



Development of parallel auditory thalamocortical pathways for two different behaviors

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Auditory thalamocortical connections are organized as parallel pathways that originate in different divisions of the medial geniculate body (MGB). These pathways may be involved in different functions. Surprisingly little is known about the development of these connections. Here we review studies of the organization and development of auditory thalamocortical pathways in the pallid bat. The pallid bat depends primarily on passive hearing of prey-generated noise for localizing prey, while reserving echolocation for general orientation and obstacle avoidance. In the inferior colliculus (IC) and the auditory cortex, physiological studies show that noise and echolocation calls are processed in segregated regions. Injection of retrograde tracers in physiologically characterized cortical sites show that the ventral division of the MGB (MGBv) projects to the cortical region selective for noise. The cortical region selective for echolocation calls receives input from the suprageniculate (SG) nucleus in the dorsal MGB, but not from the MGBv. Taken together, these studies reveal parallel IC–MGB–cortex pathways involved in echolocation and passive listening. There is overlap of thalamocortical pathways during development. At 2-weeks postnatal, when the bat begins to exhibit adult-like hearing thresholds, the SG projects to both noise- and echolocation call-selective regions. The MGBv, as in adults, projects only to the noise-selective region. The connections become adult-like only after 2-months postnatal. These data suggest that parallel auditory thalamocortical pathways may segregate in an experience-dependent fashion, a hypothesis that remains to be tested in any species.

Keywords: auditory cortex, medial geniculate body, auditory development, plasticity, parallel pathways, echolocation, thalamocortical

PARALLEL AUDITORY THALAMOCORTICAL PATHWAYS

A hallmark of sensory system organization is parallel pathways. In the auditory system, auditory nerve fibers diverge into different divisions of the cochlear nucleus setting the stage for multiple ascending pathways. At the thalamocortical level, parallel pathways arise from three distinct divisions of the medial geniculate body (MGB; for review see Winer et al., 2005). These parallel pathways are purported to serve different functions in hearing. Thalamocortical connections include a tonotopic pathway through the MGBv (ventral division) which forms the cortical substrate for functional subregions within isofrequency contours (reviewed in Read et al., 2002). The pathway through the MGBm (medial division) is implicated in multisensory integration and learning (Wepsic, 1966; Edeline and Weinberger, 1992; Bordi and LeDoux, 1994; for review see Hu, 2003). The non-tonotopic pathway through the MGBd (dorsal division, Raczkowski et al., 1976; Andersen et al., 1980; Morel and Imig, 1987; for review see Imig and Morel, 1983; Rouiller, 1997) may be involved in representing complex sounds (Aitkin and Dunlop, 1968).

The hypothesis that parallel thalamocortical pathways are involved in different aspects of hearing is supported by studies in three species of bats from different families: the mustached bat (*Pteronotus parnellii*; Mormoopidae family), the horseshoe bat (*Rhinolophus rouxi*; Rhinolophidae family) and the pallid bat (*Antrozous pallidus*; Vespertilionidae family). The mustached bat

auditory cortex contains a primary auditory cortex (A1) with a tonotopic map (reviewed in Suga, 1989). This map, as in all other species examined, receives input from the MGBv. Some specializations for echolocation, such as over-representation of dominant harmonic frequencies of the echolocation call, are present in this pathway, but within the context of tonotopic representation. Information relevant to target distance is represented by delay-tuned combination-sensitive neurons. A large delay-tuned area is located dorsal to A1. Smaller delay-tuned areas are present ventral to A1. Injections of different tracers in different delay-tuned areas label separate regions in the MGB indicating that multiple delay-tuned areas are present in the thalamus (Pearson et al., 2007; also see Wenstrup, 1999 for physiological studies with a similar conclusion) and that each thalamic delay-tuned area representation projects independently, and in parallel, to a cortical delay-tuned area (Pearson et al., 2007). Particularly, labeling was found in the rostral pole nucleus. Although it remains debated whether the rostral pole nucleus should be part of the dorsal division, the lateral part of the posterior group or an extension of the ventral division (Morel and Imig, 1987; Wenstrup et al., 1994; Lee et al., 2004; Pearson et al., 2007), it is clear there are parallel pathways conveying information from the MGB to the cortex.

