Cryptic species and hybridisation in corals: challenges and opportunities for conservation and restoration

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Abstract

shallow water coral reefs only ?

36 Conservation and management of coral reef ecosystems will depend on accurate assessments 37 of reef-building coral species diversity. However, the true diversity of corals may be obfuscated by the presence of cryptic species, which are likely much more pervasive than is 38 39 currently recognised. Additionally, cryptic species may sometimes hybridize, resulting in gene 40 introgression between species. Here, we investigate the prevalence of cryptic coral species the only 41 via a structured literature review and find that over 50% of population genomic studies show 42 evidence for divisions within taxonomically recognised species and that such closely-related 43 taxa are often linked by gene flow. We find that cryptic taxa frequently segregate by environment, especially depth, and may differ by phenotypic characteristics including 44 resilience to heat stress. This hidden biodiversity creates challenges for coral conservation 45 and restoration planning that are not well appreciated, including hiding true population 46 declines, biasing estimates for species' phenotypic breadth, overestimating the resilience of 47 48 species to stressors, yielding uncertainty in evolutionary dynamics inferred from past studies, and creating reproductive barriers that may limit mating between local and translocated 49 50 corals. Increasing awareness that coral cryptic species with incomplete species boundaries 51 are common and building this expectation into conservation and restoration plans is an 52 important pathway forward. Rich opportunities for interdisciplinary collaboration among 53 coral and speciation biologists could fill key knowledge gaps relevant to conservation. We 54 detail recommendations for best practice and strategies for identifying cryptic taxa and 55 hybrids and urge their consideration in all future studies on corals.

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1 Introduction: Hidden dimensions of coral biodiversity pose conservation 57 58 challenges

Coral reefs are highly biodiverse and productive ecosystems (Reaka-Kudla, 1997; Fisher et al., 59 2015) that substantially contribute to human well-being (Moberg & Folke, 1999; Adey, 2000). 60 61 Yet, reef-building corals are imperilled by rising temperatures and other anthropogenic 62 stressors worldwide (Hughes et al., 2017; Knowlton et al., 2021; Souter et al., 2021). Thus, 63 there is great urgency to inventory coral biodiversity and to deepen knowledge regarding the 64 processes that create and maintain this diversity to guide conservation and restoration 65 actions.

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67 For corals it has long been recognised that morphological variation is unlikely to align well 68 with genetic variation. Molecular based phylogenetic investigations are uncovering 69 unexpected relationships and unanticipated numbers of distinct taxa (reviewed by Kitahara 70 et al., 2016; Cowman et al., 2020), and similarly, multilocus population genetic surveys routinely find distinct genetic taxa within morphologically defined taxonomic species. These 71 72 so-called cryptic species (recent reported genomic examples include: Gomez-Corrales &

remove "gene" or replace by genetic, as genes are not elements that introgress

73 Prada, 2020; Underwood et al., 2020; Wepfer et al., 2020; Afiq-Rosli et al., 2021; Bongaerts 74 et al., 2021a; Feldman et al., 2021; Fifer et al., 2021; Rippe et al., 2021; Zayasu et al., 2021; 75 Adam et al., 2022; Prata et al., 2022; Rivera et al., 2022; Matias et al., 2023; Voolstra et al., 76 2023; Meziere et al., 2024) reinforce observations previously made with microsatellites and 77 few nuclear markers (such as Bongaerts et al., 2010; Souter, 2010; Ladner & Palumbi, 2012; 78 Schmidt-Roach et al., 2012; Prada & Hellberg, 2013; Boulay et al., 2014; Prada et al., 2014; 79 Warner, van Oppen & Willis, 2015; Gélin et al., 2017) and demonstrate how species 80 designations based on morphology alone can underestimate true community diversity (Fig 1). Indeed, most conservation management and restoration plans implicitly assume that coral 81 82 species are recognisable and biologically valid entities (Baums, 2008; Anthony et al., 2017; National Academies of Sciences, 2018; Colton et al., 2022). These assumptions may 83 unintentionally bias conservation and restoration efforts when such biodiversity is not 84 acknowledged. Yet, detecting and delineating distinct genetic groups presents a substantial 85 86 challenge (which we discuss in more detail in Section 2.1). Rather than adopting a specific 87 criterion for species status, we use the terms *genetic groups* or *taxa* to signify their genetic 88 distinctiveness. We avoid terms such as clade or lineage, as monophyly across all or many 89 gene trees is unlikely when divergence is recent. We use *cryptic* to signify that taxa appear 90 morphologically similar to the "untrained human eye", at least under field conditions.



to be even more convincing, suggest to add photographs taxonomic characters such a structure, to show the read ti just a matter of amateurs loc morphology

Figure 1 – Examples of closely related cryptic species. Details can be found in the original studies: *Stylophora pistillata* (Meziere *et al.*, 2024), *Porites sp.* (Starko *et al.*, 2023), and *Agaricia lamarcki* (Prata *et al.*, 2022).

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96 Further complicating taxonomic delineations, cryptic coral taxa may be linked by occasional 97 gene flow, or hybridisation. Morphologically intermediate individuals are often encountered 98 in the field (Veron, 1995; Richards et al., 2008), and many species can be crossed at least 99 under experimental conditions (Isomura et al., 2016; Chan, Peplow & van Oppen, 2019; 100 Kitanobo et al., 2022). Genomic studies confirm that hybridisation is common in corals, where 101 gene flow has been documented between distinct taxa (Cooke et al., 2020; Fifer et al., 2021; 102 Prada & Hellberg, 2021; Rippe et al., 2021; Prata et al., 2022; Matias et al., 2023; Starko et al., 103 2023; Zhang et al., 2023; Meziere et al., 2024), including historical gene flow between taxa 104 separated by over 15 million years (reviewed by van Oppen & Gates, 2006; Willis et al., 2006; 105 Mao & Satoh, 2019; González, Rivera-Vicéns & Schizas, 2021; Hobbs et al., 2021; Pinsky, Clark 106 & Bos, 2023). Cryptic genetic groups or taxa, potentially connected by limited gene flow, 107 present distinct challenges for understanding coral biodiversity. Failing to recognize cryptic 108 coral taxa and appropriately adjust interpretations can result in misleading conclusions about 109 fundamentally important aspects of biodiversity measurement, including: i) underestimating 110 species diversity, ii) overestimating within-species variances for various traits and tolerances,

111 iii) biasing inferences regarding within-species patterns of genetic diversity and population

- structure, and iv) invalidating assumptions about reproductive compatibility.
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First, biodiversity inventories typically determine the count and abundances of distinct 114 115 species. Common methods, such as field surveys, are usually based on morphological 116 identification of live organisms. If distinct taxa are not appropriately recognised and 117 delineated, then total species counts will be greatly underestimated, presenting an unstable 118 base from which to draw conclusions about coral population sizes, species extinction 119 vulnerabilities, and biodiversity valuations. For example, estimates of census population sizes 120 for present-day corals have been recently debated (Dietzel et al., 2021; Muir et al., 2022), as 121 smaller population sizes would imply greater vulnerability to extinction (Dietzel et al., 2021; 122 Muir *et al.*, 2022).

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124 Second, assessment of trait diversity is compromised if cryptic taxa are not identified, and 125 trait measurement occurs on an amalgamation of distinct taxa. The agglomerate will contain 126 more variation than the distinct groups, which will inflate estimates of both genetic and 127 phenotypic variance. This phenomenon can affect interpretation for a diverse array of traits. 128 For example, species range estimates would be upwardly biased if cryptic species are 129 geographically restricted within the broader range of a morphospecies. Environmental niche 130 breadth can also be overestimated: for instance, many coral species that were previously 131 considered depth generalists could resolve into taxa with more restricted depth distributions 132 when cryptic species are considered (e.g., Bongaerts et al., 2021a). Similarly, a presumed 133 species may appear to have a generalist phenotype but actually comprise multiple taxa that 134 are more specialized, as appears to be the case for bleaching responses among Orbicella 135 faveolata taxa (Gomez-Corrales & Prada, 2020). In summary, single-species conclusions 136 drawn from agglomerated data from multiple cryptic taxa are likely to be overly optimistic in 137 terms of coral resilience.

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139 Third, common measures of population genetics and gene flow may be inherently biased and 140 incorrect when cryptic taxa are examined as one. For example, using allele frequencies to 141 make inferences about microevolutionary dynamics without delineating separate gene pools 142 can yield incorrect results and interpretations. Lumping genetically distinct groups will inflate apparent within-population diversity (based on measures of allelic diversity or expected 143 144 heterozygosity) and will therefore cause inbreeding to be overestimated – known as the 145 Wahlund effect (as discussed in Schmidt, Thia & Hoffmann, 2023). When comparing across geographic locations, measures of population structure (notably F_{sT}) can be biased either 146 147 upwards or downwards, depending on the particular mix of cryptic taxa sampled. These 148 phenomena are neatly illustrated and discussed by Sheets et al. (2018) for Acropora 149 hyacinthus in the western Pacific (see also Pante et al., 2014; Warner et al., 2015). 150 Additionally, common metrics of population genetics such as genetic diversity, 151 differentiation, inbreeding and effective population size may be biased when hybridization

between differentiated taxa is not taken into account (Hoban *et al.*, 2022). For example,
hybridisation may inflate the measured diversity for populations that include individuals of
mixed ancestry, while gene flow or dispersal inferences may be biased in either direction.
Together, cryptic taxa and hybridisation are likely to greatly affect the accuracy of studies
aiming to assess and monitor genetic diversity.

