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Large-scale temporo–parieto–frontal networks for motor and cognitive motor functions in the primate brain

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ABSTRACT

The extent to which neural circuits and mechanisms underlying sensory, motor, and cognitive cortical functions in the human brain are shared with those of other animals, especially non-human primates, is currently a key issue in the field of comparative neuroscience. Cortical functions result from the conjoint function of different, reciprocally connected areas working together as large-scale functionally specialized networks, which can be investigated in human subjects thanks to the development of non-invasive functional and connective imaging techniques. In spite of their limitations in terms of spatial and temporal resolution, these techniques make it possible to address the issue of how and to what extent the neural mechanisms for different cortical functions differ from those of non-human primates. Indeed, 30 million years of independent evolution have resulted in significant differences between the brains of humans and macaques, which are the experimental model system phylogenetically closest to humans for obtaining highly detailed anatomical and functional information on the organization of cortical networks. In the macaque brain, architectonic, connective, and functional data have provided evidence for functionally specialized large-scale cortical networks involving temporal, parietal, and frontal areas. These networks appear to play a primary role in controlling different aspects of motor and cognitive motor functions, such as hand action organization and recognition, or oculomotor behavior and gaze processing. In the present review, based on the comparison of these data with data from human studies, we will argue that there is clear evidence for human counterparts of these networks. These human and macaque putatively homolog networks appear to share phylogenetically older neural mechanisms, which, in the evolution of the human lineage, could have been exploited and differentiated, resulting in the emergence of human-specific higher-order cognitive functions. These considerations are fully in line with the notion of “neural reuse” in primate evolution.

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1. Introduction

One long-debated key issue of comparative neuroscience is to what extent neural circuits and mechanisms underlying sensory, motor, and cognitive cortical functions in the human brain are shared with those of other animals, especially non-human primates.

The general view of the organization of the neural substrate for a given brain function has largely evolved over the years. The most accepted current view is that cortical functions are not localized to specific regions as supported by localizationism, but result from the conjoint function of different, reciprocally connected areas working together as large-scale functionally specialized networks (see, for example, Bressler & Menon, 2010; Catani, Dell'Acqua, & Bizzi et al., 2012).

There are different views on the organization of these networks and on the way in which they could operate (for reviews on this issue, see Anderson, 2010; Bergeron, 2007; Caminiti et al., 2017; Sporns, 2013). According to a modular view, functionally specialized networks are structurally and functionally distinct entities operating largely independently with one another. However, very often individual regions or even areas appear to be engaged in different functions. Furthermore, data from animal models indicate that the connectional structure of the cortex supports complex patterns of interareal interaction promoting widespread influences among cortical areas. Accordingly, any area could be responsible of specific information processing operations that could be used for different functions, taking part in different functionally specialized large-scale networks.

Thus, the identification of these networks requires multidisciplinary integration of structural and functional data in order to extract from the general pattern of cortical connectivity those connections that could mediate dynamic interactions between different areas contributing to a specific function. In this context, it is noteworthy that, as the degree of anatomical and functional selectivity varies across areas, the composition of a given network could vary based on the sets of data and criteria used for its definition.

Based on this conceptual framework it becomes clear that understanding the neural circuits and mechanisms underlying specific cortical functions requires multimodal experimental approaches aiming to define the following: i) the exact localization and extent of the areas possibly involved in a specialized network; ii) the existence of connections among these areas, and iii) their possible specific functional contribution. All together, these data can provide comprehensive pictures of large-scale functionally specialized cortical networks in terms of nodes and edges (Sporns, 2013), possible flows of information processing, and neural mechanisms from which a given function can emerge.

In recent years, the development of non-invasive functional and connectional imaging techniques has made it possible to address the definition of large-scale, functionally specialized networks in the human brain. Indeed, functionally distinct cortical sectors can be identified using functional magnetic resonance imaging (fMRI), and brain connectivity can be investigated using diffusion-weighted MRI (dMRI). Furthermore, functional connectivity MRI has recently been

proposed as a possible tool for mapping large-scale networks in the human brain.

However, a detailed definition of cortical networks in the human brain is still prevented by several limitations of these techniques. First, fMRI is limited in spatio-temporal resolution and gives indirect information of neuronal activity only at the macroscale level. Further, though multimodal techniques allow detailed post-mortem architectonic studies of the human cortex (see, Amunts & Zilles, 2015), the areal attribution of functional data obtained in living subjects can be, at best, based on probabilistic architectonic maps, which, to some extent, prevents univocal anatomic-functional correlations of experimental data. Second, recent studies in which well-known cortical pathways have been traced in macaques with dMRI have seriously questioned the technique's validity for precise in vivo tracing of point-to-point connectivity (Reveley et al., 2015; Thomas et al., 2014). Third, several observations have been made that functional connectivity is related to, but distinct from, anatomic connectivity, as it could be subserved by polysynaptic, as well as monosynaptic, anatomical circuits, and can be modulated by the task performed by the subjects or several other factors differently from structural connectivity (Biswal et al., 2010; Buckner, Krienen, & Yeo, 2013). Thus, at present, this technique does not appear capable of providing truthful comprehensive pictures of large scale functionally specialized networks as properly defined.

In spite of these limitations, the exploitation of these techniques has made it possible to make comparative observations, thus addressing the issue of how and to what extent the neural mechanisms for different cortical functions differ across different primate species. These observations are essential for assessing the extent to which detailed functional and connectional data from non-human primate studies can be used for explaining the neural mechanisms of the human brain. Indeed, 30 million years of independent evolution have resulted in significant differences between the brains of humans and macaques, which are the experimental model system phylogenetically closest to humans for obtaining highly detailed anatomical and functional information (see, e.g., Passingham, 2009; Sereno & Tootell, 2005).

Humans have brains much larger than would be expected for primates of a similar body size, and this difference appears to reflect primarily an enlargement of the neocortex and, specifically, a disproportionate enlargement of the higher-order association cortex of the frontal, temporal, and parietal lobes, relative to the primary sensory and motor areas (see, e.g., Preuss, 2011). This selective expansion is considered the neural basis for the outstanding cognitive capabilities of humans (see, e.g., Chaplin, Yu, Soares, Gattass, & Rosa, 2013; Passingham, 2009; Rilling, Glasser, Jbabdi, Andersson, & Preuss, 2012; Sereno & Tootell, 2005).

However, it appears that the association cortex expanded in a predictable manner in primate evolution (see Preuss, 2011). Indeed, the human brain is not exceptional in its cellular composition, as it contains as many neurons as would be expected for a primate brain of human size (Herculano-Houzel, 2009), and the human frontal cortex is not larger than expected for a primate brain of human size (Semendeferi, Lu, Schenker, & Damasio, 2002). Comparative observations also showed that the expansion of the cortex in

simian primates of different brain size correlates with a disproportionate expansion of some association areas typically involved in complex cognitive and behavioral functions (Chaplin et al., 2013). Furthermore, Hill et al. (2010), by comparing human and macaque cerebral cortices, found that the pattern of human evolutionary expansion is remarkably similar to the pattern of human postnatal expansion, suggesting that those association areas which disproportionately expanded in primate evolution are those that mature later in human postnatal development. Finally, recent comparative observations showed a relationship between structural interindividual variability and evolutionary expansion in the primate brain in which regions that show a higher degree of variability in a series of MRI measures of gray and white matter are those that could have evolved more recently (Croxson, Forkel, Cerliani, & Thiebaut de Schotten, 2017). Accordingly, our understanding of cognitive capabilities could be simply due to the fact that our brain has a lot more association cortex in absolute terms than do other non-human primates (Preuss, 2011).

In line with these comparative evolutionary observations, a large body of experimental evidence has been provided supporting the notion that the monkey and human brain share common plans of anatomical and functional organization of sensory, motor, and cognitive functions likely inherited from the last common ancestor shared by modern humans and macaques (Caminiti, Innocenti, & Battaglia-Mayer, 2015; Geyer, Matelli, Luppino, & Zilles, 2000; Mantini, Corbetta, Romani, Orban, & Vanduffel, 2013; Orban, Van Essen, & Vanduffel, 2004).