In the horseshoe bat, the cortical region termed dorsal cortex contains neurons that may be involved in target ranging. These neurons receive input from the MGBd (Radtko-Schuller, 2004;

Radtke-Schuller et al., 2004). The primary auditory cortex receives input from the MGBv. In both the mustached and horseshoe bat the suprageniculate (SG) nucleus, a part of the MGBd, projects diffusely to the entire auditory cortex, with a significant input to non-primary cortical regions involved in processing echolocation calls. Taken together, these data indicate that the thalamocortical pathways that serve fine frequency analysis and target distance calculation in the mustached and horseshoe bat are mostly segregated (**Figure 1A**).

Studies on the pallid bat provide evidence that parallel auditory pathways may represent sounds used in two different behaviors. The pallid bat localizes terrestrial prey by listening to prey-generated noise while reserving echolocation calls for general orientation and obstacle avoidance (Bell, 1982; Fuzessery et al., 1993). For echolocation, it uses a downward frequency modulated (FM) sweep (60 → 30 kHz). For prey localization it depends on noise transients (5–40 kHz). Its auditory cortex and inferior colliculus (IC) are organized to process FM sweeps and noise in mostly segregated regions (Fuzessery, 1994; Razak and Fuzessery, 2002). In the auditory cortex, there is a tonotopic map with frequencies from 5 to 70 kHz. Most neurons with tuning between 5 and 30 kHz respond best to noise transients. This region has been termed the low-frequency region (LFR). Most neurons with tuning between 30 and 60 kHz respond best to downward FM sweeps. This region has been termed the high-frequency region (HFR).

The majority of inputs to the LFR and HFR in adult pallid bats arise from different divisions of the MGB (**Figure 2**, schematized in **Figure 1B**). Placement of retrograde tracers in the LFR labeled neurons predominantly in the MGBv with no label in the SG (Razak et al., 2007). Placement of tracers in the HFR labeled neurons in the SG, but not the MGBv. The MGBm and parts of MGBd outside of the SG send minor projections to both the LFR and the HFR. Thus

parallel thalamocortical pathways represent sounds involved in two different behaviors, with the SG → HFR connections putatively involved in echolocation behavior and MGBv → LFR connections putatively involved with passive prey localization.

A SIGNIFICANT GAP IN KNOWLEDGE: DEVELOPMENT OF AUDITORY THALAMOCORTICAL CONNECTIONS

While the existence and functional significance of parallel thalamocortical pathways are established, very little is known about the development of these pathways. The development of these pathways between the MGB and the auditory cortex has been the subject of only one study. Gurung and Fritsch (2004) found that the MGB innervates the auditory cortex before onset of patterned sensory input. However, whether segregation of MGB-auditory cortex inputs into parallel pathways depends on sensory experience remains unknown. Research on the development of auditory pathways has focused on connections below the thalamus. Studies of connections of the midbrain (Friauf and Kandler, 1990; Gabriele et al., 2000, 2007) and lower brainstem (Leake et al., 2002) suggest that the nucleotopic connections are mostly adult-like before hearing onset. Postnatal refinement or maintenance of connections, perhaps in an activity-dependent manner, is restricted to sharpening tonotopic organization (Kim and Kandler, 2003; Franklin et al., 2006; Leake et al., 2006) and fine tuning bilateral inputs within nuclei involved in sound localization (Kapfer et al., 2002).

