Expanding the concept of social behavior to interspecific interactions

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Abstract
In pretty much any species, an individual’s survival and reproduction depends crucially on the outcome of interactions with other individuals. Key interactions may take place between individuals of the same species but also between individuals belonging to different species. However, the most accepted definition of social behavior only considers interactions between conspecifics. Here, we argue that the distinction between intra- and interspecific interactions is largely artificial and hinders the integration of the historically separately developed concepts. At the ultimate level, given that the ecological landscape of organisms is composed both by interactions with conspecifics and with heterospecifics, and both types of interactions may have evolutionary consequences. Although intraspecific interactions usually have a higher impact in fitness because in most species interactions relevant for reproduction (matting, parenting) exclusively involve conspecifics, and interactions relevant for survival are more probable between conspecifics because they share the same ecological niche, hence competing for the same resources (e.g., food, shelter), there are notable exceptions in both fitness components (e.g., heterospecific mating in parthenogenic all-female species; heterospecific brood parasitism; heterospecific aggression in sympatric species that compete for shared resources). At the proximate level, behaviors and cognitive decision-making rules used to interact with other organisms may be shared between intra- and interspecific interactions, and the mechanistic differences between conspecific social behaviors used in distinct functional domains, such as mating, aggression, or parenting, can be more expressive than those found within the same functional domain between conspecific and heterospecific behavior. Therefore, there are neither fundamental conceptual (ultimate) reasons, nor key differences in mechanisms underlying behaviors involved in conspecific vs. heterospecific interactions that support the exclusion of interspecific interactions from the conceptual framework of social behavior.

Keywords
heterospecific aggression, heterospecific mating, heterospecific alloparental care, heterospecific cooperation, social interactions
1 | INTRODUCTION

The Darwinian fitness of most animals depends to a large extent on the outcome of interactions with other individuals in multiple functional domains, including resource defense (e.g., food, shelter), mating and parenting, predation, and/or predator evasion. Typically, interactions in some of these functional domains are more frequent between conspecifics (e.g., resource defense, mating, parenting) and in other domains between heterospecifics (e.g., predator–prey interactions). Historically, the animal behavior literature exclusively considers conspecific interactions as social (Székely et al., 2010), without providing any alternative term for behavior in heterospecific interactions. However, this dichotomy between intra- and interspecific interactions is largely artificial and hinders the integration of the historically separately developed concepts in both fields. In fact, the social landscape of most species in the wild includes both conspecifics and heterospecifics and the integration of both in a common conceptual framework for the study of social interactions would be beneficial. Take the example of sympatric cichlid species of the great African lakes or South American crater Lakes with overlapping breeding seasons. Multiple species, which vary in their phylogenetic relatedness, share breeding grounds densely occupied with multiple breeding heterospecific and conspecific individuals that actively defend their breeding territories and/or offspring (Barlow, 2000; McKay, 1977). In such a scenario, territorial aggression can be directed towards (1) conspecific and heterospecific resource competitors (i.e., competing for breeding territories); (2) conspecific and heterospecific brood predators; and (3) conspecific sexual competitors (i.e., competing for mates; Lehtonen et al., 2010, 2012; McKay, 1977). Therefore, an understanding of the biology of social behavior at both ultimate (ecology, evolution) and proximate (development, physiology) in this scenario needs the integration of both conspecific and heterospecific interactions in the same conceptual framework.

The main aim of this article is to provide explicit examples in favor of a less restrictive definition of social behavior. We do not attempt to provide a new definition that includes all interspecific interaction as well, but propose that a more holistic view on interactions may improve our understanding of both ultimate and proximate causes of social behavior. Indeed, both conspecific and heterospecific interactions are characterized by the need of fine adjustment of behavioral expression as a function of the behavior of the interactant agent. Hence, they may pose similar sensory and cognitive problems that need the evolution of adaptive flexible responses to another living organism. As it stands, a meta-analysis of species discrimination in animals has shown that it is not universal and that it is dependent on the benefits and costs of responding to a conspecific vs. heterospecific that is specific to the circumstances of a given species (Ord et al., 2011). This shows that the distinction between conspecifics and heterospecifics is to some extent a human construct.

2 | A FUNCTIONAL PERSPECTIVE

From a functional perspective, Hamilton (1964) classified a behavior as social if it has the consequence $-$c on the actor and the consequence $b$ on the recipient. The lifetime consequences of two-player social interactions can be captured in a 2x2 matrix. $+/+$ has been termed either mutually beneficial behavior or cooperation, $+/-$ is defined as biological altruism, $+/-$ is defined as selfishness, and $-/-$ is defined as spite (Hamilton, 1964; Lehmann & Keller, 2006; West et al., 2007). The same 2 x 2 matrix can be used to describe the lifetime fitness consequences of interspecific interactions (Begon et al., 1990). A $+/+$ outcome has been termed mutualism, and a $+/-$ or $-/-$ outcome of interactions characterizes parasite–host and predator–prey interactions. Finally, a mutually negative effect on each other ($-/-$) has been termed competition. Hosts may be manipulated to behave altruistically ($-/+)$ toward parasites (Bakker et al., 2017); such behavior is under negative selection in hosts (and maintained due to the positive selection on manipulative strategies in parasites). Selection is expected to operate on individual strategies that differ in payoffs no matter whether an interaction partner is a conspecific or a heterospecific. Nevertheless, the strength of selection will differ depending on whether interactions are intra- or interspecific (Bergstrom et al., 2003). This is because the payoff consequences of interactions are computed in a single matrix for within-species interactions (individuals of the same gene pool are in direct competition), while for interspecific interactions, each individual competes with members of its own gene pool rather than with the interaction partner. In a football analogy, a between-species interaction is a 3-point game (if I win, I gain 3 points, and if I lose, I gain 0 points), whereas a within-species interaction is a 6-point game (the 3 points I lose are gained by a competitor). Regarding contexts, it is of course true that some types of interactions are almost exclusively observed among conspecifics (maternal, parental care) and others between heterospecifics (predation). However, as we describe below, there are various exceptions that highlight the artificiality of the conspecific/heterospecific boundaries in the definition of social behavior.

