Cytoarchitecture of the superior olivary complex of three neotropical species of bats (Noctilio leporinus, Phyllostomus hastatus and Carollia perspicillata) with different foraging behavior

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Abstract

The understanding of the echolocation by studying different auditory nuclei of echolocating bats can be an important link in elucidating questions arising in relation to their foraging behavior. The superior olivary complex (SOC) is the primary center for processing the binaural cues used in sound localization since echo locating bats rely on acoustic cues to navigate and capture prey while in flight. The present study was taken to test the hypothesis that the SOC of echolocating neotropical bats with different foraging behavior will exhibit morphological variations in relative size, degree of complexity and spatial distribution. The brains were collected from six male adult bats of each species: Noctilio leporinus (fish eating), Phyllostomus hastatus (carnivorous/omnivorous) and Carollia perspicillata (fruit eating). They were double-embedded and transverse serial sections were cut and stained with cresyl fast violet. The SOC measured as 640 ± 70 µm in the \textit{N. leporinus} bat, 480 ± 50 µm in the \textit{P. hastatus} and 240 ± 30 µm in the \textit{C. perspicillata} bat. The principal nuclei of the SOC of in all three bats were the LSO, MSO and MNTB. The MSO and LSO were very well developed in \textit{N. leporinus} bats. The MSO of \textit{N. leporinus} bat subdivided into DMSO and VMSO. The main cell type of cells present in MSO and LSO are dark staining multipolar cells in all the bats studied. The well-developed MSO and LSO of \textit{N. leporinus} bats indicate that these bats are highly sensitive to low frequency sounds and interaural intensity differences, which help these bats to forage over water by using various types of echolocation signals. The average size of SOC in \textit{P. hastatus} and \textit{C. perspicillata} bats can be attributed to the fact that these bats use vision and smell along with echolocation to forage the food.

Keywords: bats, superior olivary complex, medial superior olive, lateral superior olive.

Citoarquitetura do complexo olivar superior de três espécies neotropicais de morcegos (Noctilio leporinus, Phyllostomus hastatus e Carollia perspicillata) com diferentes comportamentos de forrageamento

Resumo

O entendimento da ecolocalização pelo estudo de diferentes núcleos auditivos de morcegos pode ser um elo importante na elucidação das inúmeras questões que surgem em relação ao seu comportamento de forrageamento. O complexo olivar superior (SOC) é o principal centro de processamento das pistas binaurais usadas na localização do som, já que os morcegos ecolocalizadores contam com sinais acústicos para navegar e capturar as presas durante o vôo. O presente estudo foi realizado para testar a hipótese de que morcegos que usam a ecolocalização para diferentes comportamentos de forrageamento irão variar na estrutura, tamanhos relativos e grau de complexidade e distribuição espacial do grupo SOC. Os cérebros foram coletados de seis machos adultos de morcego de cada espécie: Noctilio leporinus (piscívoro), Phyllostomus hastatus (carnívoro/omnívoros) e Carollia perspicillata (frugívoro). Eles foram seccionados em série e transversalmente, cortados e corados com coloração rápida cresil-violeta. O grupo SOC foi medido como 640 ± 70 µm no morcego \textit{N. leporinus}, 480 ± 50 µm no \textit{P. hastatus} e 240 ± 30 µm no morcego \textit{C. perspicillata}. Os principais núcleos do grupo SOC dos três morcegos foram o LSO e o MSO e o MNTB. O MSO e o LSO foram muito bem desenvolvidos em morcegos \textit{N. leporinus}. A MSO de \textit{N. leporinus} foi subdividida em DMSO e VMSO. O principal tipo de células presentes na MSO e LSO são as células multipolares de coloração escuro em todos os morcegos. Os MSO bem desenvolvidos e LSO de morcegos \textit{N. leporinus} indicam que estes morcegos são...
1. Introduction

The superior olivary complex (SOC) is a well-developed auditory brainstem structure in all mammals including echolocating bats. The SOC plays a number of roles in hearing including the localization of sound resources, measuring the time difference of arrival of sounds between the ears, encoding temporal features of sounds and descending modulation of cochlear nucleus (Grothe and Park, 2000). The SOC is typically located in the caudal brainstem near the facial nucleus and consists of up to 13 distinct cell groups, each contributing a unique neuronal circuit and sub serving a distinct functional role in the processing of sound (Schofield, 2002). The SOC consists of two principal nuclei, the medial superior olive (MSO) and the lateral superior olive (LSO) with established roles in hearing. The principal cell groups are flanked by a population of peri-olivary nuclei whose functional contributions are poorly understood.