DEVELOPMENT OF PARALLEL THALAMOCORTICAL CONNECTIONS IN THE PALLID BAT

The pallid bat is suited to address the development of thalamocortical connections for several reasons. First, as mentioned above, parallel pathways can be linked to representation of different behaviorally relevant sounds. Second, the ontogeny of both echolocation

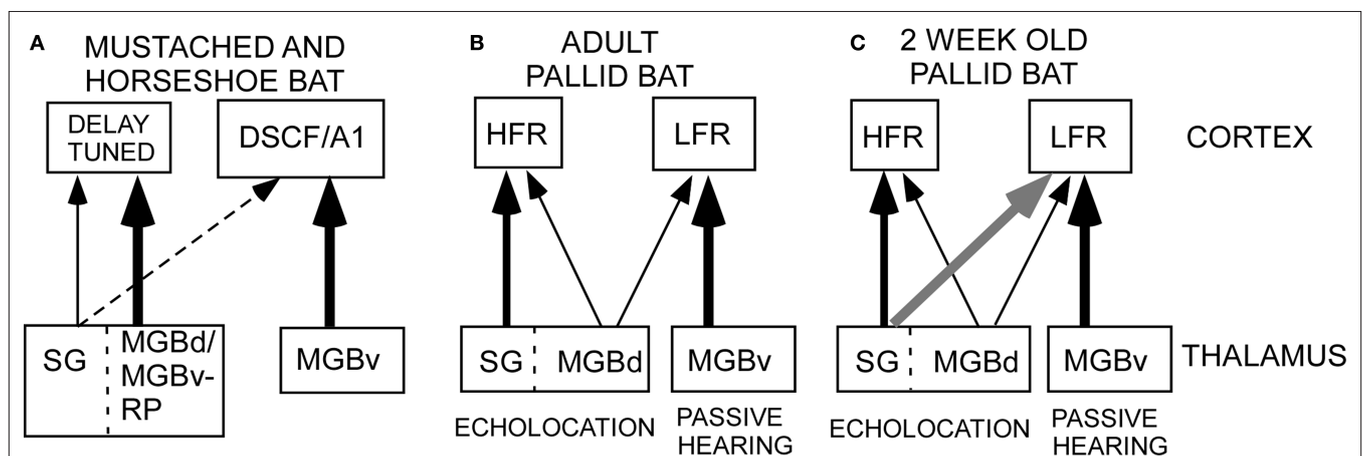
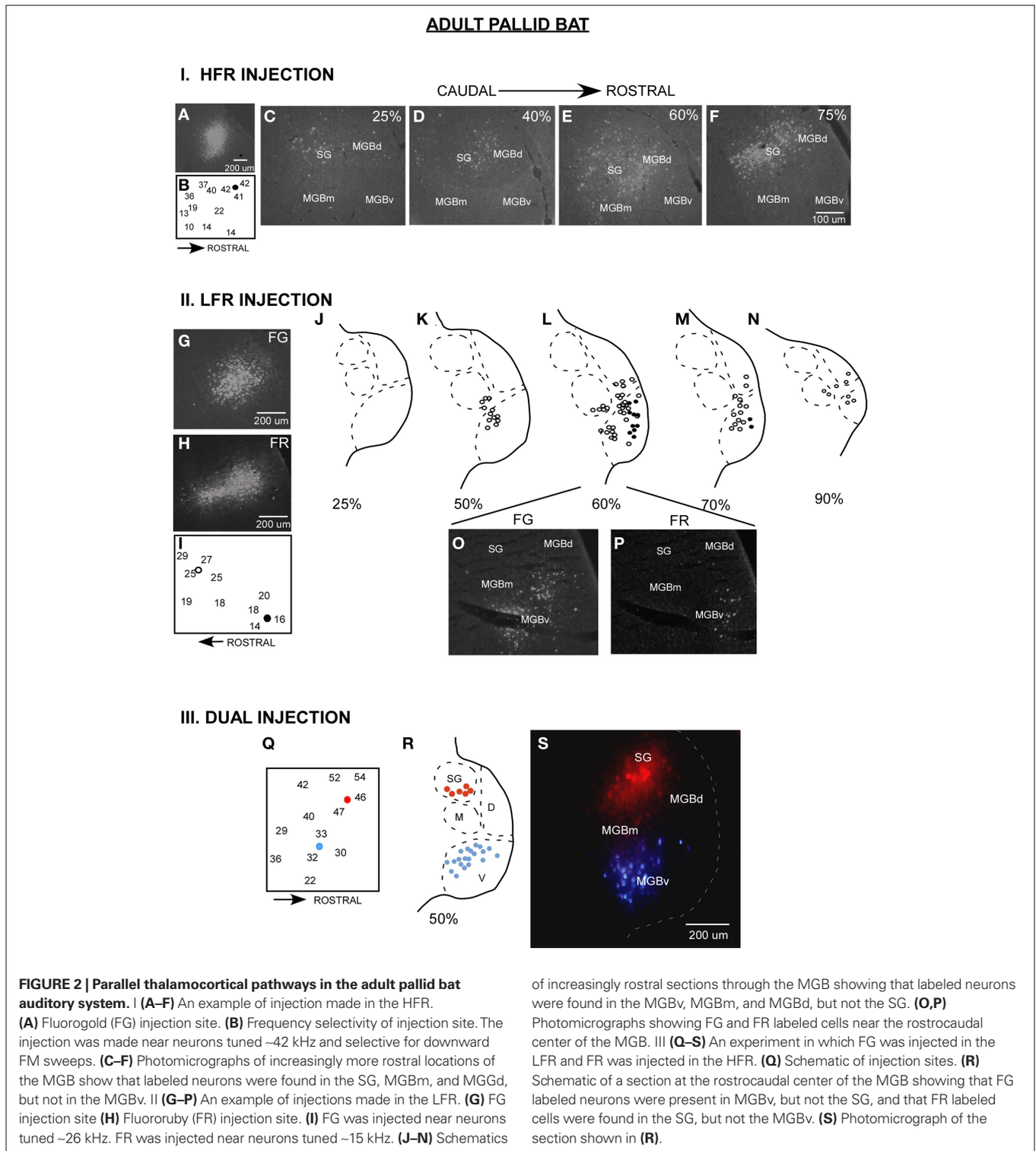


