervals starting from 1511AD. The few dated introductions before 1510AD are excluded. Black portions of the bars represent species catalogued in Long (1981), grey portions represent species not catalogued in Long (1981). The orange lines show the best-fit piecewise regression lines, with the split point (lowest mean square error) at 1860, coinciding with the appearance of Acclimatization Societies (see text).

Figure 2. The rate at which alien bird species have been introduced to Europe (number of species introduced per decade). From data in Kark et al. (2009).

Figure 3. A map of the number of alien bird species found in equal area grid cells (96.486 x 96.486 km) across the land surface of the world. The data were compiled by M. Parnell, from the maps in Long (1981), and plotted by N. Pettorelli. From Blackburn et al. (2009a).

**Figure 1**



**Figure 2**



**Figure 3**