In recent years, based on anatomical and functional data, we have provided evidence for functionally specialized large-scale cortical networks of the macaque brain involved in controlling different aspects of motor and cognitive motor functions. In the present review article, these data will be used to make comparative considerations, based on anatomical and functional data obtained in human studies. The aim is to see to what extent cortical networks and neural mechanisms identified in the macaque brain are shared with the human brain and to what extent some cognitive motor human-specific abilities could be the result of the exploitation and differentiation of neural mechanisms of the macaque brain.

2. A large-scale cortical network for controlling purposeful hand actions and for action recognition in the macaque

Fig. 1 shows the architecture of a large-scale temporo-parieto-frontal network of the macaque brain providing a possible substrate for interfacing perceptual, cognitive, and hand-related sensorimotor processes for controlling hand actions based on object identity, goals, and memory-based or contextual information. This network has been designated as *lateral grasping network* (Borra, Gerbella, Rozzi, & Luppino, 2017).

The network is centered on a parieto-frontal circuit linking the two hand-related visuomotor areas F5 and the Anterior intraparietal area (AIP), located in the ventral premotor cortex (PMv) and in the inferior parietal lobule (IPL), respectively. This circuit plays a crucial role in mediating visuomotor

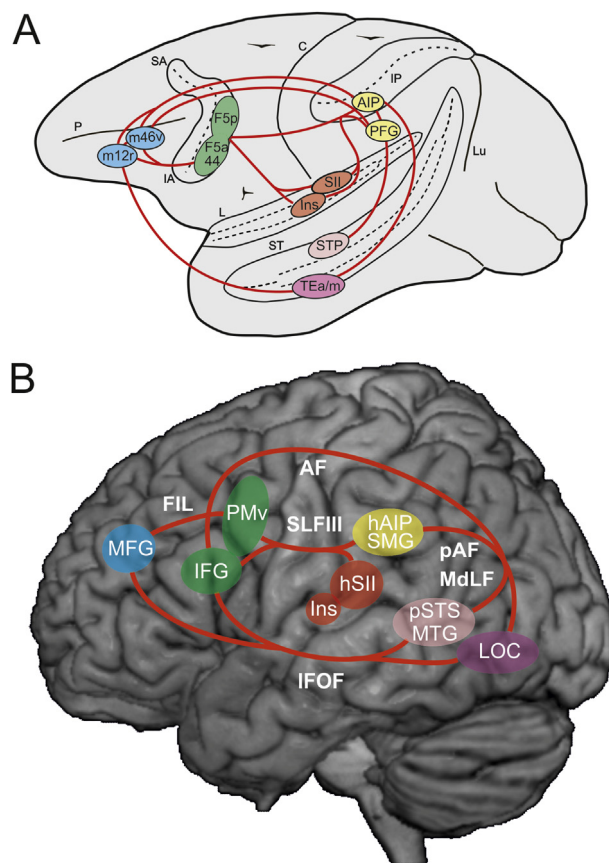


Fig. 1 – The macaque lateral grasping/action recognition network and its possible human counterpart. (A) Lateral view of a hemisphere of the macaque brain showing the main nodes of the lateral grasping/action observation network and their interconnections defined based on tract tracing connectational data. C = central sulcus; IA = inferior arcuate sulcus; IP = intraparietal sulcus; L = lateral sulcus; Lu = lunate sulcus; m12r = middle part of area 12r; m46v = middle part of area 46v; P = principal sulcus; ST = superior temporal sulcus. (B) Lateral view of a hemisphere of the human brain showing with the same color the possible homologs of the main nodes of the macaque lateral grasping/action observation network and their possible interconnections based on dMRI data. AF = Arcuate fasciculus; FIL = Frontal inferior longitudinal tract; hAIP = Human anterior intraparietal area; hSII = Human second somatosensory area; IFG = Inferior frontal gyrus; IFOF = Inferior fronto-occipital fasciculus; Ins = Insula; LOC = Lateral occipital complex; MdLF = Middle longitudinal fasciculus; MFG = Middle frontal gyrus; MTG = Middle temporal gyrus; pAF = Posterior segment of the arcuate fasciculus; PMv = Ventral premotor cortex; pSTS = Posterior superior temporal sulcus; SLFIII = Third branch of the superior longitudinal fasciculus; SMG = Supramarginal gyrus.

transformations for grasping, in which visual coding of the object's physical properties (e.g., size, shape, orientation) automatically leads to the activation of potential motor acts appropriate for hand–object interactions (Jeannerod, Arbib, Rizzolatti, & Sakata, 1995; Rizzolatti & Luppino, 2001). This process, also referred to as “affordances extraction” (see, e.g., Fagg & Arbib, 1998), primarily relies on visual coding of the object's physical properties, carried out along the occipitoparietal visual information processing pathway designated as the “dorsal visual stream” (Sakata, Taira, Kusunoki, Murata, & Tanaka, 1997). However, area AIP is also robustly connected to sectors of the inferotemporal cortex (Borra et al., 2008), located at the highest hierarchical levels of the occipitotemporal visual information processing pathway designated as the “ventral visual stream” and involved in object discrimination and recognition (see, e.g., Tanaka, 1996). One of these sectors, located in area TEa/m of the lower bank of the superior temporal sulcus, is part of a component of the ventral visual stream specifically dedicated to three-dimensional (3D) object and action processing (Orban, Zhu, & Vanduffel, 2014). These temporal connections of area AIP provide the substrate for extraction of object affordances, carried out by the AIP–F5 circuit, based on information related not only to the intrinsic properties, but also to the identity of the object target of the action. Furthermore, they could provide the access of signals related to motor and haptic representations of hand actions to the representations of object identity, thus playing a role in the neural mechanisms underlying tactile object recognition.

Area F5 is robustly connected with two other parietal areas, the visuomotor hand-related area PFG of the IPL convexity and the hand field of the higher order Second somatosensory area (SII) of the parietal operculum. In area PFG, grasping-related activity appears to be influenced by the context in which the action is performed, possibly reflecting sequential action organization according to its goal or motor intention (Bonini et al., 2011, 2012, 2010; Fogassi et al., 2005). Furthermore, the finding that area PFG grasping neurons can integrate information on both grip type and action goal suggests that this IPL area encodes information about both “how” and “why” each motor act has to be done (Bonini et al., 2012). In area SII, the presence of neurons in the hand field preferentially responsive to proprioceptive input and often responding well to active movements, especially when grasping objects, suggests this area plays a role in the somatomotor transformations for object-oriented hand actions and in haptic processing of object shapes (Fitzgerald, Lane, Thakur, & Hsiao, 2004).

The PMv area F5 and the two IPL areas AIP and PFG also host another class of visuomotor hand-related neurons—designated as “mirror neurons”—which activate during the execution of hand motor acts, as well as during the observation of similar acts done by others (Gallese, Fadiga, Fogassi, & Rizzolatti, 1996; Rizzolatti, Fadiga, Gallese, & Fogassi, 1996). This neural activity has been interpreted as the result of visuomotor transformations in which observed actions are mapped on their corresponding internal motor representations. It has been suggested that these neurons are part of an observation–execution matching system (mirror system) which is the basis for the ability to recognize others' goal-directed motor acts (see Rizzolatti, Cattaneo, Fabbri-Destro, & Rozzi, 2014). There is evidence that visual

information on observed actions can be provided to the PMv–IPL components of the mirror system through the temporal connectivity of areas AIP and PFG. Specifically, fMRI data obtained in awake macaques (Nelissen et al., 2011) showed that action observation activates cortical sectors located in both the ventral and the dorsal banks of the STS. One sector, located in the ventral bank, corresponds to the TEa/m sector connected to area AIP. Another sector, located in the upper bank of the Superior temporal sulcus (STS), corresponds to a sector of the superior temporal polysensory (STP) area connected to the PFG. Area STP is a higher-order multisensory area which integrates information within and across modalities (Baylis, Rolls, & Leonard, 1987; Bruce, Desimone, & Gross, 1981) and hosts visual neurons (see Carey, Perrett, & Oram, 1997), coding biological motion, differentiating between self-produced actions and actions made by others, and coding the intentionality of actions (Jellema, Baker, Wicker, & Perrett, 2000; Jellema & Perrett, 2003), suggesting a role in social cognition. Based on these data, Nelissen et al. (2011) suggested that visual action information, encoded in the STS, is forwarded to parietal areas of the mirror system along a TEa/m–AIP pathway which could provide visual descriptions of the type and immediate goal of hand actions made by others, and along a STP–PFG pathway which could be involved in extracting the intention behind the observed motor act. It is also noteworthy that the SII region hosts visually responsive neurons, active during the observation of human actions or objects, suggesting that this area has a role in multisensory integration for motor control and in action recognition (Hihara, Taoka, Tanaka, & Iriki, 2015).

