

Kim, Y.; Kret, M.E.; Dukes, D.; Samson, A.C.; Walle, E.A.

Citation

Kim, Y., & Kret, M. E. (2022). The emotional expressions and emotion perception in nonhuman primates. In D. Dukes, A. C. Samson, & E. A. Walle (Eds.), *The Oxford handbook of emotional development*. doi:10.1093/oxfordhb/9780198855903.013.20

Version:Publisher's VersionLicense:Licensed under Article 25fa Copyright Act/Law (Amendment Taverne)Downloaded from:https://hdl.handle.net/1887/3448114

Note: To cite this publication please use the final published version (if applicable).

Yena Kim and Mariska Kret

The Oxford Handbook of Emotional Development *Edited by Daniel Dukes, Andrea C. Samson, and Eric A. Walle*

Print Publication Date: Jan 2022 Subject: Psychology, Affective Science, Developmental Psychology Online Publication Date: Jan 2022 DOI: 10.1093/oxfordhb/9780198855903.013.20

Abstract and Keywords

The expression of emotions and their recognition in conspecifics are pivotal to social life. As Darwin postulated in his pioneering book *The expression of the emotions in man and animals*, many morphological features and functions of emotional expressions characterized in humans are homologous to those of other animals. Intriguingly, despite this early work, scientists have been skeptical about the feasibility of studying emotions in nonhumans and, therefore, the study of their emotional expressions has been limited. However, recent technological advances in neuroscience, genetics, and fine-scale behavioral analyses enable researchers to investigate human emotions in direct comparison with other animals. Throughout this chapter, the authors provide convincing evidence that nonhuman primates produce and recognize conspecific emotional expressions. Some of them, especially the bared-teeth display, are used in multiple contexts, suggesting cognitively sophisticated functions. The flexible use of emotional expressions seems to be tightly linked to species sociality, such as level of tolerance.

Keywords: emotional expression, emotion perception, evolution, primates, sociality

Adaptive Functions of Emotional Expressions and Emotion Perception

FOR decades, the concept of emotion has long been the subject of considerable debate due to differing definitions adopted across disciplines, ranging from functional (Anderson & Adolphs, 2014) to cognitive and socioconstructive (Barrett et al., 2007; Lazarus, 1991) accounts. Although the nature of emotion and to what extent we should attribute animal behavior to an emotion are still debatable (de Waal, 2011); there is a wide consensus that the ability to communicate internal emotional states is critical for social species. In this chapter, we focus on expressions of emotion and how these are perceived by conspecifics. We employ a broad definition of how emotional expressions are operationalized, which

Page 1 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

reads as follows: Any change in the face, body, or voice that is in principle perceivable by conspecifics (via visual, auditory, or olfactory channels) and that results from an altered mental state triggered by biologically relevant stimuli (e.g., a snake, an emotional conspecific) (Kret et al., 2019). Adopting this operational definition has two advantages. First, it allows the inclusion of both explicit expressions resulting from muscle activity and implicit emotion-induced changes such as piloerection and pupil size. Second, it facilitates a direct comparison between human and nonhuman primates and incorporates literature on both primatology and psychology, circumventing terminological discrepancies.

(p. 130) Primates are characterized by their gregariousness and highly developed sociocognitive capacities (Dunbar, 2003; Kappeler & van Schaik, 2002). Properly regulating behaviors in response to internal and external social stimuli is therefore critical for individuals within groups to maintain stable social relationships (Kret & Ploeger, 2015). Preston and de Waal (2002) have proposed an autonomous neural mechanism-the Perception-Action Model (PAS)—as a core mechanism which enables emotional state matching with others (de Waal & Preston, 2017; Preston & de Waal, 2002). The ability to match one's own emotional state with that of another is adaptive from both a kin and nonkin's perspective, since it allows mothers to better accommodate the needs of their offspring and unrelated group members to prepare a fight or flight response in anticipation of potential threats (Preston & de Waal, 2002; Frijda, 2016). This autonomous neural mechanism is presumed to be widespread in all animal taxa, but its complexity and the level of cognitive control presumably varies widely across species. Given the complex nature of primate sociality, primates are expected to have evolved enhanced emotion communication capacities (de Waal & Preston, 2017; Gruber & Sievers, 2019; see Clay et al., this volume).

Socioecological Constraints on Emotion Communication

Emotional expressions have evolved for various reasons. Some expressions, such as pupil dilation, may have evolved due to perceptual benefits (Mathôt, 2018), but without obvious social selective pressure, although in humans and chimpanzees these emotional cues might be picked up by others and influence their emotional state (Kret et al., 2013, 2014; for a review, see Kret, 2015). Other explicit emotional signals, such as facial or vocal expressions, often show key similairities in closely related species due to homology (Andrew, 1963; Preuschoft and van Hooff, 1995). However, specific forms and functions of such expressions are known to vary depending on socioecological conditions to which a species has adapted (for a review, see Waller and Micheletta, 2013).

Facial communication, by means of facial muscle movements, is more prominently found in diurnal than nocturnal species and in those living in large rather than in simple social groups, where close proximity or face-to-face communication is more likely to occur (Dobson, 2009a; Smuts et al., 2008). Parr and her colleagues have summarized some of the existing hypotheses describing physical and socioecological conditions that constrain or fa-

Page 2 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

vor a species' communicative repertoire (Parr et al., 2015). For example, body size constrains the length and frequency of vocalization, as well as facial mobility for facial expressions. Larger animals, compared to smaller animals, produce lower-frequency vocalizations as well as longer calls (Ey et al., 2007) and have more variable facial movements (Dobson, 2009a). These physical boundaries of expressions are tightly linked to species' capability of perceiving variable signals (Kiltie, 2000).

(p. 131) Social complexity is not only correlated with a species' general or sociocognitive capacities, but it is also known to predict the complexity of its communicative repertoire (Dunbar, 1993; Freeberg et al., 2012; Schmidt & Cohn, 2001). For example, species living in large groups have more varied communicative repertoires than those living in small groups (McComb & Semple, 2005). Moreover, tolerant species, compared to despotic species, are likely to have more variable and flexible communicative repertoires, since an error in expressing submissive signals can lead to a higher risk in species with a steep hierarchy (Ciani et al., 2012; Dobson, 2012; Maestripieri, 1999; Parr et al., 2005; Preuschoft & van Hooff, 1995; Rebout et al., 2020). For example, one such species, rhesus macaques, compared to Tonkean macaques (a tolerant species), use bared-teeth displays more unidirectionally from lower-ranking to higher-ranking individuals to display submission (Beisner & McCowan, 2014; Thierry et al., 1989). Moreover, the number of facial displays in the genus *Macaca* is found to be larger in tolerant species than despotic species (Preuschoft & van Hooff, 1995).