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158 Finally, the existence of cryptic taxa implies that there is some form of reproductive isolation 159 between taxa. Intrinsic barriers such as incompatible loci are likely to reduce gene flow among taxa in nature, so populations of a morphospecies may not be as strongly linked by gene flow 160 161 as is often assumed. In restoration contexts, intrinsic barriers could lead to translocated 162 individuals being unable to mate with local individuals and could prevent crossing in captivity 163 and/or reduce hybrid fitness (outbreeding depression). Extrinsic barriers to reproduction 164 arise when survival or fertility differs between taxa based on their surrounding environments, 165 and as a result, they inhabit different niches. This means that cryptic taxa may not be 166 ecologically or functionally equivalent. Restoration actions that involve coral outplanting (i.e., 167 fragments or sexually propagated colonies) risk mis-matching the source taxon's niche with their new destination's environmental conditions. Conversely, hybridisation between taxa 168 169 could enhance fitness or phenotypic breadth of resultant offspring (Chan et al., 2018). 170

Figure 2 outlines some possible consequences of cryptic coral taxa to current restoration actions such as direct transplantation and coral gardening, as well as actions that are being actively researched for potential use in coral restoration (National Academies of Sciences, 2018; Anthony *et al.*, 2020; Hein *et al.*, 2020; Bay *et al.*, 2023).

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Interventions

Direct transplantation & coral gardening:

Transplantation of colonies or fragments from one reef to another. Coral gardening involves rearing corals in a nursery setting (land or sea-based) before transplantation to the ree

Gamete and larval capture and seeding:

Collecting gametes and larvae from the wild and releasing larvae onto other reef areas

Intraspecific breeding within reefs:

Sexual reproduction of targeted colonies and releasing progeny to increase population size (supportive breeding) and/or increase frequency of heat tolerant alleles (selective preeding).

Assisted gene flow:

Human assisted movement, with or without crossing corals in captivity, to promote gene exchange beyond the natural dispersal scale but within the current species range

Assisted migration:

Human assisted movement, with or without crossing corals in captivity, over large spatial scales within or beyond the current species range.

Hybridization among species:

Crossing species to create novel genotypes and phenotypes.

Conditionina:

Stress exposure to make corals more tolerant of future stresses

Algal-symbiont manipulation:

Replacing algal symbionts with more heat tolerant strains that may be sourced from a reef or experimentally evolved

Probiotics:

Supplementing an established community of microbes with beneficial strains

Gene-editting:

Microbiome manipulation

Altering select genes to yield more heat tolerant phenotypes

Equivalent terms for interventions: "Managed selection for environmental tolerance

Plntraspecific breeding between reefs; outcrossing within species Experimental evolution

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Figure 2 - How cryptic coral taxa and hybridisation could affect coral reef restoration actions that aim to preserve biological diversity, counter population declines, and promote resilience to climate change biological adaptation. Terminologies follow (van Oppen *et* through al., 2015: National Academies of Sciences, 2018; Hein et al., 2020; Bay et al., 2023) and are not necessarily mutually exclusive. For example, assisted migration could be undertaken on fragments (direct transplantation), larvae, or via the progeny of captive, sexually propagated corals. In all graphics, the coral colony shown as a larger size indicates greater fitness (survival, reproductive output, thermal tolerance, etc) leading to competitive dominance.

185 As outlined above, cryptic taxa and hybridisation have substantial implications for 186 understanding and predicting the ecological and evolutionary dynamics of corals, and thus 187 will be essential for making informed conservation and restoration decisions into the future. In this review, we critically assess the prevalence of hidden biodiversity among corals. By 188 corals, we refer to benthic Anthozoans including Scleractinians (hard corals) as well as 189 Octocorallians (soft corals). We focus primarily on population *genomic* studies (supported by 190 191 whole genome or reduced representation sequencing of genome-wide variation) because 192 thousands of loci are often required to detect recent evolutionary delineations between taxa. 193 This paper builds upon important earlier reviews and syntheses of cryptic taxa (Richards, 194 Berry & van Oppen, 2016) and hybridisation in corals (van Oppen & Gates, 2006; Willis et al.,

Potential consequences arising from cryptic species or hybridisation

Taxon-environment mismatch

Cryptic taxa differ in niche and other phenotypic attributes such that transplanted fragments, colonies or host genotypes that do not match the local environment perform poorly. This may include different responses to stress

Reproductive isolation



Mating success between cryptic taxa is reduced relative to me-taxon matings. This can prevent transplants from mating with cal colonies and can diminish crossing success for interventions same that rely on sexual reproduction

Competitive dominance of transplanted taxon



If the transplanted taxon has greater growth, survival, reproductive output, etc. then the transplanted taxon may outcompete the local taxon. potenitally leading to the loss of locally adapted genetic diversity. However, increased coral cover may be viewed as a net benefit.

Competitive dominance of hybrids



Hybrids are expected to show more extreme phenotypes than their parental taxa: if hybrid colonies have greater growth, survival, reproductive output, etc. then outplanted hybrids may outcompete the local taxon, potenitally leading to the loss of locally adapted genetic diversity. However, increased coral cover may be viewed as a net benefit.

Host-microbe mismatching



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Beneficial microbes, including dinoflagellate symbionts, may confer benefits to some cryptic host taxa but not others.

Gene-manipulation mismatching

Editted genes may result in alleles that benefit some cryptic host taxa but not othe



2006; Hobbs *et al.*, 2021; Pinsky *et al.*, 2023) that primarily focused on studies with few
markers (microsatellites, few sequenced loci, or allozymes).

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198 Here we explore how evolutionary processes in corals can and should shape conservation and 199 restoration priorities and actions. Specifically, we: 1) develop criteria for recognising cryptic 200 taxa from population genetic data, 2) describe and evaluate surveyed genomic literature 201 against these guidelines to show that cryptic taxa frequently exist in sympatry, 3) determine 202 whether cryptic divergence in cnidarian hosts is accompanied by divergence in their 203 photosynthetic symbionts, 4) explore the environmental factors segregating cryptic taxa, 5) 204 review examples of hybridisation and/or gene flow between taxa, and 6) discuss how corals 205 could be exciting model systems for speciation and adaptation studies. The main text is 206 supported by text boxes that, A) provide a worked example of delineating coral taxa, B) 207 demonstrate how cryptic taxa are commonly overlooked in coral experiments, and C) outline 208 best practices for designing studies when cryptic taxa are likely to be encountered.

209 2 Closely related coral taxa are common in sympatry and frequently connected by 210 gene flow

To gauge the prevalence and impacts of cryptic coral taxa and hybridisation, we evaluate findings from a structured literature search alongside a broader literature review. Although our primary intent is to draw inferences from more informative genome-based studies, population genomic studies for corals are limited both in their number and in scope. Thus, we also include some discussion of noteworthy results based on microsatellite surveys and also review microbial symbiont diversity because of their relation to cryptic host taxa (under Sections 2.2 and 2.3).

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219 **2.1** Detecting cryptic coral species using criteria based on reproductive isolation

220 Many coral taxa potentially inhabit the "grey zone" of the speciation continuum (de Queiroz, 221 2005; Roux et al., 2016). That is, genetically distinct groups may coexist within dispersal range, 222 yet these groups may not have all the contingent properties of distinct 'species' (sensu de 223 Queiroz, 2005), including complete reproductive isolation. However, for groups with 224 overlapping ranges to remain genetically distinct, reproductive barriers of some form must 225 be present. Reproductive barriers between groups may occur due to differences in 226 microhabitats or spawning times resulting in infrequent fertilisation, producing offspring with 227 phenotypes that do not match microhabitats, or other intrinsic genetic incompatibilities. 228 When reproductive barriers are not complete, the outcome of interbreeding between groups 229 is controlled by the relative strength of selection (which promotes divergence), and the scale 230 of gene exchange (which promotes homogenisation). Even low levels of gene exchange 231 between otherwise distinct groups will cause alleles to be shared. Allele sharing will also be 232 prevalent if divergence time between groups is recent whereby groups share ancestral

this paragraph would benefit from some key references

variation. Thus, allele sharing is expected to be common between cryptic coral taxa that arerecently diverged (with or without ongoing gene flow).

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236 Population genomic surveys have the power to detect subtly differentiated genetic groups 237 where allele frequency differences between groups are small. The statistical power of 238 common methods is determined by the number of loci examined as well as the extent of 239 genetic covariance among loci, where distinct populations exhibit non-random associations 240 of alleles at various loci across the genome (i.e., *linkage disequilibria*). While physical linkage 241 on a chromosome alone will cause covariance among loci, genome-wide covariance also 242 arises as a direct consequence of population structure (reflecting distinct gene pools subject 243 to independent outcomes of genetic drift and selection). For example, reproductively isolated 244 taxa or geographically separated populations will be differentiated across loci due to genetic 245 drift and selection. In contrast, gene flow among them will erode both their allele frequency 246 differences and linkage disequilibria. Statistical power for identifying small allele frequency 247 differences and linkage disequilibria in empirical surveys is increased by sampling many 248 individuals and many loci, where subtle differences between groups can be missed when few 249 individuals or few loci are sampled. For example, two co-occurring genetic groups of 250 Montastraea cavernosa were clearly delineated using thousands of loci and yet ambiguous 251 with 9 microsatellite loci (Sturm *et al.*, 2020).