All these parietal areas and the anterior sector of area F5 (F5a) are also differentially connected with specific sectors of the areas 46v and 12r, located in the ventrolateral prefrontal cortex (VLPF). The prefrontal cortex is a large, heterogeneous region considered, as a whole, to be critically involved in the so-called “executive functions”, a term that, in general, refers to those mechanisms by which behavioral performance is optimized in situations requiring cognitive processes (see Tanji & Hoshi, 2008). Recent data provided evidence of rostrocaudal connective gradients in the VLPF, in which the caudal part is primarily connected with inferior parietal and prearcuate oculomotor areas, the middle part with parietal and frontal sensorimotor areas, and the rostral part primarily with other prefrontal areas (Borra, Gerbella, Rozzi, & Luppino, 2011; Gerbella, Belmalih, Borra, Rozzi, & Luppino, 2010, 2013). Functional studies have indeed shown that cells active in tasks requiring oculomotor responses (e.g., Averbeck, Sohn, & Lee, 2006; Boch & Goldberg, 1989; Ichihara-Takeda & Funahashi, 2007) and the execution of arm/hand responses (Bruni, Giorgetti, Bonini, & Fogassi, 2015; Hoshi, Shima, & Tanji, 2000, 1998; Requin, Lecas, & Vitton, 1990; Simone, Rozzi, Bimbi, & Fogassi, 2015) tend to be located more caudally and more rostrally in the caudal VLPF, respectively. In the context of the *lateral grasping network*, area 46v could be involved in selecting, monitoring, and updating object-oriented hand actions based on behavioral goals and guiding rules and current, memorized, or working memory information on motor programs. Furthermore, area 12r, which is robustly connected with the same sector of area TEa/m which is connected to area AIP (Borra et al., 2011), is a possible substrate for integration of the ventral visual stream with sensorimotor hand-related

information in the prefrontal cortex. Thus, in the intermediate part of area 12r, the retrieval, retention, and manipulation of information on objects or hand–object interactions could be finalized to the control of object-oriented hand actions and to tactile object recognition. The finding that, in this VLPF hand-related sector, there are neurons which respond to the observation of goal-directed actions also suggests this sector's participation in the action observation–execution matching system (Simone, Bimbi, Rodà, Fogassi, & Rozzi, 2017).

Finally, all the various parietal, premotor, and VLPF hand-related areas are connected with a specific sector located relatively dorsally in the dysgranular insula (see Borra, Gerbella et al., 2017). This sector appears to overlap, at least in part, with an insular zone from which intracortical microstimulation evokes hand movements (Jezzini, Caruana, Stoianov, Gallese, & Rizzolatti, 2012). This specific insular sector is a possible source of signals related to internal states (Ibañez, Gleichgerrcht, & Manes, 2010) modulating the control of hand actions.

Based on these data, a model has been proposed in which motor programs of “potential” hand motor acts are first activated in area F5 as a consequence of fast visuomotor transformations, and then selected based on behavioral goals, contextual information, and memorized information on object identity and properties (Borra, Gerbella et al., 2017). The selected hand motor acts can then be put into action through the robust connections of the posterior sector of F5 (F5p), from which intracortical microstimulation evokes hand movements at relatively low current thresholds, with the hand field of the primary motor area F1 (Borra, Belmalih, Gerbella, Rozzi, & Luppino, 2010). There is evidence that the projections from F5p to F1 could provide the substrate for generating the various muscle synergies (movements) represented in F1 which are necessary for executing the motor act selected at the level of F5 (Cerri, Shimazu, Maier, & Lemon, 2003; Prabhu et al., 2009; Shimazu, Maier, Cerri, Kirkwood, & Lemon, 2004; Umiltà, Brochier, Spinks, & Lemon, 2007). However, F5p is also a source of projections to the brainstem and the spinal cord, suggesting a contribution by this area to the generation and control of hand movements in parallel with the hand field of F1 (Borra et al., 2010).

The lateral grasping network, as shown in Fig. 1, is not exhaustive. First, there is connective evidence for the participation in this network of other components, such as the granular frontal opercular area GrFO. Second, there are areas which, based on their premotor connectivity, appear to contribute almost equally well to more than one network, suggesting a more general role in motor control. One of these areas is the medial premotor area F6 (pre-SMA), involved in higher-order aspects of motor control (see, e.g., Picard & Strick, 2001; Nachev, Kennard, & Husain, 2008), which could play a role in forwarding signals which transform potential actions into actual movements and determine movement onset, and in controlling the temporal organization of motor programs (see Ridderinkhof, Forstmann, Wylie, Burle, & van den Wildenberg, 2011; Rizzolatti et al., 2014). This area also hosts neurons selectively encoding others' actions and neurons showing activity increase associated with another's errors, suggesting involvement in action recognition (Yoshida, Saito, Iriki, & Isoda, 2011, 2012).

3. Cortical networks for explorative and communicative oculomotor behavior in the macaque

In addition to the lateral grasping network, there is evidence for another large-scale temporo–parieto–frontal network, in which the various nodes are linked through “dorsal” temporo–parieto–frontal and “ventral” temporo–frontal pathways, which could play a crucial role in controlling some aspects of oculomotor behavior.

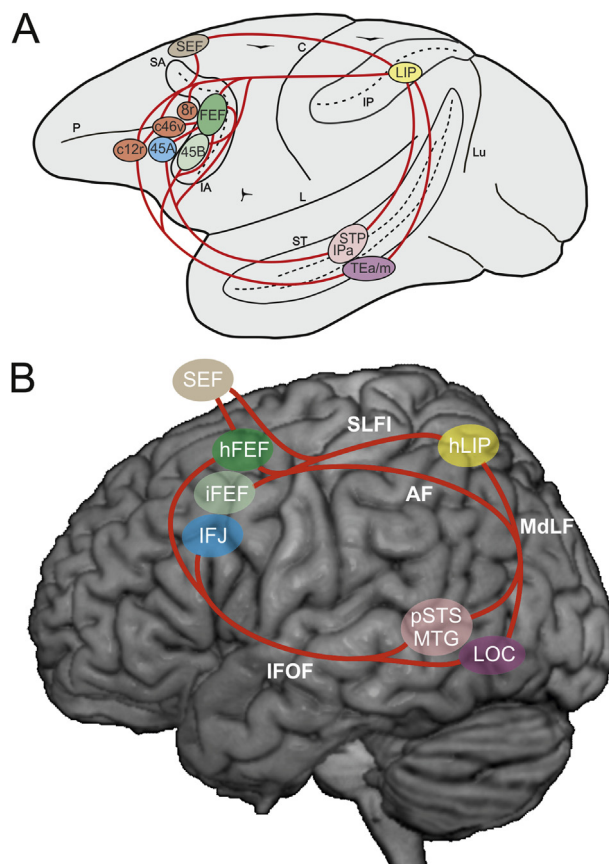


Fig. 2 – The macaque network for explorative and communicative oculomotor behavior and its possible human counterpart. (A) Lateral view of a hemisphere of the macaque brain showing the main nodes of the explorative and communicative oculomotor network and their interconnections defined based on tract tracing connective data. Temporal connections of the SEF are not shown. c12r = caudal part of area 12r; c46v = caudal part of area 46v. Other abbreviations as in Fig. 1. (B) Lateral view of a hemisphere of the human brain showing with the same color the possible homologs of the main nodes of the macaque explorative and communicative oculomotor network and their possible interconnections based on dMRI data. The Supplementary Eye Field (SEF) is located on in the medial wall of the hemisphere. hFEF = Human frontal eye field; hLIP = Human lateral intraparietal area; iFEF = Inferior frontal eye field; IFJ = Inferior frontal junction; SLFI = First branch of the superior longitudinal fasciculus. Other abbreviations as in Fig. 1.