The aforementioned studies have increased general knowledge about species' communicative repertoires. However, the link between species' sociality and their capacity of emotion communication has been mostly left untouched. Few studies have thus far compared closely related species in terms of emotional expression (Beisner & McCowan, 2014; Flack & de Waal, 2007; Preuschoft, 1995; Preuschoft & van Hooff, 1995). If emotional expression and emotion perception are indeed tightly linked to a species' communicative repertoire, then we may hypothesize that the complexity of, and the sensitivity to, emotions are directly linked to a species' social characteristics. For example, highly tolerant species living in complex social groups may have higher flexibility and variability in expressing emotions, as well as higher sensitivity to detect emotional expressions (Parr et al., 2005; Waller & Micheletta, 2013). However, without testing this possibility in combination with different emotional expressions and by comparing different species directly with each other under the same conditions, the existence of such species' differences remains speculative.

Behavioral Studies of Emotional Expression in Nonhuman Primates

More than a century ago, Darwin proposed the evolutionary continuity of emotional expressions and corresponding functions in human and nonhuman animals (Darwin, 1872). Ekman later experimentally tested the universality of emotional expressions across different cultural populations in humans and suggested that some emotions are basic and have

Page 3 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

discrete evolutionary functions which constitute a set of fundamental human emotional states important to regulate social life (Ekman, 1992). Although the classification of discrete emotions and their scalability are controversial (Anderson & Adolphs, 2014; Barrett, 1998; Devidze et al., 2006; McNaughton & Corr, 2004; Pfaff et al., (p. 132) 2005; Russell, 1980, 2003), the dominant view is that there are at least some emotions including, for instance, disgust, which are expressed similarly across different cultures (e.g., Izard, 1994) and are present in nonhuman primates (e.g., Berridge, 2000). In this section, we discuss several emotional expressions that have been most widely described in the nonhuman primate literature, and discuss their communicative value.

Among the different expressions of emotion, fear-induced responses have received most attention in both human and nonhuman primate emotion research (Cook & Mineka, 1989, 1990; LoBue & DeLoache, 2008; Weiss et al., 2015). Interestingly, most behavioral research on how primates respond to fear-inducing external stimuli, such as predators, has focused on vocalizations, in terms of linguistic properties, such as referential signaling (Seyfarth et al., 1980). However, the detection of a fear-inducing stimulus often provokes facial expressions (Parr et al., 2005). These facial expressions fundamentally share universal morphological and physiological characteristics which function to increase sensory vigilance, such as increased eye aperture and flared nostrils (Susskind et al., 2008). A growing number of studies on fear in primates, ranging from a phylogenetic analysis of facial movements to several cognitive and neuroimaging experiments (Cook & Mineka, 1989, 1990; Kalin et al., 2001, 2004; LoBue & DeLoache, 2008; Preuschoft & van Hooff, 1995; Weiss et al., 2015), have indeed supported the view that the expression and perception of fear have evolutionary origins in a predatory defense mechanism which later served a social function, such as tension reduction or conflict avoidance (Öhman, 1986, 2009). For example, naïve primates, when exposed to snake-related objects, exhibit avoiding behaviors (Weiss et al., 2015). These seemingly innate behavioral responses associated with snakes also become informative to other conspecific members (Cook & Mineka, 1990). The responses to social threat in primates show similar behavioral patterns to those observed in response to snakes, and the amygdala is known to mediate both predator-related and social threat-related fear responses (for a review, see Öhman, 2009). Therefore, the expression of fear is expected to be highly conserved across species and to have a conspicuous communicative signal. Given the space available for this chapter, we confine our discussion to the visual domain.

The bared-teeth display is one of the most frequently observed facial expressions in nonhuman primates, regardless of social structure or complexity, and most often in response to fear or aggression (marmosets: Stevenson & Poole, 1976; several species of macaques: de Waal & Luttrell, 1985; Thierry et al, 1989; mandrills: Bout & Thierry, 2005; orangutans: Liebal et al., 2006; chimpanzees: Waller & Dunbar, 2005; bonobos: de Waal, 1988). Although there are morphological similarities in the bared-teeth display across species, the frequency and function of its usage, as well as the contexts in which it is expressed, are known to vary. In some species, this expression has been ritualized to express subordinance (Flack & de Waal, 2007; Maestripieri & Wallen, 1997), benign intentions (Waller & Dunbar, 2005), and even affiliation and friendship (Bout & Thierry, 2005), which seem

Page 4 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

to be linked to sociocognitive characteristics, such as tolerance (Dobson, 2012; Thierry et al., 2000). In humans, it has been ritualized into the social smile (Van Hooff, 1972), with multiple social purposes (Martin et al., 2017). Although the involved facial muscles can vary slightly across contexts, using a similar expression (p. 133) in multiple contexts suggests behavioral flexibility and perhaps even the existence of a cognitive capacity for top-down control of emotional expressions (Kret et al., 2019).

Unlike the emotional expressions that have evolved in life-threatening situations, the expression of positive emotions is presumed to be less likely to have a specific pattern of behavioral reactions, since it is less critical to an individual's survival and therefore more susceptible to variation over the course of evolution (Fredrickson, 1998; Tooby & Cosmides, 1990). Nonetheless, one of the positive emotional expressions-the relaxed openmouth display or play face-considered to be homologous to human laughter, has been widely reported across many nonhuman primate species (Davila-Ross et al., 2008, 2015; Palagi, 2006; Pellis & Pellis, 1996; Van Hooff, 1972; Van Hooff & Preuschoft, 2003; Waller & Dunbar, 2005). The relaxed open-mouth display is mostly observed in a play context, such as tickle play (Bard et al., 2014; Palagi, 2008), wrestling (Maestripieri & Ross, 2004; Petit et al., 2008), or ritualized play fighting (Palagi et al., 2007; Palagi & Mancini, 2011). Although it is unclear whether the relaxed open-mouth display is highly conserved across species due to some physical or physiological functions, it is generally acknowledged that it originates from the ritualized play bite, signaling nonaggressive intent (Parr et al., 2015; Poole, 1978; Van Hooff, 1972), and has been reported across mammalian species (Henry & Herrero, 1974; Pal, 2010; Poole, 1978).

Infant chimpanzees as young as 4 weeks of age engage in tickle play which often accompanies this expression (Bard et al., 2014). Due to the reciprocal nature of play interactions, developing immatures learn how to signal and read play intentions from facial and bodily expressions (Burghardt, 2005; Fagen, 1993; Pellis & Pellis, 1996). Through practice, youngsters become competent in decoding cooperative or competitive signals from interaction partners, a critical skill in social living (Pellis & Pellis, 2017). In humans, positive emotions such as joy, interest, contentment, and love, are suggested to build an individual's physical and cognitive capacities by broadening the momentary thought-action repertoire (Broaden-and-Build Model of Positive Emotions: Fredrickson, 1998). Similarly, a study in nonhuman primates has shown that species' involvement in social play correlates with the relative volume of the amygdala and hypothalamus (Lewis & Barton, 2006). This indicates that producing and processing emotional signals during play are adaptive for the development of sociocognitive skills (see Zaharia et al., this volume; Veiga et al., this volume). Humans laugh not only during social play, but also in daily conversations and even without a social partner (e.g., reading comics, watching a comedy, thinking about something funny). Laughing without a social partner may reflect their genuine emotional state (see Mireault, this volume). Interestingly, similar to the social smile, laughter in humans also serves more cognitively sophisticated functions, such as signaling friendly intentions to a stranger (for a review, see Gervais & Wilson, 2005). It would be interesting to investigate whether the relaxed open-mouth display in nonhuman pri-

Page 5 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

mates similarly occurs exclusively during solo play or also outside of the play context, and whether this relates to sociality.