2.1 | Heterospecific aggression

Direct competitive interactions are common between heterospecific organisms, and aggressive behaviors are often used in such interactions (Peiman & Robinson, 2010). Heterospecific aggression is often related to competition for resource use when there is an overlap in the ecological niche (e.g., food: cutthroat trout, Oncorhynchus clarkii vs. coho salmon, O. kisutch, Sabo & Pauley, 1997; shelter (dens): red foxes, Vulpes vulpes vs. arctic foxes, Alopex lagopus, Tannerfeldt et al., 2002), and can be as intense and costly as conspecific aggression, therefore impacting Darwinian fitness with putative ecological and evolutionary consequences (Grether et al., 2013; Peiman & Robinson, 2010).
At the ecological level, heterospecific aggression results in niche partitioning through the regulation of the strength of the species interference leading to phenotypic displacement in traits that affect heterospecific interactions (e.g., habitat preferences, fighting ability, competitor recognition; Grether et al., 2009, 2013). In this scope it is interesting to note that the expression of heterospecific aggression can be flexible, with more aggressive responses directed toward heterospecific opponents that pose a higher threat. For example, the South American arrow cichlid (*Amphilophus zalisus*) adjusts its territorial defense toward heterospecific intruders according to their breeding status (breeding individuals receive lower level of aggression than non-breeders; Thus, heterospecific interactions should also be considered as a source of selection pressure shaping plasticity of aggressive behavior (Lehtonen et al., 2010).

Heterospecific aggression can be adaptive, if it increases fitness through the acquisition of limited resources, and evolve independently of intraspecific aggression. Peiman and Robinson (2010), using a simulated intrusion test, found that heterospecific aggression was greater in sympathy than in allopatry for species where resource overlap occurred and that it was not correlated with intraspecific aggression. Moreover, they also found that aggression was higher with increased resource overlap. Interestingly, there is also evidence for heritable variation in heterospecific aggression in brook sticklebacks (*Culnea inconstantis*), where aggressive responses toward heterospecific competitors (nine-spine stickleback, *Pungitius pungitius*) are higher in sympatric than in allopatric populations (Peiman & Robinson, 2007), which further supports its evolvability. Heterospecific aggression may even influence intraspecific evolutionary dynamics. In *Amphilophus sagittae*, a South American cichlid, a color polymorphism occurs with gold and dark morphs present in the population.
These color morphs elicit differential heterospecific aggression from their territorial neighbors of another cichlid species (*Hypostomus nicaraguensis*), which are more aggressive toward gold than dark morphs (Figure 1a). Thus, heterospecific aggression potentially contributes to the evolutionary equilibrium among these color morphs (Lehtonen et al., 2015).

Together these results support an adaptive role for heterospecific aggression. Apart from its role in competition for resources, heterospecific aggression may also act as a way to avoid hybridization (see section below), and selection for increased heterospecific aggression may also be associated with reduced costs of wasted reproductive effort on heterospecifics (Peiman & Robinson, 2010).

### 2.2 Heterospecific reproductive behavior: mating and parenting

Even social interactions that occur in the scope of reproduction, intuitively the most obvious reason for a distinction between intra- and interspecific interactions, may occur between species. A classic example of heterospecific mating involves Amazon mollies (*Poecilia formosa*), an all-female unisexual fish species that reproduces through gynogenesis, which is a form of parthenogenesis that requires the presence of sperm of a male from closely related species to trigger embryogenesis without the genetic contribution of sperm DNA for the zygote (Schlupp, 2005). In this system, selection for heterospecific mating occurred in both sexes, since males of the closely related species benefit from mating with heterospecific females because of mate-choice copying decision-rules of their conspecific females (Sclupp et al., 1994; Figure 1b). Several maladaptive cases of interspecific sexual interactions have been described in the literature from a wide range of species (Groning and Hochkirch, 2008). Nevertheless, these other cases of reproductive interference may have ecological and/or evolutionary consequences, leading to the species exclusion or character displacement, and are particularly relevant in the scope of invasive species because endemic species can be threatened by sexual interactions with the new introduced species (Groning and Hochkirch, 2008). In the case of rare successful hybridization, heterospecific mating may facilitate rapid evolutionary radiations, as recently described for Lake Victoria and Lake Malawi cichlids, where ancestral hybridization provided the genetic variation that subsequently became recombined and gave rise to a flock of new species (Meier et al., 2017, 2019; Svardal et al., 2020).

