### 4.15 On the Evolution of the Frontal Eye Field: Comparisons of Monkeys, Apes, and Humans

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#### Abstract

This chapter compares the location, extent, and boundaries of the frontal eye field (FEF) in prosimians, New World monkeys, Old World monkeys, apes, and humans. We are particularly interested in understanding how the location of FEF in humans relates to that in other primate species. We consider whether FEF should be understood as a collection of distinct areas or as a single area with multiple modules that can be of different sizes (and locations) in different species according to lifestyle and habitat. We also consider how the size and location of FEF is a matter of operational definition with resolution specified by the various techniques applied, making comparison across species, especially humans, challenging. Locating FEF is challenged further by the variation of sulcal morphology across species and individuals. Nevertheless, in all investigated primate species FEF appears to be located at the transition zone between premotor area 6 and prefrontal area 8 in a cortical region with significant sulcal variation where it is bordered caudally (in premotor cortex) and rostrally (in prefrontal cortex) by areas also contributing to visually guided gaze behavior. We highlight current areas of uncertainty. In particular, lacking information about precise boundaries, we do not know whether the size and modularity of FEF scales across species with overall neocortical area, or with the complexity of visual and other sensory systems, or with changes in associated subcortical circuits, or with the complexity of the musculature used for visually guided behaviors, or with the complexity of cognitive control over gaze behavior.

#### 4.15.1 Overview

Originally considered only an ocular motor area, interest in frontal eye field (FEF) has increased markedly with its recognition as an anatomical hub region, which is critically involved in covert and overt orienting in monkeys and humans (reviewed by Squire et al., 2013; Schall, 2015). Thus, FEF is associated with motor, sensory, and cognitive processes, which are mediated in association with neighboring cortical areas.

Ferrier (1874) described an area in the dorsal portion of the arcuate sulcus of anesthetized monkeys where electric stimulation elicited movements of the eyes and head in the direction opposite the stimulated hemisphere. These are referred to as contraversive movements. This finding has been replicated in multiple species including prosimians, New World monkeys, Old World monkeys, apes, and humans. Förster (1926) described the area from which he could evoke eye movement as the “frontales Augenfelds,” which was translated into “frontal eye fields” (Davidoff, 1928; Förster and Penfield, 1930).
Before proceeding, we introduce our nomenclature. This chapter describes the location of the FEF assessed with a variety of methods. For efficiency we use phrases such as "FEF is located ..." when we should more clearly state "A region activated during saccades ..." or "A region from which saccades were elicited ..." We emphasize this here to alert the reader that the location and boundaries of the FEF(s) are uncertain, operational, and have rarely been compared systematically across methods.

Today, the majority of research on FEF is done with macaque monkeys and humans, and this chapter, like previous reviews (Paus, 1996; Tehovnik et al., 2000; Amiez and Petrides, 2009; Vernet et al., 2014; Percheron et al., 2015) is animated by an apparent discrepancy in the location of FEF between monkeys and human. In both macaque and human the FEF is located in a sulcus rostral to the central sulcus, known either as arcuate or precentral (see Connolly, 1936, 1950). All authors agree that FEF in monkeys is located in the dysgranular region of caudal prefrontal cortex commonly included in Brodmann's area 8; however, most locate FEF in humans in Brodmann's agranular area 6. We address this apparent enigma by highlighting the following observations: (1) particular similarities and differences in eye-, head-, and body-orienting behavior across species, (2) variation of frontal sulcal morphology across species and individuals, (3) the location of FEF at a cytoarchitectonic transition zone between agranular area 6 and granular area 8 in a cortical region with greater than average variation of sulcal pattern, and (4) the embrace of FEF caudally (in premotor cortex) and rostrally (in prefrontal cortex) by areas also contributing to visually guided gaze behavior. These observations are framed by an appreciation of the varying technical limits and spatial resolution of methods used to locate FEF in monkeys, apes, and humans, and of the functional and anatomical position of FEF in a surrounding network of areas contributing to orienting behavior that includes but is not limited to eye movements.

4.15.2 Gaze Control and Coordination in Prosimians, Monkeys, Apes, and Humans

We briefly review key similarities and major differences in the organization of the eye and head across species. The structure of the neural circuits producing eye movements will naturally vary with differences in the nature of movement execution dictated by the size and configuration of the eye, head, body, and habitat.

A search for homologies among primates in the cortical and subcortical organization of eye movements is encouraged by the many similarities in gaze behavior observed between monkeys, apes, and humans. Members of each species produce each of the different types of eye movements—fixation maintained by gaze-holding vestibular and visual reflexes, which is interrupted by rapid gaze shifts and slow pursuit eye movements to direct vision in three dimensions. In numerous quantitative respects the eye movements of macaque monkeys and humans are more similar than different in execution but can differ in planning and context (eg, Martinez-Conde and Macknik, 2008; Baizer and Bender, 1989; Einhäuser et al., 2006; Berg et al., 2009; Shepherd et al., 2010). In a wide range of testing conditions, the advanced cognitive control of eye movements is indistinguishable between humans and macaque monkeys (eg, Munoz and Everling, 2004; Camalier et al., 2007; Hanes and Schall, 1995; Hanes and Carpenter, 1999; Nelson et al., 2010). Consequently, the macaque has been a faithful source of insights for human clinics of neurology (eg, Leigh and Zee, 2015) and psychiatry (eg, Driscoll and Barr, 2016). Comparisons of the eye movements of chimpanzees, gorillas, and orangutans with those of humans also have noted numerous similarities with particular differences (Kano et al., 2011; Kano and Tomonaga, 2009, 2011). Primates in general are visual creatures with particular ocular, muscular (eg, Blumer et al., 2016), and neural adaptations including devotion of a large proportion of the brain to visual processing.

Primates execute gaze shifts to scrutinize objects with the high acuity visual resolution afforded by the area centralis or fovea. There are qualitative differences in the organization of central retina between nocturnal and diurnal primates related to their gaze behavior that may be reflected in differences in the ocular motor network. With the exception of the nocturnal owl monkey and Strepsirrhini, all Haplorrhini, including the tarsier, possess a fovea (Wolin and Massopust, 1967). The absolute size of the fovea is comparable across diurnal primates, and a higher acuity of vision is achieved by increasing the size of the eye; for example, human eyes are about four times larger than marmoset eyes (Finlay et al., 2008). On the other hand, galagos and owl monkeys have a rudi-

In a surrounding network of areas contributing to orienting behavior that includes but is not limited to eye movements.

The unique visual abilities of primates are conferred primarily by a retina equipped with a specialized central region of high resolution that is associated with a sophisticated neural system producing large variety, range, and coordination of eye movements to gather visual information. The large oculomotor range that primates possess appears to be evolutionarily recent. The human oculomotor range spans about ±50 degrees of visual angle (Guitton and Volle, 1987; Stahl, 1999) and is nearly matched by that of the macaque monkey (Tomlinson and Bahra, 1986) and baboon (Marchetti et al., 1983). In contrast, the oculomotor range of more distantly related primates such as prosimians (Shepherd and Platt, 2006) or the New World squirrel monkey (McCrea and Gdowski, 2003; Heiney and Blazquez, 2011) and marmoset (Mitchell et al., 2014) is limited to <15 degrees. Limited ocular motility is commonly reported in nonhuman primates with small heads and poorly developed fovea, such as tree shrew (Remple et al., 2006) and cat (Guitton et al., 1984). Larger gaze shifts are accomplished with a combined rotation of the head and eyes. What qualifies as "larger" varies across species. The greater inertial mass of the larger heads of apes and humans makes head movements more energetically costly, slower, and socially revealing. Thus, while smaller prosimians and New World monkeys naturally make gaze shifts of more than about 5 degrees with a head rotation.
too (e.g., McCrea and Gdowski, 2003), humans naturally make gaze shifts of more than about 20 degrees with a head rotation too. Moreover, relative to apes and monkeys, the human eye is positioned further forward in the orbit, which expands the effective visual field with horizontal eye movements achieving larger abductions (Denison et al., 2015a). Such unique orbital morphology is adaptive for bipedal locomotion in a habitat of open planes (Denison et al., 2015b). Horizontal eye movements are more common for terrestrial species compared with arboreal species (Kobayashi and Kohshima, 2001).