As shown in Fig. 2, this network is centered on a parieto-frontal circuit linking two visually responsive oculomotor areas: the lateral intraparietal (LIP) area located in the lateral bank of the intraparietal sulcus just caudal to area AIP and the frontal eye field (FEF) located in the anterior bank of the arcuate sulcus. This circuit plays a crucial role in visuomotor transformations for controlling saccadic eye movements and in the orientation of spatial attention (see Lynch & Tian, 2006; Wardak, Olivier, & Duhamel, 2011). The FEF, defined as the arcuate bank sector from which intracortical micro-stimulation evokes saccades at relatively low current thresholds, displays a topographic organization in which smaller and larger amplitude saccades are evoked from more ventral and more dorsal sites, respectively (Bruce, Goldberg, Bushnell, & Stanton, 1985). The FEF is also connected with the supplementary eye field (SEF), located rostrally in the dorsal premotor cortex, and with several caudal prefrontal areas, including areas 8r, 45B, and 45A, and the caudal sectors of areas 12r, 46v, and 46d (Gerbella et al., 2010; Huerta, Krubitzer, & Kaas, 1987; Schall, Morel, King, & Bullier, 1995; Stanton, Bruce, & Goldberg, 1993, 1995). All these caudal prefrontal areas/sectors are, in turn, connected to the SEF and, except for caudal 12r and 45A, to area LIP (Borra et al., 2011, Borra, Gerbella et al., 2017; Gerbella et al., 2010, 2013). Thus, all these areas appear involved in the oculomotor frontal system. Furthermore, the FEF and the various caudal prefrontal oculomotor areas are provided with an access to brainstem oculomotor centers (Borra, Gerbella, Rozzi, & Luppino, 2015).

As for the parieto-frontal circuitry involved in controlling hand actions, the visuomotor transformations for controlling oculomotor behavior also appear to rely not only on input from dorsal visual stream areas, but also on input from both the lower and the upper banks of the STS, which appear to differentially distribute in the parieto-frontal oculomotor circuitry. Specifically, area LIP, the ventral part of the FEF, areas 45B and caudal 12r are connected to several inferotemporal sectors of the lower bank of the STS and of the inferotemporal convexity cortex (Blatt, Andersen, & Stoner, 1990; Cavada & Goldman-Rakic, 1989; Gerbella et al., 2010; Schall et al., 1995; Stanton, Bruce, & Goldberg, 1995). One of these sectors, located in the ventral bank of the STS, is just caudal to the sector involved in the lateral grasping network. This sector is part of the ventral visual stream component specifically dedicated to 3D object and action processing (Denys et al., 2004; Nelissen et al., 2011) and also activates during the execution of visually guided eye movements (Ward, Bolding, Schultz, & Gamlin, 2015). Accordingly, it is possible to define a large-scale temporo-parieto-frontal network involving area LIP, ventral FEF, area 45B, and caudal TEa/m, where visuospatial dorsal visual stream information and ventral visual stream information on objects and actions could be used for guiding small-amplitude saccades.

Indeed, area LIP, the ventral part of the FEF, and area 45B host neurons showing shape selectivity, encoding non-spatial attributes of the stimuli, and activating during the observation of two-dimensional (2D) shapes, thus likely reflecting input from ventral visual stream areas (Peng, Sereno, Silva, Lehky, & Sereno, 2008; Sereno & Maunsell, 1998; Toth & Assad, 2002). Furthermore, fMRI data have revealed area 45B activation for the observation of objects, faces, and actions

(Denys et al., 2004; Nelissen, Luppino, Vanduffel, Rizzolatti, & Orban, 2005; Tsao, Schweers, Moeller, & Freiwald, 2008). Though the functional properties of area 45B still remain to be fully elucidated, fMRI (Premereur, Janssen, & Vanduffel, 2015) and 2-deoxyglucose (Moschovakis et al., 2004) data have shown activation during the execution of saccades, fitting well with the proposed affiliation of this area with the oculomotor frontal system, as indicated by its connectivity pattern. It has proposed that area 45B is a “pre-oculomotor” area involved in guiding the exploration of visual scenes for the perception of objects, actions, and faces (Gerbella et al., 2010). Other caudal prefrontal oculomotor areas connected to LIP and the ventral part of the FEF (8r, caudal 46v), as well as caudal 12r, could play a role in executive functions aimed at the control of small saccades and could contribute to the suggested role of area LIP in representing salience maps combining visual information with cognitive factors, such as behavioral context, task difficulty, or reward information (Wardak, Vanduffel, & Orban, 2010).

Connectional data provide evidence for a further partially overlapping large-scale oculomotor network involving area STP and the fundal STS area IPa, which are connected to area LIP, the dorsal part of the FEF, the SEF, and area 45A (Blatt et al., 1990; Cavada & Goldman-Rakic, 1989; Gerbella et al., 2010; Huerta & Kaas, 1990; Luppino, Calzavara, Rozzi, & Matelli, 2001; Saleem, Miller, & Price, 2014; Stanton et al., 1995). As mentioned above, area STP is involved in processing various forms of biological motion, including the direction of gaze and head, which may be relevant in understanding where the conspecifics are fixating (Carey et al., 1997; Jellema et al., 2000; Mistlin & Perrett, 1990) and is involved in the integration of audiovisual communication signals (Barraclough, Xiao, Baker, Oram, & Perrett, 2005; Chandrasekaran & Ghazanfar, 2008; Dahl, Logothetis, & Kayser, 2009). Reversible inactivation of the posterior STP severely disrupts gaze-following behavior (Roy, Shepherd, & Platt, 2014). Furthermore, in area LIP, there are neurons which become more active both while directing attention toward a region of space and while observing other monkeys doing the same (Shepherd, Klein, Deaner, & Platt, 2009). These “mirror” oculomotor responses likely reflect input from the STS and suggest that area LIP plays a role in sharing attention with others (Shepherd, 2010). Area 45A is a caudal VLPF area involved in the multisensory processing of communication stimuli (Diehl & Romanski, 2014; Romanski & Averbeck, 2009; Sugihara, Diltz, Averbeck, & Romanski, 2006), and it activates during action and face observation (Kuraoka, Konoike, & Nakamura, 2015; Nelissen et al., 2005; Tsao, Schweers et al., 2008), suggesting it has a role in communication behavior. Moreover, this area is robustly connected to the dorsal part of the FEF, is a source of projections to subcortical oculomotor structures (Borra et al., 2015), and is activated during the execution of eye movements (Premereur et al., 2015).

Thus, there is connectional evidence for a large-scale oculomotor network involving areas STP-IPa, LIP, 45A, and dorsal FEF, which could provide the substrate for the role of gaze position and eye movement in social behavior, thus contributing to an understanding of the social intentions of other individuals (Ghazanfar, Nielsen, & Logothetis, 2006; Shepherd, 2010). Moreover, the rostral part of area 46d, which is connected to area STP, dorsal FEF and area 45A (Borra, Ferroni

et al., 2017), could play a higher-order executive role in this “social oculomotor” network. In this context, it is noteworthy that other oculomotor areas, such as the SEF, appear to be involved in both these described networks and that nodes of these two networks, together with other areas, could participate in large-scale networks involved in other aspects of oculomotor behavior.

