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Human ability to adapt and survive is largely based on interpersonal skills, in particular on the ability to understand and to cooperate with others. To preserve this precious capacity, social cognition relies on many mechanisms: among these, empathy and Theory of Mind are the most important ones. Although both Theory of Mind and empathy allow human beings to understand others through the representation of their mental or of their emotional states, respectively, research in the field of neuroscience has shown that the abilities of mentalizing and empathizing are associated with the activation of different neuro-cognitive circuits. On the other hand, recent studies based on naturalistic paradigms have shown that the neural systems involved in sharing and mentalization experiences play complementary roles during complex social interactions.

## Method

The present review has the purpose to discuss some features, differences and overlaps in the concepts used to define empathy and Theory of Mind: this issue will be addressed both from a psychological and a neuro-biological point of view. We chose to prepare a narrative review ( [Green et al., 2006](#B97) ; [Pan, 2008](#B144) ) because it is an interpretative-qualitative form of accounting about research ( [Ferrari, 2015](#B74) ): it allows synthesizing the findings of literature about a given topic and expand our knowledge about it ( [Grant and Booth, 2009](#B96) ).

The methodological approach ( [Egger et al., 2001](#B66) ) consisted in the collection of articles available in the international literature through electronic databases such as PROQUEST, PSICARTICLES, and PSYCINFO, published from 1995 until 2018. We used Scopus to verify the scientific relevance of the journals where these documents were published (as indicated in [Montoya et al., 2018](#B139) ). The literature search has been carried out using various combinations of the following keywords, for example: “ theory of mind,” “ mentalizing,” “ role-taking,” “ perspective taking,” “ emotional contagion” and “ empathic concern.”

The titles and abstracts of each identified article have been examined and a subset of them has been selected based on relevance: the latter was judged in terms: (1) of the number of citations gained in other articles of the same field; and (2) of the impact-factor of journals in which they have been published. Other criteria of inclusion have been: articles in Italian or English language, published in peer-reviewed journals, with the exclusion of any University Degree Thesis. At the end of the process, a total of 98 scientific articles have been included.

Another purpose of the present review was to address the issue of animal modeling. We run a PubMed search with exactly the same keywords as above, with the addition of “ rat” or “ animal model” or “ preclinical” and some review articles were returned, supporting the idea that empathy-like phenomena, which are not limited to humans but widespread across mammals, also can be studied in laboratory setting. Further support to such notion can be found elsewhere ( [Mogil, 2012](#B138) ; [Panksepp and Panksepp, 2013](#B145) ; [Keum and Shin, 2016](#B114) ). Of course, it is not easy to state that laboratory rodents can “ feel” a conspecific as a “ feeling being.” To deliberate whether this may happen, and to what extent, is perhaps a matter of philosophy and is beyond our current purposes. We decided to limit our literature search to those test paradigms in which the following criteria are met: (1) the tested experimental subjects are more than just one; and (2) the behavior observed is not just a reaction to spontaneous behavior expressed by (an)other subject(s).

The second criterion was intended to exclude all classical social interactions. Many tests are performed with laboratory rodents to measure e. g., affiliative/playful profiles, aggressive/competitive traits, sexual attraction and approach. However, in all these cases, the behavior shown by one subject does not require this subject to perceive other subjects as “ agents that feel.” Or, alternatively, it is not essential for the one subject to capture the feeling felt by the other subject. In other words, many social behaviors in animals can well be expressed as an instinct or occurring *via* automatic routines. We rather sought to focus on those (more complex) behaviors where a focal subject, in order to foresee which next action to perform in favor of the conspecific, needs to grasp and interpreter its mood; alternatively, the reaction of the focal subject is depending on perceiving the conspecific’s action as “ deliberate” or “ causal.”

The above reasoning implies that phenomena of “ contagion” are beyond the present purposes. There are quite simple settings where a demonstrator is placed when suffering from social stress or pain, and the observing subjects are evaluated. Usually, the latter quickly begin to show a response that is similar to the observed one (see e. g., [Carnevali et al., 2017](#B39) ; [Laviola et al., 2017](#B118) ). These situations resemble the vicarious experiences of pain or emotions described in humans in paragraph 3. 1; yet, it can be questioned whether this represents a real form of metacognition. The fact that a rodent becomes more prone to stress or pain, when observing another conspecific already showing such signs, does not imply that the subject is recognizing the other as a “ subject that can feel stress or pain.” We are aware of the limitations related to the work with rodents: all internal states can only be inferred from what animal subjects do. A logical consequence of this assumption is that we can explore what rodents do in favor of each other, in search of abilities reminescent of mentalization, adopting protocols similar to those described for humans. In paragraph 3. 2, we are interested in paradigms whereby animal subjects can perform actions in favor of (an)other conspecific(s). These actors—given the situation—are then recognized as “ agents” who may either promote a helpful action or benefit from that action. According to these premises, test paradigms are only taken into account if they allow situations such as “ helping” or “ donating.” The asymmetry is implicit, in that one subject will act as a “ donor” while other(s) will simply be the recipient(s). Yet, from the side of the donor, the willingness to donate may be modulated by the “ mood” (i. e., the inner state) of the recipient, or at least by the “ empathic” perception (of the recipient by the donor): some recipients may be eliciting more (or less) donations, in the same donor, meaning that empathy is involved.

On the other hand, rodents are also capable of a range of higher-order social skills, like cooperation and reciprocity: these imply that the “ other” is somewhat perceived as “ someone” that can act prosocially (or not) and perform (or not) actions to reciprocal benefit. It is assumed that such a reciprocal kind of prosocial behavior is reinforced only if benefit to *another* subject is perceived as still rewarding by the *one* subject.

## Theoretical Framework and Constructs

Despite being an important concept for social life, it is not easy to define the word “ empathy.” This term has been used by philosophers, sociologists, ethologists, psychologists and neuroscientists in different ways. The greek etymology is from the verb “ en-pathein” that means “ feeling from inside.” In the common language, the meaning of empathy is associated to the capacity of sharing emotional states with other people: this involves the ability of getting into their point of view and understanding their own way to evaluate things and situations.

In his book “ Zero degrees of empathy,” [Baron-Cohen (2011](#B9) , p. 11) defines empathy as “ the ability to understand what the other one is thinking or feeling, and to react to his/her thoughts and feelings with the appropriate feeling and—hence—deliberation. This definition includes the correct recognition of emotional and mental states of the other, as well as reacting with the consequent motivational drive.

A central nucleus of the debate concerning the structure of empathy addresses, more specifically, the investigation of all components that are necessary for its definition. The research on empathy has been divided for very long time in two different perspectives: one that considers empathy as a purely affective experience (affective empathy), and another one that considers empathy in its cognitive nature (cognitive empathy). The theories that give priority to the affective aspects in defining empathy were mainly shared by social and clinical psychologist, and represented the only possible approach until the middle years of the last century.

Psychoanalysts and psychotherapists ( [Rogers, 1959](#B157) ; [Kohut, 1984](#B117) ) share a perspective according to which empathy can be described like a semi-voluntary process of emotive activation, while for other authors it is innate: this process involves the sharing of emotional experiences by others. According to this approach, to empathize with someone means: (1) to share the emotions that the other one lives; or (2) experiencing the same emotion, although vicariously. Such a position gives importance to the affective aspects of empathy.

In the social field, the empathic experience has been defined by [Allport (1937)](#B6) as an immediate sharing of the mental state of the other, or an “ immediate sympathetic response,” followed by instinctively assuming the posture and the facial expression observed in the other. An important contribution, useful to study the conscious mechanisms of empathic sharing, came from [Mead and Mind (1934)](#B132) . He defined empathy as the ability to “ get in the role of the other.” Mead’s definition has been a fundamental pillar for all authors that, starting from the 50s–60s, decided to study the cognitive process of thought and empathy.