The eyes may be a window of the soul for humans, but gaze direction is camouflaged in other primates by coloration of the sclera resembling surrounding skin color. Comparing the external features of eyes from 88 primate species, humans are the only primate with a white sclera and the largest amount of exposed sclera relative to the outline of the eye (Kobayashi and Kohshima, 1997). This morphology affords a clear visual marker of gaze direction that facilitates communication (Kobayashi and Kohshima, 1997, 2001). Gaze following is observed across primate species (Rosati and Hare, 2009; Shepherd, 2010), but most often it utilizes the orientation of the head. The reliance on the direction of the eye seems to be specific to humans (Tomasello et al., 2007). In summary, the specialization of the fovea and the oculomotor range delimit the visual ability of primates and rationalize the amount of cortex dedicated to visual processing.

4.15.3 Variability of Sulci in the Primate Frontal Cortex

In every primate species including humans FEF is located in a sulcus (or dimple) immediately rostral to the central sulcus. In humans this is referred to as the precentral sulcus, but in monkeys, the arcuate sulcus. Mingazzini (1888) first coined the term arcuate sulcus ("solco arcuato") to describe the inferior precentral sulcus in the Old World monkey, which others had previously called "sillon coube" by Gromier or "sillon lypsoïde" by Broca (see Hervé, 1888).

The location of FEF in standardized coordinates for the individual species can vary in proportion to variability of the morphology of this sulcus. The high variability of sulcal patterns in humans is well known, and a variety of patterns can be identified across individuals (Ono et al., 1990). The precentral sulcus is interrupted into superior and inferior segments in over 50% of individuals; three segments are found in about 30% of individuals, and four segments are found in fewer than 5%. About 30% of individuals also exhibit a longitudinally oriented marginal precentral sulcus, reminiscent of the precentral dimple in macaques. The major superior and inferior segments develop at different times. In more than 90% of individuals the superior precentral sulcus connects with the superior frontal sulcus, and the inferior precentral sulcus connects with the inferior frontal sulcus. None of these tendencies are the same across hemispheres.

The first reference to a sulcus praecentralis referred only to the inferior precentral sulcus (IPrCS) (Ecker, 1869). According to Eberstaller (1890), the superior precentral sulcus was first described as a separate sulcal entity by Jensen (1871). Recent studies have further subdivided these sulci—Germann et al. (2005) distinguished three parts of the IPrCS: a dorsal and a ventral part, and a so-called horizontal extension separating the other two parts. Also, the superior precentral sulcus was subdivided into a ventral and a dorsal part, separated by the superior frontal sulcus (Amiez et al., 2006).

Naturally, the location of FEF and surrounding areas will vary as a function of systematic and random variability of sulcal patterns. In fact, as detailed in the following, Amiez et al. (2006) conducted a subject-by-subject analysis of the locus of eye movement–related functional activity revealed in relation to the detailed morphology of the precentral and superior frontal sulci. A focus of activation associated with saccadic eye movements was located in the ventral branch of the superior precentral sulcus in both hemispheres. A second focus has been found in the dorsal part of the inferior precentral sulcus. Imaging during a hand response selection task revealed activation focused in the dorsal branch of the superior precentral sulcus close to the caudal end of the superior frontal sulcus. Activation in primary motor cortex was focused in the precentral knob (or Broca’s pli de passage moyen). The relationship of FEF to the bordering premotor cortex is elaborated in the following paragraphs.

Compared to humans, the cerebral sulcal pattern is much less variable across Old World monkeys (Cercopithecidae) (Falk, 1978). The frontal lobe consists of an arcuate sulcus rostral to the central sulcus and a longitudinal principal sulcus (known also as sulcus rectus) rostral to the arcuate sulcus (Connolly, 1936, 1950; Walker, 1940). The arcuate sulcus consists of a superior (horizontal) limb or branch and an inferior (vertical); at the genu or curve, typically located caudal to the end of the principal sulcus, some individuals exhibit a posterior extension of the sulcus referred to as a "spur." Thus, the shape and extent of the arcuate sulcus in monkeys varies. However, no one has systematically characterized the variation of its sulcal morphology. For example, the prevalence of the arcuate spur is unknown in various macaque species. Therefore, we investigated the incidence of an arcuate spur in a sample of 235 hemispheres obtained from nine direct observations before histological processing. 132 magnetic resonance (MR) images, and 109 figures from the literature. This sample consists of 221 left hemispheres and 161 right hemispheres; for 147 brains both hemispheres were available. Samples were obtained from Macaca fascicularis (n = 19), Macaca mulatta (n = 162), Macaca nemestrina (n = 7), and Macaca radiata (n = 21). We could determine the gender of 54 male and 7 female monkeys. Table 1 summarizes the findings. In about one-third of both left and right hemispheres a large spur was present, another one-third exhibited a small spur, and the remaining one-third exhibited no spur. Thus, two-thirds of hemispheres exhibit an arcuate spur. In about two-thirds of hemispheres the spur was symmetric in appearance in both hemispheres.

These values may underestimate the prevalence of an arcuate spur because we found a spur more often in our sample of histological (89%) and MR (86%) images as compared with figures in the literature (57%). We cannot evaluate how faithfully authors portray cortical sulcal patterns. Also, even though the sample of female macaques was small, we found no clear difference in
prevalence between females (71%) and males (67%). An arcuate spur was most common in *M. mulatta* (75%) and *M. nemestrina* (71%), only slightly less common in *M. radiata* (67%), and least common in *M. fascicularis* (53%).

These modest findings have several implications. First, atlases based on averages of macaque brains (e.g., Frey et al., 2011; Rohlfiing et al., 2012; Calabrese et al., 2015) imply that sulcal patterns are more regular and uniform across individuals than they actually are. Such averaged atlases underestimate the length of the spur observed in many individuals by representing only a small spur (Frey et al., 2011) or none at all (Calabrese et al., 2015; Rohlfiing et al., 2012). Likewise, atlases based on a single individual will misrepresent the incidental presence or lack of a spur as standard anatomy (e.g., Saleem and Logothetis, 2012).

Second, sulcal patterns can serve as reliable landmarks to identify the location of cortical areas. For example, in macaques the location and shape of the arcuate sulcus reliably predicts the location of FEF. Also, the arcuate spur has also been identified as a boundary between functionally and anatomically distinguished dorsal and ventral premotor areas (e.g., Gabernet et al., 1999).

Third, a variety of mechanisms for cortical folding have been proposed—differential cell proliferation, differential tangential expansion, radial intercalation, axon tension, radial glia, and so on (see for review Striedter et al., 2015; see also Mota and Herculano-Houzel, 2015; Tallinen et al., 2016). These data cannot distinguish among these alternatives, but they offer an opportunity for further exploration. Finally, variation of structure most likely announces variation of function. The common lack of symmetrical occurrences of the spur represents clear evidence for cerebral asymmetry in macaques. While others have reported a rightward bias in the length of the arcuate sulcus in macaques (Sakamoto et al., 2014; Imai et al., 2011), some have reported no asymmetry (Falk et al., 1990; Heilbroner and Holloway, 1989). Whether such asymmetry is systematic and functionally meaningful requires a larger sample. However, behavioral and anatomical evidence for some degree of cerebral asymmetry in macaques has been reported (e.g., Hamilton and Vermeire, 1988; Heilbroner and Holloway, 1989; Falk et al., 1990). Perhaps an arcuate spur relates to individual hand preferences (e.g., Lehman, 1980; Mangalam et al., 2014). In any case, when quantified in the MRI atlases of macaque brains, the magnitude of deformation needed to co-register the set of macaque brains varies across the cortex and was noticeably elevated in the region of the spur (Frey et al., 2011; Calabrese et al., 2015). Does this greater variability in the organization of this region provide greater opportunity for natural selection? Further research is needed to determine how the variation of sulcal morphology relates to the location and boundaries of FEF and neighboring areas.

### 4.15.4 Location of FEF Across Primate Species

#### 4.15.4.1 Scandentia

Together with Dermoptera, for which very little is known, Scandentia is the closest mammalian order relative to Primates (Perelman et al., 2011). Most of their visual specializations evolved independently of primates. They inhabit tropical forests of southern Asia. Electrical stimulation mapping of the (small) frontal lobe has not found a discrete FEF, although stimulation of sites in frontal motor areas did elicit eye blinks (Lende, 1970; Rempel et al., 2006; Baldwin et al., 2016). This may not be not surprising, because tree shrews have neither a specialized focus of high resolution in the retina nor much range of eye movements. However, they are very visual creatures that orient readily with head and body.