The MSO and LSO are a major site of convergence of information arising from both cochlear nuclei (Glendenning et al., 1985). The MSO neurons are binaural, most sensitive to low-frequency sounds (at or below 5 kHz) and encode interaural timing differences (Spitzer and Semple, 1995) and LSO neurons are binaural and sensitive to interaural intensity differences (Sanes, 1990). The LSO is most prominent in animals with excellent high-frequency hearing, especially those utilizing echolocation (Zook and Casseday, 1982; Glendenning and Masterton, 1998). Recently however, it has been speculated that the size of the LSO is more indicative of the animal’s overall hearing range rather than sensitivity to high-frequency sounds (Moore, 2000).

Many bats use echolocation for orientation in space and for detecting and capturing prey in total darkness (Genoud et al., 1990; Webster et al., 1992; Bailey et al., 1992). The echolocating bats use different echolocation strategies based on their foraging environment. The vast majority of echolocating bats emit very short (0.5-5.0 ms) frequency modulated (FM) sweeps, covering a large proportion of their hearing range. The FM echoes returning from flying insects carry information about the range and location of those targets. In contrast to the FM bats, other bats such as horseshoe bats and mustached bats independently developed an echolocation strategy using a combination of a long constant frequency component (CF), and a brief FM sweep. The use of CF-FM echolocation calls allows these bats to segregate different aspects of information. The echo from the FM sweep is used for ranging and localization, while Doppler shifts in the echo from the CF component carry information about the relative velocity of the bat to a target. In addition, the amplitude and frequency modulations imposed on the echo of the CF component by an insect’s wing beating allow the bat to identify specific insect species (Von der Emde and Schnitzler, 1990) and to distinguish fluttering prey from non-fluttering background, even in a densely cluttered environment.

The similarities and differences that will be exhibited in the SOC of bats with different echolocation strategies can be useful in understanding the principal problems of what aspects of sound different SOC nuclei process and what some of the underlying mechanisms and circuitry patterns are in the SOC. The information on the comparative analysis of structure and cytoarchitecture of SOC associated with echolocating bats with different foraging strategies seems to be lacking.

The three species of Neotropical bats viz., Noctilio leporinus Linnaeus 1758, Phyllostomus hastatus Pallas 1767 and Carollia perspicillata Linnaeus 1758 were chosen based on their differing foraging behavior for the present study. The living range of these bats stretches from Mexico to Northern Argentina and also includes most Caribbean islands. The N. leporinus belongs to the Nectarionidae family whereas both P. hastatus and C. perspicillata belongs to Phyllostomidae family. The N. leporinus bats lives mostly around well-watered lowland and coastal areas as well as river basins. They eat small fish in both fresh and salt water but they need calm water surfaces in order to detect ripples. P. hastatus bats are omnivores, feeding on flowers and pollen, but also insects and small vertebrates, forage on open and forested regions. C. perspicillata bats are mainly frugivorous. However, they may feed on insects and sometimes pollen, which forage on moist evergreen and dry deciduous forests. Therefore, the present study is undertaken to provide comparative structural and cytoarchitectural details of the SOC of three species of bats with different foraging behavior.