Starting from the 60s, the word “ empathy” has gained a mainly completely cognitive meaning: authors have turned the object of their studies into the role of cognition and of thought in the empathic responses. Even previous studies (see [Dymond, 1949](#B63) ) introduced the term of “ role-taking” to indicate the comprehension of another’s point of view. When used in this way, the term “ empathy” can be considered as a synonymous of the “ assumption of role” or of “ perspective taking.” Following this line of thought, [Hogan (1969)](#B103) defined empathy as the ability of understanding the condition and the mental state and the subjective experience of others. In this formulation, empathy is a purely cognitive skill that does not require feeling (simultaneously) others’ emotional states.

This specific vision of empathy is shared by the theory about metacognition of mind, as defined by [Gallagher and Frith (2003](#B82) , p. 77): this concept refers to the ability to attribute mental states (such as beliefs, emotions, desires, intentions) to ourselves and to others as well as to forecast, based on these inferences, future behavior in ourselves and by others. From the 80-s, both the affective and the cognitive components of empathy, have been considered as co-occurring to generate an empathic response. Authors have started to talk of multi-factorial models of empathy.

According to the just-mentioned visions, empathy phenomena would only be possible thanks to the integrated action of both affective and cognitive components. The latter intervenes and contributes at very different levels, also depending on the complexity of the situation and on the level of analysis operated by the subject. This multi-factorial concept comprises different levels of “ sharing,” from the primitive and automatic forms (similar to emotional contagion) to the most evolved forms.

According to [Feshbach (1983)](#B75) , empathy consists of three elements: the ability of discriminate and correctly recognize the emotions expressed by the other; the ability to assume the perspective and the role of the other; self-modulation, by adhering to the necessities of the other, following a “ moral” directive gained through empathy. [Hoffman (1987](#B102) , p. 479), one of the most important authors in this field, has defined empathy as an affective response that is “ most appropriate to the situation of another person than to own one.” Feeling empathy does not mean to experience exactly what the other lives, but to understand and/or to share in a vicarious way the emotions and needs he feels. According to such multi-factorial models, empathy is defined as a process that involves and integrates cognitive and affective components; in fact, it requires the comprehension of the mental and emotional state of the other as well as the ability to respond in the most adaptive way.

Starting from the work of [Hoffman (1987)](#B102) and [Batson et al. (1991)](#B16) , “ empathy” has been defined as “ an affective response that is identical or very similar to what the other person experiences or is about to experience in a specific context” and “ a response that is triggered by the understanding of the mental state of another person.” To capture these multi-faceted concept, [Davis (1980)](#B53) proposes a multi-dimensional definition of empathy and a new tool for its measure, termed IRI (Interpersonal Reactivity Index).

In order to realize an integration between cognitive and emotional factors within a single definition, we can cite the definition of empathy formulated by [Bischof-Köhler (1991)](#B20) . For this author, empathy is “ the ability of understanding others through the direct sharing and immediate experience of their affective state, where the subject that empathizes keeps the consciousness that his current affective state is indeed mirroring the affective state of the other.” This definition entails a couple of fundamental advantages: on the one side, it allows differentiating empathy from most primitive forms of emotional contagion; on the other hand, it allows distinguishing the “ true” empathy from the more cognitive psychological processes for understanding others (ranging from perspective taking to the Theory of Mind).

Similarly, according to [Ickes (1997)](#B105) , empathy is a multi-dimensional construct: it can be defined as a complex form of psychological inference in which observation, memory, knowledge, and reasoning are combined together, in order to penetrate the thoughts and feelings of others. The definition by Ickers effectively captures the multi-dimensional nature of empathy and clearly refers its cognitive aspect, namely the comprehension of mental states of the other. In other words, such definition fully comprises the ability of mentalization. The question of the relative overlap between the concepts of empathy and mentalization remains, here, a still-open theoretical point as well as a subject of further study.

As seen above, the cognitive component of empathy consists in the comprehension of a situation from the point of view of another person. This implies taking into account that any other person reacts to one given situation based on his/her beliefs, goals and intentions that can well be different from our own ones. This ability requires skills described by the “ Theory of Mind”: when written with capital “ T” and “ M,” this phrasing indicates processes underpinning specific aspects of the meta-cognitive knowledge. Specifically, we refer here to the ability to recognize ourselves and the others as thinking entities. This means to attribute, to ourselves and to others, specific mental states (e. g., emotions, wishes, beliefs and intentions) and predict future behavior, based on this information.

The origins of the theory of metacognition are linked to work by two primatologists ( [Premack and Woodruff, 1978](#B148) ) who investigated the ability of chimpanzees to estimate the behavior of a human actor in an experimental paradigm. Some years ago, taking up the ideas of [Premack and Woodruff (1978)](#B148) and [Wimmer and Perner (1983)](#B192) set up an experimental paradigm intended to evaluate the ability of a baby to understand that different people can have a different thought relative to the very same situation. The resolution of this task by children provided the most convincing proof of the capacity to understand the mental state of others.

Different authors during recent years have focused their attention on the ontogeny of the ability to assign a mental state to the others, trying to offer a clear definition of “ metacognition of mind.” Despite many authors do agree on the developmental steps that bring to a mature development of a Theory of Mind, different positions still exist when trying to identify the mechanisms and fundamental process subserving these mentalization abilities. The theoretical investigators proposed that a child develops the knowledge on the (own and others’) mind simply on the basis of experience ( [Botterill, 1996](#B26) ; [Gopnik, 1996](#B95) ). This theory supports that ontogeny of the mentalization skills goes along with the mental progress that the child realizes in other fields of knowledge (e. g., communication and language) and depends on the maturation of his/her abilities of representation and symbolization. Studies on linguistic acquisitions in the 1st 2 years of life demonstrate that the onset of interactional communication, the use of communicative gestures and of gaze in joint attention can be considered as precursors of the mentalization ability, allowing to understand the intentional behavior and the mental state of others ( [Pizzuto and Capobianco, 2008](#B147) ; [Capobianco et al., 2017](#B36) ).

In particular, the onset (around 9–12 months) and the use of deictic gestures and of gesture-speech combinations are very important predictors of early development for both Theory of Mind and empathy in children. When a child points by alternating the gaze between an object and the adult, (s)he demonstrates that (s)he already understands an early level of the Theory of Mind: the agentivity of adults. In other words, (s)he knows that the adults’ behaviors can be modified according to his/her purposes. With further development, the child will use the pointing not only to require but also to share an object with the adult (“ Declarative Pointing,” see e. g., [Capobianco and Cerniglia, 2017](#B32) ; [Bartolomeo et al., 2018](#B14) ).

Many studies observe that the gesture-speech combinations encode different information, which represent an important portion of children’s production and play a significant role during the transition from one- to two- and multi-word speech. Recent results on 12 typically developing Italian children, observed monthly between 10 and 25 months, have highlighted that both combination types show a significant developmental pattern during the 2nd year of life, each providing specific contributions in language and communication development. Overall, this entails understanding of mental states and developing a typical interaction and a secure attachment between the mother and child ( [Capobianco and Cerniglia, 2018a](#B34) , [b](#B35) ; [Cimino et al., 2018](#B200) ).

According to “ the theory of simulation” ( [Goldman, 1992](#B93) ; [Harris, 1996](#B100) ; [Tomasello, 1999](#B187) ), the comprehension of the mind of others is not founded on a complex psychological knowledge of beliefs and wishes of others; rather, it is rather linked to the ability assume others’ standpoint. Accordingly, to deduce the mental state of others would consist in the simulation of the world as experienced by the other. An empirical support to this theory has come from the discovery of mirror neurons: it has been shown that the same neural circuit, involved in the control of action and subjective experience of emotions, is active even when we observe the same actions and emotions while they are accomplished and expressed by others. Starting from this point, [Gallese and Goldman (1998)](#B84) have supposed that the human ability of reading the mental states of others can have evolved rooting on the mirror neurons circuit in primates.

The approach by [Baron-Cohen (2011)](#B9) supports—instead—an instinctual vision for the Theory of Mind and affirms the existence of “ specific, genetically-determined and autonomously-functioning mechanisms.” For Baron-Cohen, these “ modules” always develop according to a maturation process in which environmental influence is minimized. The development of the comprehension of others’ mind is determined by the activation of a module underpinning the Theory of Mind, termed (toMM, Theory of Mind Module). Such module helps to represent the comparison between one’s own mental state and the others’ mental states, related for instance to the representation of things. Empirical evidences are given by studies on children affected by autism, or other diseases that seem to be characterized by a deficit of those modules.