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<th>Spur size</th>
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<td>Large</td>
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<td>Symmetric</td>
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#### 4.15.4.2 Strepsirrhini: Prosimians

Most prosimians are nocturnal animals. As noted in the earlier section, the retina lacks a clear fovea and instead has only a rudimentary area centralis. Their relatively poor distance vision is compensated by a long snout ending in a moist and touch-sensitive surface, similar to that of dogs. The wet nose provides a conduit for pheromones into the vomeronasal organ.

Prosimians resemble other primates and are distinct from other mammals in the presence of a clear granular prefrontal cortex rostral to the agranular motor cortex. Therefore, prosimians are a good model for the evolution of FEFs, because their frontal cortex is subdivided into a granular and agranular portion, but they do not rely as much on the visual system as simians do.
4.15.4.2 Lorisoidea (Galago)

Galagos are nocturnal primates living in low forests. Arboreal animals, they move by quadrupedal walking and leaping. Galagos have relatively large eyes with a rudimentary fovea supporting relatively low acuity vision (DeBruyn et al., 1980; Stone and Johnston, 1981). Among primates they rely least on vision. Because of the relatively large size of the eyes, galagos have a limited range of eye movements and compensate for this with more head movements to explore novel objects (Rogers et al., 1993; Cantalupo et al., 2002). Although their form of locomotion might suggest that they rely on good eye–hand coordination for their agile leaping, they primarily use olfactory and auditory information. Galagos have large ears with great mobility that are continuously moving. As insectivores, they rely on sound to localize their prey (Charles-Dominique, 1977).

Electrical stimulation of a small region medial to the anterior frontal sulcus evokes contraversive eye movements (Wu et al., 2000; Fig. 1). This very restricted representation of eye movements was compensated by a larger representation of ear movements in the cortex caudal to FEF (Fogassi et al., 1994). While the more rostral region had cytoarchitectonic characteristics of FEF seen in other species with a granular layer 4 and medium-to-large densely packed layer 5 pyramidal neurons, the caudal region appeared more dysgranular (Preuss and Goldman-Rakic, 1991). Like FEF in macaques the rostral region is connected with the multimodal division of the mediodorsal nucleus and with the intermediate and superficial layers of the superior colliculus (Markowitsch et al., 1980; Preuss and Goldman-Rakic, 1991).

Curiously, a dorsomedial eye field corresponding to the supplementary eye field has not been found (Wu et al., 2000). Research with macaque monkeys and humans has led to the general conclusion that the supplementary eye field contributes to high level, executive control of gaze behavior (e.g., Schall and Boucher, 2007). If correct, the absence of a supplementary eye field in galagos could be understood in light of the stimulus-bound simplicity of galago behavior relative to that of macaques and humans. Regardless, this reminds us that the number and organization of cortical areas mediating gaze control need not be equivalent across species.

4.15.4.3 Haplorrhini: Simians

Simians (anthropoids) are all diurnal with the exception of owl monkeys. They have less reliance on olfaction and are primarily visual animals with a well-developed fovea supporting high acuity vision and a suitably matched ocular motor system supporting the control of eye movements in coordination with head, limb, and body movements.

4.15.4.3.1 Platyrrhini (New World Monkey)

Platyrrhini are small to medium size monkeys with flat noses, which distinguishes them from Old World monkeys. Other than howler monkeys, they lack the typical trichromatic vision of the Old World monkeys.

Figure 1  Map of eye and body movements elicited by intracortical microstimulation in Galago garnetti. The frontal eye field (FEF) is highlighted red, located rostral to premotor cortex. Smooth pursuit eye movements were evoked at more caudal sites, and saccadic eye movements, more rostral. Curiously, a supplementary eye field was not found. Reproduction from Wu, C.W., Bichot, N.P., Kaas, J.H., 2000. Converging evidence from microstimulation, architecture, and connections for multiple motor areas in the frontal and cingulate cortex of prosimian primates. J. Comp. Neurol. 423, 140–177.
4.15.3.1.1 Callitricidae (Marmoset)

Marmosets are the smallest New World monkeys. Those used in research, Callithrix jacchus, naturally live in open forest habitats. Marmosets are very agile and active, moving with quadrupedal walking and leaping. Claw-like nails allow them to cling to trees while foraging. The retina of the marmoset has a well-developed fovea comparable to that found in other diurnal primates (Finlay et al., 2008; Franco et al., 2000). However, marmosets have the smallest eyes among the primates (Wolin and Massopust, 1967) affording less spatial acuity (Finlay et al., 2008). With lightweight heads, marmosets can shift gaze with rapid head movements and so have correspondingly smaller oculomotor range (Mitchell et al., 2014). Marmosets also display head-cocking, rapid head rotations along the longitudinal axis, for visual exploration (Kaplan and Rogers, 2006).

With a lissencephalic (smooth) cortex and an opportunity for genetic studies, marmosets have gained new popularity in research. A review by Bakola et al. (2015) summarizes knowledge about the frontal motor system. Although the frontal cortex of marmosets is overall a smooth surface, some individuals exhibit an arcuate dimple. Eye movements with and without head movements are elicited from the rostral frontal lobe with electrical stimulation (Mott et al., 1910; Blum et al., 1982; Fig. 2). Such movements were also elicited by stimulation of the rostral frontal lobe in prosimian lemurs (Mott and Halliburton, 1908). Subsequent research with marmosets has verified visual and ocular motor connectivity of these regions (Reser et al., 2013; Lyon and Kaas, 2001; Spatz and Tigges, 1972; Rosa et al., 2009; Krubitzer and Kaas, 1990; Collins et al., 2005). Areas 8aV and 45, which might correspond to the frontal visual (FV) area described by Krubitzer and Kaas (1990), show similar patterns of connections with

parietal and extrastriate visual areas. This observation suggests that FEF in the marmoset spans area 8 and perhaps partially area 45 (Reser et al., 2013). The cytoarchitecture of the FEF region is characterized by larger neurons in layer 5, higher myelin density, and a granular layer 4 (Burman et al., 2006; Krubitzer and Kaas, 1990), similar to other species examined. The locally elevated density of myelin was also found with structural MRI techniques (Bock et al., 2009). Information about the neurophysiology of FEF in particular or prefrontal cortex in general in marmosets is lacking. However, an fMRI study reported elevated BOLD signal within the location of area 8aV when visual stimuli were presented (Hung et al., 2015).

Eye movements with and without head movements were also elicited by stimulation caudal to the FEF (Mott et al., 1910); sites eliciting pinna movements were not found, contrasting with prosimian lemurs (Mott and Halliburton, 1908). In marmosets, an additional caudal area 8 (area 8C) was described at the junction between dorsal and ventral premotor cortex (Paxinos et al., 2012; Burman et al., 2014, 2015; Bakola et al., 2015), which is distinguished by denser myelination. This area has a connectivity pattern like area 6Va, but lacks some of the somatosensory connections and instead receives input from area 8aV and is interconnected with visual areas (Bakola et al., 2015; Burman et al., 2015; Palmer and Rosa, 2006; Rosa et al., 2009). Burman et al. (2015) consider area 8C as part of the premotor network rather than as a prefrontal area because it lacks a distinct layer 4. However, they also suggest that this region might be specialized for visually guided movements and suggest that it might correspond to a region where Preuss et al. (1996) evoked eye and neck movements in the owl monkey. Marmoset area 8C might correspond to the PMtv spur described by Cabernet et al. (1999) for the macaque monkey as is described in the following section. Burman et al. (2015) point out that area 8C might be similar to area 6Va involved in head movements. Because marmosets compensate for their limited oculomotor range with rapid head movements, an area linked with FEF that is involved in directed head movements is not surprising.

4.15.4.3.1.2 Aotidae (Owl Monkey)

Owl monkeys are nocturnal. They live in higher elevations in forests and tropical environments of Central America and northern South America (Groves, 2001). They feed upon small fruit, insects, leaves, and flowers. They are able to snatch flying insects out of the air or off of a branch, unlike foragers or tool users who exploit holes and spaces in the tree bark (Wright, 1989). Even though the nocturnal environment would be a disadvantage for animals dependent upon vision, owl monkeys have been able to thrive in the niche. They possess large eyes without a fovea and a nearly pure rod retina (Woollard, 1927; Jacobs et al., 1993). These structural features provide for better low-light spatial resolution that improves their night vision, but leaves the owl monkey with monochromatic vision.