FIGURE 1 | Schematic of thalamocortical pathways in adult mustached, horseshoe and pallid bats and young pallid bat. (A) In the horseshoe bat, the MGBd projects mainly to regions dorsal to A1 that contain the combination-sensitive neurons used in target distance computation. The MGBv projects to A1. In the mustached bat, the rostral pole nucleus projects to the delay-tuned areas. It is not clear if the rostral pole nucleus should be considered a part of the ventral or dorsal MGB. Therefore, the term “MGBd/MGBv-RP” is used in this schematic. In both species, the SG, considered a part of the MGBd, projects diffusely but more to the non-primary cortex than primary cortex. **(B)** In adult

pallid bats, the HFR involved in echolocation behavior receives input from the SG and the MGBd, but not the MGBv. The LFR, involved in passive localization, receives input from the MGBv, but not the SG. Based on response selectivity in the auditory cortex, the SG → HFR pathway is involved in echolocation behavior while the MGBv → LFR pathway is involved in passive localization of prey-generated noise. **(C)** In a 2-week-old pallid bat pup, however, the pathways overlap. This is because the SG projects to both the LFR and HFR. The pup MGBv, as in adults, does not project to the HFR. Thus anatomically segregated pathways arise through postnatal refinement of initially overlapping connections.



behavior (Brown et al., 1978) and functional organization of auditory cortex (Razak and Fuzessery, 2007) has been studied. This provides a behavioral framework during development to compare anatomical and physiological data. The overall functional organization of auditory cortex, including frequency representation, is adult-like early in development (Razak and Fuzessery, 2002, 2007).

An exception is the prevalence and anatomical distribution of functionally bimodal neurons (Razak et al., 1999). These neurons appear to receive input from both pathways, and have two discrete tuning curves tuned to frequencies used in echolocation and passive sound localization. In adults, these bimodal neurons are found near the interface of the LFR and HFR. In pups, however, they are

more widely distributed (Razak and Fuzessery, 2007), suggesting that projections from the MGBv and SG may show greater overlap in their cortical targets. We tested this hypothesis by retrograde tracing of MGB inputs from physiologically identified injection sites in the LFR and HFR of auditory cortex.