Finally, we have to mention the integrated socio-constructive approach, supported by authors like [Bruner and Feldman (1993)](#B28) . This interactionist and relational approach refuses the classical individualistic perspective in favor of a Theory of Mind that develops in a context of sequences of social interactions. Research by [Fonagy and Target (1996)](#B79) shares such an integrated approach. According to them, mentalization is defined as the ability to understand the interpersonal behavior in terms of shared mental states. This is a fundamental key for the organization of the Self; such affective regulation is acquired in the field of the early relationships of attachment, while all the psychological processes consequent to the ability to mentalize are indicated with the term “ reflective function.”

## Neurobiological and Physiological Correlates

### Endocrine and Hormonal Correlates

International research on the neurobiological and physiological bases of empathy agrees on its biological substrate, in particular, supported by the endocrine system ( [Preston and de Waal, 2002](#B149) ; [Yildirim and Derksen, 2012](#B196) ; [Gonzalez-Liencres et al., 2013](#B94) ). In this regard, recent studies highlighted some interesting neurobiological elements related to sex ( [Davis, 1983](#B54) ; [Baron-Cohen et al., 2001](#B12) ; [Derntl et al., 2010](#B57) ). In fact, several studies showed that women usually get higher results in tasks and social activities in which empathy (both cognitive and emotional) is involved, while men are more likely to suffer from problems characterized by low (or no) empathy, such as psychopathy for instance ( [Cale and Lilienfeld, 2002](#B31) ).

Such evidence can be better understood considering the presence of higher levels of testosterone in men compared to women; this difference could partly explain the variability in empathy related to sex. In fact, affective empathy results to be influenced by the activity of the amygdala, whose functioning, in turn, depends on testosterone levels ( [Terburg et al., 2009](#B185) ). The association between testosterone levels and empathy deficits seems to play an important role in autistic disorder which, according to one theory, is even attributable to fetal testosterone ( [Lutchmaya et al., 2002](#B127) ; [Baron-Cohen et al., 2005](#B10) ; [Chapman et al., 2006](#B44) ).

For a better understanding of the neurobiological mechanisms of empathy, it is important to highlight that, in addition to testosterone, other elements are found to play a relevant role: these include, for instance, the release of glucocorticoid under stressful conditions *via* the hypothalamus-pituitary-adrenal (HPA) axis ( [Tennes and Kreye, 1985](#B184) ; [Adam and Gunnar, 2001](#B4) ; [Stallings et al., 2001](#B181) ; [Nakayama et al., 2007](#B141) ; [Booth et al., 2008](#B24) ; [Barraza and Zak, 2009](#B13) ; [Buchanan et al., 2012](#B30) ).

Oxytocin (OXT) is a nonapeptide with a long evolutionary history and a well-established role in animal and human social cognition. Brain regions clearly associated with social cognition include amygdala, insula, medial prefrontal cortex (mPFC), anterior cingulate cortex (ACC), temporoparietal junction, superior temporal sulcus (STS) and inferior parietal lobule (IPL). Research ( [Mariscal et al., 2019](#B129) ) suggests that children with autism spectrum disorder (ASD) may suffer reduced empathy, as measured by impaired yawn contagion, compared to typically developing (TD) children. Blood concentrations of the “ social” neuro-peptide OXT positively predicted contagious yawning. These findings suggest that OXT identifies a biologically defined subset of children with ASD, who exhibit reduced empathy as measured by impaired contagious yawn.

Another study ( [Luo et al., 2019](#B126) ) tested the hypothesis that empathic neural responses to others’ pain are more flexible in G/G than A/A carriers (rs53576) of the oxytocin receptor gene (OXTR). They found that G/G carriers showed greater neural responses, over the frontal cortical region, to painful vs. neutral face expressions. In contrast, A/A carriers showed significant empathic responses only to faces with shared racial and group identity. These findings provide electrophysiological evidence for interactions between OXTR and intergroup relationships about empathy for others’ suffering. Another study ( [Acevedo et al., 2019](#B3) ) used magnetic resonance imaging (MRI) to visualize brain areas responsive to happy or sad faces: evidence shows that OXT increases the ability to recognize emotional expressions in faces. Moreover, some studies found a specific pharmacological effect of OXT administration on the recognition of happy faces, while others found a specific effect for the recognition of fear ( [Domes et al., 2007](#B60) ; [Schulze et al., 2011](#B162) ). Intranasal application of OXT has been shown to increase emotional, but not cognitive empathy, potentially by an amygdala-dependent mechanism ( [Luminet et al., 2011](#B125) ). The “ Reading the Mind in the Eyes Test” (RMET) requires the ability to detect emotional states from pictures that are exclusively depicting the eye region of individuals ( [Baron-Cohen et al., 2001](#B12) ): this test covers socially cognitive functions, namely mentalizing. In this specific paradigm, administration of 24 IU of OXT improves performance ( [Fischer-Shofty et al., 2010](#B77) ). This effect is specifically observed in individuals with a baseline of higher alexithymia or reduced empathy, and therefore points once again to the role of OXT in empathic abilities.

### The Neurobiological Bases of Empathic Ability

Studies in the field of biological basis of empathy have identified a series of neuronal systems that support the ability of subjects to enter into relationships, both between adult subjects (romantic relationship) and between parents and children (parenting relationship). In particular, embodied simulation and mentalization refer to the ability of subjects to observe, and care for, the other. In support to the relevance of these evidence, recent studies have shown that sensitive parenting influences the psychological development and well-being of children during development ( [Ammaniti et al., 2004](#B7) ; [Tambelli et al., 2015](#B183) ).

Many studies in the literature show the complementarity of mother-children neuro-biological systems ( [Lenzi et al., 2013](#B119) ). For example, studies have shown that parents are willing to take care of their children in response to the global characteristics of her/his face, which acts as a particularly attractive configuration-gestalt ( [Darwin, 1872](#B52) ; [Glocker et al., 2009](#B92) ; [Kalikow, 1983](#B111) ; [Stern, 1977](#B182) ). This seems to be linked to the tendency in adults to protect a child and help her/him to survive, transmitted during the course of evolution.

Several areas of the brain have been recognized as a biological substrate of empathy ( [Gallese and Goldman, 1998](#B84) ; [Carr et al., 2003](#B40) ; [Lenzi et al., 2013](#B119) ). Among them, the parieto-frontal cortical circuit assumes a crucial role: its activation, when a subject observes the action of another subject, demonstrates the biological basis of the ability to reflect the behavior of others and the related emotional content (see [Rizzolatti and Craighero, 2004](#B153) ; [Rizzolatti and Sinigaglia, 2010](#B154) ).

As anticipated above, the neuroscientific research on empathy is highly influenced by discoveries of a group led by Giacomo Rizzolatti who, in the middle of the 90s, described a population of neurons in the pre-motor area F5 of the macaques’ brain, afterwards called mirror neurons ( [Gallese et al., 1996](#B90) ; [Rizzolatti et al., 1996](#B155) ). These were activated when the ape performed an action with the hand but also when the ape observed the same action performed by another one. This discovery showed the presence, in the primates’ brain, of a transduction mechanism: when apes observe the others’ actions, mirror neurons have been demonstrated to be elicited also in terms of anticipating the corresponding motor control. As such, this mirror system transforms what they see in immediate knowledge on what to do.

Transcranial magnetic stimulation (TMS) data in humans have been used to infer the role of mirror neurons. [Fadiga et al. (2005)](#B69) suggest that, when a subject observes another individual while producing some action, (s)he is somewhat stimulated to activate the same muscles. [Gallese and Goldman (1998)](#B84) suggested that the mirror neurons’ system can constitute the neurobiological basis which underlies our ability to understand the others and to simulate automatically their possible actions and intentions. There is a wide neuroscientific literature, nowadays, supporting the idea that simulative processes could constitute a basis of our social brain and our ability to empathize with other people ( [Preston and de Waal, 2002](#B149) ; [Gallese et al., 2004](#B91) ). The mechanism that enables the sharing of a mental state with other people has been extended to the ability of sharing feelings and sensations ( [Gallese et al., 2004](#B91) ). To support this hypothesis, neuroimaging studies have investigated the brain activity associated to different empathic responses in the field of contact, smell and pain. These studies will be briefly reviewed here.