Owl monkeys are lissencephalic anterior to the central sulcus, only rarely with an arcuate dimple. FEF was identified with microstimulation at low currents around the arcuate dimple, and because of the smaller prefrontal FEF appears located relatively more rostral (Fig. 2; Gould et al., 1986; Huerta et al., 1986, 1987). Caudal to FEF is an expanded region where head movements are represented (Gould et al., 1986). Gould et al. (1986) describe another eye field that is located more dorsal and also has a caudally adjacent region representing head movements and likely corresponds to SEF of macaque monkeys (Preuss et al., 1996). Huerta et al. (1986, 1987) also characterize an FV field located ventral to FEF. Saccadic eye movements could be evoked with higher currents in area 8B—referred to it as dorsal oculomotor area—as well as in the dorsal premotor cortex (Stepniewska et al., 1993; Preuss et al., 1996). Stimulation in the cortex surrounding FEF at higher currents also leads to ear, eyelid, or vibrissa movements (Huerta et al., 1986).

FEF in owl monkeys shows a connectivity pattern comparable to squirrel monkeys and macaque monkeys with connections to the upper temporal lobe, the dorsomedial visual area, and posterior parietal cortex as well as the superior colliculus and the lateral band of the mediodorsal nucleus (Kaas and Morel, 1993; Weller et al., 1984; Krubitzer and Kaas, 1993; Kaas et al., 1977; Collins et al., 2005; Huerta et al., 1986, 1987). However, the FEF region of owl monkeys apparently lacks strong connections with the middle temporal visual area that are found in diurnal monkeys (Huerta et al., 1987; Weller et al., 1984; but see Cerkevich et al., 2014). A connectivity pattern similar to FEF is shared by the more ventral IV area.

4.15.4.3.1.3 Saimiriinae (Squirrel Monkey)

Squirrel monkeys live in Central and South America in the midlevel tangled layers of branches, vines, and trees of tropical forests. Squirrel monkeys use all four limbs to move but will occasionally walk short distances on two limbs. Squirrel monkeys predominantly eat fruits and insects. They rarely reach for insects in motion, but prefer finding insects on leaves. They engage in long periods of visual observing behavior (Glickman and Sroges, 1966); however, they tend to show notably less visual investigative behavior relative to other primates (DuMond, 1968; Glickman and Sroges, 1966; but see Haude and Ray, 1974). Male squirrel monkeys are dichromatic, while females can be either dichromatic or trichromatic. The eye movements of squirrel monkeys are in general comparable to macaque monkeys, but they have a much narrower oculomotor range of about 10–15 degrees horizontally (Heiney and Blazquez, 2011; McCrea and Gdowski, 2003). Interestingly, the oculomotor range is larger for the vertical axis (about 25 degrees); this asymmetry contrasts with terrestrial species in which horizontal eye movements are more frequent and of larger average amplitude. Like other monkeys, squirrel monkeys communicate with visual signals and social behaviors, including facial expressions, as well as signaling with their ears.

In squirrel monkeys, gyral variations of the frontal lobe range from no sulcus to a small inferior arcuate dimple to a small arcuate sulcus (Emmers and Akert, 1963; Blum et al., 1982; Huerta et al., 1987). FEF was identified with electric stimulation near the dimple, with considerable variation between individuals (Huerta et al., 1986; Fig. 2).
FEF of squirrel monkeys is connected with visual areas in the superior temporal sulcus including area MT in addition to subcortical structures paralleling macaques (Huerta et al., 1987; Tigges et al., 1981).

4.15.4.3.1.4  Cebidae (Capuchin)

Capuchin monkeys are considered to be the most intelligent of New World monkeys, possessing a relatively large brain relative to their body size. They have coordinated movements between hands, tail, feet, and mouth. They perceive objects, movements, and surfaces in much the same way humans do. They have a large repertoire of visual signals and gestural behaviors to communicate, including submissive grins, raising eyebrows, head tilt, play face, threat face, and lip smacking. Capuchins are found commonly in northern parts of Argentina but are also distributed around North and South America. They live in different regions of trees—some very high and others in the lower regions. They spend most of their waking hours foraging for a variety of food types. They eat a greater variety of food types than any other type of monkeys. They exhibit rudimentary tool use, for example, using stones to access food.

The capuchin (Cebus apella) shows a clearly defined gyral pattern (Connolly, 1936, 1950), resembling that of macaque monkeys with an arcuate sulcus that is usually accompanied by a principal sulcus (Sanides, 1970; Tian and Lynch, 1997).

FEF as identified with microstimulation is found along the anterior bank of the arcuate sulcus (Lynch et al., 1994) at a location comparable to that of macaque monkeys. It shows a comparable connectivity pattern as described for other monkeys, with connections to MT and to the posterior parietal cortex as well as subcortically to the superior colliculus and the mediodorsal nucleus of the thalamus (Tian and Lynch, 1997; Leichnetz and Gonzalez-Ruiz, 1996; Rosa et al., 1993; Lynch et al., 1994).

In addition to FEF in the anterior bank of the arcuate sulcus, two more eye fields have been characterized in Cebus. One is located dorsomedially and likely corresponds to the supplementary eye field of macaque monkeys (Tian and Lynch, 1995) and another one is located in the posterior bank of the superior limb of the arcuate sulcus (Tian and Lynch, 1996). In contrast to FEF where microstimulation evokes saccadic eye movements, stimulation at the later more dorsomedial location results in smooth eye movements. This location might correspond to the smooth pursuit zone described for macaque monkeys (Gottlieb et al., 1993, 1994); however, in macaque monkeys it is in the posterior bank of the arcuate sulcus adjacent to FEF and not located dorsomedially as in Cebus (Fig. 2). This illustrates that a common function can be mediated by cortical areas in different relative locations across species.

4.15.4.4  Cercopithecidae (Old World Monkey)

4.15.4.4.1  Macaca (Macaque)

Macaque monkeys are the most common nonhuman primate used in research to gain insights into human brain function (Passingham, 2009). Hence, the most detailed knowledge about the organization and location of the primate FEF has been obtained from various macaque species (M. mulatta, M. fascicularis, M. radiata, M. nemestrina, Macaca sinica, and Macaca fuscata). Given the relatively minor known differences in the brain organization across macaque species, many studies have used more than a single species or some do not specify clearly what macaque species was used.

Macaque are generally acknowledged as terrestrial, but some (e.g., M. fascicularis) are also arboreal. Macaques are mostly quadrupedal animals; however, they also are very dexterous in using their hands for visually guided reaching and grasping behavior. Macaques use saccadic and smooth pursuit eye movements to explore the visual field, including vergence movements to explore both far and near locations. Macaques are social animals that usually live in hierarchical groups. Gaze direction is a behaviorally relevant cue, and macaques show gaze-following behavior as directed by the head direction. Gaze direction is also an important signal in the social structure; direct staring at an animal higher in the hierarchy is an offensive behavior that could trigger retribution. Accordingly, macaques covertly observe other animals.

Following the groundbreaking experiments of Fritsch and Hitzig (1870) demonstrating that electric stimulation of certain locations in the cerebral cortex evoked specific body movements in a dog, Ferrier, (1874) carried out a thorough mapping to localize motor functions in the cortex of macaque monkeys. Based on his maps of the frontal cortex, Ferrier located a region in which stimulation evoked contraversive movements of the eyes and head as well as other ocular movements such as vergence and blinks (Fig. 3). Numerous subsequent investigators mapped locations eliciting orienting movements in macaque monkeys resolving over time the effects of anesthesia, eye movement recording sensitivity, electrode size, and electrical current levels (Horsley and Schäfer, 1888; Mott and Schäfer, 1890; Russell, 1894; Levinsohn, 1909; Smith, 1949; Crosby et al., 1952; Henderson and Crosby, 1952; Wagman et al., 1957, 1958, 1961; Robinson and Fuchs, 1969; Bruce et al., 1985). The evolution of FEF location across these studies has been reviewed elsewhere (Schall, 1997). The current definition of FEF is the region where currents less than 50 µA evoke short latency saccades, which restricts FEF in the rostral bank and fundus around the genu of the arcuate sulcus. Within this region saccade amplitude, but not direction, is topographically organized; shorter (<2 degrees) saccades are represented ventrolaterally, and progressively longer saccades (15–20 degrees) are represented dorsomedially.