Within the scope of reproductive behaviors, also parental care interactions can be observed between heterospecifics. The occurrence of brood care parasitism, in which organisms rely on others from a different species to raise their offspring, has been described in birds, insects, and one species of fish (Davies, 2000; Polacik et al., 2019; Sato, 1986). Typically, the brood parasites manipulate the behavior of the heterospecific host using brood mimicry, inducing it to express alloparental behavior toward their young (Davies, 2000; Stevens, 2013; Figure 1c). By adopting this behavior, brood parasites avoid the costs associated with parental care, such as nest building and time and energy investment in feeding the young, allowing them to increase their investment in foraging and reproduction with associated fitness benefits. Brood parasitism typically results in evolutionary arms races between brood parasites and their hosts leading to the coevolution of both species. In birds, where obligatory interspecific brood parasitism is more common, it evolved independently seven times and occurs in approximately 1% of species, and different stages of coevolution can be observed (Davies, 2000; Kilner & Langmore, 2011; Sorenson & Payne, 2005).

As a result of the arms race described above, both brood parasites and their hosts have evolved a set of behaviors and cognitive abilities that are clearly of the social domain. For example, female brood parasites are able to track the nesting progress of hosts and to choose nests or to time their egg laying accordingly to ensure optimal development of their young (Davies, 2011). Brood parasite females also behave secretly to approach the host nests without being noticed and evolved rapid egg laying, and in some species (e.g., great spotted cuckoos, *Clamator glandarius*), males may lure the hosts away from the nest, allowing the female the opportunity for egg laying (Davies, 2000, 2011). In response, hosts have evolved behavioral responses that reduce parasitism, such as mobbing behavior toward brood parasites, nesting in more concealed locations, and adopting defensive nest architectures (e.g., narrow entrance tubes; Davies, 2000, 2011). Also, parasite chicks evolved behaviors to increase the parental care provided by the hosts that range from ejecting host eggs/chicks from the nest in order to avoid competition of food provisioning, to increased begging abilities to claim an unfair share of food when competing with host chicks (Davies, 2000, 2011).

### 2.3 Heterospecific cooperation

There are numerous cases in which individuals belonging to different species cooperate with each other. As it stands, such mutualisms provide the foundation for the world as we know it, starting from endosymbionts allowing the evolution of eukaryotic cells to the diversity of flowering plants depending on pollinators and root symbionts (Bronstein, 1994, 2015).

A particularly interesting example of interspecific cooperation that is explicit on communication and behavioral coordination is the cooperative hunting between groupers of the genus *Plectropomus* and various partner species (Bshary et al., 2006; Vail et al., 2013, 2014). The groupers use strong acceleration to try to capture prey, which in turn typically tries to escape into crevices that are inaccessible for the groupers. The groupers’ partner species can access such hiding prey: Moray eels move through crevices, octopus insert their arms, and Napoleon wrasse may crush the reef structure. Groupers use two signals to elicit joint hunting. A rapid shaking of the head while in horizontal position serves to initiate a hunt with moray eels, while a headstand shake serves to attract partners to a crevice where prey is hiding (Bshary et al., 2006). The latter signal fulfills five key criteria for referential gestures (Vail et al., 2013). According to field observations in the Red Sea, following a grouper in hunting
mode for 120 min yields a 70–80% chance of observing a signaling event, and a single collaborative hunting event with a moray eel can last up to 93 min. The understanding of groupers when they need a partner, and their ability to choose between partners of different quality, rivals the abilities of chimpanzees in similar laboratory experiments (Melis et al., 2006; Vail et al., 2014).

As another example, the genus Labroides evolved from coralivorous ancestors into the most sophisticated cleaner fish on coral reefs (Cowman et al., 2009; Figure 1d). Their interactions with "client" fish have been used to demonstrate various key concepts developed to explain interspecific cooperation. Cleaners prefer to eat client mucus rather than ectoparasites (Grutter & Bshary, 2003), which creates conflicts. In order to obtain a good cleaning service, clients use punishment, partner switching, and assess the cleaners' interactions with other clients to then seek or avoid a cleaner (Bshary & Grutter, 2005, 2006). In turn, cleaners are more cooperative if observed by bystanders (Pinto et al., 2011) and reconcile with clients that responded to a cheating act with aggression (Bshary & Würth, 2001). Cleaners track their clients' behavior and act, for example, on clients' likelihood to swim off if not serviced immediately (Triki et al., 2019). Interactions with clients also affect conspecific interactions, with males punishing females for cheating joint clients, adjusting the severity of punishment to the stakes (Raithani et al., 2010, 2012). As cleaners obtain almost 100% of the food from about 2000 cleaning interactions per day (Côté, 2000; Grutter, 1995), their performance in these interspecific social interactions is obviously under strong selection.

### 2.4 | Heterospecific social groupings

Heterospecific social groups, such as mixed-species fish shoals, bird flocks, and mammalian herds, are common in nature across a range of aquatic and terrestrial ecosystems (Harrison & Whitehouse, 2011; Lukoschek & McCormick, 2000; Paijmans et al., 2019; Sridhar et al., 2013; Stensland et al., 2003). Adaptive explanations for heterospecific grouping have claimed the similar fitness benefits to those proposed for single-species groupings, including reduced predation risk, due to collective vigilance and/or predator confusion effect, enhanced foraging efficiency, and more efficient movement in the environment, due to increased aerodynamics or hydrodynamics (Goodale et al., 2020; Paijmans et al., 2019). These benefits have to out weight the costs associated with social grouping, namely increased competition for food and increased detection by predators. However, differences between species in any of these components will give rise to species-specific cost-benefit trade-offs in heterospecific grouping. For example, feeding competition is lower between species than within species, and niche overlap in resource exploitation will determine interspecific competition. Similarly, species with different sensory modalities and/or perceptual abilities will complement each other's efficiency in the detection of shared predators (e.g., Meise et al., 2019; Schmitt et al., 2014; Stears et al., 2020). In principle, these conditions favor the formation of mixed-species groups that optimize each other's grouping benefits (Terborgh, 1990). Recently, a model has been proposed which explains patterns of species associations in mixed-species groups based on net benefits to individuals resulting from the multiple trade-offs of differences between species, as well as from the trade-off arising from overall group size (Paijmans et al., 2019). Hence, although the literature on grouping behavior has mainly focused on single-species groups (for a review, see Ward & Webster, 2016), the adaptive value of grouping behavior is better understood if heterospecific grouping is also considered.

