Taking cues from ecological and evolutionary theories to expand the landscape of disgust 1 2 3 Authors: *Love, A.E^a., *Heckley, A.M^b., Webber, Q.M.R^a. 4 * These two authors contributed equally to this work 5 6 Affiliations: ^aDepartment of Integrative Biology, University of Guelph, Guelph, Ontario, 7 Canada; ^bDepartment of Biology and the Redpath Museum, McGill University, Montreal, 8 Quebec, Canada 9 10 Corresponding authors: 11 Allegra Love: allegra@uoguelph.ca Alexis Heckley: alexis.heckley@mail.mcgill.ca 12 13 14 ORCID ids: Love: 0000-0002-4637-6850; Heckley: 0000-0002-5644-3535; Webber: 0000-0002-15 0434-9360 16 17 Author Contribution Statement: Conceptualisation: AL, AH, OW (equal); Writing - Original draft: AL, AH (lead); QW (supporting); Writing - Review and editing: AL, AH, QW (equal); 18 Supervision: QW (lead). 19 20 21 Acknowledgements: We would like to thank Kiyoko Gotanda for her thoughtful comments on 22 an earlier version of this manuscript. 23 24 Funding: AMH and AEL were supported by Natural Sciences and Engineering Research 25 Council (NSERC) Postgraduate Scholarships – Doctoral. QMRW was supported by an NSERC 26 Discovery Grant. 27 28 Data Availability Statement: This paper is a review and has no associated data. 29 30 31 **Abstract:** Behavioural avoidance of parasites in the environment generates what is known as the 32 33 "landscape of disgust" (analogous to the predator-induced "landscape of fear"). Despite the 34 potential for improving our inference of host-parasite dynamics, three limitations of the 35 landscape of disgust restrict the insight that is gained from current research: 1) many host-36 parasite systems will not be appropriate for invoking the landscape of disgust framework, 2) 37 existing research has primarily focused on immediate choices made by hosts on small scales, 38 limiting predictive power, generalizability, and the value of the insight obtained, and 3) relevant

39 ecological and evolutionary theory has yet to be integrated into the framework, challenging our ability to interpret the landscape of disgust within the context of most host-parasite systems. In 40 this review, we explore the specific requirements for implementing a landscape of disgust 41 42 framework in empirical systems. We also propose greater integration of habitat selection and 43 evolutionary theories, aiming to generate novel insight, by exploring how the landscape of 44 disgust varies within and across generations, presenting opportunities for future research. Despite interest in the impacts of parasitism on animal movement and behaviour, many unanswered 45 46 questions remain.

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Keywords: Co-evolution; disease ecology; habitat selection; host-parasite dynamics; natural
selection; parasite avoidance

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1. Introduction

51 Individual organisms interpret cues from their environment to evaluate risks and inform 52 decisions to increase fitness. Animal behaviour can change based on perceived associations between a given cue and potential fitness costs or benefits, in time leading to possible 53 54 behavioural adaptation and predictable responses [1,2]. When cues are associated with the risk of 55 infection with micro- or macro-parasites (hereafter 'parasites; [3]), the perception of cues can result in feelings akin to the human feeling of disgust [4–6] and can elicit behavioural avoidance 56 57 of infectious agents [5-12]. The distribution of cues in an environment and the resulting spatial 58 distribution of organisms has been termed the 'landscape of disgust' [4], analogous to the predator-induced "landscape of fear" in which organisms respond to predator-associated cues 59 60 (sensu [13]). By describing the cues that hosts encounter, and thus may interpret and respond to, 61 the landscape of disgust is a tangible and interactive property of host-parasite systems. The 62 landscape of disgust framework has motivated increasing research on the ecological impacts of 63 parasitism in nature [4,14,15], in particular, non-consumptive effects (i.e., impacts to hosts beyond direct consumption, including changes to behavioural and developmental traits, among 64 65 other traits; [16–18]).