Similar to fear-induced facial expressions, the expression of disgust also has an adaptive function to inhibit sensory exposure, and therefore is commonly considered to be conserved across species (Susskind et al., 2008). Interestingly, humans make (p. 134) disgusted facial expressions even in socially immoral contexts which are similar to expressions following, for instance, the smell of rotten food. It has therefore been suggested that the expression of moral disgust originates from the same behavioral mechanism to avoid biological contaminants (Chapman et al., 2009). However, only a couple of studies thus far have investigated the expression of disgust, and its communicative value in nonhuman primates has rarely been explored (Berridge, 2000; Steiner & Glaser, 1984), In one study by Berridge (2000), researchers gave infants of humans, rats, and multiple species of nonhuman primates different tastes. One of these substances had a bitter taste, and the resulting facial expression was strikingly similar across species. Sarabian and her colleagues explored the disgust response in multiple species of nonhuman primates (Sarabian et al., 2017, 2018; Sarabian & MacIntosh, 2015). In their studies, the primates not only sensed potential biological contaminants through visual cues, but also through olfactory and tactile cues. Furthermore, certain nonhuman primates were able to discriminate parasite-infected individuals from olfactory cues and adjust their grooming behavior accordingly (Poirotte et al., 2017). However, how primates express aversion, and whether these putative expressions are perceived by others, should be further examined.

The specific characteristics of the facial expression of anger in nonhuman primates are not well documented, except in a few great ape species (e.g., bulging lips in chimpanzees and lip press in bonobos: van Hooff, 1971; de Waal, 1988), perhaps due to the lack of conspicuity in the expression of these emotions, or the large variability within or across species. Anger-related emotional expressions are often termed as agonistic or aggressive displays without descriptions of emotional states. Aggressive displays have been generally described in terms of an individual's exaggerated bodily expressions, rather than the facial expressions (Nishida et al., 1999). Since body size often correlates with physical strength, nonhuman primates often show exaggerated body movements with hair erection or use of objects, such as branches, to appear more threatening (Fessler & Gervais, 2010; Nishida et al., 1999). Furthermore, sexually isomorphic species living in small groups with lower levels of agonistic interactions, such as gibbons, may rely more on other behavioral signals, such as vocalizations, to signal aggressive intent (Raemaekers et al., 1984). Therefore, it is likely that the communicative signal of anger takes many different behavioral forms across and within species (Andersson, 1980).

There have been many anecdotal reports of nonhuman primates showing grief or sadness about the death of conspecifics, especially of their offspring. The most frequently reported behavior is the prolonged carrying of a dead infant which is often accompanied by inspection and grooming, and sometimes cannibalism (Anderson, 2016; Gonçalves & Biro, 2018; Watson & Matsuzawa, 2018). It is too early to draw conclusions on whether there are highly conserved facial or other behavioral features within or across primate species

Page 6 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

in expressing grief or sadness. However, it is noteworthy that responding to other's distress or grief is considered to be closely linked to empathy (Preston & de Waal, 2002; see Clay et al., this volume). Therefore, further comparative studies on these putative emotional expressions and perception of them should (p. 135) follow, with a view to enriching our understanding of the origins of human hyper cooperativeness and prosociality.

Experimental Studies of Emotion Perception in Nonhuman Primates

There are far fewer studies concerning the perception of emotional expressions in nonhuman primates than there are concerning how those emotions are expressed. Although it is possible to infer how nonhuman primates perceive emotional expressions by analyzing subsequent behaviors of the recipient, only controlled experiments can address the specific nature of the mechanisms underlying emotion perception (e.g., the capacity to categorize emotional expressions, the sensitivity to graded or variable signals of emotions, the role of expression modality on emotion perception). There have been a number of experimental approaches to investigating emotion perception in nonhuman primates, ranging from touchscreen-based tasks, such as matching-to-sample and dot-probe tasks, to eye-tracking studies (Kano & Tomonaga, 2010b; Kret et al., 2016; Parr & Heintz, 2009; van Berlo et al., 2020). Whereas the matching-to-sample task is widely used to test nonhuman primates' ability to categorize emotional expressions, dot-probe and eye-tracking tasks have been used to test attentional biases toward emotional expressions.

Studies using the matching-to-sample and similar paradigms have found that nonhuman primates can reliably discriminate and categorize images of conspecific emotional expressions (chimpanzees: Kano et al., 2008; Parr, 2001, 2003, 2004; Parr et al., 1998, 2008, 2009; crested macaques: Micheletta et al., 2015; tufted capuchin monkeys: Calcutt et al., 2017; for a review, see Nieuwburg et al., 2021). It is noteworthy that whereas tolerant species, such as crested macaques, showed error patterns influenced by functional similarities of facial expressions, chimpanzees, a despotic species, showed error patterns influenced by facial feature similarities (Micheletta et al., 2015; Parr et al., 1998). These results indicate that a species' tolerance may predict their capacity to decode facial expressions, but this assertion warrants further investigation.

The dot-probe task, a reliable paradigm to test implicit attentional biases (van Rooijen et al., 2017), has also produced concordant results in rhesus macaques, showing that bared-teeth displays (a negative expression in this species) captured attention faster than neutral expressions. In contrast, positive facial expressions did not capture macaques' attention (Lacreuse et al., 2013; Parr et al., 2013). Studies on great apes using the dot-probe paradigm have produced mixed results. Whereas bonobos showed an attentional bias toward emotional scenes compared to neutral scenes (Kret et al., 2016; van Berlo et al., 2020), chimpanzees did not (Kret et al., 2018; Wilson & Tomonaga, 2018). It is unclear whether this discrepancy can be explained by the species' different sociality or methodological differences (e.g., the use of color versus monotone pictures, or face (p. 136) and

Page 7 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

isolated body pictures versus rich emotional scenes) between these four studies. To test whether a species' sociality accounts for the difference in the attentional bias toward emotions, it would be necessary to measure this in closely related but socially different species, such as chimpanzees, bonobos, and humans, with either a set of systematically well-controlled stimuli or a very large stimulus set where low-level differences between unique stimuli are averaged out within the different emotional conditions they constitute (e.g., Kret et al., 2016).