One of the first affective state that (as shown by neuroscientists) implies a form of empathic comprehension is the *disgust* . The neurons of the anterior insula (AI), a portion of brain cortex are activated when the subject shows disgust to olfactory and gustatory stimuli. Studies ( [Wicker et al., 2003](#B191) ) have shown that the AI is activated not only in presence of these stimuli but also when we are seeing other subjects that show facial expressions of disgust. This mirror role of the insula has been confirmed with neuroimaging and clinical studies: it has been observed that a lesion of the insula not only causes an impairment in the abilities of feeling disgust but also impedes the ability to recognize the facial expression of disgust by others ( [Gallese, 2001](#B85) ). According to Rizzolatti and Sinigaglia, this result suggests that the real comprehension of disgust by others is not based on an inferential cognitive process but on a mirror empathic process. The studies on touch have shown similar evidence for the activation of the somato-sensory cortex, involved in the elaboration and in the perception of touching. This happens both when subjects are touched by others but also when they observe other people that are touched ( [Keysers et al., 2004](#B115) ).

Very interesting results were obtained, by researchers, on the capacity of empathy to feel others’ pain. It has been posited that a complex brain circuit called “ matrix of pain” is involved in the experience of pain. The matrix of pain includes the primary and secondary somato-sensory cortex, the cerebellum, motor and premotor cortices. The affective knot of the matrix includes the ACC and the insular cortex (IC): these are phylogenetically ancient regions, classically considered components of the limbic system and of the visceral brain ( [Papez, 1937](#B146) ; [MacLean, 1949](#B128) ). [Singer et al. (2004)](#B174) measured the activation of such neural structure in 16 women to which the painful stimuli (slight electric shocks on the hand) were given in a first moment; in a second moment, they watched the administration of similar painful stimuli to their fellow subjects. Some brain areas, such as the front of the insula, the ACC, the cerebellum and the brainstem, activated both during one’s own experience of pain and in the perception of the pain in others; instead, other areas as the primary and secondary somato-sensory cortices, activated just by own pain. The authors concluded that empathic experience of others’ pain consists in sharing their subjective experience of pain, without sharing the sensorial elements.

A recent neuro-imaging study ( [Novembre et al., 2015](#B142) ) shows that the social pain involved the posterior insular cortex, a brain area traditionally associated with the elaboration of physic pain: the experience of social pain would occur both when we feel it in first person but also when we experience it in an empathic way. Another indication of a neural activity linked to pain, in the absence of physical painful stimulation, came from a neuro-imaging study ( [Osaka et al., 2004](#B143) ). In an experimental paradigm, the recruited subjects listened to onomatopoeic words in Japanese that evoked pain; subsequent to this an increased magnetic-resonance signal within the ACC was observed, which suggests a real affective representation of that pain ( [Singer and Fehr, 2005](#B172) ).

Importantly, research on the above neurobiological correlates of human empathy has significantly contributed to the development of a bio-behavioral synchrony model ( [Feldman et al., 2012](#B73) ) positing that parental (mature) psycho-physiological system externally regulate the infants’ immature system by coordinating and modulating biological and behavioral processes during social contact. It has been shown that multiple neuro-physiological systems (e. g., hearth rate coupling, brain-to-brain synchrony in neural oscillations, endocrine system) are anchored in behavioral synchrony and enhance the moments of attunement and concordance between parents and children during verbal and non-verbal interactions ( [Levy and Wagner, 2011](#B120) ; [Cerniglia et al., 2018](#B43) ).

In line with this perspective, evolutionary theories, psychoanalytic theories of affect, developmental theories, social learning theories and attachment theory, all agree that parental empathy is the key component of sensitive parenting and, therefore, of children’s social adaptation ( [Abraham et al., 2018](#B2) ). Several crucial capacities in children (e. g., affect regulation, socialization, cognitive functioning, symbolic competence) are predicted and supported by their parents’ empathic ability ( [Feldman, 2007](#B72) ), which enables offspring to feel secure, show compassion towards others, and recognize their own thoughts and feeling ( [Fonagy et al., 2007](#B80) ). By modulating physiological response to stressing signals and experiences (e. g., modulating the release of cortisol in children), sensitive and emphatic parenting supports emotion regulation processes in children ( [Knafo and Plomin, 2006](#B116) ; [Muratori et al., 2011](#B140) ), and in general is associated with a positive development of both cognitive and affective empathy in children ( [Miklikowska et al., 2011](#B136) ; [Soenens et al., 2016](#B177) ). In fact, a positive caregiving can facilitate the emotional growth of children and adolescents: they will become able to consider the point of view of their peers, to care for their experiences, and to engage in prosocial behaviors ( [Hastings et al., 2007](#B101) ). Many authors have highlighted the link between difficulties in empathy and difficulties in the relationship between parents and offspring, with particular reference to the children’s problems in maintaining a visual contact and joint attention with their parents ( [Dadds et al., 2014](#B49) ; [Billeci et al., 2019](#B19) ). Following these considerations, several clinical psychologists and researchers have linked children’s empathy deficits to negative experiences with caregivers, including forms of maltreatment. Empirical data about these aspects are not conclusive: some children and adults who have experienced maltreatment show normal or even high emotional capacities, whereas others display emotional and cognitive deficits ( [Dadds et al., 2012](#B50) ). The fact that data collected in this field of studies are not homogeneous highlights the particular complexity of the construct of empathy that can be influenced by genetic and environmental variables ( [Dadds et al., 2009](#B51) ).

Most importantly, human capacity of empathizing is crucial for all social interactions, not only those between children and their parents ( [Eisenberg and Miller, 1987](#B67) ). Bio-behavioral synchrony, the coordination of biological and behavioral processes between attachment partners during social contact, is a critical component of all human attachments (parental, romantic, friendship, and fellow–human interactions), although synchronous interactions experienced during early sensitive periods are expressed in later attachments throughout life. Dopamine striatal neurons are suggested to support this synchrony because their temporal sensitivity may allow humans differentiating reward that stems from self and/or partner, anticipating predictable social interactions, and drawing reward from the matching of self and partner’s actions ( [Báez-Mendoza and Schultz, 2013](#B8) ). Recent studies tested the neural basis of humans’ multiple attachments by measuring brain responses to auditory, visual, or multimodal stimuli of the attachment target (e. g., infants to their parents, romantic partners, or friends). It has been suggested that the VTA (ventral tegmental area)/ventral striatum ignites the brain’s experience of attachment and imbues it with reward and vigor, and that human attachments integrate subcortical with multiple cortical reward- and socio-cognitive-related networks ( [Lieberwirth and Wang, 2016](#B122) ). On one side, the fronto-striatal connections of the reward circuit, particularly the ventromedial prefrontal cortex (vmPFC), the orbitofrontal cortex (OFC), and the ACC seem to support human capacity of establishing and maintaining intimate relationships by sustaining bio-physiological and psychological synchronicity and empathy ( [Abraham et al., 2016](#B1) ). On the other side, the embodied simulation/empathy network, which includes the insula, the ACC, the inferior frontal gyrus (IFG), the IPL, and the supplementary motor area (SMA), is suggested to be critical for grounding a “ shared world” in the brain underpinning the human capacity to build and maintain attachments ( [Frith and Frith, 2006](#B81) ). Besides, human bonds also rely on higher-order (mentalizing) cognitive processes, supported by fronto-temporal–parietal structures, specifically the STS, the posterior cingulated cortex, the temporo-parietal junction (TPJ), the temporal pole, and the mPFC.