2. Material and Methods

For the present study, six adult male live bats of each of the three species, N. leporinus, P. hastatus and C. perspicillata were collected. The bats were weighed and anaesthetized by using xylazine 2 mg/kg and ketamine 10 mg/kg intramuscularly. The research protocol was approved by the institutional ethical committee. Immediately after euthanasia, the brains of the bats were removed, weighed and placed in 10% formal saline. The brains were manually processed and double embedded (Gibbons et al., 2013a, b). The tissues were then blocked and coronal sections at 10 μm were made by using the rotary microtome MT 960. The sections were stained using Cresyl Fast violet. The size, shape and orientation of the cells were analyzed with the aid of the Olympus BX51 system microscope and the digital images were taken with the help of Olympus DP71 microscope digital camera.
3. Results

3.1. Superior olivary complex of the Noctilio leporinus bat

At the most caudal level, the SOC first becomes apparent at the level of facial nucleus (see Figure 1A). The facial nucleus was easily distinguished by a large cell group containing homogenous population of relatively large multipolar neurons. Lateral to the facial nucleus, the SOC comprised of mainly dense-staining multipolar neurons measuring between 12.5 to 17.5 µm in diameter. In addition, few light-staining, round cells with a diameter of 7.5 µm were present. Dense-staining, oval cells measuring 5 µm in diameter were also present but very few in numbers. The SOC measured 500 ± 55 µm at its widest point, medio-laterally. Overall, the SOC extends rostrally the mid-pons and measured 640 ± 70 µm in length.

Progressing rostrally, the caudal third of the SOC divided into two main portions: medial superior olive (MSO) and lateral superior olive (LSO) (see Figure 1B). The LSO is the larger of the two. It contained a folded region medially and this allowed for the further differentiation of LSO into two portions. The LSO measured 850 ± 70 µm in height and 640 ± 85 µm in width and MSO measuring 100 ± 14 µm in height and 400 ± 38 µm in width. The cells in this portion were large dense staining multipolar cells, measuring between 17.5 and 25 µm in diameter.

At the level of the motor nucleus of the trigeminal nerve, the middle third of the MSO divided into two distinct nuclei as the ventral medial superior olive (VMSO), and dorsal medial superior olive (DMSO) (see Figure 1C). The VMSO appeared like a horse-shoe appearance at this level (see Figure 1D). At this level, the LSO measured...
350 ± 25 μm in both height and width; while the DMSO measured 50 ± 6 μm in height and 290 ± 33 μm in width and the VMSO, which appeared horse shoe-shaped, 70 ± 9 μm in height and 290 ± 33 μm in width. The cells in all three divisions in this region were found to be mainly large, dense-staining, elongated neurons, measuring between 17.5 and 25 μm in diameter (see Figure 1F). As the SOC proceeded rostrally, only the DMSO was predominantly seen and it comprised the same large multipolar, dense-staining cells. However, the number of small round cells increased.

The rostral third of the SOC, at the level of the inferior colliculus, also revealed the three divisions (LSO, VMSO and DMSO) but the VMSO, no longer assuming a horseshoe appearance (see Figure 1E). The VMSO measured 290 ± 33 μm in width and 90 ± 7 μm in height; while the DMSO measured 350 ± 31 μm in width and 70 ± 5 μm in length and the LSO, 320 ± 37 μm in width, 200 ± 23 μm in height. The LSO, DMSO and VMSO were no longer present at the level of the commissure of the inferior colliculus. The mean body and brain weight of this bat were 48.1 ± 3.5 g and 6.93 ± 0.47 g respectively.

3.2. Superior olivary complex of the Phyllostomus hastatus bat

The SOC first appeared at the level of the facial nucleus and placed lateral to the facial nucleus (see Figure 2A). The caudal third of the SOC divided into LSO and MSO with both areas comprising similar cell types (see Figure 2B). The stromal pattern differentiated one portion from the other. It comprised large, dense-staining, multipolar cells, measuring between 15 and 22.5 μm in diameter (see Figure 2B and F).