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Despite the potential value of the landscape of disgust framework, three key limitations restrict the applicability of the landscape of disgust in natural systems. First, in many natural hostparasite systems, the landscape of disgust does not exist because potential hosts will not always behaviourally avoid infection risks [5,6]. Although parasite infection risks are ubiquitous in nature, the costs of parasitism vary widely - ranging from infections with limited fitness impacts to infections that cause mass mortality as they spread throughout populations [19]. In addition to 73 direct negative effects on survival, parasites can have also indirect fitness effects by increasing 74 variability of reproductive success [20]. Application of the landscape of disgust relies on the 75 existence of cues and the ability for potential hosts to recognize and respond to those cues [1,8], 76 which would not be present for many asymptomatic infections. For infections that have strong 77 fitness impacts, and therefore selection for preventing infection, detectable cues may allow for 78 selection of behavioural avoidance. Infections with no detectable cues may result in selection for 79 alternative strategies to prevent fitness costs such as increased resistance (the ability to limit or 80 prevent infection after contact with a parasite) or tolerance (the ability to reduce fitness 81 consequences of infection) [21]. Second, in systems where the landscape of disgust does exist, 82 empirical research has focussed on small spatiotemporal scales. Although small-scale studies 83 remain critical for improving mechanistic understanding of disgust, limited research at larger scales results in a knowledge gap about how parasite avoidance scales up to landscapes. 84 Increasing the spatiotemporal scales of the landscape of disgust is indeed critical for increasing 85 86 the value of the framework; we expect complexities introduced at higher scales may seem likely to create challenges for interpreting the findings of empirical work. We suspect the first two 87 limitations are rooted in the basis of our third argument: the current landscape of disgust 88 89 framework is limited by a lack of integration of relevant ecological theory (e.g., habitat selection theory: [22]) and evolutionary theory (e.g., foundational evolutionary forces driving adaptation: 90 91 [23]). Incorporating these broad bodies of literature will enhance robustness and ease 92 interpretability for those seeking to investigate the landscape of disgust in natural settings and at 93 larger scales.

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95 Our review aims to illustrate how critical insight could be gained by expanding the landscape of disgust interpretation to include habitat selection theory (Section 2: Landscape of disgust within 96 97 a generation; Figure 1) and evolutionary theory (Section 3: Landscape of disgust across generations; Figure 1). To provide context for this review, we provide background knowledge of 98 99 the landscape of disgust (Box 1), and a visual aid to show how aspects of the landscape of 100 disgust could change within and across generations (Box 2). We finally provide examples of 101 outstanding questions and suggestions on how to empirically approach researching these topics 102 to expand our understanding of the landscape of disgust (Section 4: Conclusions and future 103 directions; Box 3).

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2. Landscape of Disgust within a Generation

106 Habitat is a location in environmental space, defined by a set of conditions (e.g., temperature), 107 resources (e.g., food), and risks (e.g., predators or parasites) [24], with habitat selection being the 108 process through which animals differentially use habitats relative to their availability at a given 109 population density to maximise fitness [25]. Habitat selection theory provides a foundation for assessing how the landscape of disgust will be shaped by factors that vary within the lifetime of 110 111 an individual. In this section, we first discuss how the landscape of disgust will be shaped by the 112 dynamic effects of parasite transmission, host density and host sociality (Section 2.1). Next we 113 consider ways to increase the spatial and temporal scales at which the landscape of disgust is 114 assessed within a generation (Section 2.2).

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116 *2.1 Density dependence and sociality*

117 Habitat selection, by definition, is density dependent [26,27]. Following the Ideal Free 118 Distribution and density-dependent habitat selection theory, animals should select habitat to 119 maximise fitness relative to the availability of habitat, resulting in varying population densities 120 among habitats in proportion to the fitness value of each habitat [25,28]. Density-dependent 121 habitat selection theory therefore provides a null expectation for how animals select habitat 122 within the context of the conditions, resources, and risks that make up their environment. 123 However, the conceptual link between density-dependent habitat selection and parasitism is 124 lacking; an absence which is striking given that density is often associated with parasite 125 transmission risks [29,30]. 126 127 An increase in host population density is predicted to result in more social contacts and therefore 128 higher parasite transmission, and this effect has been observed in several meta-analyses (e.g., 129 [31,32]). However, transmission does not always increase with increasing host density owing to 130 the encounter-dilution effect [33], or because of increased (natural) selection for individual-level 131 avoidance behaviours (e.g., [34]). Associations between host density and parasite infection risk can change over time (e.g., [35]). For example, increased density surrounding waterholes in dry 132 133 seasons substantially increases the prevalence of oral-faecal parasites, causing a tight association 134 between seasonality and parasite prevalence [36] (see Figure B1). Notably, it is not solely host 135 conspecific density that can impact risk, but heterospecific host density can also impact risk [37]. 136 To provide important context to the landscape of disgust, both host social behaviour and density-137 dependent habitat selection must be incorporated into estimates of the landscape of disgust 138 (Figure 1).