The macaque FEF is well known to be highly visually responsive with pronounced modulation according to the allocation of attention and other cognitive demands (reviewed by Squire et al., 2013; Schall, 2015). Visually responsive neurons are found in cortical areas extending rostral to FEF. Indeed, the map of saccade amplitude is paralleled by a map of visual receptive field eccentricity described in the prearcuate gyrus (Suzuki and Azuma, 1983).
A region related to slow tracking pursuit eye movements is located at the fundus of the arcuate sulcus immediately caudal to the principal sulcus (MacAvoy et al., 1991; Gottlieb et al., 1994; Fukushima et al., 2002; Tanaka and Lisberger, 2002; Fig. 3). Low-intensity electrical stimulation of FEF in head-free monkeys also elicits contraversive head movements and natural eye-with-head gaze shifts (Tu and Keating, 2000; Knight and Fuchs, 2007; Monteon et al., 2010, 2013). Medial to FEF in the dorsomedial cortex is an anatomically separate and functionally distinct supplementary eye field (Schlag and Schlag-Rey, 1987; Schall, 1991). Along the superior limb of the arcuate is a region that represents orienting movements of the pinna (Bon and Lucchetti, 1994; Lucchetti et al., 2008; Lanzilotto et al., 2013) with a gradual transition from eye movements and ear movements that also correspond to the eccentricity of localized events (Barbas and Mesulam, 1981; Lanzilotto et al., 2013). Finally, neurons in the rostral convexity of the arcuate sulcus are active during vergence and accommodation for perception of objects at different distances (Gamlin and Yoon, 2000).

The location of FEF has a distinctive cyto- and myeloarchitecture (Walker, 1940; von Bonin and Bailey, 1947; Stanton et al., 1989; Preuss and Goldman-Rakic, 1991). Not unexpectedly, it is distinguished by a concentration of large pyramidal cells in layer 5 higher than surrounding areas. It has a thin granular layer 4 distinct from the agranular area 6 at its caudal boundary but noticeably thinner than the rostrally bordering area referred to now as 8Ar. Area 8Ar should probably be considered as functionally distinct from FEF; it may correspond to the area FV that has been distinguished from the heavily myelinated part of FEF in owl monkeys based on patterns of connectivity with prefrontal visual areas (Weller and Kaas, 1987; Krubitzer and Kaas, 1990). The medial aspect of FEF has relatively fewer large pyramidal cells in layer 3 and a loosely organized granular layer and has more connectivity with cortical areas representing the peripheral visual field. The dorsomedial border of FEF transitions into area 8B. The ventral aspect of FEF has a higher density of larger pyramidal cells in layer 3 and has more connectivity with cortical areas representing the central visual field. The ventrolateral border transitions into area 45. The medial and lateral parts of FEF in the arcuate sulcus are also distinguished by connectivity; they are connected with many common but several distinct cortical areas (Schall et al., 1995; Markov et al., 2014). The lateral segment, which is responsible for generating short saccades, receives visual inputs from the foveal representation in retinotopically organized areas, from areas that represent central vision in inferotemporal cortex and from other areas having no retinotopic order. Meanwhile, the medial segment, which is responsible for generating longer saccades coupled with head and also pinna movements, is innervated by the peripheral representation of retinotopically organized areas, from areas that emphasize peripheral vision or are multimodal and from other areas that have no retinotopic order or are auditory. Convergence from the dorsal and ventral visual processing streams occurs in lateral but not in medial FEF. These differences of connectivity must have corresponding differences of functionality, but they are unknown at this time.

The location of FEF in macaques has also been investigated with a variety of neuroimaging techniques. One approach used 2-deoxyglucose imaging to locate activated areas in both the anterior and posterior banks of the arcuate sulcus, the prearcuate convexity, dorsomedial frontal, caudal principal and periprincipal, anterior cingulate, and even some of the orbitofrontal cortex (Moschovakis et al., 2004; Savaki et al., 2015). The relation of these diverse cortical areas to eye movement production was reinforced by retrograde transneuronal labeling by rabies virus that was injected into the lateral rectus muscle (Moschovakis et al., 2004; Fig. 3D).

FEF has also been localized using fMRI with macaques. Early studies identified an involvement of FEF in visual motion processing in awake behaving monkeys (Vanduffel et al., 2001). Subsequent primate fMRI studies characterized involvement of FEF in visually guided saccade tasks (Koyama et al., 2004; Baker et al., 2006; Premereur et al., 2015), memory guided saccade tasks (Kagan et al., 2010), antisaccade tasks (Ford et al., 2009), visual search tasks (Wardak et al., 2010), and spatial attention tasks (Caspari et al., 2015). The connectivity found in resting state fMRI of the macaque FEF corresponds very well to the connectivity pattern found with anatomical tract tracers (Babapoor-Farrokhran et al., 2013).

Finally, the location of FEF in the arcuate sulcus has been confirmed through inactivation (e.g., Sommer and Tehovnik, 1997) and lesion studies demonstrating temporary impairments in eye movements (Schiller et al., 1980, 1987; Keating, 1993; Morrow and Sharpe, 1995; Heide et al., 1996) and in eye–head movements (van der Steen et al., 1996). Inactivation of FEF also impairs visual search (Latto, 1978; Wardak et al., 2006).

While there is no disputing the location of the FEF in the rostral bank of the arcuate sulcus of macaques, multiple studies have described visual- and saccade-related function in areas surrounding the arcuate sulcus. For example, visually responsive, eye movement, and eye position–related neural activity is found in premotor cortex caudal to the arcuate sulcus around the arcuate spur (Boussaoud, 1995; Boussaoud et al., 1993, 1998; Fuji et al., 1998, 2000). Neural activity recorded in this general region is also modulated during tasks requiring visually guided selection of targets for action (e.g., Cisek and Kalaska, 2005; Song and McPeek, 2010). Also, measurements of 2-DG uptake during production of saccades of various directions and amplitudes defined two maps, one running between area 44 and ventral area 6, and the other spanning areas 8 and 45 in the anterior bank of the arcuate sulcus and extending onto the prearcuate convexity (Savaki et al., 2015). Finally, fMRI measurements in macaque monkeys reveal saccade-related activation in areas 45A, 45B, and 46 outside the conventional FEF (e.g., Premereur et al., 2015).

Thus, numerous studies in macaques agree on the core location of FEF in the rostral bank of the arcuate sulcus, surrounded by areas associated with pursuit and vergence movements. Many other studies demonstrate that neural activity associated with visual responses and eye movement production is found in areas surrounding FEF both caudally in premotor cortex and rostrally in prefrontal cortex. Thus, defining the boundaries of “FEF” in the macaque is a matter of specifying criteria.
### 4.15.4.4.2 Papio (Baboons)

The frontal lobe of baboons resembles that of macaques although with somewhat more complexity in the relationship of the arcuate and principal sulci and the presence of more dimples or partial sulci (Connolly, 1936, 1950).

Less is known about the properties of FEF in baboons. Confidence in its location in the rostral bank of the arcuate sulcus, though, has guided studies of the effects of FEF lesions (e.g., Pribram, 1955). Some evidence from cytoarchitecture and connectivity reinforces this confidence (Watanabe-Savaguchi et al., 1991).

### 4.15.4.5 Hominoidea

The evolution of visually guided reaching and grasping is closely associated with primate ecology and the need to move through the forest canopy. Apes and monkeys that use brachiation rely upon vision to select, reach for, and secure their grip onto appropriate branches. These actions are often rapid and coordinated with other self-generated movements that strongly affect the visual input, such as the motion of the body, turning of the head, and frequent saccades. Since a miscalculation can have fatal consequences, primates have evolved under a strong selective pressure for accuracy in this domain. The origins of precise manual behavior may stem from specific aspects of the feeding behaviors of early primates. One influential hypothesis holds that early primates were nocturnal predators who were aided by a wide field of stereoscopic vision for catching insects in their hands (Cartmill, 1992).

An alternative view is that precision reaching and grasping evolved for the purposes of extracting small fruits from terminal branches of angiosperms (Bloch and Boyer, 2002; Sussman et al., 2013).

#### 4.15.4.5.1 Pongo (Orangutan), Gorilla (Gorilla), and Pan (Chimpanzee and Bonobo)

Relative to the Cercopithecidae brain, major changes have occurred in the cerebral cortex of hominids, especially in the frontal lobe (Connolly, 1936, 1950). Apes have no sulcus resembling the arcuate sulcus of Cercopithecidae (e.g., Bogart et al., 2012). The chimpanzee frontal lobe consists of a prominent superior precentral sulcus and an inferior frontal sulcus, both with multiple limbs. The inferior precentral sulcus branches off of the inferior frontal sulcus. Ventrolaterally is found a fronto-orbital sulcus. Differential studies carried out on a large number of chimpanzee brains show considerable individual variation, particularly in the inferior precentral region (Sherwood et al., 2003).