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253 We focus on two general approaches for detecting subtle differentiation between genetic 254 groups: ordination-based analyses and model-based clustering. Ordination-based analyses, such as principal components analyses (PCA), principal coordinates analysis, 255 256 multidimensional scaling, etc., describe multidimensional relationships among entities and 257 are based on (and visually represent) the genetic covariance matrix (Patterson, Price & Reich, 258 2006). Model-based clustering analyses – typified by admixture detection analyses such as 259 STRUCTURE (Pritchard, Stephens & Donnelly, 2000) and ADMIXTURE (Alexander, Novembre 260 & Lange, 2009) – partition groups (K) based on associations among alleles and loci. Ordination 261 and model-based clustering approaches are unsupervised machine learning methods that are 262 valuable for exploring relationships between individuals (that is, without pre-assigning 263 individuals to "populations", as is required by F-statistics and other population-level metrics). 264 These methods are some of the most common and routinely employed methods in genomic 265 surveys and provide complementary insights into spatial patterns of genetic diversity. We 266 omit results from supervised methods, such as discriminant analysis, that maximize variance 267 using user-assigned groupings (see Thia, 2022 for extended discussion).

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We define taxa as distinct groups of individuals (genotypic clusters) within named species that
 maintain their distinctiveness even when their ranges overlap, therefore they have the

271 opportunity to mate (in line with Mallet, 1995). This definition aligns to many species 272 concepts (de Queiroz, 2005), under the assumption that some form of reproductive isolation 273 must be occurring. Instances where distinct genetic groups are found together (i.e., sympatric 274 within the scale of dispersal distance) provide the strongest circumstantial evidence for some 275 degree of reproductive isolation between groups, as reduced gene flow due to dispersal 276 restrictions cannot be the primary cause of genetic divergence in these cases. These "taxa" 277 might be considered species, depending on the definition of species employed (de Queiroz, 278 2005); as our focus is not taxonomy but evolutionary processes and their implications, the 279 term "taxon" is more flexible. Thus, we focus on evidence for sympatric cryptic coral taxa. 280 When distinct genetic groups are geographically separated (*i.e.*, allopatric), then divergence 281 may solely reflect dispersal barriers to gene exchange and therefore are not informative for 282 inferring reproductive isolation.

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We propose three requirements for identifying and delineating taxa using ordination and model-based clustering:

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1) Distinct genetic groups occur in sympatry relative to their dispersal ability.

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 2) Ordination (e.g., PCA) clusters these distinct genetic groups based on genotypes of individuals and/or model-based clustering indicates that individuals belong to separate groups.
- 3) When genetic distance between sympatric individuals is greater than the
 genetic distance between allopatric individuals. This is evidenced by
 divergence between groups across the primary ordination axis and/or at low
 hypothesized group numbers (K values) as compared to axes or groups that
 describe geographic structure.
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296 To determine if published genomic surveys of corals typically test for and find evidence for 297 cryptic taxa following the above criteria (see Textbox 1 for worked example), we searched the 298 Web of Science for published papers displaying graphical results from ordination (PCA, 299 primarily) and/or model-based clustering analyses (Structure and ADMIXTURE). We focus on 300 studies that genotyped individual cnidarian genomes for 1000s of loci or more and sampled 301 from two or more geographic locations (see Appendix for details of literature search). We 302 additionally collected information on depth ranges and symbiont composition where 303 reported. The literature search uncovered 41 papers describing results for 31 species, where 304 some papers included multiple species and some species were studied across multiple papers, 305 yielding a total of 51 unique records (available as supplemental data). Although we did not 306 restrict the search by sampling depth, none of the recovered records included species deeper 307 than mesophotic depths.

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In our review of published studies, we apply the above criteria (1-3) conservatively by only looking for sympatric differentiation i) along the first principal component axis (referred to as 'PC1') in ordination analyses, and ii) for model-based clustering examining outcomes when individuals were assigned to only two groups (K=2). We consider individuals sampled ≤ 10 km apart as being broadly sympatric since dispersal distances for many coral species are unknown.

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316 As shown in Fig. 3, over half of the coral population genomic studies (23 records out of 39 317 that could be evaluated against all three criteria) showed evidence for distinct cryptic taxa. While we might expect genetic differentiation to be greater among brooding corals that have 318 319 less innate dispersal ability relative to broadcast spawning corals (Knowlton, 2001), the 320 relative proportions of sympatric versus non-sympatric groups did not differ between 321 brooding and broadcast spawning corals (P> 0.2 with Fisher's exact test using either 322 ordination or clustering). Criterion 3 is based on genetically distinct groups co-occurring. This 323 evidence is stronger when this groups co-occur across many geographic locations. Thus, we 324 investigated whether pairs of genetically distinct coral groups were repeated across multiple 325 sites using model-based clustering results based on author-selected K values. Across studies, 326 it was common for cryptic groups to be sympatric at multiple sampled sites (Fig. 3). This 327 observation strengthens the conclusion that closely-related, but distinct taxa, can co-occur 328 over extensive geographic areas and implicates some degree of reproductive isolation 329 maintaining the distinctiveness of each group. In summary, cryptic taxa are common features 330 in population genetic studies of corals.

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332 Our literature review shows that cryptic taxa are prevalent, implying that many conclusions 333 related to biodiversity, species traits, and population genetic structure based at the 334 morphospecies level are likely to be inaccurate based on the reasons discussed previously 335 (see Section 1). Even when genomic data to evaluate cryptic taxa are presented, five studies 336 that we assessed failed to include any mention of cryptic species or taxa, despite their results 337 showing evidence for genetically distinct groups. Additionally, some of the studies reviewed 338 here computed summary statistics such as heterozygosity, or F-statistics using all individuals 339 from the sampling locations, despite evidence for cryptic taxa co-occurring within locations. 340 We surmise that inferences regarding the absence of potential cryptic taxa based on analyses using few loci (including microsatellite studies) are likely lacking in terms of the power to 341 342 detect recently differentiated taxa, and therefore many published studies – including studies 343 published by authors of this review – may inadvertently base conclusions on heterogeneous 344 mixes of cryptic taxa.

Class	Family	Species	Ordination	Model-based clustering	Number sympatric locations ^a
Hexacorallia	Acroporidae	Acropora aspera	•	•	
		Acropora digitifera			
1.15		Acropora downingi			
		Acropora hyacinthus			2
		Acropora palmata			
		Acropora tenuis			1
		Isopora brueggemanni.			2
C C C	Agariciidae	Agaricia fragilis			3
	0	Agaricia grahamae			6
		Agaricia lamarcki		•	4
	Pachyseridae	Pachvseris speciosa			2
And the second	Poritidae	Porites lobata		•	
		Porites sp.			
	Euphylliidae	Galaxea fascicularis			-
	Astrangiidae	Astrangia poculata			
	Merulinidae	Orbicella faveolata			
		Platvovra daedalea			
and the second	Montastraeidae	Montastraea cavernosa			
	Pocilloporidae	Pocillopora acuta			
		Pocillopora damicornis			2
		Pocillopora meandrina.			4
		Pocillopora verrucosa			
		Stylophora pistillata			
Octocorallia	Coralliidae	Corallium iaponicum			-
	Nephtheidae	Dendronephthya austra	lis		
	 Studies that me Studies that do 	et criteria 1-3 for cryptic ta not meet criteria 1-3 for cr	axa ryptic taxa	^a Dash indicate	s that evaluation is not possible

Figure 3 - For many genomic studies of corals, the greatest axis of genetic differentiation defines groups that are sympatric. Results by species are summarized as either meeting or not meeting the three criteria for cryptic taxa under the strictest definition where sympatric differentiation is aligned to the first axis (ordination) or K = 2 (model-based clustering). Some studies had both ordination and model-based clustering, and some species were included in more than one study, thus multiple points can appear against each species. Studies that did not present ordination or model-based clustering are not shown. The number of locations where distinct groups were sympatric was based on K clustering, using the K value preferred by the authors, where a dash indicates that this value could not be inferred. From the 51 studies examined, 39 presented results that could be evaluated against criteria 1-3.

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357 **2.2** No clear patterns for symbionts associated with cryptic coral host taxa

358 An important aspect of coral biodiversity is the diverse microbial community living within the 359 cnidarian host, where mutualistic relationships with endosymbiotic dinoflagellates from the 360 family Symbiodiniaceae are known to affect whole organism physiology (LaJeunesse et al., 361 2018). The specificity of coral-symbiont pairings is influenced by host-symbiont genetic 362 interactions, host reproduction, symbiont transfer mode, and environment (see Baker, 2003; 363 Davies et al., 2023; Turnham et al., 2023 for further readings). While symbiont evolution is 364 not the focus of this review, joint consideration of coral hosts and their microbes is relevant 365 in understanding the specificity of associations that can affect traits, local adaptation, and 366 restoration actions.