4. Possible human counterpart of the macaque lateral grasping/action recognition network

In early functional imaging studies and in others since then (Binkofski et al., 1998; Culham et al., 2003; Ehrsson et al., 2000; Johnson-Frey, 2004; Toni, Rushworth, & Passingham, 2001), it has been shown that the execution of object-oriented hand actions activates two cortical zones located in the frontal lobe and in the IPL, respectively. The frontal zone is located mostly in the ventral part of the precentral gyrus and also extends rostrally into the inferior frontal gyrus (IFG), involving Brodmann's architectonic area 44 (Brodmann, 1909), corresponding to the caudal part of the language-related Broca's region. The IPL zone corresponds to both the rostral part of the lateral bank of the IPS and the supramarginal gyrus (SMG). Based on their location, these two zones have been considered as the possible human counterparts of the macaque PMv and IPL areas of the *lateral grasping network*, respectively. Several studies have provided further evidence for these proposed homologies. Specifically, Forna et al. (2018) showed that short-train or single-pulse electrical stimulation of the cortical surface along the dorso-ventral extent of the PMv is effective in evoking hand, orofacial, and combined orofacial and hand motor responses from the dorsal, ventral, and intermediate part of it, respectively, providing evidence for a somatotopic arrangement of this region similar to that of the macaque F5. Furthermore, transcranial magnetic stimulation (TMS) studies showed an interaction of the dorsal part of the PMv with the primary motor cortex, modulated during grasping execution, and that a virtual lesion of this PMv sector impairs grasping execution (Davare, Andres, Cosnard, Thonnard, & Olivier, 2006, 2008, 2009). These data provide support for the possible homology between this human PMv sector and the posterior subdivision of the macaque area F5 (F5p). Other studies showed that the lateral bank of the IPS also activates during surface orientation discrimination and subsequent related spatial adjustment of finger position (Shikata et al., 2003) and is involved in 3D shape processing from disparity (Georgieva, Peeters, Kolster, Todd, & Orban, 2009) and in coding intrinsic object properties (Monaco, Sedda, Cavina-Pratesi, & Culham, 2015). Furthermore, this region hosts neurons selectively tuned for motor imagery of specific hand shapes (Klaes et al., 2015) and TMS studies have shown that virtual lesions of this zone affect hand shaping, scaling of grip force (Dafotakis, Sparing, Eickhoff, Fink, & Nowak, 2008; Davare, Andres, Clerget, Thonnard, & Olivier, 2007), and online adjustments of goal-directed hand actions (Rice, Tunik, & Grafton, 2006; Tunik, Frey, & Grafton, 2005). Finally, there is an increase in effective connectivity between this region and PMv when grasping small objects (Grol et al., 2007) and a reduction in

PMv–M1 interactions during grasping preparation after a virtual lesion of this hand-related zone of the IPS (Davare, Rothwell, & Lemon, 2010). These data have provided strong support for the possible homology between this anterior intraparietal hand-related sector (human area AIP) and the macaque area AIP.

Execution of object-oriented actions (Gazzola & Keysers, 2009; Grèzes & Decety, 2001) also activates the SMG, which appears to be involved in comparing predicted and actual sensory input during object manipulation and updating of sensorimotor memories (Jenmalm, Schmitz, Forssberg, & Ehrsson, 2006). Furthermore, TMS of this region (but also of the caudal IFG) affects planning of sequential goal-directed hand actions in which object grasping is embedded in actions with different goals (Tunik, Lo, & Adamovich, 2008). The human rostral IPL hosts a cluster of architectonic areas which, based on dMRI observations, appears to share several connective features with the macaque rostral IPL areas (Caspers et al., 2011, 2013, 2006; Ruschel et al., 2014). One of these areas, area Pft, has been considered the putative homolog of the macaque area PFG (Caspers et al., 2011).

Finally, there is evidence that the human parietal operculum hosts two architectonically distinct somatosensory areas, designated as OP1 and OP4 (Eickhoff, Amunts, Mohlberg, & Zilles, 2006; Eickhoff, Schleicher, Zilles, & Amunts, 2006; Eickhoff, Grefkes, Zilles, & Fink, 2007). Data from dMRI studies suggested connectivity of these two areas with rostral IPL and PMv areas and with Broca's region (Eickhoff et al., 2010). Functional imaging data showed that this region activates during both tactile stimulation (Burton, Sinclair, Wingert, & Dierker, 2008; Disbrow, Roberts, & Krubitzer, 2000; Eickhoff, Grefkes, Zilles, & Fink, 2007) and movement execution (Gazzola & Keysers, 2009; Hinkley, Krubitzer, Nagarajan, & Disbrow, 2007) and is involved in tactile object recognition (Reed, Shoham, & Halgren, 2004). Furthermore, TMS studies provided evidence for a causal role of this region in the haptic working memory of object properties and grasping motor programs (Cattaneo, Maule, Tabarelli, Brochier, & Barchiesi, 2015; Maule, Barchiesi, Brochier, & Cattaneo, 2015). These data provided clear support for the proposed homology between this human opercular region (human SII) and the macaque SII region (Eickhoff, Amunts et al., 2006; Eickhoff, Schleicher et al., 2006; Eickhoff et al., 2007).

Though technical limitations still prevent reliable dMRI definition of point-to-point anatomical connectivity in the human brain, several studies have provided evidence for anatomical connectivity between the rostral IPL and the PMv/IFG (Hecht et al., 2013; Ramayya, Glasser, & Rilling, 2010; Rushworth, Behrens, & Johansen-Berg, 2006; Schubotz, Anwender, Knösche, von Cramon, & Tittgemeyer, 2010), which, as in macaques, is largely supported by the third branch of the superior longitudinal fasciculus (SLF; Schmahmann et al., 2007; Thiebaut de Schotten et al., 2012). In sum, in the human brain, there is a possible parieto-frontal circuitry linking the PMv/IFG with the rostral IPL and parietal operculum, which very likely represents the human counterpart of the macaque parieto-frontal circuitry at the core of the lateral grasping network. These possible homologies appear even more plausible considering that the PMv/IFG, the human area AIP, the SMG (especially area Pft), and the human SII all

activate during the observation of goal-directed hand actions, thus also suggesting involvement of this possible parieto-frontal circuitry in action recognition (human mirror system), as in macaques (Caspers, Zilles, Laird, & Eickhoff, 2010; Fogassi & Simone, 2013; Gazzola & Keysers, 2009; Rizzolatti et al., 2014).

In recent years, evidence has been accumulated indicating that, as in macaques, human visuomotor processing for selecting and controlling hand actions carried out in area AIP is influenced by processing which takes place in the ventral visual stream areas (see van Polanen & Davare, 2015).

Comparisons between the human and the macaque temporal cortex based on merely topological criteria are complicated by the great expansion of language-related and other higher-order associative areas in the evolution of the human lineage. Comparative fMRI observations have provided clear evidence for human homologs of the macaque visual extrastriate and adjacent temporal areas, which, however, appear to be located more posteriorly and medially than their macaque counterparts (for reviews on this issue, see Orban et al., 2004, 2014). Specifically, there is evidence for a cortical sector located ventral to the human homolog of the motion sensitive Middle temporal (MT) area, in the posterior inferior temporal and the fusiform gyrus (the lateral occipital complex, or LOC) in which different sites are specifically active during the visual processing of shapes, faces, and actions (e.g., Bell, Hadj-Bouziane, Frihauf, Tootell, & Ungerleider, 2009; Denys et al., 2004; Jastorff & Orban, 2009; Kanwisher, McDermott, & Chun, 1997; Malach et al., 1995; Tsao, Moeller, & Freiwald, 2008). Based on comparative fMRI observations, Denys et al. (2004) proposed that the LOC could be the human homolog of the lower bank of the STS (area TEa/m) and the laterally adjacent IT convexity cortex in macaques.