Similar to the dot-probe task, eye-tracking methodology can further elucidate whether the animal is drawn toward, or avoids looking at, certain emotional expressions. For example, rhesus macaques preferred to look at neutral faces over aggressive faces when they were in a stressful condition, but displayed an opposite tendency when they were in an enriched condition (Bethell et al., 2012). Studies on chimpanzees and orangutans have shown the apes' sustained attention toward negative emotional expressions compared to positive and neutral expressions (Kano & Tomonaga, 2010a; Pritsch et al., 2017). Although further work is necessary, the likely explanation of the difference between them would be related to the steepness of hierarchy, as attending to negative expressions in rhesus macaques may elicit substantial stress compared to chimpanzees and orangutans.

There have been a number of experimental studies investigating neural and physiological mechanisms underlying emotion processing in nonhuman primates. Similar to the findings in humans, studies in chimpanzees have shown brain lateralization when processing emotional expressions (Parr & Hopkins, 2000) and specific event-related potential (ERP) waveforms elicited by affective pictures (Hirata et al., 2013). Moreover, peripheral temperature changes, such as nasal temperature drop, were also found in nonhuman primates when processing negative emotional expressions (Dezecache et al., 2017; Kano et al., 2016; Nakayama et al., 2005). In nonhuman primates, the neuropeptide hormone oxytocin also seems to modulate the sensitivity to emotional expressions and regulate emotional behaviors (Chang & Platt, 2014; Crockford et al., 2013; Dal Monte et al., 2014; Parr et al., 2013, 2018). Genetic variation of oxytocin and vasopressin receptor genes found in chimpanzees and bonobos further suggests that behavioral and perceptual differences in these species are closely linked to their genetic and physiological differences (Staes et al., 2014). Therefore, it is most parsimonious to assume that there is evolutionary continuity in both emotional behaviors and their processing in humans and nonhuman primates.

Discussion and Future Directions

In this chapter, we aimed to shed light on the socioecological selective pressures that have shaped emotion communication by reviewing the literature on emotional expression and emotion perception in nonhuman primates. Indeed, a handful of studies suggest that the way species produce and use emotional expressions is largely influenced (p. 137) by these pressures. In particular, social complexity and tolerance seem to be positively correlated with the variability and complexity of species' emotion communication (Dobson,

Page 8 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

2009b, 2012; Rebout et al., 2020; Thierry et al., 1989; Preuschoft & van Hooff, 1995). However, throughout the review, we have encountered more limitations than concrete empirical findings to link emotions and species' socioecological characteristics in the nonhuman primate emotion literature. In this section, we discuss the preliminary results from which we draw our conclusions, and further outline the limitations and, most importantly, possible future directions of research.

First, some of the facial expressions of emotions, such as bared-teeth and open-mouth displays, are conserved across primate species in terms of morphological characteristics. However, the frequency, flexibility, and function of expressions vary in relation to species' social characteristics. With the exception of a few studies (Dobson, 2012; Thierry et al., 2000), there is a lack of systematic investigation of emotional expressions across closely related species. Moreover, comparative studies on the contexts in which these are expressed in relation to species sociality are virtually absent. It is of critical importance that this gap is addressed, since the expression of emotions in multiple contexts with various functions will likely reveal species' capacities to use and decode emotional signals, as well as the evolutionary trajectory that shaped species' sociality.

Second, observational studies have shown that nonhuman primates are, in general, able to express and recognize emotions. However, unlike observational studies, experimental studies on emotion perception in nonhuman primates are scarce. In particular, we do not know whether certain species are more sensitive than others to particular modalities, a particular valence or intensity of emotional cues, and whether this can be explained by their specific sociality. Since different methodologies often lead to different results, even in the same species, it has to be noted that experimental stimuli should be prepared and presented under ecologically valid, yet controlled conditions.

Third, studies of emotional expressions and perception in nonhuman primates mostly concern facial expressions. Studies in humans, however, have shown that people can be as good at recognizing bodily expressions of emotion as they are at recognizing facial expressions (De Gelder, 2009; Kret et al., 2013). In this regard, studying the expression and perception of emotions through different modalities with regard to species' sociality and their environment (e.g., dense forest or open savannah) could help construct a comprehensive evolutionary framework of emotion.

Finally, studies on the perception of subtle emotional cues have almost never been explored in nonhuman primates. For example, only one study has tested the perception of pupil size in humans and chimpanzees (Kret et al., 2014). A growing body of research in humans suggests that these subtle emotional cues (e.g., pupil size, blushing, tears during sadness or laughter, signs of sweating or goosebumps) play a significant role in cooperative decision making and social bonding (Behrens & Kret, 2019; Prochazkova et al., 2019, 2021; for a review, see Prochazkova & Kret, 2017). Therefore, studying the expression and perception of these subtle emotional cues would provide a promising avenue for future emotion research.

Page 9 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

(p. 138) Conclusion

In broad terms, species' sociality is tightly interconnected with perception, cognition, communication, and behavior (Dobson, 2009b; Dunbar, 1993; Freeberg et al., 2012; Kano et al., 2018). We therefore presume a link with emotional capacities, too. Here, we suggest that species' social complexity and level of tolerance determine the flexibility and complexity in expressing and perceiving emotions. Specifically, we predict that species living in complex and tolerant social environments have higher flexibility and variability in expressing emotions, as well as higher sensitivity to detect emotional expressions in their group mates. We encourage future studies to test this hypothesis in closely related but socially distinct species, such as chimpanzees and bonobos. Studying their expression and perception of emotional expressions would fill the gap between species' sociality and emotional capacities, and thereby help to construct a full evolutionary picture of emotions.

References

Anderson, D. J., & Adolphs, R. (2014). A framework for studying emotions across species. *Cell*, *157*(1), 187–200.

Anderson, J. R. (2016). Comparative thanatology. Current Biology, 26(13), R553-R556.

Andersson, M. (1980). Why are there so many threat displays? *Journal of Theoretical Biology*, 86(4), 773–781.

Andrew, R. J. (1963). The origin and evolution of the calls and facial expression of the primates. *Behaviour*, 20(1/2), 1–109.

Bard, K. A., Dunbar, S., Maguire-Herring, V., Veira, Y., Hayes, K. G., & McDonald, K. (2014). Gestures and social-emotional communicative development in chimpanzee infants. *American Journal of Primatology*, *76*(1), 14–29.

Barrett, L. F. (1998). Discrete emotions or dimensions? The role of valence focus and arousal focus. *Cognition & Emotion*, *12*(4), 579–599.

Barrett, L. F., Mesquita, B., Ochsner, K. N., & Gross, J. J. (2007). The experience of emotion. *Annual Reviews in Psychology*, 58, 373-403.

Behrens, F., & Kret, M. E. (2019). The interplay between face-to-face contact and feedback on cooperation during real-life interactions. *Journal of Nonverbal Behavior*, 43(4), 513–528.