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Among the population genomic studies that we surveyed, 43% genotyped dinoflagellate symbionts alongside the coral hosts. Several studies (e.g., Howells *et al.*, 2016; van Oppen *et al.*, 2018; Gomez-Corrales & Prada, 2020; Bongaerts *et al.*, 2021a; Prata *et al.*, 2022; Buitrago-Lopez *et al.*, 2023; Starko *et al.*, 2023) used internal transcribed spacer (*ITS*) sequences to characterize the community of symbiont lineages living within a colony, aligning with traditional marker-based methods (Davies *et al.*, 2023) for symbiont characterisation. Some studies also used incidentally recorded symbiont sequences retrieved from whole-colony
sequencing (either reduced representation or shotgun whole genome sequencing) to make
inferences about symbionts, including reconstructing organelle diversity (Bongaerts *et al.*,
2017; Forsman *et al.*, 2017; Gonzalez-Zapata *et al.*, 2018; Cooke *et al.*, 2020; Bongaerts *et al.*,
2021a; Matias *et al.*, 2023; Zhang *et al.*, 2023) or summarizing symbiont genomes with *k*-mer
analyses (Zhang *et al.*, 2023).

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381 A central issue for coral adaptation and diversification is the extent to which symbiotic 382 associations are flexible. Brooding corals are expected to have less flexible symbiont 383 associations because they often exhibit vertical symbiont transmission from the maternal 384 colony (Knowlton, 2001; Johnston, Cunning & Burgess, 2022; Turnham et al., 2023). However, 385 in our review, only one brooding coral (Stylophora pistillata) that was genotyped for both 386 symbionts and hosts (Buitrago-Lopez et al., 2023) preventing adequate comparisons among 387 reproductive modes. Symbiont types were commonly found to be shared across cryptic host 388 lineages for S. pistillata (Buitrago-Lopez et al., 2023), for broadcast spawners (O. faveolata: 389 Gomez-Corrales & Prada, 2020; Pachyseris speciosa: Bongaerts et al., 2021a; Pocillopora 390 verrucosa: Buitrago-Lopez et al., 2023; Porites lobata: Rivera et al., 2022; Porites sp.: Starko 391 et al., 2023) for mixed mode Pocillopora damicornis (van Oppen et al., 2018), and for taxa 392 with unknown modes of reproduction (Agaricia fragilis: Bongaerts et al., 2017; Agaricia 393 *lamarcki*: Prata et al., 2022). Some symbiont types appeared specific to cryptic taxa (O. 394 faveolata: Gomez-Corrales & Prada, 2020; P. speciosa: Bongaerts et al., 2021a; P. verrucosa: 395 Buitrago-Lopez et al., 2023; massive Porites sp.: Starko et al., 2023; P. damicornis: van Oppen 396 et al., 2018), whereas other sets of cryptic taxa were not associated with specific symbiont 397 types (Agaricia fragilis: Bongaerts et al., 2017; Agaricia lamarcki: Prata et al., 2022; S. 398 *pistillata*: Buitrago-Lopez et al., 2023). Thus, some symbiont strains may be specialized, as 399 evidenced by symbiont lineages only associating with one host group; however, this pattern 400 is not always a characteristic accompanying cryptic cnidarian taxa.

401

Moving beyond single marker genotyping of symbionts (Davies *et al.*, 2023; Ishida *et al.*, 2023;
Zhang *et al.*, 2023) may provide better resolution of host-symbiont associations. For example,
Rivera et al. (2022) found that symbiont identities among *P. lobata* did not align to host taxa
using ITS genotyping but were concordant when using higher resolution SNP-based analyses.
Additional studies are needed, however, to determine whether alternative methods
harnessing whole-genome data to summarize symbiont diversity will reveal more nuanced
patterns than ITS based results.

409

410 **2.3 Depth** and microhabitats can segregate cryptic taxa

For coral hosts, the existence of distinct genetic groups in sympatry implies that differentiation may be maintained by strong natural selection arising from local environments (Richardson *et al.*, 2014). Such strong selection and genotype-by-environment associations may merely reflect pre-existing reproductive isolation (Bierne *et al.*, 2011), although 415 divergent natural selection can also contribute to reproductive isolation (Schluter, 2001; 416 Rundle & Nosil, 2005). For marine species, including corals, depth often delineates closely-417 related species (Knowlton, 1993) and could drive ecological speciation in corals (González et 418 al., 2020). In coral reef communities, depth is a predominant structuring aspect, with distinct 419 species turnover between shallow (approximately < 30 m) and mesophotic (approximately 420 30-150 m) depth zones Importantly, these transitions exist across communities because 421 many environmental factors covary with depth, such as light intensity and spectrum (Lesser, 422 Slattery & Leichter, 2009), temperature (Kahng et al., 2019), nutrients (Leichter, Stokes & 423 Genovese, 2008), water flow (Muir et al., 2015), as well as disturbance frequency and severity 424 (Bongaerts & Smith, 2019), creating highly contrasted habitats often only meters apart. The 425 strongest environmental differences are within the first few meters due to the exponential 426 decay of light in the aquatic environment – the most important environmental factor for light-427 dependent scleractinians. Thus, it is highly likely that recently diverged taxa will also be 428 partitioned by depth, matching patterns of species turnover.

429

430 Among the population genomic studies surveyed, cryptic taxa differed in abundance by 431 depth-associated habitats. Eleven studies to date undertook extensive sampling over 432 replicated depth-associated habitat contrasts (i.e., more than one site >10 km apart with two 433 depth habitats sampled within each site). Replicated differentiation by depth was found for 434 Agaricia fragilis (i.e., shallow vs. mesophotic, Bongaerts et al., 2017), Agaricia lamarcki (i.e., 435 shallow vs. mesophotic, Prata et al., 2022), Isopora brueggemanni (i.e., lagoon vs. slope; 436 Thomas et al., 2019), Pocillopora damnicornis (i.e., flat vs. slope, van Oppen et al., 2018) and 437 Montastraea cavernosa (i.e., shallow vs. mesophotic, Sturm et al., 2022) but not for Agaricia 438 grahamae (i.e., upper vs. lower mesophotic, Prata et al., 2022), Stephanocoenia intersepta 439 (i.e., shallow vs. mesophotic, Bongaerts et al., 2017), Acropora digitifera (i.e., lagoon vs. slope, 440 Thomas *et al.*, 2019), *Agaricia undata* (i.e, shallow vs. upper mesophotic and upper vs. lower 441 mesophotic, Gonzalez-Zapata et al., 2018). Thus, differentiation by depth frequently 442 discriminates cryptic coral taxa.

443

444 The most extensive investigations of differentiation by depth have been for Seriatopora 445 hystrix (brooder) on the Great Barrier Reef, Australia and Eunicea flexuosa (broadcast 446 spawner) in the Caribbean. Although most of the studies discussed below pre-date genomic-447 scale genotyping and thus rely on small numbers of microsatellite loci for making genetic 448 inferences, they are noteworthy because they meticulously documented small scale depth 449 and habitat segregation of host genotypes and also used reciprocal transplantation 450 experiments to further support evidence for depth-associated adaptive differentiation. 451 Bongaerts et al. (2010) showed that genetically distinct groups of S. hystrix were associated 452 with reef zone and depth position (backreef at 2 m, shallow reef at 6 m, deep reef at ~27 m; 453 locations within 200 m distant horizontally). For the gorgonian E. flexuosa, Prada & Hellberg 454 (2013) demonstrated that individuals from <5 m and >20 m depths were distinct and joined 455 by a narrow hybrid zone (< 200 m horizontally). This depth separation was likely maintained

456 by selection against juveniles (Prada & Hellberg, 2014). Reciprocal transplantation supported 457 evidence for local adaptation in both cases (Bongaerts et al., 2011; Prada & Hellberg, 2013). 458 Together, these two examples (S. hystrix and E. flexuosa) show that cryptic coral taxa could 459 comprise distinct depth-specific ecotypes maintained by selection and furthermore illustrate 460 the utility and feasibility of using experiments to strengthen evidence for local adaptation 461 across small spatial scales. There is an unrealised opportunity to combine future genomic 462 investigations with manipulative experiments to advance the understanding of speciation and 463 adaptation (discussed further in Section 3).

464

465 While depth has been one of the most investigated environmental axes of local genetic 466 differentiation in corals, cryptic taxa can be associated with other habitat differences (and 467 sometimes are evident with few markers (Warner et al., 2015)). In one example, broad 468 physiological differences between A. hyacinthus taken from two reef pools less than 500m apart but varying in natural thermal variability (coined "highly" and "moderately" variable 469 470 pools (reviewed by Thomas et al., 2018); were partially explained by the relative abundance 471 of four cryptic species (identified using ordination of SNP-based genotypes) that differed in 472 abundance between the two pools (Rose et al., 2017). Similarly, substantial differences in 473 skeletal morphologies and gene expression were found between P. lutea colonies sampled 474 from a mangrove lagoon versus colonies from a reef approximately 500 m away (Scucchia et 475 al., 2023); however, these phenotypic differences also aligned with genome-wide 476 differentiation measured by transcriptome-derived SNPs, suggesting that mangrove and reef 477 populations could simply be different taxa (applying the criteria in Section 2.1).