Indeed, grasping objects based on the processing of pictorial depth cues increases the activity of the human area AIP and its functional connectivity with the PMv and the lateral occipital complex (LOC) (Verhagen, Dijkerman, Grol, & Toni, 2008). Furthermore, when planning object-oriented actions, there are activity patterns in the LOC reflecting the type of hand action (Gallivan, Chapman, McLean, Flanagan, & Culham, 2013), and the organization of visual object representations in this region reflects action-related properties of the objects (Bracci, Cavina-Pratesi, Ietswaart, Caramazza, & Peelen, 2012; Bracci & Peelen, 2013; Mahon et al., 2007; Peelen et al., 2013). Finally, the visual and haptic coding of objects activates the human area AIP (Grefkes, Weiss, Zilles, & Fink, 2002) and a part of the LOC (Amedi, Jacobson, Hendler, Malach, & Zohary, 2002; James et al., 2002; Reed et al., 2004), providing evidence for the multimodal representation of objects in the human ventral visual stream and suggesting an interaction between area AIP and the LOC for tactile object recognition (Lacey, Tal, Amedi, & Sathian, 2009; Tal & Amedi, 2009).

In the human temporal cortex, rostral to MT and dorsal to the LOC, there is a region including the posterior STS (pSTS) and middle temporal gyrus (MTG) involved in multisensory processing and responsive to diverse types of biological motion (Allison, Puce, & McCarthy, 2000; Frith & Frith, 2007), which appears to be the putative homolog of the macaque area STP (Beauchamp, Yasar, Frye, & Ro, 2008). Biological motion processing in this region appears to mostly concern

kinematic aspects, whereas, in the LOC, it appears to mostly concern configuration changes of the observed actions, suggesting, as in macaques, a dual-stream processing of action observations (Jastorff & Orban, 2009). These sectors of the pSTS/MTG and LOC are considered the major source of visual action information for the human mirror system (Caspers et al., 2010; Grosbras, Beaton, & Eickhoff, 2012; Molenberghs, Sale, & Mattingley, 2012; Rizzolatti et al., 2014).

In the human brain, there is a conspicuous fiber system linking the human rostral IPL with the posterior part of the superior, middle, and inferior temporal gyri, which has been identified as the posterior segment of the arcuate or of the SLF (Catani, Jones, & ffytche, 2005; Martino et al., 2013; Wu, Sun, Wang, Wang, & Wang, 2016). Connectivity between the Superior temporal gyrus (STG) and SMG could also be supported by the middle longitudinal fasciculus (Makris, Zhu, Papadimitriou, Mouradian, & Ng, 2016). Thus, it is possible that human temporo-parietal connectivity includes components equivalent to the macaque pathways connecting TEa/m with area AIP and area STP with area PFG, and that the human temporal cortex includes areas which take part in what is possibly the human counterpart of the macaque lateral grasping/action recognition network.

To our knowledge, there is no clear evidence for activation of prefrontal areas other than the IFG during the mere execution or the observation of hand-object interactions. However, there is clear clinical, electrophysiological, and imaging evidence for the involvement of the human middle frontal gyrus (MFG)—for the most part, the putative homolog of the macaque ventral area 46 (Petrides, 2005)—in different aspects of the executive control of motor behavior (Goldenberg & Spatt, 2009; Haaland, Harrington, & Knight, 2000; Rowe, Stephan, Friston, Frackowiak, & Passingham, 2005) including hand actions. Indeed, TMS over this region affects free selection of hand actions (Hadland, Rushworth, Passingham, Jahanshahi, & Rothwell, 2001) and modulates the excitability of the primary motor cortex, showing temporally and spatially selective interaction between these two areas (Hasan et al., 2013). Furthermore, functional imaging evidence showed involvement of the MFG during the preparation of contralateral and ipsilateral hand actions (Gallivan, McLean, Flanagan, & Culham, 2013). Finally, the MFG displays visual object-related activation (Denys et al., 2004), activates during texture recognition (Stylianou-Korsnes, Reiner, Magnussen, & Feldman, 2010), and also appears to be involved in tactile object recognition (Lacey, Flueckiger, Stilla, Lava, & Sathian, 2010; Reed et al., 2004; Savini et al., 2010). Cieslik et al. (2013) have suggested a rostrocaudal subdivision of the MFG into two distinct subregions in which the caudal one is characterized by functional connectivity with bilateral intraparietal sulci, including the location of the human area AIP and appears to be more strongly related to action execution and working memory. The second and the third branches of the SLF connecting the IPL with the frontal lobe (Makris et al., 2005; Thiebaut de Schotten et al., 2012) and the frontal inferior longitudinal tract connecting the precentral gyrus with the MFG (Catani, Dell'Acqua & Vergani et al., 2012; Rojkova et al., 2016) could represent the possible substrate for the participation of this region in the putative human counterpart of the lateral grasping/action recognition network. Furthermore, the inferior fronto-occipital

fasciculus, linking temporal areas (including the LOC and caudal temporal areas) with prefrontal areas (including the MFG) (Sarubbo, De Benedictis, Maldonado, Basso, & Duffau, 2013; Thiebaut de Schotten et al., 2012) could provide the substrate for a connectivity equivalent to the macaque inferotemporal connectivity with hand-related VLPF areas. However, note that in the human brain there is a component of the arcuate fasciculus (long direct segment, Catani et al., 2005), directly connecting the temporal with the frontal cortex, for which there is no equivalent in the macaque brain (Rilling et al., 2008).

Functional data have also provided evidence for a putative human homolog of the hand-related sector of the macaque insula, which activates during the execution and observation of hand actions with a vitality form, suggesting a modulation of the cortical circuits for controlling hand actions according to the internal state of the individual (Di Cesare et al., 2014, 2015). Observations based on dMRI have provided evidence for the connectivity of the insula with the PMv, the IFG, the MFG, and the IPL (Cerliani et al., 2012; Di Cesare et al., 2018; Ghaziri et al., 2017).

Finally, the rostral part of the medial premotor cortex, based on architectonic and functional data, has been considered the homolog of the macaque area F6/pre-SMA (Geyer et al., 2000; Nachev et al., 2008; Picard & Strick, 1996; Zilles et al., 1996). As in the macaque, this area appears to play a more general role in several higher-order aspects of motor control (see Geyer, Luppino, & Rozzi, 2012; Nachev et al., 2008). Specifically, recent evidence showed that the human pre-SMA could play a role, together with the IFG, in the neural mechanisms underlying response inhibition (Angelini et al., 2015; Aron, Behrens, Smith, Frank, & Poldrack, 2007; Swann et al., 2012). A bundle of fibers connecting the pre-SMA and the rostral SMA with the PMv/IFG, designated as the frontal aslant tract (Catani, Dell'Acqua & Vergani et al., 2012; Rojkova et al., 2016), is likely the substrate for this interaction.

In sum, there is robust evidence that, in the human brain, there is a set of potentially linked parietal, temporal, and frontal areas, which, as a result of their topology and functional properties, appear very likely to form a human counterpart of the macaque lateral grasping/action observation network.

However, this same set of cortical nodes, or at least part of it, also appears to be involved in cognitive abilities unique or almost unique to humans, such as higher-order aspects of organization of object-oriented actions, including tool use (Johnson-Frey, Newman-Norlund, & Grafton, 2005; Peeters, Rizzolatti, & Orban, 2013, 2009; Ramayya et al., 2010), imitation, and imitation learning (Buccino et al., 2004). Specifically, functional studies have shown that tool use action planning, execution, and observation and tool observation and naming activate a set of cortical regions of the left hemisphere, including the LOC and posterior MTG in the temporal cortex, the anterior IPS and SMG in the parietal cortex, and the PMv/IFG and MFG in the frontal cortex, which largely overlap with the putative human lateral grasping/action observation network (Brandi, Wohlschlagler, Sorg, & Hermsdorfer, 2014; Choi et al., 2001; Johnson-Frey et al., 2005; Moll et al., 2000; Peeters et al., 2009, 2013). Based on comparative observations in humans and macaques, Peeters et al. (2009, 2013) have concluded that the left rostral SMG also includes an

evolutionarily new human-specific zone specifically devoted to tool use, which could have differentiated from phylogenetically older rostral IPL hand-related areas.