139

140 The landscape of disgust does not only exist for uninfected individuals avoiding infectious 141 agents. Infected individuals with clinical symptoms are also a part of the landscape of disgust 142 (Figure B1a) and their behaviours affect how uninfected and infected conspecifics interact and 143 the extent to which they can avoid infection risks in the environment. For instance, parasites can impact social behaviour to facilitate further infection (e.g., increased shoaling in infected fish; 144 145 [38]) or alternatively, parasite infection can maintain social behaviour even as host movement 146 and habitat selection change [39]. Conversely, infected individuals may not be accepted into 147 social groups (e.g., guppy, *Poecilia reticulata*, shoals avoid infected conspecifics) [40], reducing 148 the risk for all individuals in the group and altering the structure of the landscape of disgust. 149 Indeed, whether an animal is solitary or social can impact the behavioural defences employed 150 against parasites, including avoidance [10]. In some cases infected conspecifics with infection-151 associated pathologies are not avoided. For example, eastern water dragons (Intellagama 152 *lesueurii*) do not avoid conspecifics infected with a lesion-causing fungus unless the severity of 153 the infection is severe, presumably because the benefits of sociality outweigh the costs of most 154 infections [41].

155

Social behaviour contributes to the spatial structuring of populations [42], which in turn affects the placement of one type of infection risk (i.e., direct transmission from conspecifics) on the landscape of disgust. Although infection risk of directly transmitted parasites (i.e., parasites that require direct contact between hosts for transmission; [43]) is generally higher in large social groups [32,44], social behaviours, such as allogrooming and social learning of parasite cues, can offset costs of living in large groups [45,46]. Further, animals can modify their movement to reduce infection risks for themselves or group-mates. For instance, primates are thought to cycle 163 through sleeping groves and defecate in specific areas to reduce parasite transmission [47,48], 164 bats avoid recolonization of recently used roosts with potential for infection [49], and badgers 165 (Meles meles) moderate sett usage based on infection risk [34]. The landscape of disgust can also 166 be shaped by social hosts mitigating risk by controlling where they generate risks (e.g., latrines 167 or defecation behaviour). Taken together, the interactive effects of host sociality, habitat 168 selection, density, and parasitism are multifaceted and can contribute to a dynamic landscape of 169 disgust within the lifetime of an individual or group of individuals. It is well known that sociality 170 and density operate on various biological scales [42], suggesting that the interactions with the 171 landscape of disgust will vary from small to larger scales.

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173 2.2 Expanding spatiotemporal scales

174 Habitat selection theory proposes that habitat decisions are hierarchical [50,51]; animals select 175 habitats first at larger spatial scales and then make smaller-scale decisions within habitats. 176 Habitat selection is considered at four scales: first order (the geographical area used by a 177 species), second order (the home range of an individual or group), third order (the resource selection decisions made by individuals within their home range), and fourth order (an 178 179 individual's immediate decision making, often related to foraging decisions) [52]. Most 180 landscape of disgust research focuses on third and fourth order scales. Intake maximisation is the 181 most heavily studied driver of habitat selection in the context of landscape of disgust (e.g., [53– 182 55]). Some food sources, such as carcasses and faeces, are high risk for a host to acquire 183 parasites. As a result, they should be avoided behaviourally while foraging at the fourth order of 184 habitat selection [56,57]. Quantifying behavioural avoidance at the fourth order could be done 185 using cafeteria style experiments to measure "give up density" (a metric used to indicate when an animal quits harvesting from a patch) [13]. Within the third and fourth orders of habitat selection
we would expect individuals to typically avoid areas or resources where parasite-associated cues
are perceived (e.g., [58,59], Figure 1). However, in natural systems risk-free habitat may not
exist, leading to trade-offs in habitat selection. For example, individuals typically favour taking
parasite risks over predation risks [60], though not always (e.g., additive avoidance responses;
[61]).

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193 Connecting ecological processes across the orders of habitat selection has become a hallmark of 194 empirical and theoretical habitat selection studies. Because the landscape of disgust should 195 correlate with infection risk for a given parasite, variation in this risk across large spatiotemporal scales might predict avoidance behaviour and generate broad-scale patterns in finer scale 196 197 avoidance behaviour [62]. Broad-scale variation in parasite infection risks could emerge owing 198 to climatic gradients like temperature or precipitation that can predict parasite prevalence at 199 population or species levels [63]. Temperature can also affect host immune function [64], 200 potentially changing the effectiveness and benefits of avoidance behaviour at the third and fourth 201 orders of habitat selection. At the third order of habitat selection, some animals can use 202 behaviours such as migration to seasonally avoid infested habitats and reduce parasite prevalence 203 as they traverse diverse climatic conditions [65]. These effects on avoidance could be most 204 dramatic in unfavourable environments, such as at species ranges limits where host condition 205 may be lower [66], affecting trade-off dynamics between avoiding fitness costs associated with 206 parasite infection and satisfying other needs, such as foraging or mating. Drivers of broad-scale 207 patterns could also depend on community composition. Diversity varies at large spatial scales, 208 with the most diverse communities often occurring closer to the equator [68], potentially