478

479 It is important to note that the structure and composition of coral-associated microbial 480 communities also often vary along environmental gradients, including those associated with 481 depth and disturbance (Klaus et al., 2007; Bongaerts et al., 2013; Howells et al., 2013; Quigley 482 et al., 2022), and likely help mediate coral holobiont adaptation to environmental stress. 483 Endosymbiotic dinoflagellates (LaJeunesse et al., 2018) enable certain coral species to reside 484 within specific light environments, where mesophotic and shallow depth coral hosts often 485 harbour distinct symbiont strains, whereas depth-generalists appear to be able to host many symbiont strains (Bongaerts et al., 2013). Consistent with these previous findings, several of 486 487 the studies examined here reported greater spatial or environmental partitioning among symbionts as compared to hosts (e.g., Astrangia poculata: Aichelman & Barshis, 2020; P. 488 489 verrucosa: Buitrago-Lopez et al., 2023; S. pistillata: Buitrago-Lopez et al., 2023; Platygyra 490 daedalea: Howells et al., 2016; Acropora tenuis: Matias et al., 2023; A. lamarcki: Prata et al., 491 2022; but not so for A. digitifera: Zhang et al., 2023). Intriguingly, Starko et al. (2023) 492 demonstrated that a distinct symbiont community associated with one cryptic taxon of 493 massive Porites sp. changed following a heatwave, such that the post heatwave composition 494 better matched the symbiont communities living in the other two cryptic taxa. Thus, 495 symbionts may change to track local environments (Baker, 2003), although this flexibility may 496 differ among host taxa (Quigley et al., 2022).

497

498 Whereas differentiation by depth and habitat appear to be fairly common in corals (and their 499 associated microbes), sampling strategies for many coral genomic studies are surprisingly 500 underpowered in their ability to detect genetic differentiation by depth or habitat. Among the 501 population genomic studies examined here, 25% failed to report sampling depth (or any other 502 relevant habitat, including our own work, e.g. Matias et al., 2023). Presumably, most of the 503 genotyped corals across the studies reviewed here were sampled on SCUBA from < 30 m and 504 were sampled from a similar depth range across all sites. Among studies that did report depth, 505 many sampling regimes had depth confounded with geography where each location was 506 sampled at a single depth. A minority of studies (21%), however, implemented a structured 507 sampling design where the same depth was sampled at more than one location (Fig. 4). More 508 complete reporting on depth and other microenvironmental attributes alongside sampling 509 study designs that replicate environmental contrasts are needed to advance our 510 understanding of how cryptic taxa, reproductive barriers, heterogeneous environments, and 511 natural selection interact to shape coral biodiversity.



Figure 4 - Summary of depth sampling schemes for studies that reported depth and sampled at more than one depth. Numbers indicate the number of distinct locations that were sampled per depth. Locations were considered distinct if the nearest locations were depth contrasts (e.g., adjacent sites sampled at 5 and 15 m were considered as two locations); otherwise, same depth locations within 10 km were collapsed as a single location. Dotted lines connect locations from the same study and thick grey lines indicate the sampling range (as reported by authors). Citations are as follows: a) (Thomas *et al.*, 2019); b) (Bongaerts *et al.*, 2017); c) (Prata *et al.*, 2022); d) (Gonzalez-Zapata *et al.*, 2018); e) (Bongaerts *et al.*, 2021); f) (Shilling *et al.*, 2023); g) (Rippe *et al.*, 2021); h) (Aichelman & Barshis, 2020); i) (Drury *et al.*, 2020); j) (Sturm *et al.*, 2020); k) (Sturm *et al.*, 2022); l) (Aurelle *et al.*, 2022); m) (van Oppen *et al.*, 2018); n) (Meziere *et al.*, 2024).

524 **2.4 Cryptic taxa can differ in their susceptibility to thermal stress**

525 Limited but growing evidence shows that phenotypes of closely related cryptic coral taxa can 526 differ with respect to environmental tolerances, which may affect where they can live (i.e., 527 their niche, including depth) and how they might respond to climate change and other 528 anthropogenic stressors. For example, under experimental conditions, one genetic group of 529 A. hyacinthus showed greater resistance to bleaching conditions under experimental trials 530 compared to the three other groups, matching within reef microhabitat distributions of these 531 taxa in American Samoa (Rose et al., 2021). Likewise, heat resilience and survival may differ 532 between cryptic taxa in the wild. By monitoring tagged colonies during an extreme heatwave, 533 Starko et al. (2023) found that mortality was significantly higher for one of three massive 534 Porites taxa on Kiritimati. In Palau, cores from P. lobata showed differences in past growth 535 rates and stress banding between four cryptic taxa – indicative of taxa-specific differences 536 over long time periods (1970-2014), with the two most heat tolerant taxa predominating at 537 the warmest location (Rivera et al., 2022). Similarly, survival through a heatwave differed 538 among Pocillopora taxa distributed across Moorean reefs (Burgess et al., 2021). These studies 539 illustrate that cryptic taxa are not necessarily interchangeable, especially with respect to heat 540 stress responses and survival over both short and long-time frames.

541

542 In Textbox 2, we investigate whether thermal biology studies are attuned to the coral cryptic 543 taxon "problem". If cryptic coral taxa consistently differ in their phenotypes, including 544 response to thermal stress, then experimental outcomes need to be evaluated with respect 545 to taxonomic identity. Yet, we find that only 8% of such studies included genotyping that 546 could identify cryptic taxa, suggesting that many studies could be inadvertently evaluating 547 multiple taxa and thereby supporting incorrect or biased conclusions.

548

549 **2.5 Gene flow and introgression link taxa across divergence histories**

550 Interbreeding between genetically distinct groups of corals could elevate genetic diversity 551 and contribute to adaptation. Hybridisation in corals has long been suspected (van Oppen & 552 Gates, 2006; Willis et al., 2006), but studies using few genetic markers often lack the 553 resolution to appropriately investigate hybridisation in the context of recently diverged taxa, 554 where genetic similarities can result from either shared ancestral diversity or gene flow. If 555 hybrids interbreed with parental species (backcrossing), interspecific recombination leads to introgression of genetic material from one lineage into the other. Thus, by analysing 556 557 thousands of genomic SNPs, genomic studies can often resolve whether genetic similarities 558 are due to gene flow and sometimes identify genomic regions that have experienced high or 559 low levels of gene flow as well as identify where selection may have shaped introgression 560 patterns (Taylor & Larson, 2019).

561

562 Coral population genomic studies often find individuals with possible hybrid ancestries, but 563 only two studies included explicit tests for recent hybridisation (such as tests implemented in 564 NewHybrids (Anderson & Thompson, 2002)) that probabilistically assign individuals as 565 putative first-generation and early backcrosses. Hybrid individuals were found among 566 Agaricia taxa (Prata et al., 2022) but not among S. pistillata taxa (Meziere et al., 2024). Other 567 studies have identified likely hybrid individuals based on the proportion of assignment to 568 different groups from model-based clustering outputs (e.g. Cooke et al., 2020; Kitchen et al., 2020; Bongaerts et al., 2021a; Fifer et al., 2021; Rippe et al., 2021; Rivera et al., 2022; Matias 569 570 et al., 2023). Using a majority background group assignment score of < 0.75 for distinct 571 sympatric genetic groups as an indication of possible recent hybridisation, we identified 572 potential hybrid individuals for 21 of the 34 species surveyed; however, these mixed ancestry 573 individuals are based on original author preferred K groups and therefore do not necessarily 574 reflect admixture between partially reproductively isolated groups (see Textbox 3 for further 575 discussion). Nonetheless, the prevalence of individuals with mixed ancestry suggests that 576 hybridisation could be ongoing for many taxa.

577

578 Historical gene flow between divergent species can be an important aspect of their 579 evolutionary history (Shinzato et al., 2015; Voolstra et al., 2023) even without early 580 generation hybrid individuals being frequent. Unique signatures of historical ("ancient") gene 581 flow between taxa have been demonstrated by contrasting population genetic demographic 582 models using alternative speciation scenarios (e.g., no gene flow versus periodic or ongoing gene flow) (Gutenkunst et al., 2009; Beaumont, 2010; Sousa & Hey, 2013; Fraïsse et al., 2021) 583 584 thereby resolving the contributions of shared ancestral polymorphisms and introgression to 585 shared genetic variation among taxa. Where demographic modeling has been applied to 586 corals, divergence time estimates among cryptic taxa ranged between a few thousand years 587 (e.g., ~10k between A. digitifera: Zhang et al., 2023) and up to 9 million years (i.e., Galaxea 588 fascicularis: Wepfer et al., 2020). However, most studies reported recent divergence times 589 ((~0.2-1 million years: Cooke et al., 2020; Bongaerts et al., 2021a; Fifer et al., 2021; Matias et 590 al., 2023; Meziere et al., 2024) that predate Holocene reef configurations. This suggests that 591 divergence began before species shifted into their present-day ranges. Additionally, several 592 studies found strong support for models of speciation with ongoing or episodic gene flow 593 between genetically differentiated and sympatric and/or depth associated cryptic coral taxa 594 with low-to-moderate levels of genetic exchange (e.g., Wepfer et al., 2020; Fifer et al., 2021; 595 Rippe et al., 2021; Prata et al., 2022; Tsuchiya et al., 2022; Matias et al., 2023; Starko et al., 596 2023; Zhang et al., 2023; Meziere et al., 2024) (Prada & Hellberg, 2021). These findings 597 suggest that recent divergence with gene flow is common across a range of phylogenetic and 598 functional coral diversity and is geographically widespread.