Apart from these studies, a particular type of cell has been also observed, called the von Economon’s neurons that are present only in the ACC and in the AI of humans. [Allman et al. (2011)](#B5) believes that this neuron could constitute a neural basis for mechanisms underpinning knowledge about the one’s own and others’ roles and interactions within complex social settings. Thus, the same structures that play a decisive role in the representation of own affective states (like AI, ACC) seem to have a decisive role in the comprehension of the affective state by others.

During the last 10 years, many studies with functional neuro-imaging on healthy adults have focused on investigation of the ability to mentalize ( [Ruby and Decety, 2003](#B158) ; [Singer, 2006](#B173) ). The results have shown the existence of a neural network underpinning the ability to understand and predict the behavior of others, possibly through the assignment to them of mental states. This network includes three brain areas that are greatly involved: TPj and temporal poles (TP), upper posterior temporal sulcus (pSTS) and, above all, the mPFC ( [Farrow et al., 2001](#B71) ). Given the heterogeneity of methods, and of the demographic characteristics of participants recruited in these studies, it is remarkable that the results converge on the same areas of the brain. According to the vision currently shared in literature, the three brain area supporting the Theory of Mind play different functions: the STS could be responsible for detecting and recognizing the biological movement of others, as the direction of gaze, the reading of lips and the movement of body, hands and lips; the TP, associated to memory processes, give the semantic context in which stimuli are elaborated; finally, the mPFC is implicated in the subsequent analysis of such stimuli and produce explicit representations of one’s own and others’ mental states (see [Fletcher et al., 1995](#B78) ; [Gallagher et al., 2000](#B83) ; [Cuff et al., 2016](#B47) ).

## Emotions vs. Cognition (or Both)?

### The Subcomponents Within Theory-of-Mind

Although the topics dealt with by the theory about metacognition of mind have gained attention in cognitive neuroscience during the last 20 years, only recently the experimental research started to examine subcomponents of such complex activity. Therefore, it is important to be aware that the different roles of its neural correlates are currently investigated, at least for what concerns the prefrontal cortex (PFC). [Shamay-Tsoory et al. (2009](#B167) , [2010)](#B168) proposed a model for the Theory of Mind in which two separated systems were involved: one elaborates the inferences on beliefs and intentions of others (Cognitive Theory of Mind) and the other realizes an inference of emotions and feeling by other people (Affective Theory of Mind).

A series of studies on lesions ( [Shamay-Tsoory and Aharon-Peretz, 2007](#B166) ) have shown that patients with lesions to the vmPFC have a deficit in tasks that involve an affective component, as the comprehension of the affective state hidden behind an ironic comment in a non-affective context. These observations support the hypothesis that vmPFC could be necessary for the affective empathy but not for the cognitive one. This area has many connections with regions involved in the affective elaboration (amygdala, AI and TP). Thus, it may integrate cognitive and affective information. A support to this hypothesis comes from the studies on patients with psychosis (see [Shamay-Tsoory et al., 2010](#B168) ): a deficit in empathy was shown in correspondence to dysfunctions of the amygdala and of the OFC.

One neuroimaging study explored the overlapping neural correlates of cognitive and affective Theory of Mind, hypothesized by Shamay-Tsoory. The study (see [Sebastian et al., 2011](#B164) ) has inquired into the relationship between the affective and cognitive subcomponents with illustrations that predicted separate intentional conditions for the cognitive vs. affective mind, plus a condition to control for physical causality. Results have shown that both cognitive and affective capability of inferring others’ mental states activate brain areas, such as the posterior STS, the TPj, precuneus, and temporal lobes, thus confirming the model ( [Shamay-Tsoory et al., 2010](#B168) ).

An important contribution about PFC’s different roles in Theory of Mind was given by a recent study ( [Xi et al., 2011](#B194) ), who used a complex task, the Faux Pass Test ( [Baron-Cohen et al., 1999](#B11) ), that comprehends both affective and cognitive subcomponents. This test puts participants in the position of reading a story in which the statement-making character does not realize that he should not have expressed such affirmation and that listeners could have felt offended. Subsequently, participants are given six questions, for evaluating their ability to recognize the Faux Pass as well as their empathic capabilities and mental representation.

Results of the study evidenced that patients having vmPFC lesions did not recognize the gaffe correctly (affective component of the test). Conversely, patients with dorsal/lateral PFC lesions recognized the gaffe correctly but did not manage to infer the other’s mental state (cognitive component of the test). The affective mind also involved the vmPFC, providing a validation of the hypothesis that affective Theory of Mind requires additional elaboration compared to cognitive Theory of Mind: vmPFC’s activation reflects the integration of cognitive and affective information.

### Differences and Overlaps Between Empathy and Theory-of-Mind

In neuroscientific and cognitive literature, there is often a clear difference between the Theory of Mind and empathy. On one side, empathy concerns our ability to share affective states with others; on the other hand, Theory of Mind represents our ability to interpret their mental state, their intentions and beliefs ( [Blair et al., 1996](#B21) ). The distinction between the two concepts is supported by the evidence of social impairments in people with psychosis: indeed, people with a diagnosis of schizophrenia or under drug-induced psychosis show inability to empathize but they do not show any deficit in mentalizing. The dichotomy between empathy and Theory of Mind distinguishes between intentions and beliefs, on one side, and emotions on another side. Rational vs. affective domains of life are indeed the two poles if such a dichotomy.

According to this model (see also above, “ Endocrine and Hormonal Correlates” section), the affective empathy, after a sensory perception of emotions in others, occurs through the same neural mechanisms that would be activated and are responsible for the emotional experience in first person; thus, empathy would be based on mirroring processes like emotional contagion, shared pain, recognition of emotions. The cognitive facet of empathy, instead, consists of the ability to “ infer”—using cognitive skills—the emotional state of others: according to the model by Shamay–Tsoory, mentalization involves both cognitive and affective processes, and specifically, affective empathy refers to the possibility of inferring others’ emotions.

With reference to the studies of Shamay–Tsoory, empathy is composed by two components, affective and cognitive, while the Theory of Mind is referred to our ability to get inferences on others’ beliefs but also emotions. By functional neuro-imaging [Schnell et al. (2011)](#B161) investigated neural correlates of “ cognitive” empathy as defined by [Preston and de Waal (2002)](#B149) . These authors highlighted how this mechanism of emotional representation is still relatively unexplored and connects to the (cognitive) domain of the mentalization system.

A differential study was realized with illustrated stories of faux beliefs in first and third person, in which all direct signals depicting affective states (like the face’s expression of all characters of the story) have been deleted. Results confirmed the hypothesis that cognitive empathy involves, at the same time, the neo-cortical network of mentalization and the limbic areas involved in the experience of emotion in first person. This implies that cognitive empathy entails a “ simulated” generation of an affective state through activation of the limbic system, in the absence of an external (direct) emotional signal.

The general involvement of simulation in mentalization is moreover supported by studies of [Lombardo et al. (2009)](#B124) . Limbic simulation can explain the mirroring of others, i. e., elaboration of affective empathy; mentalization, in its affective and cognitive subcomponents, could support cognitive empathy. A reduction of empathy, according to this model, could be caused by a deficit of either mirroring (limbic simulation) or affective inference (mentalization), that are mediated by different neural systems.

From a neurobiological point of view, when humans are required to solve cognitive tasks that imply empathy or Theory of Mind, different neural structures are activated. Shamay-Tsoory’s theory distinguishes two different capacities for the comprehension of others. Such two forms of comprehension are interconnected, from both a psychological and a neural viewpoint, since neo-cortical or limbic brain areas often interact with each other.

[Völlm et al. (2006)](#B190) studied this overlap (in neural correlates of empathy and Theory of Mind) and found that the comprehension of others’ emotional states required the involvement of other interconnected areas, in particular, the amygdala and the orbital–PFC.

### The Full Interaction Between Empathizing and Mentalizing

According to the model proposed by Shamay–Tsoory, our human interpersonal relationships in normal social circumstances would activate, independently, both a mirroring of the emotional response (emotional empathy) and a cognitive evaluation of the others’ affective mental state (cognitive empathy; [Shamay-Tsoory et al., 2009](#B167) ). Several evidence seem to confirm the existence of such different pathways, acting in parallel and relatively autonomous, allowing the full experience of empathy ( [Eres et al., 2015](#B68) ; [Valk et al., 2016](#B188) ).