Details about injection method, sizes, and definition of divisions of the MGB have been addressed in detail in the original papers (Razak and Fuzessery, 2007, 2009). Although it was relatively easy to distinguish the SG from rest of the MGBd and MGBv, the boundary between MGBd and MGBv was less obvious based on Nissl or neutral red stains. However, luxol fast blue, neutral red stained sections show that MGBv stains weakly for myelin, whereas the MGBd contains fibers that run in a dorsomedial to ventrolateral direction. The boundary between the dorsal and the ventral divisions was drawn based on myelin staining. The injection sizes are noted in Razak and Fuzessery (2007). A comparison of adult and pup brain photomicrographs shows similar injection sizes for fluororuby and flurogold (Figures 2 and 3).

There is greater overlap in thalamocortical connections during early development (Razak et al., 2009). Injections of retrograde tracers in the LFR labeled neurons in both the MGBv and the SG (Figure 3, schematic in Figure 1C). The input from the SG to LFR was prominent from the caudal half of the MGB, and was at least as strong as the input from the MGBv to the LFR based on the number of labeled neurons. However, as in adults, there was no input from the pup MGBv to the HFR. An overlap of pathways was seen in P60 bats, but not in P150 bats, indicating that the refinement is occurring during this time period. Pallid bats obtain adult-like hearing sensitivity ~2-weeks postnatal. Pallid bats begin to fly and use echolocation in flight around 5–6 weeks (Brown et al., 1978) and are fully weaned around 12 weeks. Thus the refinement occurs a few weeks after flight onset, and more than 6 weeks after onset of adult-like hearing. These data show that the parallel thalamocortical pathways seen in adults emerge during postnatal development and may involve refinement of overlapping connections well after the bat begins to experience patterned input.

WHY PARALLEL PATHWAYS FOR NOISE AND FM SWEEPS?

We have suggested that the parallel pathways for noise and FM sweep processing observed in the IC, MGB, and cortex are an adaptation for gleaning behavior in the pallid bat (Barber et al., 2003). As the bat hunts, it receives both echoes from the flight path and prey-generated noise from the ground. The bat has to segregate these two streams of temporally overlapping inputs. The noise and FM sweeps used in prey localization and echolocation differ in spectral (low versus high frequency), temporal (noise versus sweep), and spatial (ground versus flight path) qualities. These differences should enable the bat to segregate the two sounds (Bregman, 1990). In addition, the parallel processing with strong filters (noise versus FM sweep selectivity) will provide a substrate for additional segregation of the two auditory streams. During development, the onset of adult-like hearing thresholds may not be the critical experience for initiation of segregated pathways. The experience with dual processing of echoes and terrestrial prey-generated noise may be more important in initiating segregated pathways; therefore refinement occurs when the bat begins to hunt. Such experience-dependent plasticity has been described previously in the barn owl midbrain (Bergan et al., 2005).