### 2.5 | Intraspecific predation

Although predation is commonly seen as an interspecific interaction, intraspecific predation (aka cannibalism) is widespread in the animal kingdom, from invertebrates (e.g., arachnids, insects) to all vertebrate classes (Elgar & Crespi, 1992; Fox, 1975; Polis, 1981). It has been shown to play a significant role in population dynamics and in the competitive relationships of organisms, since it can reduce population size and alter age structure in face of shortage of resources or high population density (e.g., Van Buskirk, 1989; Fincke, 1994; Lloyd, 1968; Park et al., 1965; Wagner & Wise, 1996). Cannibalism, including filial cannibalism, is nutritionally beneficial, increasing survival, reducing developmental time and increasing fecundity, hence having an effect on the fitness of cannibals (FitzGerald, 1992; Fox, 1975; Manica, 2002; Matsumoto et al., 2018; Sonleitner & Guthrie, 1991; Spence & Carcamo, 1991; Tschinkel, 1993). Such fitness benefits may provide a selective advantage that leads to the evolvability of cannibalism (e.g., Eickwort, 1973; Stevens, 1989). Filial cannibalism, when organisms eat one's own offspring, is a sub-type of intraspecific predation that also has clear fitness implications. It is particularly common in fish where it has been regarded as an adaptive trade-off between present (i.e., offspring survival) and future (i.e., feeding in order to increase nutritional status) reproductive success (FitzGerald, 1992; Manica, 2002). However, some recent evidence suggests that, at least in some cases, filial cannibalism does not have nutritional benefits but is rather a way to end the parental phase and restart courtship, suggesting that this is infanticide rather than cannibalism (e.g., Matsumoto et al., 2018).

In conclusion, intraspecific predatory behavior may have fitness implications that are analogous to those of other intraspecific social processes such as dispersal, spacing behavior, or dominance hierarchies. Its effects can hence rival the effects of interspecific predation, blurring the distinction between social behavior and predator-prey interactions.

### 3 | A MECHANISTIC PERSPECTIVE

Conspecific and heterospecific interactions impose similar selective pressures on sensory and cognitive mechanisms that regulate social behavior. In order to be successful in both types of interactions, individuals need to be able to recognize relevant interactants.
and discriminate them into meaningful categories (e.g., potential mate, competitor, cooperator/mutualist, predator) and to use adaptive decision-making rules in order to adjust a flexible behavioral response accordingly. A key question is to what extent the functional categories of behavior observed in intraspecific interactions (i.e., aggression, mating, parenting) are homologous (i.e., share the same proximate mechanisms) to those observed in heterospecific interactions. In this respect, two scenarios are possible for the evolution of the proximate mechanisms (genes, neural circuits, cognitive processes) underlying a behavior used in both conspecific and heterospecific interactions: (1) They evolved first in a conspecific context and were later co-opted to be used in interspecific interactions, when species first evolved in allopatry and later became in contact, or (2) a shared mechanism evolved simultaneously toward both conspecifics and heterospecifics, when species evolved sympatrically.

3.1 | Social recognition and discrimination

In order to respond adequately to social interactions, animals have to discriminate interactants into functional categories. A major recognition category that has been considered in evolutionary behavioral ecology is species recognition, given its importance for the differential expression and response to mating behaviors among closely related species, hence playing a key role in restricting gene flow and consequently in behavioral isolation and speciation. Hence, animals produce a variety of species-specific stereotypic behaviors (signals) in different sensory modalities (e.g., acoustic, chemical, visual) to communicate with each other in the scope of intraspecific social interactions, and their sensory/perceptual mechanisms are tuned to respond to these signals (aka conspecific recognition; e.g., pheromones in insects and mammals, Coyne et al., 1994, Brennan & Kendrick, 2006; Lebreton et al., 2017; body coloration in fish, Seehausen & van Alphen, 1998, Couldridge & Alexander, 2002, Williams & Mendelson, 2011; vocalizations in amphibians and birds, Gerhardt, 1994, Toledo et al., 2015, Louder et al., 2019). Therefore, animals are usually able to discriminate between conspecific and heterospecific social signals and these conspecific signals are categorically represented in the nervous system. Thus, one possibility for the occurrence of heterospecific interactions is that the same cognitive and sensory systems may be used in signal recognition toward heterospecifics of closely related species that phenotypically resemble conspecifics (Ord et al., 2011), as supported by a literature review that found heterospecific aggression to be more pronounced among congeneric heterospecific individuals, than toward more distantly related genera (Peiman & Robinson, 2010).