In line with a “ theorethical/deductive” approach, people use mind deductions to attribute intentional states to others; that is, they adopt a fully theoretical-reasoning approach for the comprehension of others’ mental state, relying onto a mechanism of mentalization. According to the “ Theory of Simulation,” instead, we attribute mental states through a process of intuitive mirroring. This perspective emphasizes the use of shared representations: thanks to the innate faculties of observation-execution, the emotion felt by others is shared empathically. The system of mirror neurons would be particularly suited for providing a simple mechanism of motor empathy, like for imitations and emotional contagion. In an independent way, the deductive processes would be activated to infer the perspective and the mental state of others: it would require the ability of distinction between self-else and recruit the autobiographic memory. Therefore, a final cognitive empathic response would be generated, involving—in parallel—the cognitive mind (mPFC, STS, TP) and the affective mind (vmPFC, OFC). On the contrary, an emotional (“ instinctive”) empathic response is simply guided by simulation and involves the same regions that mediate the first-person emotional experience (amygdala, insula).

Even if the emotional and cognitive components of empathy work autonomously, every empathic response would activate the same components, as a function of different variables including social context, the level of stress ( [Jackson et al., 2005](#B108) ) and the perceived similarity between the observing subject and the observed person ( [Mitchell et al., 2005](#B202) ). Therefore, although both the affective and cognitive components of empathy can work independently, in any natural social situation every empathic response involves both components ( [Zaki and Ochsner, 2012](#B197) ). We can assume that the balanced activation of such networks is fundamental for an adaptive social behavior. As the system of mirror neurons seems to be of a lower level, one could think that its activation could represent a prerequisite for the activation of the more complex mentalization system.

[Van Overwalle and Baetens (2009)](#B189) have suggested that the mirror neurons and the mentalization systems are rarely active at the same time: it is possible that the mirror-neuron system provides a fast and intuitive input to the mentalization system. But social interactions themselves represent the privileged condition to explore the potential connection between the two systems. Social interactions are in fact characterized by a continuous exchange of roles between actors; this exchange, of course, not only the comprehension of the other’s action but also the capability to infer other’s intentions. To the purpose of studying this dynamics, which seems to require a connection between mirror neurons and mentalization systems, [Sperduti et al. (2014)](#B178) realized a study in which two subjects underwent functional magnetic resonance while involved in a reciprocal game of imitation of each other’s hands. The two participants could observe each other’s hand movements and they were free to choose whether to imitate each other’s movements or to freely produce some gestures at any moment. To produce a harmonic exchange of roles, each subject needed to anticipate the other’s intentions. Results have evidenced a relationship between the functioning of mirror neurons and mentalization systems. During active (imitating) and passive (to be imitated) “ imitation” conditions, when compared to a passive observation condition, functional neuroimaging has shown a major activation in the IFG concurrently with ventral and dorso-medial prefrontal cortices and with the ACC.

In sum, the mirror-neuron system is involved into the preparation of one’s own actions and into the simulation of the other’s actions, while the mentalization system is likely involved in higher-level cognitive processes, that allow the anticipation of the other’s intentions, a fundamental step in the co-regulation of reciprocal action. A recent review ( [Catmur, 2015](#B42) ) has expanded on the role of mirror neurons and mentalization systems: mirror neurons would be majorly activated in the perception of action, a process that could be necessary but not sufficient for the full comprehension of the intention. Another recent research on comprehension of others’ intentions ( [Isoda, 2016](#B107) ) supports the idea that mirror-neuron system is activated by others’ visible actions, and predicts physical consequences in terms of scopes, while mentalization system would be mainly involved into the forecasting of others’ intentions. Interestingly, impending actions would be detected regardless whether others’ current actions can be observed directly or not. Isoda suggests the hypothesis that the two systems strictly collaborate to judge the meaning of the others’ actions.

A speculative conceptual model, that allows to comprise the most critical aspects of social cognition, includes formulating statements about others’ intentions and comprehending the meaning of their actions. [Tidoni and Candidi (2016)](#B186) evidence the difficulties of demonstrating that mirror neurons and mentalization areas contribute separately to the perception of actions and to the comprehension of others’ intentions. Another recent and interesting research ( [Kanske et al., 2016](#B112) ) has further looked into the relationship between emotion sharing and the comprehension of others’ mental state during social interactions. People with higher levels of empathy do not necessarily possess a greater mentalization capability. In highly emotional situations, though, empathic sharing could inhibit the activity that is bound to mentalization and therefore degrade the mentalization performance.

Data would, therefore, suggest the existence of two distinct social systems interacting with each other rather than a unitary construct of social comprehension. The AI takes on a more instinctive and visceral reaction about what’s happening to others, with a role in modulating current processes within TPJ, such as the undertaking of the others’ point of view or reasoning about the others’ mental state. In a very recent study with functional magnetic resonance, [Buades-Rotger et al. (2017)](#B29) have investigated the role of brain areas associated with aggressive or avoidant responses to interpersonal provocation. The participants are put in situations where there is choice of facing or avoiding an opponent with a high (HP) or low (LP) level of provocation. The brain areas containing mirror neurons have shown a major activation when the participant decided to face the opponent, while mentalization circuits were more involved in the avoidance decision. Moreover, avoidance of HP opponents was associated to a major activation of amygdala and to a slightly lower activation of mentalization regions (IFG, TPJ), compared to avoidance of an LP opponent. Such results would suggest that avoidant responses are associated to a minor assumption of perspective and could represent the outcome of a major anticipation of threatens. Data also suggests that the two systems play a convergent role in decisions which regard response to provocation. However, there is a necessity of causal evidence from new social-interaction studies, proposing different conditions under which each one of them may activate the other one.

### Empathy and Psychopathology

Research shows that many psychopathological forms are characterized by difficulties in empathy. In particular, [Smith (2006)](#B175) proposed a distinction into four empathy disorders. These disorders are: *cognitive empathy deficit disorder* (CEDD), characterized by low cognitive empathy and high affective empathy; *emotional empathy deficit disorder* (EEDD), characterized by low affective empathy and high cognitive empathy; *general empathy deficit disorder* (GEDD), characterized by low abilities in both cognitive and affective empathy; *general empathy surfeit disorder* (GESD), characterized by a high level of both cognitive and affective empathy.

The CEDD type of empathy disorder is associated with ASD and Asperger’s syndrome, both of which are characterized by difficulties in understanding the mental and emotional state of a subject ( [Capps et al., 1993](#B38) ; [Smith, 2009](#B176) ). But several clinicians and researchers do not agree on the presence of a unique deficit of empathy at an emotional level in autistic subjects. Also, many authors do not agree with the understanding of autism based on the Smith model, which seems to be too rigid and limiting if considering a greatest variability of phenotypic manifestations within the autistic spectrum ( [Shamay-Tsoory et al., 2002](#B165) ; [Minio-Paluello et al., 2009](#B137) ). Moreover, in this regard, several studies showed the presence—in autistic subjects—of difficulties in empathy even at the lowest levels of cognitive processing, with errors in the simple identification and recognition of facial expressions ( [McIntosh et al., 2006](#B131) ; [Clark et al., 2008](#B45) ).

The CEDD category of empathic disorder has been associated not only with autism but also with other disorders, including fronto-temporal lobar degeneration (FTLD), bipolar disorder and borderline personality disorder (BPD; [Harari et al., 2010](#B99) ; [Cusi et al., 2011](#B48) ; [Mier et al., 2013](#B135) ). In addition, the other typologies of Smith’s classification have also been associated with various disorders. More specifically, psychopathy and antisocial personality disorder (ASPD) seem to correspond to EEDD, because the subjects understand the emotions of others but are not able to socialize with them (see [Dolan and Fullam, 2004](#B58) ; [Decety and Ickes, 2009](#B56) ; [Schwenck et al., 2011](#B163) ), whereas schizophrenia seems to be categorized as GEED, as the subjects show difficulties in understanding the mental and emotional state of others from both an emotional and cognitive level ( [Bora et al., 2008](#B25) ; [Haker and Rössler, 2009](#B98) ).