Classic studies used electrical stimulation of the cortical surface under anesthesia to map sites eliciting movements of the body and eyes in chimpanzees (Grünbaum and Sherrington, 1901; Leyton and Sherrington, 1917; Hines, 1940; Dusser de Barenne et al., 1941; Bailey et al., 1950), orangutan (Beevor and Horsley, 1890; Leyton and Sherrington, 1917), and gorilla (Grünbaum and Sherrington, 1901; Leyton and Sherrington, 1917). Leyton and Sherrington (1917) mapped body and eye movements elicited by electrical stimulation of the cortical surface in 22 chimpanzees, 3 orangutans, and 3 gorillas. In chimpanzees, they showed that conjugate contraversive ocular rotations were elicited at a range of sites around the middle and inferior sulci with some sites eliciting eye opening as well (Fig. 4). Most of the rotations were horizontal, but some sites elicited rotations with upward or more often downward components. A few sites elicited convergence movements. They noted that higher currents were needed to elicit eye movements as compared to limb movements from the precentral gyrus. They further showed that contraversive ocular rotations were not a primary movement from the precentral gyrus and occurred only associated with head movements.

In a sample of 38 chimpanzees Bailey et al. (1950) located regions from which ocular deviations could be elicited in 11. Their results indicate two general regions from which ocular deviation was evoked with greatest frequency. One was located ventrolaterally straddling the inferior frontal sulcus, and the other was located dorsomedially ventral to and straddling the superior frontal sulcus. In most of the chimpanzees, eye movements were elicited much more rostral than the precentral sulci, and in many cases stimulation of the cortex rostral to the regions from which ocular deviations were elicited resulted in no movements. When explored, stimulation of the cortex caudal to the eye fields often elicited skeletal movements and other eye movements such as eye blinks and pupil size changes. The dorsomedial locations eliciting ocular rotations most commonly were located in area FC, which is described as transitional between agranular and prefrontal cortex having a slight granular layer. The ventrolateral locations appeared most commonly located in the weakly granular area FC and possibly extending into granular areas FDr and FDp. By the way, note that the summary figure offered by the authors seems not to most accurately portray their findings because it identifies an eye movement region more ventral than was observed (Fig. 4C).

In orangutan and gorilla, Leyton and Sherrington (1917) found sites eliciting ocular rotations similarly situated about the middle frontal sulcus (Figs. 5 and 6). Thus, the site most clearly identified with FEF is located comparably in chimpanzee, orangutan, and gorilla, distant from primary motor cortex and centered rostral to the inferior principal sulcus around the inferior frontal sulcus.

#### 4.15.4.5.2 Human

Readers of this chapter need no introduction to the importance of visually guided eye movements coordinated with ongoing manual, locomotor, and social behavior. They will also know that the human cerebral sulcal pattern is noticeably elaborated from that of apes and appreciate how the significant individual gyral variation, sometimes even between hemispheres, can obscure descriptions of the location of cortical areas. We should also remind the reader that different investigators have arrived at somewhat different descriptions of the organization of cortical areas (Fig. 7). In considering which cortical areas in the human are homologous

Figure 5  Location of frontal eye field based on electrical stimulation in the orangutan. From Leyton, A.S., Sherrington, C.S., 1917. Observations on the excitable cortex of the chimpanzee, orangutan, and gorilla. Q. J. Exp. Physiol. 11, 135–222.
to areas in the brains of nonhuman primates, the FEF provides an interesting test case. The novice is surprised how far caudal FEF is until appreciating how much frontal lobe is in the human brain.

FEF has been located in humans using electrical stimulation in various forms for many years. To summarize, most studies have evoked ocular rotations by stimulation of the caudal end of the middle frontal gyrus, but results disagree about the extent of surrounding cortex that also elicits eye movements and whether it extends to the skeletal motor representation in the precentral gyrus. Dejerine and colleagues located a region, which they called “area D,” from which conjugate deviation of the head and eyes were evoked (Dejerine and Roussy, 1906; Dejerine, 1914). Förster (1931, 1936) identified FEF as the region from which contraversive ocular rotation was elicited at the caudal end of the middle frontal gyrus (Fig. 8) in a region designated 8abγ by Vogt and Vogt (1926) (Fig. 7A). Förster also reported coordinated contraversive eye, head, and trunk rotation following stimulation of a dorsomedial region designated 6aβ, corresponding to Dejerine’s area D. Subsequently, Penfield and coworkers evoked eye movements over a wider area of frontal cortex, extending caudally onto the precentral gyrus; however, most sites were rostral to the precentral sulcus at a location Penfield identifies as Vogt’s 8abγ (Penfield and Boldrey, 1937; Rasmussen and Penfield, 1947). Subsequently, Lemmen et al. (1959) reported conjugate and divergent eye movements from stimulation of the posterior end of the middle frontal gyrus rostral to sites in the precentral gyrus that elicited limb and lip movements (Fig. 9); these authors noted a particular absence of eye movements elicited by stimulation of the precentral gyrus.

Later studies used subdural electrode arrays implanted over the frontal lobe of patients and found FEF in a zone rostrally contiguous with the motor cortex representation of head and forelimb (Godoy et al., 1990). More recent work has located FEF in the posterior part of the middle frontal gyrus, just below the intersection of the precentral sulcus and the superior frontal sulcus (Blanke et al., 2000; Blanke and Seeck, 2003; Lobel et al., 2001; Mälea et al., 2002; Yamamoto et al., 2004; Thurtell et al., 2009; Kaiboriboon et al., 2012; Montemurro et al., 2016; Fig. 10). Some studies have also located a more lateral region associated with eye movement production close to the surface of the precentral gyrus (Lobel et al., 2001). The reader should note that these electrical stimulation studies typically can explore only the cortical surface not buried in sulci and that spatial localization is difficult to interpret because of current spread influencing remote sites.

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**Figure 6** Location of frontal eye field based on electrical stimulation in the gorilla. From Leyton, A.S., Sherrington, C.S., 1917. Observations on the excitable cortex of the chimpanzee, orangutan, and gorilla. Q. J. Exp. Physiol. 11, 135–222.

**Figure 7** Frontal lobe areas identified by Vogt and Vogt (1926) (A) and Brodmann (1909) (B). Reproduced from (A) Vogt, C., Vogt, O., 1926. Die vergleichend-architektonische und die vergleichend-reizphysiologische Felderung der Großhirnrinde unter besonderer Berücksichtigung der menschlichen. Naturwissenschaften 14, 1190–1194; (B) Brodmann, K., 1909. Localisation in the Cerebral Cortex. Springer, US.
FEF has also been localized using transcranial magnetic stimulation (e.g., Wessel and Kömpf, 1991; Müri et al., 1991; Olk et al., 2006; Neggers et al., 2007). Transcranial magnetic stimulation does not elicit ocular rotations, so the influence is inferred through indirect measures of influence on saccade latency or accuracy. Such studies point toward the caudal end of the middle frontal gyrus at the intersection of the superior frontal sulcus and the superior precentral sulcus. Some investigators also locate FEF relative to the motor hand area (e.g., Ro et al., 1999), on average 5 cm lateral of the sagittal midline and 3–4 cm rostral of each subject’s motor hand area.

Human FEF has been located through specific deficits in eye movements observed after focal lesions (reviewed by Pierrot-Deseilligny, 1994). Early studies noted gaze deficits with damage involving large parts of frontal cortex; however, more recent studies of patients with more restricted lesions have converged on the precentral sulcus ventral to the superior frontal sulcus (Rivaud et al., 1994; Fig. 11).
Human FEF has been described in numerous PET and fMRI studies (reviewed by Grosbras et al., 2005; see also Luna et al., 1998; Petit et al., 1997, 1999; Neggers et al., 2012; Ford et al., 2005; Amiez et al., 2006; Kastner et al., 2007; Ikkai and Curtis, 2008; Derrfuss et al., 2012; Thakkar et al., 2014). Execution of saccadic eye movements is accompanied by relatively strong and consistent bilateral activation in the superior precentral sulcus and by relatively weaker and less consistent activation in the superior segment of the inferior precentral sulcus. Pursuit eye movements are accompanied by activation in the superior precentral sulcus occupying a smaller region that tends to be inferior and slightly lateral to that occupied during saccades, closer to the fundus. Within individuals, the activation in both precentral sulci is restricted to the banks and does not extend onto the surface convexity, even though maps of average activation give that appearance. High-resolution imaging locates the most active voxels on the rostral bank of the superior precentral sulcus. The dorsal activation is located at the junction of the precentral sulcus and the superior frontal sulcus.