Perceptual filters for conspecific signals have been well studied in acoustic communication in birds where conspecific vocalizations elicit differential behavioral responses that are paralleled by heightened neural activity in distinct nuclei of the avian auditory forebrain (Louder et al., 2019), often based on innate predispositions toward conspecific vocalizations (Araki et al., 2016; Hauber et al., 2013; Long et al., 2002; Whaling et al., 1997). Experience with heterospecific vocalizations during critical periods in development (e.g., zebra finch nestlings raised with Bengalese finch tutors) may lead to a learned preference and expression of heterospecific song, which is accompanied by a specialization of auditory cortex neurons for the tutor song, through tuning of spectrotemporal receptive fields of the neurons (Moore & Woolley, 2019) and a differential transcriptional responses suggestive of release of inhibition toward the heterospecific stimulus (Louder et al., 2018). While such tutor experiments described above represent an artificial situation where developmental mechanisms have not been under selection, a natural situation where nestlings are raised by heterospecific tutors is offered by brood parasites. In the pin-tailed whydah (Vidua macroura), an obligatory brood parasite, adults still recognize and express a behavioral preference for their own host parent species vocalizations, paralleled by an increased neural response within the auditory forebrain homologous to the one observed in phylogenetic related non-parasitic species that are raised by their biological parents (Louder et al., 2016). Studies in another obligatory brood parasite, the brown-headed cowbird (Molothrus ater), suggest that this is apparently made possible through an inborn predisposition to a password-like conspecific chatter call, which not only identifies conspecifics but triggers social learning of other conspecific traits, including song (Hauber et al., 2001). It has recently been shown that the chatter call induces increased expression of the immediate early gene ZENK, used as a marker of neural activity, in the caudal medial nidopallium (NCM) in juveniles, whereas exposure to either conspecific or heterospecific songs elicits higher gene expression in an experience-dependent manner (Lynch et al., 2017). Thus, in parasitic cowbirds the NCM has been evolutionarily co-opted to selectively recognize a species-specific password chatter, allowing juvenile cowbirds raised by different host species to avoid imprinting on heterospecifics, while recently experienced song is still represented in the caudal medial mesopallium (CMM).

While species identity is of great biological relevance in a reproductive context, it is expected that selection may have acted irrespective of species boundaries to promote the recognition of relevant interactant categories in other ecological contexts, like competition for resources. Indeed, it has been shown in experimental settings that species that live in sympathy with heterospecific competitors display higher aggressive responses toward them than allopatric ones, suggesting that heterospecific competitor recognition was enhanced in sympathy (Grether et al., 2009). Furthermore, differential aggressive responses have been reported toward heterospecific members of the same competitor species that pose different levels of threat, suggesting an ability to discriminate heterospecifics into different threat categories. For example, neotropical cichlid fish that share breeding grounds with heterospecifics are more aggressive toward heterospecific nonbreeding individuals, that pose a threat to their offspring, than to heterospecific breeders, using body coloration as a recognition cue of heterospecific breeding status (Lehtonen et al., 2010, 2015). Interestingly, when exposed to manipulated territorial intrusions individuals of allopatric populations do not show
an adjustment of their aggressive response in relation to heterospecific color markings that are associated with different levels of threat (Lehtonen et al., 2016), highlighting the importance of coevolution and learning in the categorization of heterospecific interactants. Interestingly, the same social signal can be differentially perceived by the different sexes of another species. In tungará frogs (Physalaemus pustulosus), the calls of the heterospecific (Physalaemus petersi) elicit a vocal response in males but fail to elicit a mating phonotaxis response in females (Bernal et al., 2007; Ryan & Rand, 1995), and this sex difference in the behavioral response is mirrored by the neural response. Indeed, the activity of the midbrain auditory region, as measured by the expression of an immediate early gene (egr-1), is elevated in males in response to both conspecific and heterospecific calls, whereas in females, it is only elevated in response to conspecific calls (Hoke et al., 2008). Therefore, it seems that the perceptual system of males and females can be shaped by different selection pressures for the discrimination or not of relevant social categories: for males, both conspecific and heterospecific males represent competitors for mating territories, whereas for females, only the conspecific call is relevant for sexual selection.

3.2 Social learning and memory

The ability to successfully interact with other behavioral agents, either conspecific or heterospecific, relies to a large extent on flexible behavioral responses that require learning and memory (Taborsky and Oliveira, 2012). In the scope of social interactions, individuals can learn from observing others rather than by costly trial-and-error strategies (aka social learning; Dall et al., 2005; Danchin et al., 2004; Heyes, 1994). This form of learning is widespread in animals, from invertebrates to mammals (e.g., Brown & Laland, 2003; Galef & Giraldeau, 2001; Galef & Laland, 2005; Leadbeater & Chittka, 2007). Conspecifics have been traditionally seen as the primary source of information for social learning (Hoppitt & Laland, 2013). However, heterospecifics can be as valuable as information sources as long as they share the same ecological needs (Avarguès-Weber et al., 2013). In fact, heterospecific social learning has been demonstrated for food location, habitat choice, and predator evasion across a wide range of species (Table 1 for selected examples). A peculiar case of social learning that blends intraspecific and heterospecific aspects involves juvenile cleaners learning from adults how to interact with various client types (Truskanov et al., 2020). From the many examples available in the literature, three patterns are worth noting. First, the diversity of examples shows that the occurrence of heterospecific social learning is taxonomically as widespread as conspecific social learning, ranging from invertebrates to mammals, hence not suggesting more complex cognitive demands. Moreover, social information transfer may occur across widely different taxa. For example, Augabries flat lizard (Platysaurus capensis), which feed on ripe figs, use birds, that also feed on figs, flocking in fig trees for food location (Whiting and Greeff, 1999), whereas yellow-casqued hornbills (Ceratogymna elata), which are vulnerable to predation by crowned eagles (Stephanoaetus coronatus) but are not preyed on by leopards (Panthera pardus), respond to sympatric Diana monkey eagle alarm calls but not to the leopard alarm call (Rainey et al., 2004). Secondly, individuals of other species can be as good demonstrators as conspecifics (e.g., bumblebees: Dawson & Chittka, 2012; birds: May & Reboreda, 2005; lizards: Damas-Moreira et al., 2018), and more efficient than non-social cues, suggesting that organisms may have a predisposition to social cues when learning about food sources (Dawson & Chittka, 2012). Finally, the fact that phylogenetically closely related species show marked differences in the use of heterospecific information (e.g., Coolen et al., 2003; Magrath et al., 2009; Magrath and Bennett, 2012; Kitchen et al., 2010; Avarguès-Weber et al., 2013) suggests that heterospecific social learning only emerges if it is under selection (e.g., high predation risk).