Figure 2. Superior olivary complex of the Phyllostomus hastatus bat. (A) transverse section of the brainstem at the level of the facial nucleus. 1. Cochlear nucleus, 2. Caudal cerebellar peduncle, 3. Medial vestibular nucleus, 4. Fourth ventricle, 5. Facial nucleus, 6. Superior olivary complex; (B) the first (1) and second (2) divisions of the caudal third of the superior olivary complex; (C) transverse section of the brainstem at the level of the trigeminal nucleus. 1. Cochlear nucleus, 2. Trigeminal nucleus, 3. Fourth ventricle, 4. Superior olivary complex; (D) the three (3) divisions of the middle third of the superior olivary complex. 1. Lateral superior olive (LSO), 2. Dorsal medial superior olive (DMSO), 3. Ventral medial superior olive (VMSO); (E) transverse section of the brainstem at the level of the nucleus of the trapezoid body. 1. Fourth ventricle, 2. Dorsal medial superior olive (DMSO), 3. Nucleus of the trapezoid body; (F) the cells found in the superior olivary complex.
At the level of the trigeminal nucleus, the middle third of the SOC measured 970 ± 78 µm in height and all three divisions, LSO, DMSO and VMSO were now apparent (see Figure 2C and D). At the rostral third at the level of nucleus of the trapezoid body, only the DMSO division was seen (see Figure 2E). No divisions of the superior olivary complex were visible at the level of the inferior colliculus. The total length of SOC measured 480 ± 50 µm in length from rostrocaudally. The mean body and brain weight of this bat were 73.24 ± 7.25 g and 8.57 ± 0.67 g respectively.

3.3. Superior olivary complex of the Carollia perspicillata bat

At the level facial nucleus, the caudal third of the SOC was observed immediately lateral to the nucleus of trapezoid body. It divided into two: lateral and medial portions (see Figure 3A). Both divisions contained medium and large multipolar cells measuring between 12.5 µm and 22.5 µm in diameter as well as small round cells measuring average diameter of 5 µm.

The divisions of the middle third of the SOC were very indistinct only at the level of the middle cerebellar peduncle (see Figure 3B). The rostral third of the SOC was located at the level of the inferior colliculus (see Figure 3C) and comprised mainly large multipolar cells, measuring between 20 to 25 µm in diameter. The total length of SOC measured 240 ± 30 µm from rostrocaudally. The mean body and brain weight of this bat were 14.4 ± 2.1 g and 3.74 ± 0.75 g respectively.

4. Discussion

The nuclei involved in audition tend to be larger in animals that echolocate and in those with an excellent sense of hearing, compared to other animals (Reis and Erhart, 1979; Casseday et al., 1988). The other auditory nuclei like inferior colliculus, medial geniculate body and cochlear nuclear complex shown significant differences in their size in N. leporinus, P. hastatus, and C. perspicillata bats (Gibbons et al., 2013a; b; Adogwa et al., 2014). SOC displays a significant interspecies variation, being largest in bats and rodents and smaller in primates (Grothe and Park, 2000). The SOC is very well developed in all the bats studied so far. In the present study, the SOC was measured as 640 ± 70 µm in the N. leporinus bat, 480 ± 50 µm in the P. hastatus and 240 ± 30 µm in the C. perspicillata bat. The body and brain weight do not reflect proportionately on the size of the SOC in the present study as the body and brain weight of the P. hastatus measured as 73.24 ± 7.25 g and 8.57 ± 0.67 g respectively which is higher than the N. leporinus 48.1 ± 3.5 g and 6.93 ± 0.47 g, but the length of the SOC in P. hastatus (480 ± 50 µm) was lesser than N. leporinus (640 ± 70 µm).

The principal nuclei of the SOC of in all three bats in the present study were the lateral superior olive (LSO) and the medial superior olive (MSO), which is similar to the mustached bats (Covey and Casseday, 1995). The principal nuclei of the SOC project to the central nucleus of the inferior colliculus with the lateral superior olive projecting bilaterally and the others projecting to the inferior colliculus ipsilaterally. The different divisions of

Figure 3. Superior olivary complex of the Carollia perspicillata bat (A) the brainstem at the level of the facial nucleus of the C. perspicillata bat. 1. Cerebellum, 2. Pyramids, 3. Nucleus of the trapezoid body, 4. Superior olivary complex, 5. Cochlear nucleus, 6. Facial nucleus; (B) the brainstem at the level of the middle cerebellar peduncle. 1. Cerebellum, 2. Nucleus of the trapezoid body, 3. Fibres of the trapezoid body, 4. Superior olivary complex, 5. Cochlear nucleus, 6. Middle cerebellar peduncle; (C) the brainstem at the level of the inferior colliculus. 1. Cerebellum, 2. Inferior colliculus, 3. Fibres of the trapezoid body, 4. Nucleus of the trapezoid body, 5. Superior olivary complex.
the superior olivary complex received inputs from either the ipsilateral or the contralateral cochlear nucleus (Zook and Casseday, 1982).