Amiez et al. (2006) conducted a subject-by-subject analysis of the locus of eye movement–related functional activity revealed in relation to the individual morphology of the precentral and superior frontal sulci (Fig. 15). A focus of activation associated with saccadic eye movements was located in the ventral branch of the superior precentral sulcus in both hemispheres. A second focus has found in the dorsal part of the inferior precentral sulcus. Imaging during a hand response selection task revealed activation focused in the dorsal branch of the superior precentral sulcus close to the caudal end of the superior frontal sulcus. The relative magnitude of activation in the superior and inferior precentral sulci varies with the nature of the eye movement (eg, saccade or pursuit, in light or dark) and task demands (eg, prosaccade or antisaccade) (Fig. 12A; eg, Neggers et al., 2012). Medial and lateral

**Figure 9**  Red highlights sites from which cortical stimulation resulted in conjugate deviation of the eyes. Stimulation at site 5 had no effect. Stimulation at more caudal sites elicited movements of the indicated parts. From Lemmen, L.J., Davis, J.S., Radnor, L.L., 1959. Observations on stimulation of the human frontal eye field. J. Comp. Neurol. 112, 163–168.

**Figure 10**  Red highlights location where conjugate eye movements were elicited with the lowest currents using subdural electrode arrays. From Blanke, O., Spinelli, L., Thut, G., Michel, C.M., Perrig, S., Landis, T., Seeck, M., 2000. Location of the human frontal eye field as defined by electrical cortical stimulation: anatomical, functional and electrophysiological characteristics. Neuroreport 11, 1907–1913.
foci are also observed during covert attention tasks as well (Beauchamp et al., 2001). Furthermore, two regions recruited in auditory attention tasks were reported to be interdigitated with these two visual attention regions (Fig. 12B; Michalka et al., 2015).

The interpretation of the location of BOLD activation during eye movements can be complicated by other factors. First, eye blinks are commonly associated with saccades (e.g., Evinger et al., 1994; Gandhi, 2012). Since the original electrical stimulation mapping it has been clear that cortical regions caudal to the arcuate sulcus in monkeys produce other ocular movements including blinks. Thus, some of the BOLD activation during eye movement tasks could arise from regions producing eye blinks. However, an fMRI study of spontaneous eye blinks in macaque monkeys did not report activation in or around FEF, calling into question whether this is an actual confounding factor for FEF localization (Guipponi et al., 2015). Second, saccades may be associated with neck contractions to rotate the head (e.g., Goonetilleke et al., 2015). Thus, some of the BOLD activation during eye movement tasks could arise from premotor cortical areas that contribute to head movements (e.g., Preuss et al., 1996). Third, in the macaque areas rostral and caudal to the arcuate sulcus have visually responsive neurons, which are modulated by attention tasks. If the human FEF is similarly surrounded, then BOLD activation would occupy a larger region than FEF proper. Also, as noted earlier, in the macaque areas caudal to the arcuate sulcus include neurons active during saccade tasks. We suggest alternative hypotheses concerning the relation of the monkey and human FEF and premotor region in the following section.

4.15.5 Is FEF Located Differently Across Species?

In this section we consider from multiple perspectives whether the FEF can be regarded as homologous across primate species, including humans.

4.15.5.1 Comparative Neuroimaging

Several studies compare directly macaque and human fMRI data (Koyama et al., 2004; Baker et al., 2006; Ford et al., 2009; Kagan et al., 2010). For example, Koyama et al. (2004) compared fMRI activation patterns in humans and macaques performing visually guided saccades. BOLD activation in macaques was found in the rostral bank of the arcuate sulcus, and also in premotor cortex caudal to the arcuate sulcus. In humans a large domain of activation occupied the banks of the precentral sulcus. The authors propose that the peak of activation near the junction of the precentral and the superior frontal sulci corresponds to the FEF in monkeys. They observed another focus of activation in the inferior precentral sulcus (Fig. 14). Resting-state fMRI connectivity of FEF has been compared in macaques and humans (Hutchison et al., 2012; Babapoor-Farrokhran et al., 2013). These studies have found consistent functional connections of the FEF with frontal and parietal cortical areas in humans and macaques (Fig. 14). However, the analysis also revealed more lateralization of connectivity of the region identified as FEF in humans than in monkeys.

The homology or at least comparability of the cortical region in and rostral to the arcuate sulcus in macaques and the superior precentral sulcus in humans has also been revealed in quantitative functional connectivity maps (Sallet et al., 2013; see also Goulas et al., 2012). Of note, in macaques a particular hot spot of connectivity with this arcuate region was caudal to the arcuate, dorsal to a spur in the region overlapping that associated with visually guided saccades described previously.
4.15.5.2 Comparative Architecture

As noted earlier, the FEF in macaque monkeys is centered in the rostral bank of the arcuate sulcus at the caudal end of prefrontal cortex. The cytoarchitecture of the area is included as Brodmann’s area 8, generally. Likewise, the FEF mapped in other monkeys is recognized as being located in granular prefrontal cortex. The FEF in apes also appears to be in prefrontal cortex, although we have less specific information about the cytoarchitecture of the regions mapped. However, many authors describe the human FEF as being located in Brodmann’s agranular area 6. Here we address this apparent discrepancy.
Figure 13  Lateral view of the human cerebral hemispheres, showing the localization of functions in the cerebral cortex. The numbers list Brodmann’s (1909) cytoarchitectonic areas. Note that the domain for “ocular adverse movements” is identified as area 8. From von Kleist, K., 1934. Gehirnpathologie. J.A. Barth, Leipzig.

The literature on the location of FEF in humans has referred exclusively to Brodmann’s cytoarchitectonic map (1909). Of course this is not the only or the last description, nor does it correspond to modern descriptions in many respects (Zilles and Amunts, 2010; Nieuwenhuys, 2013). In Brodmann’s map area 6 occupies a very large amount of the frontal lobe, but contemporaneous as well subsequent maps by other investigators subdivide Brodmann’s area 6 into many more areas. For example, myeloarchitectonic studies have distinguished the caudal end of the middle frontal gyrus as distinct from surrounding areas (e.g. Nieuwenhuys et al., 2015). While other authors locate the caudal end of the middle frontal gyrus in area 6 (Sarkissov et al., 1955), it has also been labeled area FB (von Economo and Koskinas, 1925), area 4s (von Bonin, 1949), the boundary of FA and FB (Bailey and von Bonin, 1951), and 8zy by (Vogt and Vogt, 1926). Penfield, with Förster, described the majority of stimulation sites eliciting eye movements as being in 8zy (Penfield and Rasmussen, 1950), and the exceedingly elaborate map of von Kleist (1934) identifies ocular adverisive movements with Brodmann’s area 8 (Fig. 13).

Of note, the structure of the cortex occupied by FEF has been reevaluated more recently. Human FEF can be distinguished from surrounding areas by MRI myelin mapping (Glasser et al., 2016). In addition, a recent anatomical study reexamined the architecture of this region using modern chemoarchitectonic methods in postmortem tissue from six subjects (Rosano et al., 2003; see also Schmitt et al., 2005). The histological structure of the superior precentral sulcus was distinct from adjacent rostral and caudal regions. A thin granular layer 4 was observed in sections labeled with neuronal nuclear protein (NeuN), and the nonphosphorylated neurofilament triplet protein (NNFP). Also, clusters of large, intensely immunoreactive pyramidal cells were located in deep layers 3 and 5. In sections labeled for calcium-binding proteins, the two walls of the sulcus were characterized by higher density of calretinin-labeled interneurons, lower density of calbindin-labeled pyramidal neurons, higher density of calbindin-labeled interneurons in layers 2–3, and higher density of large parvalbumin-labeled interneurons in deep layer 3. These histological features resemble the macaque FEF more than agranular area 6. These immunohistochemistry methods highlighted distinctions across this cortical region that are obscured in Nissl-stained section. Based on this analysis of cytoarchitectural, myeloarchitectural, and histochemical studies, one can conclude that Brodmann’s description is unlikely to be correct because it misplaced the caudal boundary of area 8 of humans rostrally.