599

An emerging observation among diverse metazoans is that gene flow between closely-related species is variable across genomes (Ravinet *et al.*, 2017) due to local adaptation or reproductive incompatibilities that reduce gene flow for selected regions (Bay & Ruegg, 2017; Martin & Jiggins, 2017). In cases where variable introgression rates were tested explicitly using demographic modeling of coral divergence, models including heterogeneous gene flow rates received the highest support (Fifer *et al.*, 2021; Rippe *et al.*, 2021; Starko *et al.*, 2023; Meziere *et al.*, 2024), providing evidence for genomic barriers to gene flow in the sympatric taxa studied and supporting the contention that some degree of reproductive isolation helps maintain genetic cohesion of these interbreeding taxa. For example, in *S. pistillata*, more divergent taxa had a higher proportion of their genomes experiencing reduced gene flow compared to the less divergent taxa, implying that islands of differentiation become wider as speciation proceeds (Meziere et al., 2023). These findings are consistent with morphologically similar taxa being at various stages of the divergence process (Roux *et al.*, 2016).

613

614 Low levels of gene flow can contribute to adaptation via introgression (Martin & Jiggins, 615 2017), where alleles derived from a different species can affect a specific beneficial trait (e.g. 616 resistance to hypoxia at high altitude in humans: Huerta-Sánchez et al., 2015; and winter coat 617 colour in hares Giska et al., 2019). In corals, a genomic region (approximately 220 kb) that 618 appear to contribute to increased bleaching-tolerance for one A. hyacinthus taxon relative to 619 other cryptic A. hyacinthus taxa was likely acquired through hybridisation with A. millepora 620 (Rose *et al.*, 2021). This finding is consistent with some evidence suggesting that historical 621 introgression events may have coincided with Acropora range expansions, with possible links 622 to ecological opportunities associated with their diversification (Mao, Economo & Satoh, 623 2018). While these studies implicate a role for hybridisation in adaptive evolution, and the 624 possibility that large chromosomal variants may be important for adaptation in the context 625 of gene flow, there have been no comprehensive investigations of adaptive introgression in 626 corals to date.

627

628 **2.6** *Implications*: investigations that do not test for cryptic taxa risk biased estimates of 629 *ecologically important traits*

630 The preceding review of population genomic studies shows that cryptic coral taxa are 631 common (Section 2.1) and may be adapted to and associated with different 632 microenvironments – especially depth (2.3). These cryptic taxa, however, are often linked by 633 sharing symbiont strains (2.2) and via some gene exchange (2.5). Therefore, cryptic taxa may 634 be distinct in terms of ecology, physiology, and evolution, but how to describe and delineate 635 taxa is not obvious, as there may not be complete divisions or obvious diagnostic characteristics. Even with access to genomic-scale genotyping, taxonomic resolution is 636 637 affected by sampling and open to interpretation. It is clear, however, that gross morphology 638 assessed by humans under field conditions is unreliable for recognising closely related taxa, 639 as over 50% of studies in our literature search show genetic evidence for cryptic taxa (Fig. 3). 640 Simply put, any coral investigation that does not include genotyping, risks treating a 641 heterogeneous mix of partially reproductively isolated taxa as a single species. To get a sense 642 of how extensive this issue could be for other fields in coral biology, in Textbox 2 we show 643 that only 8% of coral experiments on thermal tolerance include genotyping that could identify 644 cryptic taxa. Although it is outside the scope of this paper to examine the thousands of papers 645 in coral biology that examine phenotypes including niches and geographic distributions, we 646 would expect that most experimental and ecological studies are not combined with

647 genotyping in a way that can detect cryptic taxa, if they are present. Given that unaccounted 648 cryptic taxa can bias estimates of species' traits (alongside estimates of biodiversity and 649 evolutionary dynamics - Section 1), inferences from many studies need to be viewed with 650 scepticism.

651

652 3 Corals as untapped systems for studying speciation and adaptation in a changing 653 world

Although cryptic species and gene flow (hybridisation) between closely related coral taxa are
more common than previously thought, corals present largely overlooked systems for
studying speciation and adaptation. Such studies could bring new insights to processes
generating coral biodiversity and would also clarify biological attributes of corals that may
sway conservation management decisions and strategies.

659

660 The emerging consensus that closely-related coral taxa are frequently sympatric at coarse spatial scales yet segregate by depth or other micro environmental characteristics aligns well 661 662 with models of ecological speciation (Rundle & Nosil, 2005). Furthermore, the presence of 663 distinct cryptic taxa in close geographic proximity suggests that selection may be very strong (a high selection to migration ratio: Richardson et al., 2014)). To what extent cryptic taxa differ 664 665 phenotypically or in terms of competitive ability is largely unknown, although differences in 666 bleaching susceptibility among some cryptic taxa suggest differing vulnerabilities to climate 667 change (Section 2.4). As sessile organisms, corals are well-suited to manipulative experiments 668 such as common garden or reciprocal translocation designs. Additionally, their clonal nature 669 means that genetically identical fragments from the same colony can be exposed to differing 670 treatments, offering rich opportunities to combine experiments with genomic analyses to 671 holistically investigate the interactions between taxon identity, phenotype, and environment 672 (Pinsky et al., 2023; Richards et al., 2023).

673

674 Divergence dates between cryptic taxa often pre-date Holocene reef configurations, implying 675 that old standing genetic diversity is spread across contemporary reefs that are characterized 676 by spatially complex yet replicated microhabitats and environmental gradients. Thus, corals 677 would be ideal for investigations that explore the genetic mechanisms of parallel divergence 678 (e.g. analogous to fishes that have colonized post-glacial lakes: Rougeux, Bernatchez & 679 Gagnaire, 2017; De-Kayne et al., 2022). Such investigations would also provide insights on the 680 geographic distribution of standing genetic variation, which may be under increasing selective 681 pressure due to pervasive environmental changes. For example, knowing whether 682 geographically distant populations do or do not share alleles for advantageous traits can guide 683 decisions regarding the utility of assisted migration. This is because evolutionary rescue is only worth considering if donor and recipient populations differ in functional standing genetic 684 685 diversity.

686

- 1t is likely that endogenous barriers to reproduction suppress gene flow between taxa to some extent. Analyses to date support evolutionary genomic models that allow different genomic regions to be more or less permeable to gene flow, consistent with chromosomal inversions or other structural variants contributing to reproductive isolation (see section 2.5). Future studies that use chromosomal resolution genotyping will be critical to forming a deeper understanding of how species boundaries are maintained (e.g., Leitwein *et al.*, 2020) and can guide decisions on assisted migration or choosing broodstock for selective breeding (Fig. 2).
- 695 Individual colonies with genotypes consistent with recent hybrid ancestry have been noted 696 (Section 2.5). To date, these individuals have largely been treated as curiosities and not 697 subject to focused study. Studies of hybridization and hybrid zones also provide important 698 insights on speciation and adaptation (Hewitt, 1988) and would yield key background 699 information to evaluate suggestions that hybrid corals are viable in nature and could be used 700 in restoration. Potential restoration interventions based on hybridisation rest on the 701 supposition that hybrid corals differ in phenotypes from parental species through some 702 combination of hybrid vigor or transgressive segregation. Promisingly, first generation (F1) 703 crosses among Acropora species have showed that some crosses survived as well as or better 704 than parentals in some conditions (Chan et al., 2018). Aside from the transect studies of Prada 705 & Hellberg (2014), we know of no systematic attempt to map the spatial and environmental 706 distributions of hybrids relative to parental taxa, and yet this knowledge can yield important 707 insights into natural selection and dispersal of cryptic taxa (Barton & Hewitt, 1985). Mapping 708 hybrids would also allow restoration biologists to easily locate naturally-occurring hybrid 709 corals with a wide diversity of genotypes that could be used as broodstock in restoration 710 methods that rely on sexually-produced offspring (Table 1).
- 711

712 Throughout this review, we have focused primarily on the cnidarian component of coral 713 genomes to document evidence for cryptic species and hybridization. However, in considering 714 how future studies could build on these observations to better understand speciation and 715 adaptation processes, it will also be important to integrate genetic analyses of microbes -716 especially symbiotic dinoflagellates – with those of the cnidarian host. An exciting line of 717 investigation would be to try to understand the co-evolutionary dynamics of hosts and 718 symbionts in reference to environmental adaptation and speciation, where environmental 719 heterogeneity likely exerts direct selection on both corals and their symbionts (i.e. the coral 720 holobiont) and indirect selection via host-symbiont genetic interactions. 721

722

4. Conclusions

In this review, we demonstrate that cryptic coral taxa are extremely common and are oftenconnected by gene flow. Many previous studies have emphasized the importance of such

cryptic coral taxa (Gomez-Corrales & Prada, 2020; Bongaerts *et al.*, 2021a; Burgess *et al.*,
2021; Feldman *et al.*, 2021; Rippe *et al.*, 2021; Zayasu *et al.*, 2021; Prata *et al.*, 2022; Rivera *et al.*, 2022; Matias *et al.*, 2023; Pinsky *et al.*, 2023; Starko *et al.*, 2023; Voolstra *et al.*, 2023;
Meziere *et al.*, 2024) and here we use a systematic examination of published population
genomic studies to document that cryptic taxa are indeed widespread across coral families
(Fig. 3).