Furthermore, there is evidence that imitation or imitation learning of hand actions activates temporal, rostral IPL, and PMv/IFG regions involved in action observation and, especially for imitation learning, the MFG (Buccino et al., 2004; Caspers et al., 2010; Higuchi, Holle, Roberts, Eickhoff, & Vogt, 2012; Rizzolatti et al., 2014; Vogt et al., 2007). Based on comparative dMRI observations, Hecht et al. (2013) have suggested that stronger and more extensive connectivity of the SMG with the pSTS/MTG and LOC regions involved in shape and action coding, as well as stronger connectivity between the SMG and PMv/IFG, differentiate the cortical mirror system of humans from that of macaques and could have contributed to the emergence of the role of this system in imitation and imitation learning (see also Rizzolatti et al., 2014). Furthermore, the differentiation of the MTG areas responsible for storing conceptual and semantic information about tools and of the rostral SMG sector devoted to tool use, and their interconnectivity, could have contributed to the emergence of tool use from a specialization of neural mechanisms for controlling hand–object interactions shared with macaques (Orban & Caruana, 2014; Peeters et al., 2013, 2009; Ramayya et al., 2010).

5. Possible human counterpart of the macaque networks for explorative and communicative oculomotor behavior

A large number of human functional studies have shown that tasks requiring the execution of visually- or memory-guided saccades and/or shifts of attention in the visual field activate several foci located in the posterior parietal and frontal cortex (e.g., Alvarez, Alkan, Gohel, Douglas Ward, & Biswal, 2010; Corbetta et al., 1998; Curtis & Connolly, 2008; Dieterich et al., 2009; Koyama et al., 2004; Petit, Clark, Ingeholm, & Haxby, 1997; Petit & Haxby, 1999). In the posterior parietal cortex, one of these foci, observed in virtually all studies, is located dorsally and posteriorly in the medial bank of the IPS in a sector which has been designated as the dorsal IPS medial (DIPSM, see Orban et al., 2004) and is usually referred to as the “parietal eye field” (PEF). Based on different lines of functional evidence, there is a general consensus that this sector corresponds to the macaque area LIP (Orban, 2016). The finding that the possible human equivalent of the macaque area LIP is located in the medial, and not in the lateral bank of the IPS, is an example of the possible differences in the topology of equivalent areas between the macaque and human brains, which, in this case, could be accounted for by the disproportionate increase of the human IPL. Some studies (e.g., Corbetta et al., 1998; Koyama et al., 2004) have also described additional foci, one of them located more rostrally in the IPS, whose possible macaque equivalent still remains to be verified. In the frontal cortex, one focus constantly observed in all studies is located within the dorsal part of the precentral sulcus at the junction with the superior frontal sulcus. There is unanimous consensus that this sector corresponds to the macaque PEF. However, the exact location of this field tends to vary across different studies. According to a

high-resolution fMRI study, this field is located in the anterior bank of the precentral sulcus and corresponds to a distinct chemoarchitectonic area (Rosano et al., 2002, 2003). An additional, constantly observed focus is located in the medial frontal gyrus and is considered to correspond to the macaque SEF. Furthermore, several studies have provided evidence for at least one additional frontal oculomotor field located ventral to the human FEF in the precentral sulcus and usually referred to as the “inferior” FEF (Amiez & Petrides, 2009; Corbetta et al., 1998; Derrfuss, Vogt, Fiebich, von Cramon, & Tittgemeyer, 2012; Heide et al., 2001; Koyama et al., 2004; Luna et al., 1998; Mort et al., 2003). Though varying in location across studies, this field has been usually attributed to Brodmann area 6 (Brodmann, 1909) and a homology with a possible oculomotor premotor field of the macaque brain has been then suggested (Amiez & Petrides, 2009; Koyama et al., 2004). However, evidence for a postarcuate oculomotor field in the macaque brain is not univocally supported by fMRI data (Baker, Patel, Corbetta, & Snyder, 2006; Koyama et al., 2004; Premereur et al., 2015), nor is it supported by electrophysiological studies. Furthermore, the attribution of this field to Brodmann area 6, does not necessarily imply that it is actually a premotor field located within architectonic area 6. Indeed, the human FEF is also located within the limits of Brodmann area 6 but, as reviewed above, is a distinct architectonic granular area (Rosano, Sweeney, Melchitzky, & Lewis, 2003) and a homolog of a macaque prefrontal area (area 8/FEF). It is noteworthy that all the above mentioned studies have not considered that, in the macaque caudal prefrontal cortex, there are several oculomotor areas, including area 45B, which are located just ventral to the FEF and, as reviewed in Section 3, activate during the execution of saccadic eye movements. Finally, Patel et al. (2015) found that, in human subjects, visuospatial attentional tasks activated, in addition to the FEF, two other fields, one apparently located ventral to the inferior FEF in the precentral sulcus and the other located in the inferior frontal sulcus. Additional comparative observations are needed in order to examine the possible homologies between the human and the macaque frontal oculomotor systems in the light of data showing a multiplicity of caudal oculomotor prefrontal areas in the macaque.

All together, these data provide evidence for a parieto-frontal circuit corresponding to the macaque LIP-FEF circuit involved in oculomotor control and in a dorsal attention network for controlling spatial and featural attention (Corbetta & Shulman, 2002). The First branch of the superior longitudinal fasciculus (SLFI) could provide the substrate for the connectivity in the human brain between the PEF and the FEF, as suggested by Thiebaut de Shotten et al. (2011) and, possibly, between the PEF and other frontal oculomotor fields. In fact, dMRI evidence for connectivity between the Superior parietal lobule (SPL) and inferior frontal areas has been provided by Hecht et al. (2013).

Functional studies have also suggested the interaction of ventral visual stream areas with the parieto-frontal oculomotor circuitry. For example, Preston, Guo, Das, Giesbrecht, and Eckstein (2013) have shown that, in a visual search task, the LOC appears to play a role in coding the contextual location of objects and features in real scenes, and they suggested that information on the likely location of the targets could be relayed from the LOC to the parieto-frontal

oculomotor network for directing attention to the contextually relevant location.

Furthermore, there is evidence that parietal and frontal oculomotor areas are also involved in the neural mechanisms underlying gaze perception and joint attention, together with temporal areas (Bristow, Rees, & Frith, 2007; Grosbras, Laird, & Paus, 2005; Hooker et al., 2003; Nummenmaa & Calder, 2009; Williams, Waiter, Perra, Perrett, & Whiten, 2005; see also; Shepherd, 2010). Specifically, the involved temporal regions include the posterior STS/MTG specialized for perceiving social signals mediated by biological motion including gaze shifts (Allison et al., 2000; Blakemore, Winston, & Frith, 2004; Caruana et al., 2014; Marquardt, Ramezanpour, Dicke, & Thier, 2017) and the fusiform/LOC regions specialized for face processing, which can be modulated by the configuration of the gaze (George, Driver, & Dolan, 2001). In the frontal lobe, in addition to the FEF, some studies have observed activation in a more ventral region involving the IFG (Hooker et al., 2003) or the junction between the inferior frontal sulcus and the precentral sulcus (inferior frontal junction – IFJ, Bristow et al., 2007; Grosbras et al., 2005; Nummenmaa & Calder, 2009). It is noteworthy that a cortical zone located at the IFJ was found to activate during the observation of faces only when the eyes were not masked (Chan & Downing, 2011). These data suggest homology of this region with the macaque area 45A.

In the human brain, there is evidence, based on dMRI and fiber dissection observations, for a temporal connectivity of the SPL. Specifically, these connections appear to involve, in the temporal cortex, the rostral and caudal regions located in the STG, MTG, and Inferior temporal gyrus (ITG), including the fusiform gyrus and, in the SPL, the caudal part corresponding to Brodmann's area 7 (Makris et al., 2013, 2016, 2009; Hecht et al., 2013; Kamali, Sair, Radmanesh, & Hasan, 2014; Wang et al., 2013; Wu et al., 2016). These connections appear to run through the MdLF and a component of the Arcuate fasciculus (AF)/SLF (Makris et al., 2013, 2016, 2009; Kamali et al., 2014; Wang et al., 2013; Wu et al., 2016). This temporo-SPL fiber system has been described as evolutionarily new, not present in the macaque brain (e.g., Hecht et al., 2013; Wang et al., 2013), and as a possible substrate for a role of the human SPL in action observation (Abdollahi, Jastorff, & Orban, 2013) or visuo-auditory attentional processing (Hecht et al., 2013; Makris et al., 2016; Wang et al., 2013). However, it is possible that components of this connectivity are equivalent to the temporal connectivity of the macaque area LIP. Furthermore, it is possible that components of the arcuate fasciculus and inferior fronto-occipital fasciculus linking temporal to frontal areas are equivalent to the temporal connections of the macaque frontal oculomotor areas.