This characterization seems to resolve the discrepancy. However, questions remain. As described earlier, BOLD activation is also observed in the inferior precentral sulcus, near areas 44 and 45. In nonhuman primates tested, the FEF in the arcuate sulcus is bordered ventrally by areas 44 and 45. Also, the macaque has a region caudal to the FEF in agranular premotor cortex with visual responsiveness and from which saccadic eye movements can be elicited by intracortical microstimulation. Does this area have a homologue in humans? To motivate future experimental work, we suggest for consideration the hypothesis that the “FEF” in the inferior precentral sulcus is homologous with the “FEF” in the arcuate sulcus, while the “FEF” in the superior precentral sulcus is homologous with the premotor eye movement region caudal to the arcuate sulcus of macaques. We note that this assignment is opposite that featured in a recent comprehensive mapping of human cerebral cortical areas (Glasser et al., 2016; see also Amiez and Petrides, 2009). Is it conceivable that premotor cortex in humans takes over functions that are performed by prefrontal cortex in monkeys? The emergence of language, for example, may entail such a phylogenetic displacement. We find it instructive to consider the finding that cortical control of the larynx in monkeys is located in area 6, while in humans it is located in area 4, presumably to provide for articulated speech (Simonyan, 2014).
Identification of the FEF region in the superior precentral sulcus as a premotor area can provide rationale for the findings of a recent study that compared diffusion tractography of corticostriatal pathways in humans and macaques (Neggers et al., 2015). In macaques a seed region in the rostral bank of the arcuate sulcus had connectivity primarily with the head of the caudate and also the anteromedial putamen, while a seed in primary motor cortex in the rostral bank of the central sulcus was connected with more posterior sections of caudate and mainly putamen. These observations replicate neuroanatomical tract tracing findings in macaques. In humans, though, the medial region of activation associated with saccadic eye movements was connected primarily to putamen and only a small portion of the caudate. Moreover, the region of striatum connected with the dorsal FEF in humans overlapped substantially the region of striatal connectivity with a point in primary motor cortex in the rostral bank of the central sulcus. We look forward to a future study examining the connectivity of the FEF region in the inferior precentral sulcus.

**Figure 16** A frontal cortex cladogram to illustrate the location of FEF in a range of prosimian and primate species. The diagrams are not to scale. Sulcal labels are as follows: c, central; fi, frontal inferior; fm, frontal medius; fo, frontal orbital; fs, frontal superior; h, horizontal ramus of precentral inferior sulcus; io, inferior orbital; o, orbital; ob, olfactory bulb; pci, precentral inferior; pcm, precentral medius; pcs, precentral superior; r, rectus; R, ramus ascendens fissurae Sylvii; sca, subcentral anterior; W, frontomarginal of Werneck. Note the use of the label “rectus” for the macaque and *Cebus* principal sulcus and of the labels “pci” and “h” for the macaque and *Cebus* arcuate sulcus. Two regions are highlighted in *Cebus*, macaque, chimpanzee, and human based on evidence for at least two discrete zones. Reproduced from Connolly, C. J., 1936. The fissural pattern of the primate brain. Am. J. Phys. Anthropol. 21, 301–422; Connolly, C. J., 1950. External Morphology of the Primate Brain. CC Thomas.
4.15.5.3 Comparative Sulcal Patterns

Another way to compare the location of FEF across species involves analyzing the topographic pattern of sulci (Fig. 16). The FEF in lissencephalic species is located in the rostral frontal lobe, sometimes around a cortical dimple. In monkeys with an arculate and principal (previously known as rectus) sulcus, the saccade-related FEF is consistently located in the rostral bank of the arcuate sulcus. Curiously, the smooth pursuit–related region has been found dorsal to the arculate in Cebus and at the fundus of the arcuate in macaques. Another region associated with visually guided saccades has been located immediately caudal to the arcuate sulcus around the arculate spur (when it exists).

The sulcus pattern in the apes and human is notably more complex than that in monkeys, but particular associations have been described (eg. Hervé, 1888; Owen, 1900; Connolly, 1936, 1950; Rizzolatti and Arbib, 2002; Falk, 2014). First, there is general agreement that the arculate sulcus, or at least the ventral (vertical) branch, of monkeys corresponds to the inferior precentral sulcus in apes and humans. Second, the superior frontal sulcus and superior precentral sulcus of apes and humans may be derived from the precentral dimple of monkeys. Alternatively, they may correspond to the superior (horizontal) limb of the macaque arculate sulcus. Third, early authors identified the macaque principal sulcus (also referred to as rectus sulcus) with the hominid inferior frontal sulcus, but later analyses conclude that the inferior frontal sulcus of apes and humans has no counterpart in monkeys and instead the principal sulcus of monkeys corresponds to the middle frontal sulcus of humans. Thus, the ventral frontal lobe of humans has expanded relative to apes and monkeys. Such an expansion could lead to a relatively more dorsal location of the FEF in humans relative to apes and monkeys.

These topological identifications of sulci can guide a perspective on the location of FEF in nonhuman primates and humans. Hence, if the FEF of monkeys is in the arculate sulcus, and the arculate sulcus is homologous with the inferior precentral sulcus, then does it follow that the FEF of humans is located in the inferior precentral sulcus? Such reasoning can account for the lateral BOLD activation observed in imaging studies. If this reasoning is sound and valid, then what is the status of the FEF region at the intersection of the superior frontal sulcus and the dorsal precentral sulcus? One possibility is that this dorsal region is homologous to the premotor eye field of macaques. On the other hand, if the dorsal region is correctly understood as the homologue of the macaque FEF, based on evidence reviewed previously, then what is the proper relation of the ventral FEF region? Perhaps it is a premotor eye field as suggested earlier. Confusing all of this, though, is the evidence that the eye movement region in orangutans, gorillas, and chimpanzees is located clearly rostral to the precentral sulci. We note, though, that in spite of the similar appearance of the frontal sulci of humans and chimpanzees the precentral sulcus of chimpanzees does not mark the border between premotor and prefrontal cortex as it may in humans; instead it marks the border between motor and premotor cortex (Preuss, personal communication). This can explain why the FEF is located on the middle frontal gyrus of the apes, but it cannot help us understand the apparent caudal migration of the FEF in humans. Clearly, many questions remain.

4.15.6 Conclusion

We end this chapter with more questions and cautions than conclusions. First, should the term "frontal eye field" be singular or plural? Shall we regard "FEF" (now in scare quotes to remind the reader of the uncertainty) as a collection of distinct areas that can be of different sizes (and locations) in different species? Or shall we regard "FEF" as a single area with multiple modules adapted according to the lifestyle and habitat of the respective species? Under either hypothesis the apparent location of "FEF" might appear to differ across species as one function is emphasized over another. In the macaque, for example, we have described both arculate and post-arcuate regions from which saccades are elicited with weak electrical stimulation. Interposed between these is a pursuit region. Rostral to the saccade map in the arcuate sulcus is a vergence region. The medial and lateral segments within the arcuate sulcus support very different patterns of cortical connections. More dorsomedially in the macaque is a region concerned with eye movements and orienting to sounds. Evolutionary specialization of any (or all of these) could result in elaboration of at least five or six subregions or areas.

Next, across primate species does the size (and complexity) of "FEF" scale with total neocortical area? With the area of visual cortical areas in the parietal and temporal lobes? Does it scale with changes in the subcortical network mediating orienting such as the mediodorsal thalamus, basal ganglia, and—in particular—the superior colliculus. Does it scale with the degree of elaboration of extrastriate visual areas and other sensory systems? Does it scale with the complexity of the musculature used for orienting and associated visually guided behaviors (eye, head, eye + head, eye + head + hand)? Does it scale with the complexity of cognitive control over gaze behavior (eg, dissociation of gaze from visual salience for social deception tested by an ability to make antisaccades)? We cannot answer these questions because precise boundaries of FEF are uncertain, especially in humans. For this reason, it remains essential that the human imaging studies identify the anatomical landmarks and full extent of any activation interpreted as corresponding to FEF.

The size and location of "FEF" is a matter of operational definition with resolution specified by technique. Consider, for example, mapping based on the eye movements elicited by electrical stimulation. Results differ according to state of consciousness (anesthetized or alert), electrode size (surface or intracortical), stimulation parameters (brief or long stimulation trains), method of movement inspection (simple observation or quantitative eye tracking), and inclusion of coordinated orienting movements (head free or fixed). Until common methods are applied across all species, including humans, ground truth comparisons cannot be made. Therefore, seeking convergence of criteria across methods seems the only viable approach. Ultimately, a comparative analysis of the location, architecture, and connectivity of "FEF" should be framed by an appreciation of the interaction of gaze control with the body...
size, posture, locomotion, and reaching behavior, plus the elaborated relevance of gaze for communication in monkeys, apes, and humans living in larger social groups.

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