731

732 The prevalence of cryptic coral taxa means that many accepted understandings and 733 conclusions regarding coral biology could be incorrect. In this review, we argue that hidden 734 taxonomic diversity can affect conservation thinking and, in the extreme, mislead possible 735 management decisions if ignored. In Section 1, we highlight how ignoring cryptic taxa can: 736 bias estimates of spatial biodiversity patterns; make species appear to have larger ranges, 737 trait spaces, and niches; and can skew inferences regarding intraspecific population structure 738 and gene flow in unpredictable ways. Thus, as a field, we are unable to confidently generalize 739 species distributions, ranges, and phenotypes including resilience to heat stress without 740 adequate study of potential cryptic taxa (Textbox 3). Identifying locations with high genetic 741 diversity that may harbour greater adaptive potential or inferring sites with high gene flow 742 and dispersal will crucially depend on analyses that are able to detect and account for distinct 743 taxa.

744

745 Although observations of cryptic coral taxa are frequent, our collective knowledge regarding 746 the evolutionary dynamics that enable closely-related taxa with incomplete species 747 boundaries to persist in sympatry remains limited. There is a vast potential to unite coral 748 studies with the insights and approaches into speciation and adaptation from other fields and 749 organisms. For example, partnerships between coral ecologists, physiologists, and population 750 geneticists may bridge insights into coral population microevolution to climate change, while 751 collaborations with experts from other fields may broker novel analyses and genomic-based 752 approaches to speciation and hybridisation in corals.

753

The reality, however, is that the future for corals and coral reefs is perilous (Knowlton *et al.*, 2021). Management decisions cannot wait for perfect information. Therefore, it is incumbent for evolutionary biologists to identify attributes that do or will affect coral health and evolutionary trajectories for the short-term future. An important next step will be to explore how conservation and management actions can best proceed with a newfound expectation that coral species boundaries are unlikely to be well defined – a conservation challenge that ultimately afflicts many other taxa apart from corals (Roux *et al.*, 2016).

762 **Textbox 1: Applying taxonomic delineation with reproductive isolation criteria**

We propose three requirements for identifying and delineating taxa using genomic-informed ordination and model-based clustering approaches. In the empirical example that follows, we detail how coral cryptic taxa were identified using three criteria (section 2.1) and highlight difficulties with interpretation.

767

768 Colonies of the brooding coral, Agaricia agaricites, were sampled at four sites ~10-15 km apart along southwest Curaçao, and collections were further subset to three depths (5, 10 769 770 and 20 m) at each site. Genotyping using reduced representation sequencing of 335 colonies 771 and 1,629 SNP-loci revealed distinct genetic groups co-occurring within four sampled sites. 772 This study provides a clear example of cryptic taxa identified according to criteria 1-3. 773 Furthermore, taxa occupied unique depth ranges (AA1 occurs predominantly at 20 m, 774 whereas AA2 occurred at all depth sampled) suggesting divergence of taxa by habitat (Fig. 775 T1A & C).





Figure T1 - *Agaricia agaricites* resolved into two distinct taxa and fulfil criteria 1, 2 and 3 for cryptic taxon delineation. A) PC1 resolves two sympatric groups at every sampling site. PC2 represents an example of geographic partitioning in AA2 (Site 1 vs Site 2 - 4) and therefore does not necessarily imply reproductive isolation. B) Shows that the percentage of variation explaining PC1 (14.65%) is ~5x more than PC2 (3.01%). C) Analyses using ADMIXTURE for K=2 assigned individuals to each group with high confidence (q > 0.9) and showed that AA1 and AA2 were sympatric at all sites. D) There is a significant drop in cross-validation error between K=1 and K=2, and greater log-likelihood, supporting

the selection of K = 2. All three criteria are met in delineating AA1 and AA2 cryptic taxa within A. agaricites.

788

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787

789 This example also illustrates some complications for interpreting differentiation among 790 putative taxa. The second PC axis (also mirrored in ADMIXTURE outcome for K=3 and K=4) 791 shows partitioning that largely aligns with geographic separation and so would not be 792 considered as delineating distinct taxa under our criterion of sympatry. If the variation 793 captured by the second PC axis (and K=3 and K=4) reflects geographic differentiation, 794 geographically mismatched genotypes likely reflect recent immigration (shown by dashes 795 among K=3 and K=4 in Fig. T1C). Distinguishing migrants from distinct taxa may be especially 796 difficult when sampling numbers are low. However, if immigration is high (> 1 migrant per 797 generation) and there are no barriers to reproduction then the structure between 798 populations is expected to dissipate over a few generations (Waples & Gaggiotti, 2006).

799

This example highlights the utility of PCA and cluster-based modeling methods for identifying cryptic coral taxa. However, additional subsetting and filtering steps are necessary to thoroughly scrutinize data for consistent patterns and reveal accurate groupings. To better understand the possible biases of both PCA and assignment methods, we refer readers to (Pritchard *et al.*, 2000; McVean, 2009; Puechmaille, 2016).

805

806

Textbox 2: Are coral experiments designed to detect cryptic taxa?

807 Overlooking cryptic taxa in experiments can bias interpretations of experimental results. To 808 ascertain how substantial this issue might be for coral studies, we focus on experiments 809 related to thermal tolerance as a subset of coral studies more generally. Marine heatwaves 810 have caused extensive coral mortality and bleaching events globally (Leggat et al., 2019), and 811 thus numerous coral studies have aimed to ascertain intra- and inter-specific differences in 812 phenotypic and physiological heat stress responses using experiments (e.g., common 813 gardens, reciprocal transplants, etc.) and natural heating events. Mounting evidence suggests 814 that cryptic species and hybrids display contrasting responses to heat stress (Section 2.4), and 815 so experimental results may be more accurate when taking into account potential cryptic taxa 816 identified using genomic-scale genotyping .

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The Coral Research and Development Accelerator Platform (CORDAP) database (Ortiz, Humanes & Scharfenstein, 2023) represents a curated search for papers that study thermal biology of corals. We screened the database to identify records which used genome-wide data of the coral host (i.e., coral SNP data) and those that conducted either ordination or model-based clustering (as in 2.1). We evaluated the database to determine:

- 1. The number of studies within the database undertaking marine heatwave
- 824 experiments that genotyped corals for multiple unlinked markers (i.e., created data 825 that could be used for ordination or model-based clustering).
- 826827827827828829<l
- 828 3. Whether there is evidence for cryptic taxa based on applying the criteria outlined in829 Section 2.1.
- 830

We found that very few experimental studies genotyped coral colonies - from 562 studies, 831 832 222 studies undertook marine heatwave experiments, of which 59 studies included any sort 833 of host genotyping and only 17 used high-resolution genome-wide markers such as SNPs. Still 834 fewer studies undertook either ordination or model-based clustering on their genomic data 835 (n= 9; Fig. T2). Of those nine studies, three showed indications of cryptic taxa in line with proportions of cryptic taxa in population genomic surveys (Fig. 3). It is likely that many studies 836 837 will have inadvertently sampled multiple taxa and therefore we would anticipate that 838 reported variances among individuals within studies will be greater than true variances within 839 cryptic taxa (see Section 1). This could manifest as a bias in overestimated thermal tolerance 840 breadth and thus may also mask differences in measured tolerance in comparative tests between morphospecies. The CORDAP database focuses on one group of studies, but we 841 842 would anticipate that similar issues arise across all coral experimental work that does not 843 leverage genomic-level genotyping of individual colonies.



Figure T2 – Proportion of individual studies from the Coral Research and Development Accelerator Platform (CORDAP) thermal tolerance experiment database that: (1) record genotype data capable of identifying cryptic taxa via genome-wide data of hosts, and (2) test for cryptic taxa via ordination-based analyses or model-based clustering analyses.

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Textbox 3: Best practice recommendations

Future genomic surveys of corals should be designed with the expectation that cryptic species could be encountered. This means undertaking structured and replicated sampling, reassessing field collection protocols, and testing for cryptic taxa as part of bioinformatic and population genetic analyses. Ensuring that all data and metadata are thoroughly documented
will create the best chances that future investigations can re-examine published data as novel
methods emerge and thus bypassing the need for additional fieldwork and genotyping in
some cases.

859

860 Spatial sampling at the colony level

The best evidence for discriminating cryptic taxa from population structure is when distinct taxa are observed in sympatry (or close geographic proximity) at multiple locations (Section 2.1). We acknowledge that there is an element of chance in co-sampling distinct taxa. Given that depth appears to be the most common axis of differentiation, however, researchers who are planning to sample across depth should ensure that the same depths are sampled at multiple distinct locations to enable the detection of repeated co-occurrences of distinct taxa. 867

868 Alongside structured sampling, investigators would greatly enhance their data's value (and 869 scope for future inference) by transitioning from a population sampling mindset to focusing 870 on individual sampling and seeking to capture as much environmental context as possible at 871 the colony level. For instance, "cryptic" species may in fact be morphologically distinguishable 872 based on subtle characteristics (for example, S. pistillata, Meziere et al., 2024) and therefore 873 taking comprehensive photographs that can be examined later (see, for example, Protocol for 874 Coral Collection & Curation by Proiect Phoenix: 875 https://coralprojectphoenix.org/resources/#protocols) may allow diagnostic characters to be 876 identified post hoc. For a subset of samples, it would be useful to retain larger colony fragments that would be suitable as museum voucher specimens, if permits allow. Recording 877 878 each colony's geoposition and depth can greatly support analyses based on depth (i.e. as a 879 continuous rather than categorical predictor) and space, while simultaneously could provide 880 insights on the microhabitat attributes of cryptic taxa and hybrid individuals (as in (Prada & 881 Hellberg, 2014)). A particularly exciting technology that can greatly advance this colony-882 focussed perspective is photogrammetry (Bongaerts et al., 2021b). We recognise that moving 883 the focus from coral populations to colonies will require more time, effort, and expense, but 884 the insights will be far richer.