All together, these data suggest that, as in the macaque, the human brain hosts a large-scale temporo-parieto-frontal circuitry for oculomotor control, covert shift of attention, and gaze perception. Accordingly, humans and non-human primates appear to share neural circuits and mechanisms for basic gaze-following behavior, which, in humans, could have been the foundation for more sophisticated social skills, such as mutual awareness of shared mental states (Shepherd, 2010).

6. Human-specific functions versus human-specific areas

It is largely agreed in comparative neuroscience that primates share common principles of cortical organization. Indeed, primates display similar layouts of homologous sensory, motor, and association areas organized in similar sensorimotor domains, which, to a large extent, can be identified even in prosimians (Kaas & Stepniewska, 2016). Furthermore, there are very strong similarities in the organization of the major fiber tracts connecting parietal with frontal areas (Schmahmann et al., 2007; Thiebaut de Schotten et al., 2012). Accordingly, primates appear to share common plans of the organization of sensorimotor functions, which have been likely conserved along the various lineages that have differentiated during primate evolution.

In the evolution of the lineage leading to *Homo sapiens*, there have also been substantial changes in brain size and organization, which are considered to be at the basis of some higher-order human- or almost human-specific cognitive functions. According to Darwin's theory, the evolution of complex structures is incremental, so that human specific mental functions would derive from phylogenetically older mental processes with gradual evolutionary trajectories. However, it has been proposed that human-specific mental functions reflect discontinuities pervading nearly every domain of cognition (Penn, Holyoak, & Povinelli, 2008). de Waal and Ferrari (2010) have argued that the concept of discontinuity results from the adoption of a “top-down” perspective in comparative cognitive research in which the main question is which animals possess or do not possess a given cognitive ability. Conversely, if a “bottom-up” approach is adopted, focusing on the constituent capacities underlying larger cognitive phenomena, it appears quite clear that the basic building blocks of cognition might be shared across a wide range of species, suggesting mental continuity in primate evolution (de Waal & Ferrari, 2010; Sherwood, Subiaul, & Zawidzki, 2008). In line with this view, there are theories positing that one central organizational principle of the functional structure of the brain is based on reuse of neural, often sensorimotor, circuitries for various cognitive purposes (“neural reuse theories”, see Anderson, 2010). Specifically, according to these theories “neural circuits established for one purpose are commonly exapted (exploited, recycled, redeployed) during evolution or normal development, and put to different uses, often without losing their original functions” (Anderson, 2010). Two of these theories differ on the time course over which they operate. One – the massive redeployment theory (Anderson, 2007) – concerns the evolutionary emergence of the functional organization of the brain, the other – the neuronal recycling theory (Dehaene, 2005; Dehaene & Cohen, 2007) – explains those cognitive abilities for which there has been insufficient time for specialized neural circuits to have evolved.

Taken for granted that in primate evolution neural reuse for developing progressively more complex cognitive abilities has been paralleled by a disproportionate increase in size of some cortical regions, one important issue is to what extent regional expansions have resulted in the generation of evolutionary new cortical areas.

In addressing this issue, it is important first to note what makes a cortical area. It is largely accepted that the cerebral cortex contains many distinct entities, usually referred to as “areas,” although it has been matter of debate what precisely constitutes a cortical area and what the best criteria for their definition are (see, e.g., Van Essen, 1985). In general, three main criteria, the architectonic, the connectional, and the functional, are considered most useful for the definition of a cortical area. Converging evidence, based on these criteria, is generally considered a strong argument for reliable identification and delineation of a cortical area (see, e.g., Felleman & Van Essen, 1991; Van Essen, 1985). For example, in macaques, the IPL convexity cortex has been subdivided into four distinct areas based on converging architectonic (Gregoriou, Borra, Matelli, & Luppino, 2006), connectional (Rozzi et al., 2006), and functional (Rozzi, Ferrari, Bonini, Rizzolatti, & Fogassi, 2008) evidence.

In human studies, the term “area” is very often simply used to designate cortical zones which appear to have specific functional properties. However, the functional distinctiveness of a given cortical zone does not necessarily imply that it corresponds to a distinct area as defined above. Indeed, a functionally distinct cortical zone could correspond to a module of a larger cortical area, or could extend over adjacent cortical areas which share some common functional features. Unfortunately, the issue of the areal attribution of functional data, even when addressed, is often seriously prevented by the coarseness of the architectonic maps for several brain regions. It is noteworthy, however, that, for some human cortical regions, a higher functional complexity seems not to be paralleled by a higher architectonic complexity, with respect to the corresponding regions of the macaque brain. This appears to be the case of the parietal lobe in which detailed architectonic studies have identified a number of superior parietal, intraparietal, and inferior parietal areas almost comparable to those of the macaque (Caspers et al., 2006; Choi et al., 2006; Scheperjans et al., 2008). For example, in the inferior parietal lobule, the evolutionarily new human-specific tool use SMG zone appears to be located within architectonic area PFT, which also activates during hand action execution, observation, and imitation and has been considered the possible homolog of the macaque area PFG (Caspers et al., 2011, 2010; Gazzola & Keysers, 2009; Peeters et al., 2009). Conversely, in some human cortical regions, the architectonic organization appears to be more complex than in the corresponding regions of the macaque brain. This is the case, for example, of the caudal IFG, which, based on chemoarchitectonic data, has been subdivided into several areas (Amunts et al., 2010) that could in part correspond to evolutionarily new areas not present in the macaque brain. In sum, there is no compelling evidence that evolutionarily new cognitive capacities in humans are necessarily linked to the addition of evolutionarily new cortical areas as properly defined (e.g., Preuss, 2011; Sherwood et al., 2008).

7. Concluding remarks

In the present article we have reviewed comparative observations showing that some human- or almost human-specific functions, such as tool use, imitation learning, and language,

appear to involve cortical zones of the parietal and frontal cortex which overlap or, at least, are in contiguity with the nodes of the human counterpart of the macaque lateral grasping/action recognition network. These observations suggest that neural mechanisms underlying tool use could have emerged from the exploitation and adaptation of phylogenetically older neural mechanisms, shared with macaques, underlying the selection and control of object-oriented hand actions (see, e.g., Orban & Caruana, 2014). Similarly, imitation and imitation learning could have emerged from the exploitation and adaptation of phylogenetically older neural mechanisms, shared with macaques, involved in mapping observed actions into their corresponding motor representations (see e.g., Rizzolatti et al., 2014). It has been also suggested that the neural mechanism for recognizing actions made by others (mirror mechanism) could have represented a neural prerequisite for the development of interindividual communication and, finally, of speech (Pulvermüller, 2018; Rizzolatti & Arbib, 1998). Similar considerations could be done also for the neural mechanisms underlying some higher-order aspects of social attention and interactions based on gaze perception, which appear to involve cortical nodes of the putative human counterpart of the macaque networks for explorative and communicative oculomotor behavior (Nummenmaa & Calder, 2009; Shepherd, 2010).

In sum, data reviewed above provide clear examples of the fact that the monkey and the human brain share neural circuits for sensory, motor, and cognitive motor functions, likely inherited from the last common ancestor and the neural mechanisms mediated by these circuits could have represented building blocks for the generation, based on a process of neural reuse, of higher order human specific functions. Obviously, the degree of detail of this type of comparative observations would greatly benefit from a desirable future development of non-invasive techniques for a more detailed definition of point-to-point cortical connectivity and the structural correlation of functional data.

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