Moreover, several studies in patients with brain lesions have confirmed the usefulness of distinguishing between various empathy disorders. For example, cognitive-empathy deficits were found in patients with lesions at the level of the vmPFC, while emotional-empathy deficits were found in patients with lesions in the lower frontal gyrus cortex (IFG; [Shamay-Tsoory et al., 2009](#B167) ). Still, the well-known area 44 of Broca has been identified (by neuroscientists) as neurobiological correlate of affective empathy and of the mirror neuron system, which is particularly important in sharing mental states between subjects ( [Rizzolatti, 2005](#B152) ; [Shamay-Tsoory et al., 2009](#B167) ). On the other hand, though, the association between difficulties in empathy and pathologies cannot always be univocal, as there are also a number of psycho-pathological conditions that cannot be traced back to the specific type of empathy deficit nor to the categories of the Smith’s model. This is the case, for example, of attention deficit and hyperactivity disorder (ADHD) and related worsening like e. g., conduct disorder ( [Braaten and Rosén, 2000](#B27) ).

Taken as a whole, the elements of evidence on the neurobiological foundations of empathy analyzed so far recall the need for future investigation on the sub-types of emotional and cognitive empathy, and on their characterization within different psychopathologies. This effort can turn out to be particularly useful, e. g., to organize prevention and intervention programs effectively based on the characteristics of cognitive and relational functioning of the patients.

## The Contribution by Animal Modeling

Today, the utility of animal models is recognized in neuroscience. Laboratory rodents can very easily learn how to produce a behavioral response in order to produce a given desired effect, and many “ operant” tasks have been developed. Interestingly, however, these subjects are often tested individually; alternatively, they can be tested while living in group, like e. g., within IntelliCages: however, the measured behavior usually still produces effect for the subject itself. Rodents are rarely confronted with situations in which the effect of own actions is tapping onto another subject (and vice versa). When such protocols are proposed, there are the concepts of metacognition popping up. To what extent indeed does the acting subject demonstrate, by observable actions, to select those actions based on own “ empathy”?

As far as preclinical modeling is concerned, with the term “ empathy” we mean the ability to identify the mood of other conspecifics. This is based on:

1. understanding their emotional signals, which allows to foresee in advance their intentions;

2. sharing their feelings (e. g., mirroring expressions of disgust or pain);

3. assuming their subjective perspective, i. e., role-taking.

We hereby propose that evidence of modulation in deeds by one subject based on the emotional state of the other subject in a dyad can be used to infer an empathic response. As a matter of fact, among definitions of empathy described above, it was assumed that the behavior of one subject would be influenced after getting the (correct) insight about the mood of the other subject. In other words, observable behavior may well adapt to the necessities of the other if such needs are correctly inferred. Therefore, in those paradigms whereby one subject is satisfying the needs of another conspecific, we could well modulate prosocial responses by this one subject by simply changing the conspecific or its emotional status.

Prosociality, or “ prosocial behavior,” is a construct that includes all those actions which can produce, maintain or increase the benefits (or well-being) of other individuals, often bringing lack of benefits or even disadvantages for those who perform them ( [Roche Olivar, 2002](#B156) ). For this reason, it is strongly linked to concepts of altruism and generosity that—indeed—disregard possible benefits. In naturalistic conditions as well as in the wildlife, prosocial behaviors that bring benefits to individuals related to each other are often observed: this happens because animals tend to favor the survival of other individuals who have a similar genetic background and behave so that to increase their fitness accordingly. Prosociality settings in most laboratory protocols are often measuring helpful donations, namely not related to the possibility of obtaining, with these disinterested behaviors, secondary benefits like a courtesy back in the next future.

It has long been recognized that prosocial behavior is widespread in non-human species. The first animals in which prosocial behavior was observed were bats. These nocturnal rodents normally go on excursions to get a blood meal: if an individual fails to eat, it is often fed by a conspecific who regurgitates part of her/his own meal, in favor of the fasting and hungry subject. In addition, individuals who have shown prosocial behavior in the past are fed more frequently than those who have never shown, or not yet, a prosocial behavior.

To understand how rats can be studied for prosocial choice, [Márquez et al. (2015)](#B130) developed a two-choice task in which prosocial behavior did not yield a benefit nor a cost to the focal rat. They used a double T-maze in which a focal rat controlled access to the food-baited arms both of its own maze and the recipient rat’s one. In this task, the focal rat could freely choose between one side of the maze, which yielded food only to itself (selfish choice), or the opposite side, which yielded food both to itself and to the recipient rat (prosocial choice). Rats showed a high proportion of the latter choices. By manipulating the recipient’s ability to display a preference for the baited arm, the authors found that the display of food-seeking behavior by the recipient was driving prosocial choices of the focal rat. Thus, rats provide to others access to food, in the absence of direct self-benefit.

Once established this, the next step is to ascertain whether rats can judge the entity of such donations in comparison to own reward. “ Inequity aversion” is defined as the resistance to inequitable outcomes, which occurs in humans when subjects hate inequality. In the case of lab animals, this concept implies that a focal subject is not willing to provide another subject with a reward being much greater than own one. In some instances, inequity aversion can be disadvantageous: in order to prevent another animal from receiving a superior quantity or quality of a given reward, focal animals may be willing to skip an own gain.

The apparatus exploited by [Douglas et al. (2018)](#B61) was composed of two identical operant boxes, placed side-by-side on a table: they had food cups at the center of the wall and a food-pellet dispenser connected to a lever, which released 45 mg sucrose pellets in the recessed food cup. The sample was composed of adult, male rats and a specific goal of the article was to examine how a downward shift in reward value, with food acquisition occurring in a social context, would impact individual operant responding (inequity aversion).

Animals were exposed to different conditions, including a “ together” (TOG) condition: two animals placed in the two chambers simultaneously, next to each other, could observe the other rat and hear the different tones. Following a lever press, during a trial, the subsequent outcome was a high-value reward. After the control rat completed one trial, a trial then began for the experimental rat: this schedule continued back and forth until each rat completed twenty trials. Then, a socially-negative contrast was tested in that the session began with a string of five high-magnitude rewards, followed by 15 low-magnitude rewards, for one rat. The neighbor animal continued to receive the high reward outcome. As such, the first (experimental) rat was expected to “ suffer and react” because of such inequality. To summarize the overall results, performance in this TOG condition appeared slower than the “ alone” control one, when the experimental rat completed the session with no subject nor program running in the adjacent cage.

Although this approach does not address prosocial behavior as no donation is involved, it is quite evident that the experimental rat, while observing the neighbor rat to perform lever pressing and eating the food pellets, and while considering magnitudes of this reward compared to own one, shall be supposed to express a metacognitive skill. This rat is somewhat detecting the intentional actions performed by the other rat to earn food and somewhat considering the level of reward and/or of satiety states, gained by the other compared to own one; as such, elements of mentalization and/or of empathy are somewhat involved.

As far as empathy-driven responses are concerned, some experiments were recently run to observe whether rats were motivated to help a conspecific that showed signs of stress. In particular, in the study by [Ben-Ami Bartal et al. (2011)](#B17) , a free rat was placed in an arena where a companion was trapped in a container that could only be opened from the outside. After a few sessions, the free rat learned to intentionally and quickly open the container to help a cage mate. In support of the hypothesis that the intention of the rat was indeed to release the trapped mate, it was observed that rats did not open the container when it was empty or with an inanimate object inside. In addition, rats released their cage mates even when social contact was prevented. Therefore, rats behave prosocially in response to the suffering of a conspecific: as a matter of fact, the subject shall *first* detect the suffering of the other and *then* act to help it. These studies are hence providing important evidence of the biological roots of prosociality as an empathically-motivated behavior.

[Silberberg et al. (2014)](#B171) devised another similar experiment, in which a rat could release a conspecific trapped in a tube by allowing it access to a space located quite away. This protocol was chosen to exclude that the free rat could release the conspecific just to seek for social contact. It rather emerged that free rats are empathically motivated to reduce the discomfort of the trapped rat also when they become very distant after the latter issuing the tube.