resulting in a reduction in avoidance behaviour where the dilution effect is observed (i.e.,
increased diversity is associated with decreased parasite prevalence; [67]). Considering large
scale environmental and ecological processes should strengthen inferences when predicting the
strength and variation in the landscape of disgust as it spans across these different orders of
habitat selection.

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215 In the context of avoidance behaviour, this relationship between different orders of habitat 216 selection is likely akin to a feedback loop. The presence or efficacy of avoidance at one order 217 may impact the need for avoidance at higher or lower orders. Although we expect animals to 218 place their home ranges to minimise infection risk [69], home ranges may still include areas with 219 higher risk (third order; [70]), in which case finer scale avoidance is an important adaptive 220 behaviour (fourth order; [71]). Another aspect that may affect the relevance of avoidance 221 behaviours is the predictability of risk [72]. In areas with high spatial and temporal predictability 222 in infection risk, third and fourth order avoidance behaviours should be most effective as 223 individuals can reliably change their foraging behaviour and habitat selection to prevent risk 224 exposure. In contrast, when predictability of risk in space or time is low the efficacy of fine scale 225 avoidance is lower and therefore it may be more adaptive for individuals to invest in other anti-226 parasite defences such as resistance or tolerance. Indeed, the predictability of risk could impact 227 the trade-offs for selecting habitat that has associated infection risk. If there is a high fitness cost 228 to infection and no potential for fine scale avoidance at the fourth order of habitat selection then 229 we expect avoidance at third or second order habitat selection decreasing exposure to risky areas 230 [73].

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232 We suggest the consideration of avoidance in habitat selection behaviour at different scales 233 should be a priority for future empirical work. At the first and second orders of selection, 234 biogeographic patterns of parasite avoidance behaviour can provide insight into historical 235 avoidance or the factors driving selection for these traits at smaller scales. The process of scaling 236 up the landscape of disgust from fourth order to the first order of habitat selection relies on the 237 integration of macroecological principles with existing knowledge and theory [62]. Additional 238 inference could be gained by correlating propensity for fine-scale avoidance behaviours with an 239 individual's patterns of habitat selection (third order). Similarly, estimating individual, 240 population, or species-level niche partitioning within the context of behavioural avoidance of parasites could shed light on the integration of avoidance across scales. Researchers could use 241 242 meta-analyses that combine small-scale studies of avoidance behaviour among populations or 243 species at different spatial scales (with carefully selected moderators to untangle sources of 244 variation in behavioural responses). One caveat to this approach is that it requires sufficient data 245 on avoidance, and therefore might not be immediately feasible and awaits future work. Broader-246 scale research is necessary given that small-scale processes, although providing important 247 mechanistic insight, rarely scale linearly, and emergent properties at the higher organisational 248 levels require broad-scale investigation to be identified [62].

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3. Landscape of Disgust Across Generations

Within an animal's lifetime the landscape of disgust can be static or dynamic depending on how
various mechanisms develop or continue to develop via evolutionary processes (Figure 1).
Evolution generating variation in the landscape of disgust could be reflected in many ways,
including the mechanisms that hosts use to recognize and avoid parasites, their ability to resist or

255 tolerate infections by certain parasite species ("ghosts of parasitism past"; [74]), or through non-256 behavioural avoidance (e.g., morphological adaptations [75]) [11]. Variation will affect all three 257 components of the landscape of disgust framework: actual infection risks, perceived infection 258 risks, and the threshold for avoidance behaviour. Not all variation in the landscape of disgust is a 259 product of evolution; plasticity could underlie some phenotypic variation, although plasticity can 260 also be a product of evolutionary forces [76]. Below, we discuss how the landscape of disgust 261 will be shaped by evolutionary change in the hosts (Section 3.1), focusing on evolutionary 262 processes that drive those changes (i.e., natural selection, sexual selection, gene flow and drift). 263 Next, we discuss how the landscape of disgust will be shaped by evolutionary changes in the 264 parasites, often in response to host evolution (Section 3.2).