A NOTE ON THE SUPRAGENICULATE NUCLEUS

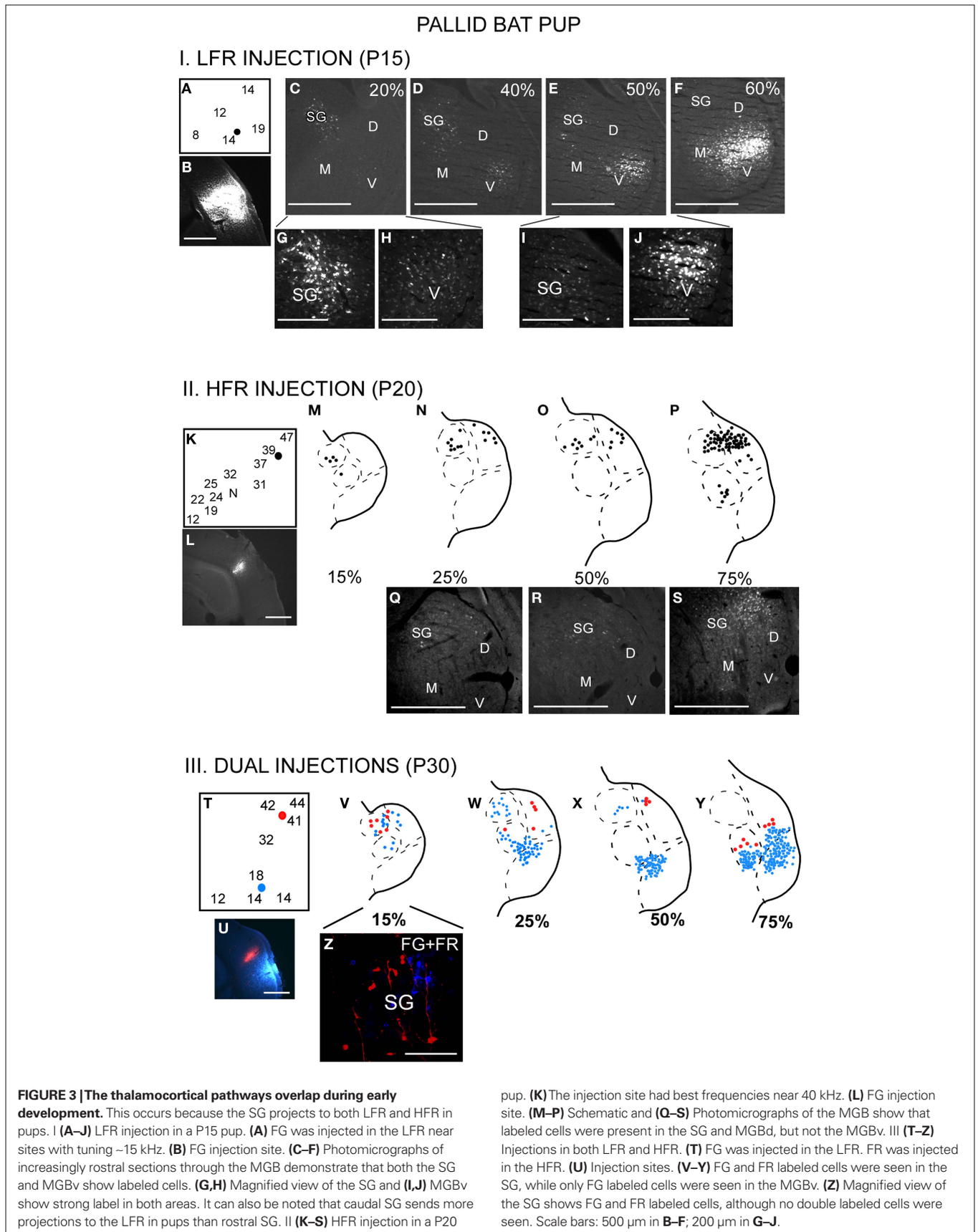
The SG appears to be a phylogenetically and ontogenetically malleable region of the thalamus. In non-chiropterans the SG is dominated by visual input from the superior colliculus (cat: Calford and Aitkin, 1983; Katoh and Benedek, 1995; rat: Tanaka et al., 1985). Less than 15% of the neurons respond only to auditory stimuli, whereas 65% of the SG neurons respond to visual stimuli alone (Benedek et al., 1997). The entire rostrocaudal extent of the SG receives input from the superior colliculus (SC) (Katoh and Benedek, 1995). Auditory inputs to the SG may arise from the nucleus of the central acoustic tract in the brainstem and the external nucleus of the IC (Henkel, 1983; Katoh and Benedek, 1995), both of which are considered to be a part of the extralemnisal auditory pathways. The SG projects primarily to non-primary auditory cortical areas, although sparse connections to the primary auditory cortex are found in many species (owl monkey: Morel and Kaas, 1992; macaque monkey: Hackett et al., 1998; tamarin: Luethke et al., 1989; dog: Malinowska and Kosmal, 2003; rat: Roger and Arnault, 1989).

In bats, the SG is dominated by auditory inputs. The SG of the mustached bat, unlike non-chiropterans, receives inputs from all frequency bands of the central nucleus of the IC (Wenstrup et al., 1994). Like non-chiropterans, the SG also receives direct input from the nucleus of the central acoustic tract (Casseday et al., 1989; Gordon and O'Neill, 2000). In addition to broad regions of the auditory cortex, the SG of the mustached bat projects to frontal cortex, which in turn projects to the SC, suggesting a role in acoustic-motor reflexes (Kobler et al., 1987). In the horseshoe bat, the SG projects to both primary and non-primary cortical fields (Radtke-Schuller et al., 2004). There is overlap in projections from the SG and the MGBv to any given cortical region, but the dorsal cortical fields (non-primary) receive most SG inputs. In the pallid bat, the echolocation pathway is primarily routed through the SG. Taken together, these studies suggest that the SG can be taken over to increase thalamic representation of species-specific dominant sensory modalities.