Beisner, B. A., & McCowan, B. (2014). Signaling context modulates social function of silent bared-teeth displays in rhesus macaques (*Macaca mulatta*). *American Journal of Primatology*, 76(2), 111–121.

Page 10 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

Berridge, K. C. (2000). Measuring hedonic impact in animals and infants: Microstructure of affective taste reactivity patterns. *Neuroscience & Biobehavioral Reviews*, *24*(2), 173-198.

Bethell, E. J., Holmes, A., MacLarnon, A., & Semple, S. (2012). Evidence that emotion mediates social attention in rhesus macaques. *PLOS ONE*, *7*(8), Article e44387.

Bout, N., & Thierry, B. (2005). Peaceful meaning for the silent bared-teeth displays of mandrills. *International Journal of Primatology*, *26*(6), 1215–1228.

Burghardt, G. M. (2005). The genesis of animal play: Testing the limits: MIT Press.

(p. 139) Calcutt, S. E., Rubin, T. L., Pokorny, J. J., & de Waal, F. (2017). Discrimination of emotional facial expressions by tufted capuchin monkeys (*Sapajus apella*). *Journal of Comparative Psychology*, 131(1), 40.

Chapman, H. A., Kim, D. A., Susskind, J. M., & Anderson, A. K. (2009). In bad taste: Evidence for the oral origins of moral disgust. *Science*, *323*(5918), 1222–1226.

Chang, S. W., & Platt, M. L. (2014). Oxytocin and social cognition in rhesus macaques: mplications for understanding and treating human psychopathology. *Brain Research*, *1580*, 57–68.

Ciani, F., Dall'Olio, S., Stanyon, R., & Palagi, E. (2012). Social tolerance and adult play in macaque societies: A comparison with different human cultures. *Animal Behaviour, 84*(6), 1313–1322.

Cook, M., & Mineka, S. (1989). Observational conditioning of fear to fear-relevant versus fear-irrelevant stimuli in rhesus monkeys. *Journal of Abnormal Psychology*, *98*(4), 448.

Cook, M., & Mineka, S. (1990). Selective associations in the observational conditioning of fear in rhesus monkeys. *Journal of Experimental Psychology: Animal Behavior Processes*, *16*(4), 372.

Crockford, C., Wittig, R., Langergraber, K., Ziegler, T., Zuberbühler, K., & Deschner, T. (2013). Urinary oxytocin and social bonding in related and unrelated wild chimpanzees. *Proceedings of the Royal Society B, 280*, 20122765.

Dal Monte, O., Noble, P. L., Costa, V. D., & Averbeck, B. B. (2014). Oxytocin enhances attention to the eye region in rhesus monkeys. *Frontiers in Neuroscience*, 8. https:// doi.org/10.3389/fnins.2014.0004

Davila-Ross, M., Jesus, G., Osborne, J., & Bard, K. A. (2015). Chimpanzees (*Pan troglodytes*) produce the same types of "laugh faces" when they emit laughter and when they are silent. *PLOS ONE*, *10*(6), Article e0127337.

Davila-Ross, M., Menzler, S., & Zimmermann, E. (2008). Rapid facial mimicry in orangutan play. *Biology Letters*, 4(1), 27-30.

Page 11 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

Darwin, C. (1872). *The expression of the emotions in man and animals*. London, UK: John Murray.

De Gelder, B. (2009). Why bodies? Twelve reasons for including bodily expressions in affective neuroscience. *Philosophical Transactions of the Royal Society B: Biological Sciences, 364*(1535), 3475-3484.

De Waal, F. B. (1988). The communicative repertoire of captive bonobos (*Pan paniscus*), compared to that of chimpanzees. *Behaviour*, 183–251.

De Waal, F. B. (2011). What is an animal emotion? *Annals of the New York Academy of Sciences*, 1224(1), 191–206.

De Waal, F. B., & Luttrell, L. M. (1985). The formal hierarchy of rhesus macaques: An investigation of the bared-teeth display. *American Journal of Primatology*, 9(2), 73–85.

De Waal, F. B., & Preston, S. D. (2017). Mammalian empathy: Behavioural manifestations and neural basis. *Nature Reviews Neuroscience*, *18*(8), 498.

Devidze, N., Lee, A. W., Zhou, J., & Pfaff, D. W. (2006). CNS arousal mechanisms bearing on sex and other biologically regulated behaviors. *Physiology & Behavior, 88*(3), 283–293.

Dezecache, G., Zuberbühler, K., Davila-Ross, M., & Dahl, C. D. (2017). Skin temperature changes in wild chimpanzees upon hearing vocalizations of conspecifics. *Royal Society Open Science*, *4*(1), 160816.

Dobson, S. D. (2009a). Allometry of facial mobility in anthropoid primates: Implications for the evolution of facial expression. *American Journal of Physical Anthropology, 138*(1), 70–81.

(p. 140) Dobson, S. D. (2009b). Socioecological correlates of facial mobility in nonhuman anthropoids. *American Journal of Physical Anthropology*, 139(3), 413–420.

Dobson, S. D. (2012). Coevolution of facial expression and social tolerance in macaques. *American Journal of Primatology*, 74(3), 229–235.

Dunbar, R. I. (1993). Coevolution of neocortical size, group size and language in humans. *Behavioral and Brain Sciences*, *16*(4), 681–694.

Dunbar, R. I. (2003). The social brain: Mind, language, and society in evolutionary perspective. *Annual Review of Anthropology*, *32*(1), 163–181.

Ekman, P. (1992). An argument for basic emotions. *Cognition & Emotion*, 6(3-4), 169-200.

Ey, E., Pfefferle, D., & Fischer, J. (2007). Do age- and sex-related variations reliably reflect body size in non-human primate vocalizations? A review. *Primates*, *48*(4), 253–267.

Fagen, R. (1993). Primate juveniles and primate play. In M. E. Pereira & L. A. Fairbanks (Eds.), *Juvenile primates* (pp. 182–196). Oxford University Press.

Page 12 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

Fessler, D. M., & Gervais, M. (2010). From whence the captains of our lives: Ultimate and phylogenetic perspectives on emotions in humans and other primates. In P. Kappeler & J. Silk (Eds.), *Mind the gap* (pp. 261–280). Springer.

Flack, J. C., & de Waal, F. (2007). Context modulates signal meaning in primate communication. *Proceedings of the National Academy of Sciences*, *104*(5), 1581–1586.

Fredrickson, B. L. (1998). What good are positive emotions? *Review of General Psychology*, *2*(3), 300–319.

Freeberg, T. M., Dunbar, R. I., & Ord, T. J. (2012). Social complexity as a proximate and ultimate factor in communicative complexity. *Philosophical Transactions of the Royal Society B: Biological Sciences, 367*(1597). https://doi.org/10.1098/rstb.2011.0213

Frijda, N. H. (2016). The evolutionary emergence of what we call "emotions." *Cognition and Emotion*, *30*(4), 609–620.