265

266 *3.1 Host evolution*

267 The most apparent process by which evolution could generate change in the landscape of disgust 268 over time is natural selection. When infection has negative fitness consequences, and where 269 heritable variation exists in the traits that affect fitness, natural selection should drive adaptation 270 to improve avoiding, resisting, or tolerating infection [7,11]. In the landscape of disgust, natural 271 selection could improve cue detection (adding resolution to the landscape of disgust; see Figure 272 B2a-b) and increase avoidance of risks (how an individual reacts to the landscape of disgust it 273 perceives). Importantly, cue detection and avoidance are likely under correlated selection, as 274 these traits go hand in hand [77]. Detection and avoidance also presumably correlate with 275 resistance and tolerance [6,11]. Populations with high resistance or tolerance may not have as 276 strong selection for avoidance behaviours, as they handle infection with a different strategy 277 [7,78,79]; in which case, high resistance or tolerance strategies could translate to a higher

278 threshold for parasite avoidance (Figure B2c) or a lower perceived risk (e.g., decreased detection 279 of parasite cues). For instance, house finches (Carpodacus mexicanus) that have stronger 280 behavioural avoidance responses invest less in immune defences [80]. Raccoon (Procyon lotor) 281 latrines tend to have high prevalence of raccoon roundworm (Baylisascaris procyonis), and 282 tolerant species (e.g., raccoons and rats) use latrines frequently, whereas intolerant species (e.g., 283 birds and small mammals) avoid them [57]. Although these 'strategies' (cue detection vs. altered 284 avoidance threshold vs resistance and tolerance) may be difficult to tease apart empirically, 285 modelling provides an avenue that could attempt to investigate the independent effects of these 286 strategies (Box 3). For instance, theory suggests that the evolutionary dynamics leading to 287 behavioural defences can differ from resistance or tolerance depending on avoidance behaviour 288 type and cost of infection [81]. Researchers could also conduct comparative analyses among 289 populations or species that occupy different environments that might select for different cue 290 detection methods, as detection mechanisms are likely highly associated with the organism's 291 ecology and the environment they inhabit [6].

292

293 Sexual selection could also generate variation in host parasite defences. For instance, more 294 vibrant or ornamented individuals are typically hypothesised to be preferred by the choosier sex 295 because they are more resistant to parasites and hence can afford to produce energetically costly 296 ornamentations [82]. When the choosier sex selects mates that are more resistant, tolerant, or 297 best at avoiding infection, variation in parasite avoidance could arise, leading to variation in the 298 landscape of disgust if subsequent generations inherit these anti-parasite defences. Increased 299 anti-parasite defences owing to sexual selection could present in the landscape of disgust similar 300 to the outcomes of natural selection described above (e.g., high resistance or tolerance could

301 result in decreased avoidance). However, natural selection could also remove individuals from 302 populations that are the most resistant, tolerant, or least effective at avoiding infection if the 303 individuals that cope best with parasite infections (i.e., that are more conspicuous) are also more 304 likely to be predated upon (e.g., [83]). The balance ("trade-off") between these two selective 305 pressures will likely impact the landscape of disgust, owing to selection acting differently on 306 host behaviour. In empirical studies this expectation that healthier and more conspicuous 307 individuals will be preferred by the choosier sex is often not met ([84]). Many hypotheses exist 308 to explain the lack of expected trade-offs, including that the association will be shaped by 309 characteristics of the host or parasite (e.g., [85]), that the association is not being investigated at 310 the appropriate scale of inference (e.g., [84]), or that other sources of selection in the 311 environment could be 'confounding' the expected association (e.g., [86]). In other words, the 312 association can be context dependent and is likely to be variable within and among populations, 313 but could nevertheless help to shape the landscape of disgust over time.

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315 Natural and sexual selection are not the only mechanisms that can generate evolutionary change. 316 Some host populations are more susceptible to genetic drift (e.g., if they are small and isolated; 317 [87]), and the associated randomness could create challenges for predicting parasite avoidance as 318 it relates to the landscape of disgust. In systems where behavioural defences are particularly 319 effective at reducing infection, it has even been suggested that immune based responses could be 320 lost through drift [7]. Additionally, in connected populations where individuals disperse, gene 321 flow could affect host-parasite dynamics [88] and the landscape of disgust over time. As an 322 example, the introduction of individuals from a different population that have not co-evolved 323 with a given parasite could reduce the extent to which the resident population is locally adapted

to those parasites, which could impact selection [89]. Specifically, gene flow could swamp out
evolved defence mechanisms, such as cue recognition or avoidance behaviours, essentially
'resetting' the landscape of disgust. In such cases, decreases in the accuracy of risk perception
(upper layer in Figure B2) or avoidance behaviours may be observed. Alternatively, the
introduction of genetic variation could shift the landscape of disgust by facilitating adaptation
and the potential for more effective anti-parasite responses to evolve.