The MSO was well-developed in *N. leporinus* than the other two bats and it further divided into dorsal medial superior olive (DMSO) and ventral medial superior olive (VMSO) in *N. leporinus* and *P. hastatus* bats whereas it was less developed in the *C. perspicillata* bats. The variations in the structure of SOC among bats were reflecting varying foraging strategies (Grothe and Park, 2000). The well-developed MSO in *N. leporinus* can be attributed to the need for high sensitivity to high frequency when these animals forage in open water surfaces and need to detect the water ripples.

In the present study, the subdivisions of MSO were referred to as DMSO and VMSO respectively as stated by Zook and Casseday (1982) in mustached bats rather than MSO and dorsal medial paraolivary olive (DMPO) as it was referred in the Mexican free-tailed bat Grothe et al. (1994). The DMPO showed varying quantities of cell-types with different projection patterns. The fusiform cells were seen in the MSO whereas multipolar cells were seen in the DMPO (Schofield and Cant, 1991; Schwartz, 1977). It was also found that in the free-tailed bat, more than 30% of the cells of the MSO sent projections to both ipsilateral and contralateral inferior colliculi, whereas these cells sent projections ipsilaterally in other animals (Grothe et al., 1994). In the present study, there was no considerable difference in cell types between these two nuclei, with dark staining multipolar cells occurring predominantly in all three bats.

At the rostral third of the SOC, the VMSO of the *N. leporinus* folded into a horse-shoe shape. No apparent evidence of this occurring in other species of bats or other animals (Rietzel and Friauf, 1998; Helfert and Schwartz, 1986, 1987). The VMSO features that were noted in this study appeared to be equivalent to those of the DMPO (Grothe et al., 1994) in Mexican free-tailed bats. This comparison is based on similar cell-types and location of the nucleus. This VMSO projects to ipsilateral inferior colliculus and receive inputs from the cochlear nuclei (Rietzel and Friauf, 1998; Grothe et al., 1994). This division of the SOC did not appear in the *P. hastatus* and the *C. perspicillata* bats. This difference is probably attributed to the fact that the *N. leporinus* bat uses echolocation as a major part of its feeding strategy (Wenstrup and Suthers, 1984; Schnitzler et al., 1994), whereas, *P. hastatus* and *C. perspicillata* use vision and smell along with echolocation to locate food (Neuweiler, 1989).

The LSO was well developed in *N. leporinus* than the other two bats in this study. The cell population was predominantly the multipolar and elongated cells, which is similar to the multipolar and banana-like cells seen in the LSO of rat, gerbil and cat. These animals use high frequency hearing excellently during foraging due to the well-developed LSO (Rietzel and Friauf, 1998; Helfert and Schwartz, 1986, 1987). Recently however, it has been speculated that the size of the well-developed LSO is more indicative of the animal’s overall hearing range rather than sensitivity to high-frequency sounds (Moore, 2000). The cells of the LSO were described as fusiform and bipolar in the ferret and the guinea pig (Rietzel and Friauf, 1998; Helfert and Schwartz, 1986). Generally, the neurons in this area may be either excitatory or inhibitory depending on whether the stimulus is received from the ipsilateral or the contralateral ear, respectively (Schofield and Cant, 1991). The well-developed LSO in *N. leporinus* used in this study can be attributed to the fact these bats with excellent high-frequency hearing and the overall hearing range than other two bats which helps the *N. Leporinus* to forage effectively over water by various using various echolocation signals. The bats belong to Phyllostomidae family use vision and smell along with echolocation while foraging whereas the bats belongs to Noctilionidae family rely heavily on echolocation for foraging (Neuweiler, 1989).

In the present study confirmed the statement of Neuweiler (1989) as the *P. hastatus* and *C. perspicillata* are belonging to the Phyllostomidae family whereas the *N. leporinus* belong to the Noctilionidae family. Among the Phyllostomidae family bats, the *P. hastatus* have better developed SOC in the present study than the *C. perspicillata*, which concurs the reports of Hutcheon et al. (2002) that the insectivorous bats rely profoundly on echolocation for the pursuit of and capture of prey than do phytophagous species.