From a proximate causal perspective, heterospecific social learning may rely on relatively simple forms of classical conditioning as in learning to associate alarm calls with a predator appearance (Avarguès-Weber et al., 2013), but also on second-order conditioning based on indirect associations (Dawson et al., 2013). This is a potential explanation for social learning about food location, where the indirect association between a location or cue (second-order conditioned stimulus) and the reward (food) is established by the prior association of foraging conspecifics/heterospecifics (first-order conditioned stimulus) with the food, as in the case of nine-spined sticklebacks (Pungitius pungitius) learning a food location after observing either conspecifics or three-spined sticklebacks (Gasterosteus aculeatus) eating in the same spatial position (Coolen et al., 2003).

Alternatively, perceptual resemblance of conspecific and heterospecific cues may trigger appropriate responses, as it has been shown for bird alarm calls and for bat distress calls (de Kort and Cate, 2001; Fallow et al., 2011; Russ et al., 2004), and perceptual resemblance should share the same neural mechanisms. In accordance with this prediction, it has been shown that playbacks of black-capped chickadee mobbing calls, which are used to recruit conspecifics and other avian species to mob perched predators (Ficken & Witkin, 1977), elicited similar patterns of forebrain secondary auditory cortex activation (i.e., CMM and NCM) as measured by ZENK expression, in both conspecific and heterospecific (i.e., mountain chickadee, Poecile gambeli) listeners. Interestingly, black-capped chickadees produce referential mobbing calls that convey the degree of threat posed by different predators (Templeton et al., 2005), and referential mobbing calls signaling a low-threat predator elicited lower activations of the auditory cortex in both chickadee species (Avey et al., 2011). Therefore, the level of activation of the secondary auditory cortex seems to reflect the level of assessed threat irrespective of source of information, hence suggesting a shared neural substrate to encode the socially transmitted information about predation risk. Spatial cognition is another key element for successful social interactions. Within the brain, spatial cognition has been related
### TABLE 1  Selected examples of heterospecific social learning

<table>
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<tr>
<th>Ecological domain</th>
<th>Observer species</th>
<th>Demonstrator species</th>
<th>Learning efficiency (referent to a conspecific demonstrator)</th>
<th>Author(s)</th>
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</thead>
<tbody>
<tr>
<td>Food location</td>
<td>Bumblebee (<em>Bombus terrestris</em>)</td>
<td>Honey bees (<em>Apis mellifera</em>)</td>
<td>(H=C) non-social cue</td>
<td>Dawson and Chittka (2012)</td>
</tr>
<tr>
<td>Food location</td>
<td>Italian wall lizard <em>Podarcis sicula</em></td>
<td>Bocage’s wall lizard <em>Podarcis bocagei</em></td>
<td>H=C</td>
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</tr>
<tr>
<td>Food location</td>
<td>Augabries flat lizard <em>Platysaurus capensis</em></td>
<td>Bird flocks of red-eyed bulbuls (<em>Pycognotus nigricans</em>) and pale-winged starlings (<em>Onychognathus nabolourop</em>)</td>
<td>n.a.</td>
<td>Whiting and Greeff (1999)</td>
</tr>
<tr>
<td>Habitat (nest-site) selection</td>
<td>collared flycatchers (<em>Ficedula albicollis</em>) / pied flycatchers (<em>F. hypoleuca</em>)</td>
<td>Great tits (<em>Parus major</em>) / blue tits (<em>P. caeruleus</em>)</td>
<td>n.a.</td>
<td>Seppänen and Forsman (2007)</td>
</tr>
</tbody>
</table>

Abbreviations: and n.a., not available; C, Conspecific; H, Heterospecific.
to the hippocampus in mammals and to homologous structures (i.e., medial pallium) in other vertebrates (Hartley et al., 2013). In rodents, the hippocampal complex (i.e., hippocampus and entorhinal cortex) contains a population of neurons (aka place cells) that respond to specific locations in the environment and another cell population (aka grid cells) which fires when an animal enters an environment with geometrically patterned locations hence providing a spatial cognitive map of the environment (Hafting et al., 2005; Hartley et al., 2013; O’Keefe, 1979). This map is also capable of processing other spatially representable information, hence providing a spatially based system (aka memory map) for encoding declarative memories (Eichenbaum, 2017; Jeffery, 2018), which is relevant for social memory. Direct evidence of the involvement of the hippocampus in mapping the social landscape has been recently demonstrated in bats, where specific place cells map the presence of other bats in the environment (Omer et al., 2018). Also in rats, the cornu ammonis 2 (CA2) area of the hippocampus has been shown to use place fields to encode information about conspecifics (Alexander et al., 2016). Thus, the hippocampus may play a more relevant role in social cognition than originally thought.