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331 *3.2 Parasite (co)evolution*

332 Evolutionary processes affect the landscape of disgust over longer time scales as frequencydependence or 'arms race' dynamics play out in host-parasite systems [11]. While the host is 333 334 "winning" the arms race, the cost of infection may be reduced due to shifts in behaviour, 335 resistance, and tolerance; however, similar evolutionary processes also occur for parasites, and 336 selection can drive variation in adaptive parasite traits (e.g., [90]). The strength of selection 337 acting on parasites is highly dependent on host defences. If host populations evolve increased 338 tolerance, parasites may not suffer substantially reduced fitness, and so natural selection acting 339 on the parasites will be weak [91]. In contrast, if host populations evolve increased resistance, 340 parasite fitness will decrease, and there will be strong natural selection acting on parasites. 341 Likewise, if host avoidance strategies are successful at reducing infection, then parasite 342 populations could decrease, reducing the strength of selection acting on hosts but increasing the 343 strength of selection on the parasites to rapidly adapt in response [11]. When selection on 344 parasites is strong, the parasites may evolve less noticeable cues, or shifts may occur in the 345 presentation of disease caused by the parasite, limiting the efficacy of avoidance behaviours for 346 reducing infection risks. For example, for some viruses, such as SARS-CoV-2, infectiousness is highest prior to the onset of symptoms ([92]; impacting the upper layer in Figure B2). In this
sense, parasites can adapt in response to host evolution to successfully infect despite their
presence in the landscape of disgust (the former scenario) or can avoid entering the landscape of
disgust in the first place (the latter scenario; at least for the pre-symptomatic period).

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352 In many cases, humans alter the movement of animal hosts and parasites, increasing interactions 353 between hosts and parasites that have no, or weak, co-evolutionary histories, which can have 354 devastating impacts on host populations [93]. The movement of hosts or parasites could 355 introduce novel parasite species to host populations, or familiar parasite species (i.e., a species 356 that the host has co-evolved with) from genetically distinct populations that the host has not co-357 evolved with [94]. Such scenarios are hypothesised to explain the success of some invasive 358 species (i.e., the novel weapons hypothesis; [95]). For example, when the American grey squirrel 359 (Sciuris carolinensis) was introduced to Europe it also introduced a parapox virus, contributing 360 to the grey squirrels' ability to outcompete Eurasian red squirrels (S. vulgaris), a host that had no 361 previous exposure to the virus [96]. Scenarios of novel host-parasite interactions highlight the 362 importance that evolutionary histories or genetic backgrounds can have in host-parasite 363 dynamics. The importance of shared evolutionary histories in shaping the landscape of disgust 364 (affecting actual risks, perceived risks, and host responses to those risks) is a promising avenue 365 for future work, both in natural contexts and with increasing anthropogenic impacts to host-366 parasite interactions.

367

368 4. Conclusions and future directions

369 In this review, we emphasise the value gained by incorporating habitat selection and 370 evolutionary theories into the landscape of disgust framework. There are many practical ways to 371 integrate concepts from ecological and evolutionary theory into the landscape of disgust that 372 could be leveraged in future work (Box 3). We recognize that determining the level of 373 information required to map actual and perceived infection risks in natural systems may be 374 difficult. The work required to quantify and map or predict the infection risks a host may 375 encounter, or the evolutionary processes acting on hosts and parasites, poses several logistical 376 barriers including: difficulty detecting parasites, the time required to collect data, and potential 377 cost or technological barriers. One solution is to use simulation tools such as Agent-Based 378 Modelling [97] to investigate how the landscape of disgust changes and how it can impact other 379 aspects of natural systems to inform future empirical work (Box 3).