Gervais, M., & Wilson, D. S. (2005). The evolution and functions of laughter and humor: A synthetic approach. *Quarterly Review of Biology*, *80*(4), 395–430.

Gonçalves, A., & Biro, D. (2018). Comparative thanatology, an integrative approach: Exploring sensory/cognitive aspects of death recognition in vertebrates and invertebrates. *Philosophical Transactions of the Royal Society B: Biological Sciences, 373*(1754), 20170263.

Gruber, T., & Sievers, C. (2019). Affective social learning and the emotional side of cultural learning in primates. In D. Dukes & F. Clément (Eds.), *Foundations of affective social learning: Conceptualising the transmission of social value* (pp. 41–66). Cambridge University Press.

Henry, J., & Herrero, S. (1974). Social play in the American black bear: Its similarity to canid social play and an examination of its identifying characteristics. *American Zoologist*, *14*(1), 371–389.

Hirata, S., Matsuda, G., Ueno, A., Fukushima, H., Fuwa, K., Sugama, K., Kusunoki, K., Tomonaga, M., Hiraki, K., & Hasegawa, T. (2013). Brain response to affective pictures in the chimpanzee. *Scientific Reports*, *3*, 1342.

Izard, C. E. (1994). Innate and universal facial expressions: Evidence from developmental and cross-cultural research. *Psychological Bulletin*, *115*(2), 288–299.

Kalin, N. H., Shelton, S. E., & Davidson, R. J. (2004). The role of the central nucleus of the amygdala in mediating fear and anxiety in the primate. *Journal of Neuroscience*, *24*(24), 5506–5515.

Kalin, N. H., Shelton, S. E., Davidson, R. J., & Kelley, A. E. (2001). The primate amygdala mediates acute fear but not the behavioral and physiological components of anxious temperament. *Journal of Neuroscience*, *21*(6), 2067–2074.

Page 13 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

(p. 141) Kano, F., Hirata, S., Deschner, T., Behringer, V., & Call, J. (2016). Nasal temperature drop in response to a playback of conspecific fights in chimpanzees: A thermo-imaging study. *Physiology & Behavior*, 155, 83–94.

Kano, F., Shepherd, S. V., Hirata, S., & Call, J. (2018). Primate social attention: Species differences and effects of individual experience in humans, great apes, and macaques. *PLOS ONE*, *13*(2), Article e0193283.

Kano, F., Tanaka, M., & Tomonaga, M. (2008). Enhanced recognition of emotional stimuli in the chimpanzee (*Pan troglodytes*). *Animal Cognition*, *11*(3), 517–524.

Kano, F., & Tomonaga, M. (2010a). Attention to emotional scenes including whole-body expressions in chimpanzees (*Pan troglodytes*). *Journal of Comparative Psychology*, *124*(3), 287.

Kano, F., & Tomonaga, M. (2010b). Face scanning in chimpanzees and humans: Continuity and discontinuity. *Animal Behaviour*, 79(1), 227–235.

Kappeler, P. M., & van Schaik, C. P. (2002). Evolution of primate social systems. *International Journal of Primatology*, 23(4), 707–740.

Kiltie, R. (2000). Scaling of visual acuity with body size in mammals and birds. *Functional Ecology*, 14(2), 226–234.

Kret, M. E. (2015). Emotional expressions beyond facial muscle actions. A call for studying autonomic signals and their impact on social perception. *Frontiers in Psychology*, *6*, 711.

Kret, M. E., Jaasma, L., Bionda, T., & Wijnen, J. G. (2016). Bonobos (*Pan paniscus*) show an attentional bias toward conspecifics' emotions. *Proceedings of the National Academy of Sciences*, *113*(14), 3761–3766.

Kret, M. E., Muramatsu, A., & Matsuzawa, T. (2018). Emotion processing across and within species: A comparison between humans (*Homo sapiens*) and chimpanzees (*Pan troglodytes*). *Journal of Comparative Psychology*, 132(4), 395.

Kret, M. E., & Ploeger, A. (2015). Emotion processing deficits: A liability spectrum providing insight into comorbidity of mental disorders. *Neuroscience & Biobehavioral Reviews*, *52*, 153–171.

Kret, M. E., Prochazkova, E., Sterck, E., & Clay Z. (2019). Emotional expressions in human and non-human great apes. *Neuroscience and Biobehavioral Reviews*, *115*, 378–395.

Kret, M. E., Stekelenburg, J., Roelofs, K., & De Gelder, B. (2013). Perception of face and body expressions using electromyography, pupillometry and gaze measures. *Frontiers in Psychology*, *4*, 28.

Page 14 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

Kret, M. E., Tomonaga, M., & Matsuzawa, T. (2014). Chimpanzees and humans mimic pupil-size of conspecifics. *PLOS ONE*, *9*(8), Article e104886.

Lacreuse, A., Schatz, K., Strazzullo, S., King, H. M., & Ready, R. (2013). Attentional biases and memory for emotional stimuli in men and male rhesus monkeys. *Animal Cognition*, *16*(6), 861–871.

Lazarus, R. S. (1991). Cognition and motivation in emotion. *American Psychologist*, 46(4), 352.

Lewis, K. P., & Barton, R. A. (2006). Amygdala size and hypothalamus size predict social play frequency in nonhuman primates: A comparative analysis using independent contrasts. *Journal of Comparative Psychology*, 120(1), 31.

Liebal, K., Pika, S., & Tomasello, M. (2006). Gestural communication of orangutans (*Pongo pygmaeus*). *Gesture*, *6*(1), 1–38.

LoBue, V., & DeLoache, J. S. (2008). Detecting the snake in the grass: Attention to fearrelevant stimuli by adults and young children. *Psychological Science*, *19*(3), 284–289.

Maestripieri, D. (1999). Primate social organization, gestural repertoire size, and communication dynamics: A comparative study of macaques. In B. King (Ed.), *The origins of language: What nonhuman primates can tell us* (pp. 55–77). School of American Research.

(p. 142) Maestripieri, D., & Ross, S. R. (2004). Sex differences in play among western lowland gorilla (*Gorilla gorilla gorilla*) infants: Implications for adult behavior and social structure. *American Journal of Physical Anthropology*, 123(1), 52–61.

Maestripieri, D., & Wallen, K. (1997). Affiliative and submissive communication in rhesus macaques. *Primates*, *38*(2), 127-138.

Martin, J., Rychlowska, M., Wood, A., & Niedenthal, P. (2017). Smiles as multipurpose social signals. *Trends in Cognitive Sciences*, *21*(11), 864–877.

Mathôt, S. (2018). Pupillometry: Psychology, physiology, and function. *Journal of Cognition*, 1(1).

McComb, K., & Semple, S. (2005). Coevolution of vocal communication and sociality in primates. *Biology Letters*, 1(4), 381–385.

McNaughton, N., & Corr, P. J. (2004). A two-dimensional neuropsychology of defense: Fear/anxiety and defensive distance. *Neuroscience & Biobehavioral Reviews, 28*(3), 285-305.