In wild animals, hippocampus size has been shown to be evolutionarily shaped by conspecific interactions, namely the spatial range of mating interactions and social group size that may cause interspecific and/or intraspecific intersexual variation (e.g., primates: Todorov et al., 2019; voles (Rodentia: Microtus): Boonstra et al., 1993; Jacobs et al., 1990; desert kangaroo rats (Dipodomys spp.): Jacobs and Spencer, 1994; Langley, 1994; deer mice (Peromyscus maniculatus): Galea et al., 1996; giant panda (Ailuropoda melanoleuca) and Asian small-clawed otter (Aonyx cinereus): Perdue et al., 2011; intertidal blenniid fish species [i.e., shanny (Lipophrys pholis) and Azorean rock-pool blenny (Parablennius parvicornis)] (Costa et al., 2011)). Once again, parasitic brood parasites offer a great system to test for interspecific hippocampus size adjustments, since they need to search for and monitor host potential nests in which to lay their eggs, hence facing a selection pressure for heightened spatial cognition. Moreover, there is interspecific variation among brood parasites in the role of males in host nest search behavior, and there are non-parasitic closely related species that can be used as a phylogenetic reference of the ancestral state. As predicted, hippocampi size is larger in brood parasitic species when compared to closely related non-parasitic species (Reboreda et al., 1996; Sherry et al., 1993), and there is a sex difference with females having larger hippocampi in species where females are not assisted by the males in searching for host nests [i.e., brown-headed cowbird (Molothrus ater) and shiny cowbird (M. bonariensis)], but not in species where both sexes search for nest [i.e., screaming cowbird (M. rufoaxillaris); Sherry et al., 1993; Reboreda et al., 1996]. Moreover, the sex differences present during the breeding season in the shiny cowbird disappear during the nonbreeding season, when there are also no sex differences in spatial behavior (Clayton et al., 1997). The sex difference in hippocampi size in brown-headed cowbirds is explained by enhanced adult hippocampal neurogenesis in females, which increases in the post-breeding season, suggesting that old memories for host nest sites may be lost through hippocampal reorganization following breeding (Guigueno et al., 2016). In summary, heterospecific interactions seem to also impose a selection regime that leads to cognitive and neurobiological specializations, as observed for conspecific interactions.

### 3.3 | Social decision-making

Social decision-making requires the integration of multimodal sensory information about social stimuli with information stored in memory regarding previous social experience in order to produce an adaptive behavioral response to the perceived social environment.

An evolutionarily conserved brain social decision-making network (SDMN) has been proposed to regulate the expression of social behavior across vertebrates (O’Connell & Hofmann, 2011, 2012). This network integrates two previously described networks: the social behavior network and the mesolimbic reward circuit (Adinoff, 2004; Goodson, 2005; Newman, 1999), with the former regulating the expression of the social repertoire of the species and the latter providing a mechanism for the reinforcement of adaptive behavioral responses. Subsequently, there have been proposals to expand this network to include also brain regions related to the sensory integration and salience assessment of social information (e.g., insular cortex for sensory integration and anterior olfactory nucleus and prefrontal cortex for social salience assessment in rodents; Johnson et al., 2017; Rogers-Carter & Christianson, 2019). Irrespective of the specific structure of this network, two key features of this conceptual proposal are that (1) social information is processed in a network of reciprocally interconnected brain nodes emphasizing distributed, interregional information processing, such that the expression of social behavior (i.e., aggression, mating, parenting, cooperation) is better reflected by the overall profile of activation across the different loci in the network rather than by the activity of a single node (Johnson et al., 2017; Teles et al., 2015); (2) different combinations of activation across nodes and variation in the strength of the connections among them have the potential to generate unlimited variation in social behavior (Cardoso et al., 2015; Goodson & Kabelik, 2009); and (3) the nodes of the SDMN extensively express receptors for steroids and neuropeptides (O’Connell & Hofmann, 2011, 2012), thus allowing the modulation of the activity of this network by hormones and neuromodulators that promote the integration of organismal state in social decision-making (e.g., adjust expression of mating behavior to gonadal maturation state).

From a mechanistic perspective, our proposal of extending the concept of social behavior to heterospecific interactions is that the SDMN must have been evolutionarily co-opted to regulate also social decision-making toward heterospecific interactants. Therefore, one can generate two predictions: (1) Similar patterns of activation of the SDMN should be observed in response to social interactions with conspecifics vs. heterospecifics in the same functional domain (e.g., response to conspecific vs. heterospecific territorial intrusion).
To the best of our knowledge, the first prediction has never been directly tested by contrasting the patterns of activity across the SDMN in conspecific vs. heterospecific interactions in the same functional domain. However, indirect evidence can be gathered by comparing the relative development of the forebrain, that harbors most of the nodes of the SDMN, between relevant social conditions.

In a recent study with cleaner wrasse, which engages in complex social interactions with heterospecific client reef fish species, the relative forebrain size correlated positively with the complexity of their intra- and interspecific social environment, for high-performing, but not for low-performing individuals, in an ecologically relevant cognitive task for heterospecific interactions (Triki et al., 2020). This suggests that cleaners with relatively larger forebrains used optimal social decision-rules for the level of social complexity they experience and that heterospecific interactions are shaping the forebrain, and putatively its SDMN, development.