380

381 Our review highlights that the landscape of disgust remains in its infancy; without an 382 understanding of how the landscape of disgust changes within and across generations, we cannot 383 fully comprehend how parasite infection risks impact host ecology. Many future avenues of work 384 remain that would complement the ideas presented in this manuscript. For instance, individual 385 variation owing to acquired immunity or plastic behavioural responses may impact how a 386 potential host interacts with the landscape of disgust [98] and should be investigated alongside 387 repeatable host behavioural defences (e.g., 'hygienic personalities': [99]). Exploring whether 388 some of the landscape of disgust concepts may be applied to other parasitism models that do not 389 have classic host-parasite dynamics would also be valuable (e.g., individuals use visual cues to 390 detect brood parasitism; [100]). Little work has explored how parasites may interact with, or 391 compensate for, the landscape of disgust. Finally, because ecology and evolution can have

reciprocal effects [23], we also suggest investigations that explore how these two processes could
interact to influence the landscape of disgust, particularly in natural systems. Clearly, many
outstanding questions regarding the landscape of disgust framework remain. We focus our
discussion above on habitat selection and evolutionary theories given that they are the focus of
the current paper, although a longer-term goal for the landscape of disgust should include
integration with other frameworks and theories to create a more holistic - and therefore even
more powerful - framework.

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734 Figure Captions and Boxes



735

736 Figure 1: Factors affecting the formation and demonstration of the landscape of disgust. 737 Here we illustrate a hypothetical system to demonstrate different factors that affect the landscape 738 of disgust both within (tan boxes on left) and across (yellow boxes on right) generations. These 739 factors include principles from habitat selection theory operating at varying spatial scales that 740 could be impacted by detection of risk and ecological context, along with evolutionary principles 741 such as selection for traits associated with the detection and avoidance of parasites, or geneflow 742 or genetic drift impacting genetic variation for selection. This is not an exhaustive representation 743 of factors impacting the landscape of disgust but demonstrates several factors that should be

considered when interpreting avoidance behaviour and making predictions in natural systems.

Figure was illustrated by Juan Aristizabal.

746

747 Box 1: Describing the landscape of disgust

748 The landscape of disgust represents behavioural responses to parasite infection risks [4].

Researchers have emphasised the role of host anti-parasite behaviours for decades [1,101], and

the landscape of disgust provides a framework to renew previous lines of research, inspire new

approaches to classic questions, and concentrate research efforts on the investigation of non-

752 consumptive effects of parasitism.

753

Host-parasite dynamics influencing the landscape of disgust can vary between species,

populations, and individuals within populations. For a landscape of disgust to exist and impact

host behaviour, there are at least three prerequisite environmental, physiological, and

revolutionary factors required. First, a cue - which can be visual, auditory, olfactory, or

mechanosensory - must be produced either directly by a parasite, the infected host, or something

associated with risk of infection [6]. For example, the avoidance of faeces which may or may not

be infested with parasites. Second, hosts must have the capacity to detect cues; the physiological

761 mechanisms to interpret and recognize the cue are required for the cue to be perceived by the

host within the landscape of disgust (e.g., detection of chemical cues in mice; [102]). Finally,

parasitism must be costly enough to impact potential host behaviour, and this change must

764 prevent or reduce infection resulting in a fitness benefit to the host [5,6]. Both cue detection and

response behaviour could be learned [45,79,103] or adaptive if sufficient time and genetic

variation are present to evolve a connection between the cue and response [104]. Selection can

act on phenotypes involved in different stages of infection, including recognizing parasite cues,

and responding to infection [105]. In natural systems these prerequisites for the landscape of

769 disgust can take on many forms; any breakdown in the connection between the cue production,

perception, and response will prevent a quantifiable landscape of disgust from forming.

771

772 Box 2: Visualising landscape of disgust within and across generations

773 We visualise the landscape of disgust as the summation of any 'perceivable' parasite infection 774 risks within an individual's spatial perception, relative to the 'active' infection risks (Figures B1, 775 B2). We primarily discuss and interpret the landscape of disgust as a two-dimensional plane, as 776 this reflects the movement of most terrestrial individuals, although in some systems a third axis 777 of movement is available to avoid parasite infection, such as in aquatic systems [106], or for 778 aerial or arboreal animals that can use vertical movement to avoid transmission (e.g., [107]). 779 Because not all parasites can be detected, and not all perceived risks warrant a response [5,6], we 780 visualise a biologically relevant threshold for infection risk that determines whether an avoidance 781 response occurs at all (represented by the horizontal line overlapping perceived risks in Figures 782 B1, B2), which parallels the "cost-benefit" structure used in other parasite avoidance and disgust 783 reviews [5,6,11]. An avoidance threshold can be affected by individual state or past experiences, 784 such as an individual's body condition or hunger levels [108] and prior or current infection status 785 [56,109]. Although we illustrate individual avoidance as binary in the context of this threshold, 786 the strength of an avoidance response can vary such that individuals may moderate their foraging 787 behaviour to decrease exposure (weaker response) or leave the area to seek lower-risk habitat 788 (stronger response). We predict the avoidance response to be correlated with the size of the 789 perceived risk (represented as height of peak in Figures B1, B2). The risk of parasite infection 790 also often exists as a gradient, and variation in parasite risk can occasionally be detected (e.g.,

[41,110]); this variation in risk is represented with simple peaks here but in natural systems there
will be variation in the gradient around the cue as well [111]. In visualising the landscape of
disgust across two timescales - within a generation (Figure B1) and across generations (Figure
B2) - we provide a visual aid to help the reader understand how ecological and evolutionary
theory fit into, and would expand, the existing framework.