Micheletta, J., Whitehouse, J., Parr, L. A., & Waller, B. M. (2015). Facial expression recognition in crested macaques (*Macaca nigra*). *Animal Cognition*, *18*(4), 985–990.

Page 15 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

Nakayama, K., Goto, S., Kuraoka, K., & Nakamura, K. (2005). Decrease in nasal temperature of rhesus monkeys (*Macaca mulatta*) in negative emotional state. *Physiology & Behavior, 84*(5), 783–790.

Nishida, T., Kano, T., Goodall, J., McGrew, W. C., & Nakamura, M. (1999). Ethogram and ethnography of Mahale chimpanzees. *Anthropological Science*, *107*(2), 141–188.

Nieuwburg, E., Ploeger, A., & Kret, M.E. (2021). Emotion recognition in nonhuman primates: How experimental research can contribute to a better understanding of underlying mechanisms. *Neuroscience & Biobehavioral Reviews*. 123, 24–47.

Öhman, A. (1986). Face the beast and fear the face: Animal and social fears as prototypes for evolutionary analyses of emotion. *Psychophysiology*, 23(2), 123–145.

Öhman, A. (2009). Of snakes and faces: An evolutionary perspective on the psychology of fear. *Scandinavian Journal of Psychology*, *50*(6), 543–552.

Pal, S. K. (2010). Play behaviour during early ontogeny in free-ranging dogs (*Canis familiaris*). *Applied Animal Behaviour Science*, *126*(3–4), 140–153.

Palagi, E. (2006). Social play in bonobos (*Pan paniscus*) and chimpanzees (*Pan troglodytes*): Implications for natural social systems and interindividual relationships. *American Journal of Physical Anthropology*, *129*(3), 418–426.

Palagi, E. (2008). Sharing the motivation to play: The use of signals in adult bonobos. *Ani- mal Behaviour*, *75*(3), 887–896.

Palagi, E., Antonacci, D., & Cordoni, G. (2007). Fine-tuning of social play in juvenile lowland gorillas (*gorilla gorilla gorilla*). *Developmental Psychobiology*, 49(4), 433–445.

Palagi, E., & Mancini, G. (2011). Playing with the face: playful facial "chattering" and signal modulation in a monkey species (*Theropithecus gelada*). *Journal of Comparative Psychology*, *125*(1), 11.

Parr, L. A. (2001). Cognitive and physiological markers of emotional awareness in chimpanzees (*Pan troglodytes*). *Animal Cognition*, 4(3-4), 223–229.

Parr, L. A. (2003). The discrimination of faces and their emotional content by chimpanzees (*Pan troglodytes*). *Annals of the New York Academy of Sciences*, 1000(1), 56-78.

Parr, L. A. (2004). Perceptual biases for multimodal cues in chimpanzee (*Pan troglodytes*) affect recognition. *Animal Cognition*, 7(3), 171–178.

(p. 143) Parr, L. A., Cohen, M., & de Waal, F. (2005). Influence of social context on the use of blended and graded facial displays in chimpanzees. *International Journal of Primatology*, *26*(1), 73–103.

Parr, L. A., Hecht, E., Barks, S. K., Preuss, T. M., & Votaw, J. R. (2009). Face processing in the chimpanzee brain. *Current Biology*, 19(1), 50–53.

Page 16 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

Parr, L. A., & Heintz, M. (2009). Facial expression recognition in rhesus monkeys, *Macaca mulatta*. *Animal Behaviour*, 77(6), 1507–1513.

Parr, L. A., & Hopkins, W. D. (2000). Brain temperature asymmetries and emotional perception in chimpanzees, *Pan troglodytes*. *Physiology & Behavior*, *71*(3–4), 363–371.

Parr, L. A., Hopkins, W. D., & de Waal, F. B. (1998). The perception of facial expressions by chimpanzees, *Pan troglodytes*. *Evolution of Communication*, *2*(1), 1–23.

Parr, L. A., Micheletta, J., & Waller, B. M. (2015). Nonverbal communication in primates: observational and experimental approaches. In *APA handbook of nonverbal communication*. American Psychological Association.

Parr, L. A., Mitchell, T., & Hecht, E. (2018). Intranasal oxytocin in rhesus monkeys alters brain networks that detect social salience and reward. *American Journal of Primatology, 80*(10), Article e22915.

Parr, L. A., Modi, M., Siebert, E., & Young, L. J. (2013). Intranasal oxytocin selectively attenuates rhesus monkeys' attention to negative facial expressions. *Psychoneuroendocrinology*, 38(9), 1748–1756.

Parr, L. A., Waller, B. M., & Fugate, J. (2005). Emotional communication in primates: Implications for neurobiology. *Current Opinion in Neurobiology*, *15*(6), 716–720.

Parr, L. A., Waller, B. M., & Heintz, M. (2008). Facial expression categorization by chimpanzees using standardized stimuli. *Emotion*, 8(2), 216.

Pellis, S. M., & Pellis, V. C. (1996). On knowing it's only play: The role of play signals in play fighting. *Aggression and Violent Behavior*, *1*(3), 249–268.

Pellis, S. M., & Pellis, V. C. (2017). What is play fighting and what is it good for? *Learning & Behavior*, 45(4), 355–366.

Petit, O., Bertrand, F., & Thierry, B. (2008). Social play in crested and Japanese macaques: Testing the covariation hypothesis. *Developmental Psychobiology*, *50*(4), 399–407.

Pfaff, D., Westberg, L., & Kow, L. M. (2005). Generalized arousal of mammalian central nervous system. *Journal of Comparative Neurology*, 493(1), 86–91.

Poirotte, C., Massol, F., Herbert, A., Willaume, E., Bomo, P. M., Kappeler, P. M., & Charpentier, M. J. (2017). Mandrills use olfaction to socially avoid parasitized conspecifics. *Science Advances*, *3*(4), Article e1601721.

Poole, T. B. (1978). An analysis of social play in polecats (*Mustelidae*) with comments on the form and evolutionary history of the open mouth play face. *Animal Behaviour, 26*, 36-49.

Preston, S. D., & de Waal, F. (2002). Empathy: Its ultimate and proximate bases. *Behavioral and Brain Sciences*, *25*(01), 1–20.

Page 17 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

Preuschoft, S. (1995). *"Laughter" and "smiling" in Macaques: an evolutionary perspective* [Unpublished doctoral dissertation]. University of Utrecht.

Preuschoft, S., & van Hooff, J. A. (1995). Homologizing primate facial displays: A critical review of methods. *Folia Primatologica*, *65*(3), 121–137.

Pritsch, C., Telkemeyer, S., Mühlenbeck, C., & Liebal, K. (2017). Perception of facial expressions reveals selective affect-biased attention in humans and orangutans. *Scientific Reports*, 7(1), 7782.