[Sato et al. (2015)](#B160) evaluated whether rats do help stressed conspecifics who are wet with water. In a first experiment, rats quickly learned to open the door, allowing the wet trapped rat to enter the safe area escaping from the water area. In addition, it was seen that rats that had previously been wet learned faster to open the door to the trapped partner than rats that had never been wet. This, probably, may happen because the former were able to recognize the unpleasant situation of the latter conspecific. In a second experiment, rats had to choose between opening the door to help the stressed partner and opening a different door to obtain a food reward. In most cases, rats would choose to help their partner before they got their food reward. These results suggest that rats behave prosocially; their intention to help may be motivated by states of emotional empathy towards stressed companions.

Also, in the rat there are evidence of reciprocity in prosocial behavior. In pioneer study, [Rutte and Taborsky (2007)](#B159) showed that the cooperative behavior of rat females is influenced by having previously received help, regardless of the identity of the partner. Rats were trained to pull a stick so to provide food to the partner. Test subjects threw it more often for an unknown partner when they had been helped than in situations when they had not previously received help. This phenomenon, called generalized reciprocity, does not require specific knowledge of the partner and may promote the evolution of cooperation between unrelated and unknown individuals. Subsequently, [Dolivo and Taborsky (2015)](#B59) have shown that rats are able to distinguish between different collaborators based on the quality of their help. Their data confirm that rat’s propensity to reciprocate aid is proportional to the quality of aid previously received by the same partner. To sum up, rats are more likely to help a conspecific they do not know but who previously helped them.

The aim was then to understand factors on which rats rely, in order to decide when they should return the aid received ( [Dolivo and Taborsky, 2015](#B59) ). In fact, even when visual information was missing, rats provided more food to partners who had previously helped them to obtain food than to those who had not. In a similar way, [Hernandez-Lallement et al. (2015)](#B201) trained pairs of rats—an actor and a partner—to decide between two alternatives that differed only in the reward given also to the partner. With the “ own reward” choice, the reward was accessible only to the actor, while the “ both reward” choice also produced a reward for the partner or for an inanimate toy, placed in an adjacent compartment. These authors found that the actors chose the “ both reward” option significantly more than the random level and more often in the condition with the real partner than in the condition with the toy. This study shows how rats are influenced by the presence of a conspecific in implementing a behavior that produces also a benefit for the other.

Last but not least, a study by [Li and Wood (2017)](#B121) determined if unrelated male laboratory rats would respond *on behalf* of a partner in an iterated sequential game. Trial-by-trial analysis of “ donor” and “ responder” responses provides evidence that rats are not following a strategy of generalized reciprocity, but are sensitive to responses of their partner in previous trials. The study revealed that, when rats were paired with their cagemate, the responses were consistent with reciprocal altruism. By contrast, when paired with another unrelated partner (a stranger), and even when paired with a maximally-generous partner (good stooge), response rates decreased, since test rats failed to adjust their donor responses to obtain more pellets. These findings, obtained with a “ repeated donation” game, seem to align with previous studies of sequential cooperative behavior in primates.

These results are consistent with direct reciprocity, whereby participants work on behalf of a partner who has worked for them previously. This suggests however that rats, when playing an iterative sequential game, are more likely to respond on behalf of a cagemate. Nevertheless, rats show direct reciprocity for a partner who previously provided a food reward and respond more rapidly for a partner who provided a preferred reward. This study proves that the “ repeated donation” game is a good protocol to model and test cooperative behavior in laboratory rats, offering the opportunity to explore the neurobiology of cooperation.

In our lab, we are currently developing some novel paradigms to identify the empathy-driven change in prosocial behavior. We developed an experimental setting whereby rats learn to push a syringe to let a food pellet drop. Once they have been trained to push in order to obtain food for themselves, the test phase is consisting of dividing the arena (by means of a grid) into two separate halves: in this way, any one rat of a dyad will be able, by pushing, to let the food pellet drop towards the other conspecific and vice versa. In such way, the both of them can eat only by reciprocal pushing, while if only one subject of a dyad pushes, only the other will eat ( [Festucci et al., 2019](#B76) ).

We hereby propose that the exact rate at which the subjects will push, within a given session, may well depend on the status of the conspecific found behind the dividing grid. Our goal is to demonstrate that such prosocial habit is sensitive to the level of e. g., stress, or hunger, of the recipient. We propose that such a modulation of the prosocial habit, if demonstrated, would imply a metacognitive step: if the donor increases donations when the recipient is stressed or hungry, then the recipient is felt by the donor as a “ subject” that may experience stress or hunger.

## Conclusions

Today, hundreds of research articles are available which studied the neural mechanism underlying human empathy. Most of the work focused on a detailed characterization of two processes: a *mirroring of the emotional response* i. e., living “ as if the same feelings or perceptions occurred to me” and a *mentalization* as a cognitive evaluation of the others’ internal state (thoughts, intentions, feelings, perceptions).

Classic works, developed from 1995 to 2000, have gathered fundamental knowledge of the neural activity underlying these two processes, who seem to be completely separated at least at first glance. Current psycho-biological models of empathy nowadays consider the two processes deeply interactive, along with prosocial motivation.

Neuroscientists have often adopted consolidated concepts used by psychologists to describe empathic processes: sharing of emotion, role taking, Theory of Mind. Some researchers define empathy referring to the multi-dimensional construct, which comprehends its dual nature, both cognitive and affective. Other researchers refer to empathy taking into account only one component, either affective (studies concerning contact, pain and smell) or cognitive ( [Völlm et al., 2006](#B190) ). Some other authors suggest that, even though cognitive empathy seems to be depending on deductive abilities also supporting Theory of Mind, it is not equivalent to Theory of Mind. Good cognitive empathic skills are likely to rely on the same underlying skills that also enable mentalization, yet cognitive empathy is concerned with the attribution of emotions as opposed to cognitions; as such, the two constructs are potentially dissociable ( [Reniers et al., 2011](#B151) ).

Some studies ( [Lombardo et al., 2009](#B124) ; [Schnell et al., 2011](#B161) ) surprisingly demonstrate that cognitive empathy leads to a “ simulated” generation of affective states through the activation of the limbic system in the absence of external emotive signals.

Additional ambiguity comes from the definition of cognitive empathy vs. affective subcomponents of Theory of Mind. Both are often similarly defined in terms of comprehension of emotions. Shamay–Tsoory is the first author who tried to clarify the situation, elaborating a model in which a specific role was defined for all of the components. Albeit pitfalls encountered in the study of empathy may be troublesome, such kind of problems naturally came to surface during initial studies of any complex psychological phenomenon. In neuroscience, simplified stimuli and tasks have been fundamental to track the roots of empathic sub-processes, and such localization was absolutely necessary to continue with the study. Social-cognitive naturalistic paradigms are becoming frequent in neurosciences and some of them have already evidenced limitations of previous visions of empathy that focused on a single process. Such researches have shown that neural systems involved with experience-based sharing and mentalization, other than being activated in separate ways, are normally co-activated when the person is subject to complex social signals.

Moreover, functional neuroimaging will not have to be the sole research approach to study biological substrates of empathy: important outcomes could well come from genetics ( [Ebstein et al., 2010](#B65) ), from pharmacology ( [Bartz et al., 2011](#B15) ), from clinical studies on patients ( [Shamay-Tsoory and Aharon-Peretz, 2007](#B166) ) and from many other techniques. Finally, new approaches will not have to be limited to the study of healthy population. For example, naturalistic studies ( [Courchesne and Pierce, 2005](#B46) ) are demonstrating that people diagnosed with autism can be identified by difficulty in integrating multiple social signals: such an anomaly may be related to interruptions of connections between neural systems. Such widening of views will allow to plan for novel interventions with specific therapies, addressed towards children, teenagers and adults with deficiencies in multiple aspects of empathy and mentalization.

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LC and LB wrote a first draft of this review. MC and FF added specific sections. SLR translated the first draft into English. RT and WA commented on the first draft and extended it. SC coordinated the whole workgroup.

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## Conflict of Interest Statement

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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