796



798 Figure B1. Visual depiction of a hypothetical change to the landscape of disgust that can occur 799 across seasons, within a generation of a host. A critical resource (water) becomes increasingly 800 scarce from wet to dry seasons, increasing density of conspecifics (animals per unit area) around 801 water sources that could harbour infection through time. The geographic landscape and 802 conspecific locations (lower layer), the actual risk of infection (middle layer), and the perceived 803 risk of infection (upper layer) are denoted as layers in each panel. From a) to b) as the resource 804 becomes less available, individuals will choose lower density areas that present less risk but still 805 provide access to the resource (where perceived risk is below threshold in upper layer). Between 806 b) and c) the threshold of avoidance (upper layer) becomes higher as the resource becomes

807 increasingly scarce. In this example, individuals will choose to increase their risks of infection







810

811 Figure B2. Visual depiction of a hypothetical change to the landscape of disgust over 812 generations. A novel environmentally transmitted parasite is introduced to a landscape via a 813 contaminated waterbody that represents the environmental reservoir; the introduction of this 814 parasite drives adaptive changes in the host population over generations. The geographic 815 landscape and conspecific locations (lower layer), the actual risk of infection (middle layer), and 816 the perceived risk of infection (upper layer) are denoted as layers in each panel. Between a) and 817 b) the host population evolves to associate the parasite cue with the environmental reservoir 818 (there is novel perceived infection risk) (upper layer). Between b) and c) the host population has 819 evolved to become more tolerant to infection by that parasite (reduction in magnitude of actual 820 risk in middle layer), reducing the perceived cost of infection; notably, there is a shift in the 821 magnitude of perceived infection risk such that the perceived risk is now lower than the 822 actionable avoidance threshold.

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Box 3: Tools for measuring and interpreting the landscape of disgust in nature.

825	1)	Mapping parasites on the landscape: To predict both the benefits of avoidance and where
826		avoidance should occur, quantifying the parasites that actually exist in a given
827		environment provides important context. It is possible to map parasite distributions when
828		they are inside hosts or outside hosts. Species distribution models (SDM) [112] and
829		resource selection functions (RSF) [113] are useful tools for estimating organismal
830		distribution through space and time. One way to estimate parasites without intermediate
831		or external stages (e.g., viruses) is to develop SDMs or RSFs for infected hosts through
832		space and time (e.g., [114]). For hosts with intermediate or external stages (e.g.,
833		nematodes or ectoparasites), we propose a multi-step approach, similar to the joint-SDM
834		approach [115], where SDMs or RSFs are developed for definitive hosts, intermediate
835		hosts, and/or for the parasites themselves while they are outside of the host.
836	2)	Agent Based Modelling of the Landscape of Disgust: Agent based models are spatially
837		explicit individual level models. The emphasis on the individual level (the agent)
838		provides an excellent opportunity to explore decision making and trade-offs when
839		perceiving risks. These models are used to investigate movement and processes from
840		individual level physiological processes (e.g., energetics; [116]) up to ecosystem level
841		processes (e.g., landscape heterogeneity; [117]). Researchers could model a landscape of
842		disgust with agents that are a source of risk (i.e., social transmission), or that create risks
843		(e.g., faecal deposition), tracking how individuals who perceive these risks avoid them
844		and how this could impact the system at varying levels (e.g., disease dynamics or nutrient
845		cycling).

3) Tracking the landscape of disgust across generations: To estimate the strength and 846 direction (e.g., stabilising, disruptive) of selection in the context of the landscape of 847 disgust, a researcher could measure any given trait (e.g., avoidance behaviour, resistance 848 849 or tolerance) and a fitness proxy (e.g., number of offspring that survive to reproductive 850 age, number of offspring produced) [23]. If the traits of interest are heritable (which can 851 be estimated in different ways, such as the traditional method of correlating offspring to 852 parental phenotypes; [118]), researchers can gain insight into whether selection on a trait (or correlated traits) will lead to evolutionary change in the landscape of disgust 853 854 [119,120].