The developmental plasticity of SG connections is illustrated by cross-modal plasticity in ferrets (Pallas et al., 1990). By appropriate lesions of retinal axon targets, visual input can be guided to the MGB. This rewiring leaves the MGBv-primary auditory connections similar to those of controls. However, an anomalous input to the primary auditory cortex arises from the dorsal thalamus, including the SG, in the cross-modal animals. This finding suggests that the SG typically receives and sends more exuberant connections during early development. These connections may be either stabilized or pruned by activity-dependent mechanisms. The data from the pallid bat thalamocortical development support the hypothesis that the SG sends exuberant projections during early development.

FUTURE STUDIES

1. The studies on pallid bat thalamocortical connections suggest postnatal refinement. However, it remains unclear if this is experience-dependent. It is possible to modify echolocation experience during development (Razak et al., 2008). Such modifications significantly alter response selectivity within the



echolocation pathway. Future studies will address how refinement of pathways proceeds in bats with altered experience with echolocation calls.

2. Although several studies have focused on development of auditory pathways below the thalamus, the pallid bat is the only animal we are aware of in which postnatal development of auditory thalamocortical connections has been examined. Therefore it remains unclear if the postnatal refinement of parallel thalamocortical pathways is a general phenomenon. In animals, such as the cat, in which the adult auditory pathways have been well characterized, studies are required on the development of these pathways. Significantly more is known about development of visual and somatosensory thalamocortical pathways.
3. A key hypothesis to emerge based on physiological and anatomical studies of the pallid bat auditory system is that in gleaning bats, parallel pathways from the IC–MGB–cortex represent different sounds used in different behaviors. This hypothesis is supported by physiological studies in the IC of *Megaderma lyra*, another gleaner bat from a different family (Rubsamen et al., 1988). Gleaning as a hunting strategy appears to have evolved independently in different families of bats. If parallel pathways are an adaptation for gleaning in the pallid bat, then a similar organization of the auditory system must be present in other gleaner bats. This hypothesis of convergent evolution of the auditory system remains to be tested. One unusual aspect of

the pallid bat thalamocortical connections is that the cortical tonotopic map is composed of non-overlapping inputs from two different divisions of the MGB (Razak et al., 2007). The presence of such an organization in other gleaners will strongly suggest a modified thalamocortical plan in gleaners compared to other mammals, including obligate echolocating bats.

4. The molecular mechanisms that make the SG ontogenetically and phylogenetically labile are unknown. Recent studies that compared FOXP2 expression between echolocating and non-echolocating bats identified the SG as one of the nuclei in which expression was higher in the echolocators (Berquist et al., SFN abstracts, 2009). Expression of FOXP2 in the MGB also changes in an activity-dependent manner (Hornig et al., 2009). FOXP2 may therefore be a part of the molecular milieu of the SG that makes it developmentally and evolutionarily more labile. Future studies comparing the expression of FOXP2 across various divisions of the MGB will be useful in determining the mechanisms of SG malleability and to address mechanisms underlying connectional and therefore functional plasticity in the thalamocortical connections.

ACKNOWLEDGMENTS

We thank T. Zumsteg for valuable comments on the manuscript and G. McLellan for programming the software required for this study. This research was supported by an NIDCD grant DC05202 and INBRE P20 RR0 16474-04.

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

Received: 15 January 2010; paper pending published: 03 August 2010; accepted: 29 August 2010; published online: 21 September 2010.

Citation: Razak KA and Fuzessery ZM (2010) Development of parallel auditory thalamocortical pathways for two different behaviors. *Front. Neuroanat.* 4:134. doi: 10.3389/fnana.2010.00134

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