(p. 144) Prochazkova, E., & Kret, M. E. (2017). Connecting minds and sharing emotions through mimicry: A neurocognitive model of emotional contagion. *Neuroscience & Biobehavioral Reviews*, *80*, 99–114.

Prochazkova, E., Sjak-Shie, E. E., Behrens, F., Lindh, D., & Kret, M. E. (2019). The choreography of human attraction: Physiological synchrony in a blind date setting. *bioRxiv*. https://doi.org/10.1101/748707

Prochazkova, E., Sjak-shie, E.E., Behrens, F., Lindh, D., & Kret, M.E. (2021). Physiological synchrony predicts attraction in a blind date setting. Nature Human Behavior

Raemaekers, J. J., Raemaekers, P. M., & Haimoff, E. H. (1984). Loud calls of the gibbon (*Hylobates lar*): Repertoire, organisation and context. *Behaviour*, *91*(1/3), 146–189.

Rebout, N., De Marco, A., Lone, J.-C., Sanna, A., Cozzolino, R., Micheletta, J., Sterck, E. H. M., Langermans, J. A. M., Lemasson, A., & Thierry, B. (2020). Tolerant and intolerant macaques show different levels of structural complexity in their vocal communication. *Proceedings of the Royal Society B: Biological Sciences*, 287(1928), 20200439.

Russell, J. A. (1980). A circumplex model of affect. *Journal of Personality and Social Psychology*, *39*(6), 1161.

Russell, J. A. (2003). Core affect and the psychological construction of emotion. *Psychological Review*, *110*(1), 145.

Sarabian, C., Belais, R., & MacIntosh, A. J. (2018). Feeding decisions under contamination risk in bonobos. *Philosophical Transactions of the Royal Society B: Biological Sciences, 373*(1751), 20170195.

Sarabian, C., & MacIntosh, A. J. (2015). Hygienic tendencies correlate with low geohelminth infection in free-ranging macaques. *Biology Letters*, *11*(11), 20150757.

Sarabian, C., Ngoubangoye, B., & MacIntosh, A. J. (2017). Avoidance of biological contaminants through sight, smell and touch in chimpanzees. *Royal Society Open Science*, 4(11), 170968.

Page 18 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

Schmidt, K. L., & Cohn, J. F. (2001). Human facial expressions as adaptations: Evolutionary questions in facial expression research. *American Journal of Physical Anthropology, 116*(S33), 3–24.

Seyfarth, R. M., Cheney, D. L., & Marler, P. (1980). Vervet monkey alarm calls: Semantic communication in a free-ranging primate. *Animal Behaviour, 28*(4), 1070–1094.

Smuts, B. B., Cheney, D. L., Seyfarth, R. M., & Wrangham, R. W. (2008). *Primate societies*. University of Chicago Press.

Staes, N., Stevens, J. M., Helsen, P., Hillyer, M., Korody, M., & Eens, M. (2014). Oxytocin and vasopressin receptor gene variation as a proximate base for inter- and intraspecific behavioral differences in bonobos and chimpanzees. *PLOS ONE*, *9*(11), Article e113364.

Stevenson, M. F., & Poole, T. B. (1976). An ethogram of the common marmoset (*Calithrix jacchus jacchus*): General behavioural repertoire. *Animal Behaviour, 24*(2), 428-451.

Steiner, J. E., & Glaser, D. (1984). Differential behavioral responses to taste stimuli in nonhuman primates. *Journal of Human Evolution*, *13*(8), 709–723.

Susskind, J. M., Lee, D. H., Cusi, A., Feiman, R., Grabski, W., & Anderson, A. K. (2008). Expressing fear enhances sensory acquisition. *Nature Neuroscience*, *11*(7), 843.

Thierry, B., Demaria, C., Preuschoft, S., & Desportes, C. (1989). Structural convergence between silent bared-teeth display and relaxed open-mouth display in the Tonkean macaque (*Macaca tonkeana*). *Folia Primatologica*, *52*(3-4), 178–184.

Thierry, B., Iwaniuk, A. N., & Pellis, S. M. (2000). The influence of phylogeny on the social behaviour of macaques (Primates: Cercopithecidae, genus Macaca). *Ethology*, *106*(8), 713–728.

Tooby, J., & Cosmides, L. (1990). The past explains the present: Emotional adaptations and the structure of ancestral environments. *Ethology and Sociobiology*, *11*(4-5), 375–424.

(p. 145) van Berlo, E., Bionda, T., & Kret, M. E. (2020). Attention towards emotions is modulated by familiarity with the expressor. A comparison between bonobos and humans. *bioRxiv*. https://doi.org/10.1101/2020.05.11.089813

Van Hooff, J. (1971). Aspecten van het sociale gedrag en de communicatie bij humane en hogere niet-humane primate [Unpublished doctoral dissertation]. Utrecht University.

Van Hooff, J. (1972). A comparative approach to the phylogeny of laughter and smiling. In R. A. Hinde (Ed.), *Non-verbal communication* (pp. 209–241). Cambridge University Press.

Van Hooff, J., & Preuschoft, S. (2003). Laughter and smiling: The intertwining of nature and culture. In F. B. M. de Waal & P. L. Tyack (Eds.), *Animal social complexity: Intelligence, culture, and individualized societies* (pp. 260–287). Harvard University Press.

Page 19 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).

van Rooijen, R., Ploeger, A., & Kret, M. E. (2017). The dot-probe task to measure emotional attention: A suitable measure in comparative studies? *Psychonomic Bulletin & Review*, 24(6), 1686–1717.

Waller, B. M., & Dunbar, R. I. (2005). Differential behavioural effects of silent bared teeth display and relaxed open mouth display in chimpanzees (*Pan troglodytes*). *Ethology, 111*(2), 129–142.

Waller, B. M., & Micheletta, J. (2013). Facial expression in nonhuman animals. *Emotion Review*, *5*(1), 54–59.

Watson, C. F., & Matsuzawa, T. (2018). Behaviour of nonhuman primate mothers toward their dead infants: Uncovering mechanisms. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 373(1754), 20170261.

Weiss, L., Brandl, P., & Frynta, D. (2015). Fear reactions to snakes in naïve mouse lemurs and pig-tailed macaques. *Primates*, *56*(3), 279–284.

Wilson, D. A., & Tomonaga, M. (2018). Visual discrimination of primate species based on faces in chimpanzees. *Primates*, *59*, 243–251.

Yena Kim

Yena Kim, Cognitive Psychology Unit, Institute of Psychology, Leiden University, The Netherlands

Mariska Kret

Mariska Kret, Cognitive Psychology Unit, Institute of Psychology, Leiden University, The Netherlands

Page 20 of 20

PRINTED FROM OXFORD HANDBOOKS ONLINE (www.oxfordhandbooks.com). © Oxford University Press, 2022. All Rights Reserved. Under the terms of the licence agreement, an individual user may print out a PDF of a single chapter of a title in Oxford Handbooks Online for personal use (for details see Privacy Policy and Legal Notice).