Regarding the second prediction, some studies on nonapeptides are available that support it. The involvement of oxytocin, whose receptors are extensively expressed in the SDMN and that is known to regulate social affiliation across a wide range of species (Donaldson & Young, 2009; Jurek & Neumann, 2018; Knobloch & Grinevich, 2014), in social bonding toward conspecifics and heterospecifics has been tested in dogs (Canis familiaris). When sprayed with oxytocin, dogs displayed more affiliative behaviors both toward their owners and toward other dogs (Romero et al., 2014), suggesting a shared causal role of this neuropeptide on conspecific and heterospecific social bonding. Moreover, exogenous administration of oxytocin to dogs also increases gazing toward owners, which elicit increased oxytocin levels in owners, consequently facilitating owners’ affiliation toward, which in turn lead to further increases in oxytocin concentration in dogs (Nagasawa et al., 2015). Arginine-vasopressin/vasotocin (AVP/AVT) is another key neurohormone regulating the modulation of several intraspecific social behaviors, such as social approach/withdrawal and aggression (Goodson & Bass, 2001; Goodson & Thompson, 2010). Research on cleaner fish (L. dimidiatus) reveals how the expression of AVT has been co-opted for interactions with heterospecific client fish. Cleaners evolved smaller and less numerous AVT-ir neurons in the gigantocellular preoptic area (gPOA) compared to non-cleaners, apparently to reduce AVT production to be able to approach the diversity of typically larger client species (Mendonça et al., 2013). Indeed, administration of AVT causes cleaners to reduce cleaning interactions (Soares et al., 2012) and to be less willing to feed against preference (Cardoso, Teles, et al., 2015), indicating that AVT makes them perceive
cleaning interactions as more competitive situations. Together, these results suggest that oxytocin regulation of affiliative behavior in dogs and AVT regulation of social interactions in cleaner fish have been co-opted for heterospecific interactions, which may have led to the coevolution of human–dog bonds and cleaning mutualistic behavior, respectively.

3.4 | Physiological responses to social interactions

Peripheral hormones and physiology both respond to and regulate conspecific social interactions as a way to promote brain–body integration in the scope of social behavior.

Two hormone families have been consistently shown to promote such phenotypic integration: androgens, in the scope of agonistic and reproductive interactions, and glucocorticoids, in the scope of the physiological challenge imposed by social interactions in general (Archer, 2006; Oliveira, 2009; Wingfield et al., 1990). Similarly, although less studied, heterospecific interactions may also elicit similar physiological responses. Indeed, in the territorial algae farmer fish (dusky gregories, Stegastes nigricans) territorial intrusions either by conspecific or by heterospecific intruders elicited similar levels of aggressive response and an increase in cortisol levels that was positively related with the levels of aggression, irrespective of the intruders’ species (Ros et al., 2014). Regarding the cellular stress response, upregulation of heat-shock protein genes (HSP70) has been shown to respond to interspecific interactions in some species (e.g., aggregating anemone, Anthopleura elegantissima; Rossi & Snyder, 2001) but not in others (e.g., porcelain crabs, Petrolisthes spp.; Gunderson et al., 2017), with the reasons for these interspecific differences not being clear. In contrast, interaction or the mere exposure to predators has a consistent effect on the cellular response in prey (Gunderson et al., 2017). Finally, red-winged blackbirds (Agelaius phoeniceus), which respond aggressively toward other conspecific blackbird males that compete for breeding territories, and toward females of the brood parasite brown-headed cowbird, also show partially similar transcriptional responses. Both heterospecific and conspecific aggressive responses triggered a downregulation of immune response genes, but only the latter elicited an increase in singing behavior, which was associated with an upregulation of genes associated with metabolic processes (Louder et al., 2020). Thus, conspecific aggression and heterospecific aggression share similar physiological responses for the similar component of the interaction (i.e., physiological stress).

Testosterone influences the expression of aggressive behavior in males of many vertebrates, especially in the context of territoriosity (Wingfield, 1994; Oliveira, 2004). Conversely, aggressive interactions induce an increase in circulating androgen levels, which has been interpreted as a way to promote a brain–body integrated flexible response to social challenges (Oliveira, 2009; Goymann et al., 2019). Interestingly, the source of androgens regulating aggressive behavior differs seasonally with gonadal androgens being the main source during the breeding season and androgens produced locally in the brain being the main effectors outside the breeding season (Soma, 2006). This suggests that mating and non-mating competitive interactions may be regulated by different androgen-based mechanisms, with the former promoting a brain–gonad integration that is not relevant in the latter. Hence, it is expected that although androgens should promote aggressive behavior in both mating and non-mating interactions, only the former should elicit a response in circulating androgens, which originate predominantly in the gonads. The available data on androgens and interspecific interactions fully support these predictions. In fish (S. nigricans) defending feeding territories, a blocker of androgen receptors (flutamide) suppresses aggressive behavior toward both intraspecific and heterospecific intruders, and neither type of intruder elicits a response in circulating androgens (Vullioud et al., 2013). Similarly, Eastern Bluebirds (Sialia sialis) respond to simulated intrusions of both conspecifics and heterospecifics (house Sparrows, Passer domesticus), and a positive correlation as found between testosterone baseline levels and the aggressiveness displayed toward heterospecific intruders (Arboleda & Khan, 2016). Finally, in sympatric competing warblers the species (Townsend’s warblers, Dendroica townsendi) that is aggressively dominant and outcompetes the other (hermit warblers, D. occidentalis) also exhibits higher androgen levels, and hybrids have intermediate androgen levels, suggesting that these hormonal differences are partially genetically based and regulate heterospecific competition (Owen-Ashley & Butler, 2004).
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