5. Conclusion

The principal nuclei of the SOC of in all three bats were the LSO, and MSO. The MSO and LSO are well-developed in *N. leporinus* bats. The MSO of *N. leporinus* bat subdivided into DMSO and VMSO. The main cell type of cells found in MSO and LSO were dark staining multipolar cells. The well-developed MSO and LSO of *N. leporinus* bats indicated that these bats were highly sensitive to low frequency sounds and interaural intensity differences, which helped these bats to forage over water by using various types of echolocation signals. The average SOC in *P. hastatus* and *C. perspicillata* bats can be attributed to the fact that these bats use vision and smell along with echolocation to forage.

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References

ADOGWA, A., SUNDARAM, V., GIBBONS, I. and ODEKUNLE, A., 2014. Cytoarchitecture of the medial geniculate body of three species of bats: *Noctilio leporinus*, *Phyllostomus hastatus* and *Carollia perspicillata*. *Annual Research & Review in Biology*, vol. 4, no. 3, pp. 460. http://dx.doi.org/10.9734/ARRB/2014/4961.

BAILEY, W.J., SLIGHTOM, J.L. and GOODMAN, M., 1992. Rejection of the “flying primate” hypothesis by phylogenetic evidence from the epsilon-globin gene. *Science*, vol. 256, no.
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MOORE, D.R., 2000. Auditory neuroscience: Is speech special? Current Biology, vol. 10, no. 10, pp. 362-364. http://dx.doi.org/10.1016/S0960-9822(00)00479-6. PMid:10837214.

NEUWEILER, G., 1989. Foraging ecology and audition in echolocating bats. Trends in Ecology & Evolution, vol. 4, no. 6, pp. 160-166. http://dx.doi.org/10.1016/0169-5347(89)90120-1. PMid:21227342.

REIS, P.F. and ERHART, A.E., 1979. The brain of the marmoset (Callithrix jacchus). Acta Anatomica, vol. 103, no. 3, pp. 350-357. http://dx.doi.org/10.1159/0000415034. PMid:107711.

RIETZEL, H.J. and FRAUFA, E., 1998. Neuron types in the rat lateral superior olive and developmental changes in the complexity of their dendritic arbors. The Journal of Comparative Neurology, vol. 390, no. 1, pp. 20-40. http://dx.doi.org/10.1002/(SICI)1096-9861(19980105)390:1<20::AID-CNE3>3.0.CO;2-S. PMid:9456173.

SANES, D.H., 1990. An in vitro analysis of sound localization mechanisms in the gerbil lateral superior olive. The Journal of Neuroscience, vol. 10, no. 11, pp. 3494-3506. http://dx.doi.org/10.1016/S0270-7146(99)80517-0. PMid:67031.

HUTCHERSON, J.M. and KIRK, A.J.W. and GARLAND, T.A., 2002. Comparative analysis of brain size in relation to foraging ecology and phylogeny in the Chiroptera. Brain, Behavior and Evolution, vol. 60, no. 3, pp. 165-180. http://dx.doi.org/10.1159/000065938. PMid:12417821.

WEBSTER, W.R., DAY, R.H., GILLIES, O. and CASSANDON, B., 1992. Spatial-frequency- contingent color aftereffects: adaptation with two-dimensional stimulus patterns. Perception & Psychophysics, vol. 51, no. 1, pp. 66-78. http://dx.doi.org/10.3758/BF03205075. PMid:1549426.

WENSTRA, J.J. and SUTHERS, R.A., 1984. Echolocation of moving targets by the fish-catching bat, Noctilio leporinus. Journal of Comparative Physiology, vol. 155, no. 1, pp. 75-89. http://dx.doi.org/10.1007/BF00610933.

ZOOK, J.M. and CASSEDAY, J.H., 1982. Origin of ascending projections to inferior colliculus in the musclete bat, Pteronotus parnellii. The Journal of Comparative Neurology, vol. 207, no. 1, pp. 14-28. http://dx.doi.org/10.1002/cne.902070103. PMid:7096636.