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886 Datasets that link genomic genotyping with ecological context at the colony level will be 887 immensely valuable for gaining insights into ecological and evolutionary processes relevant I suggest being a tat more specific 888 to conservation. To maximise this value, investigators should strive to make all facets of their on this topic, eg data open, which includes linking genotypes with all recorded metadata including metadata FAIR guidelines 889 https://doi.org/ 890 that might not be relevant to the original study (but that might be of value to others). 10.1038/ 891 Analytical pipelines also need to be fully reproducible by enabling consistency in sdata.2016.18 892 bioinformatics and analytical decisions across studies such that outcomes can be confidently 893 compared. No doubt, all this extra documentation is a substantial amount of work, and 894 therefore should be incorporated into initial project planning. Coral biologists can take 895 inspiration from plant population geneticists who have greatly advanced insights and impacts

ie "pseudocryptic" species? perhaps use that term? by sharing highly curated data sets that have been re-used and supported a myriad of
additional studies after their initial publication, for example, the IntraBioDiv (Meirmans *et al.*,
2011) dataset of 27 co-distributed alpine plant genotypes has supported numerous
reanalyses and test cases. Additionally, the genomic (and phenotypic) datasets for lodgepole
pine and spruce from the AdapTree group (https://adaptree.forestry.ubc.ca/about/scientificsummary/) have greatly advanced understandings of spatial adaptive diversity in trees.

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903 Adjusting bioinformatic pipelines and analyses

904 Bioinformatics and population genetic analyses also need to be sensitive to the possibility of 905 cryptic taxa (see also Section 2.1 and Textbox 1). Missing data thresholds and other data 906 quality filters are employed as standard practice on individuals and loci. However, sensitivity 907 of different missing data thresholds to test taxon assignment and hierarchy hypotheses are 908 not often mentioned. The more divergent groups are, the fewer sites they will share and thus 909 blanket missing data thresholds on heterogeneous samples may bias patterns. Applying 910 different missing data filters and subsetting datasets by selecting an even representation of 911 predetermined groups (from initial structure analyses) or isolating certain groups can help in 912 determining if the assignment and hierarchy of groups is stable and robust to the filters 913 selected (Pritchard et al., 2000; McVean, 2009; Puechmaille, 2016). Intermediate or admixed 914 individuals may appear as hybrid individuals, but the causes of these patterns are many, e.g., 915 unexplained variance due to geographic structure, under-sampled taxa, admixture with 916 unsampled taxa, or higher levels of missing data for some individuals. Thus, we suggest formal 917 hybrid tests are employed for clarification (e.g., NewHybrids). Investigators should be 918 transparent regarding how biases or decisions were handled when reporting groupings. We 919 suggest following advice from (Meirmans, 2015) by always reporting multiple K values when 920 using assignment methods, as clustering analyses represent a heuristic approach that is open 921 to interpretation for all biologically-sensible K values, even if an optimal K-value is selected 922 by the user-defined summary statistic. Similarly, PCA results should present the percent of 923 variation explained and include multiple axes. Ultimately, we hope that the guidelines 924 presented here can be used as a framework to detect coral cryptic taxa in future population 925 genomic investigations.

926

927 In the absence of genotypes....

928 While population geneticists are the target audience for our recommendations, any coral 929 biologist whose data interpretations could be affected by cryptic species would do well to 930 incorporate genotyping in their project planning. For instance, genotyping is largely 931 overlooked in experiments (Textbox 2). For experimental work, we propose that future 932 studies should: 1) where possible, include larger sample sizes (n > 30) to screen for cryptic 933 genetic population structure (this will ensure downstream comparisons in individual 934 phenotypic differences are not confounded by cryptic speciation); 2) follow guidelines from 935 2.1 to recognise cryptic species; 3) report initial data checking methods and results (e.g., 936 screening population structure) in publications and reports (e.g., in supplementary items) to

937 938	assist the interpretation of individual- and population-level differences; and 4) clarify definitions and conventions for terms such as "cryptic species" and establish common
939 940	terminology.
941	Acknowledgements
942	We thank S. Starko for photos of <i>Porites</i> sp.
943	
944	Data, scripts, code, and supplementary information availability
945	Data and supplementary information are available linked to this pdf at EcoEvoRxiv.
946	Conflict of interest disclosure
947 948	The authors declare that they have no financial conflicts of interest in relation to the content of the article.
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Appendix

1439 *Structured review for population genomic studies*

An initial search of Web of Science on 21/10/2022, using the search terms "(TI=(coral) OR TI=(scleractinia*) NOT TI=(fish)) AND (AB=(rad*) OR ALL=(snp*)). The search returned 802 studies. Titles and abstracts were skimmed, and many irrelevant papers were excluded. Any manuscript that appeared to contain population genomic data of scleractinians or octocorallians was then retained for manual inspection.

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1447 Each manuscript was read and evaluated by two people to ensure that the genomic data: i) 1448 pertained to the cnidarian coral host, ii) used many loci on a genomic scale (i.e., not 1449 microsatellites, not metabarcoding), iii) surveyed two or more sites, and iv) presented results 1450 that included ordination based on individual genotypes (principal components analysis, 1451 principal coordinates analysis or multidimensional scaling) and/or unsupervised model based 1452 clustering test (such as ADMIXTURE or STRUCTURE). We did not consider papers that only 1453 reported discriminant analysis of principal components, as DAPC finds the eigenvectors that 1454 best differentiate prespecified groups, whereas PCA, PCOA and MDS find eigenvalues that 1455 best capture total diversity regardless of group membership (see (Thia, 2022) for further 1456 discussion). If these four criteria were not met, the paper was excluded. For each retained 1457 paper, the two evaluators independently extracted various attributes from the study (see raw 1458 data) and reconciled any discrepancies between their scoring through discussion. Despite 1459 attempting to undertake a rigorous and inclusive search, the authors noticed that several 1460 suitable manuscripts were missing and therefore on July 18, 2023 we ran an ad hoc search in 1461 WOS based on authors that are known to be publishing on population genomics of corals 1462 (namely Barshis, DJ; Baums, IB; Bay, LK; Bongaerts, P; Cooke, I; Matz, MV; Palumbi, SR; 1463 Richards, ZT, Underwood, JN, van Oppen, MJH) and also repeated the the above search with 1464 exactly the same criteria for articles published since 21/10/2022. These new papers were 1465 evaluated as above.

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The initial search yielded 853 papers that were reduced to 27 after skimming titles and abstracts, and 25 were found to be suitable for data extraction. The ad hoc search (combining authors and new publications) initially identified 897 papers in WOS that were reduced to 16 papers once titles and abstracts were skimmed, and only 11 were suitable for data extraction.

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1472 *Heat stress studies*

1473The CORDAP database (Ortiz *et al.*, 2023) was downloaded and searched on 11/09/2023,1474searching for any records which used genome-wide data of the coral host (i.e., coral SNP data)

1475 and those which conducted clustering analyses (e.g., PCA, ADMIXTURE, or STRUCTURE). First,

records were filtered based on whether the database columns "Host genotype", 1476 "Symbiodiniaceae genotype", and "Microbiome genotype" were listed as "TRUE". Next, the 1477 1478 columns "Symbiodiniaceae_genotyping_approach", "Host_genotyping_approach" and 1479 "Microbiome genotyping approach" were interrogated and only records where the 1480 genotyping method was listed as RADseq, WGS or RNAseq were kept for further checks. The 1481 titles and abstracts of the remaining records were skimmed, and only records for which at 1482 least host genotyping was performed were included. The remaining records were read and 1483 evaluated to ensure that they i) pertained to the cnidarian coral host, ii) used many loci on a 1484 genomic scale (i.e., not microsatellites, not metabarcoding, but single nucleotide 1485 polymorphisms), iii) presented results that included ordination based on individual genotypes 1486 (principal components analysis, principal coordinates analysis or multidimensional scaling) and/or an assignment test (such as ADMIXTURE or STRUCTURE). Additionally, the number of 1487 1488 cryptic species assigned in each paper by the original authors was noted, as well as the 1489 evaluators' interpretation of the number of cryptic species based on the plots and analyses 1490 included (where possible).

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1492 The original database consisted of 562 records, many of which did not include host, 1493 Symbiodiniacae or microbiome genotyping. The initial filtering of the database for records 1494 that included some aspect of host genotyping yielded 223 results. Applying criteria i & ii by 1495 skimming abstracts and methods reduced the studies to 17 records. Of these 17 studies, nine 1496 studies included either ordination or model-based clustering analyses (e.g., PCA, 1497 ADMIXTURE). Of these, three (Rose et al., 2017; Ruiz-Jones & Palumbi, 2017; Rose et al., 2021) 1498 the criteria in Section 2.1. The remaining six studies were either ambiguous or showed no 1499 clear evidence for cryptic taxa.