Quantitative Analysis of the Superior Colliculus and the Parabigeminal Nucleus in the Hereditary Unilaterally Microphthalmic Rat*

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Summary: Quantitative morphological changes in the superior colliculus (SC) and the parabigeminal nucleus (PB) were studied in hereditary unilaterally microphthalmic rats. The mutant animals have one vestigial and one almost normally developed eyeball. The former eye completely lacks the optic nerve. The proportion of uncrossed to crossed retinal fibers in the mutant rats was estimated at about 6%. Conspicuous changes in SC were found only on the contralateral side of the abnormal eye. Relative volume of the superficial layers of SC (SCS) to the central gray matter (SGC) was decreased to 50% of the normal. However, the cell density in SCS increased up to 130–190% of the normal. The str. griseum superficiale consisted mainly of small roundish neurons in dense and irregular arrangements. Small amounts of fibers were observed in the medial 1/3 of the str. opticum, but very few in the lateral 2/3. No significant changes were found in the deeper collicular layers of the mutant rats.

Unilaterally microphthalmic PB was subdivided normally into three parts: the dorsal (PBD), middle (PBM) and ventral sub-divisions (PBv). The relative volumes of each PB subdivision on both sides had decreased to 57 to 65% of the normal, except for that of PBv on the contralateral side to the anomalous eye (26% of the normal). Cell densities of PB were slightly lowered (76–84% of the normal) in PBM and PBv, but not much in PBD (88–90% of the normal) on both sides.

Uncrossed retinal fibers of rats are reported to terminate in a restricted region of the superficial gray and optic layers of the superior colliculus (SC)⁸,¹³,²⁴,²⁶. It has been also well documented that prenatal or neonatal removal of one eye in experimental animals resulted in the marked expansion of the terminal field in the uncrossed retinal projection (see Discussion). The expanded distribution was observed not only in the superficial layers of SC (SCS) but also in the lateral geniculate nucleus of either the
experimentally induced\(^\text{23}\) or the congenital unilaterally micro-\(^\text{39}\) and anophthalmic animals\(^\text{9}\).

Tokunaga et al. \(^\text{85}\) quantitatively demonstrated a significant hypoplasia of SCS in the bilaterally microphthalmic rats. An autopsy report of the uniocular blindness revealed no morphological changes in the mesencephalic tectum\(^\text{14}\). On the other hand, remarkable reduction in the size of contralateral SC has been reported in experimental animals having one eye removal at birth\(^\text{6,7,9,15,27,29,37,41}\). However, little is known about the quantitative morphological changes in the visual centers of hereditary unilaterally micro- or anophthalmic animals. The uni- and bilaterally microphthalmic rats, which are maintained with brother-sister matings in our laboratory, lack completely the optic nerve on one and both sides, respectively, due to growth inhibition of the optic cup at the early embryonic stage\(^\text{20}\). It is of interest to find out what sort of influences could be exerted by the expanded uncrossed retinotectal fibers upon developing SC in the mutant rat.

Recently, the parabigeminal nucleus (PB) the secondary visual centers in the midbrain because of its intimate connections with SC\(^\text{2,12,42}\). It can be expected that PB in the unilaterally microphthalmic rats is also involved in some structural changes during its development under the influence of the altered SC.

In this study morphological changes of SC and PB in the unilaterally microphthalmic rats will be described, with special reference to quantitative analysis, as well as a comparison, with those in bilateral microphthalmia\(^\text{40}\).

### Materials and Methods

Nine unilaterally microphthalmic rats (Donryu), ranging from 5 to 7 months old (body weight, 250–350 g), were used with 6 normal rats from the same strain as control. Animals were anesthetized with sodium amobarbital (60 mg/kg body weight) and perfused transcardially with a brief rinse of physiological saline (100 ml/each animal) followed by 300 ml of buffered 10% formaldehyde (pH 7.4). The brain was stored in the same fixative for 1 week at 4°C, and then embedded in celloidin. Serial coronal sections were cut at 25 \(\mu m\) and every other section was stained by the Klüver-Barrera method.

In order to compare the width of the optic tract, the relative thickness of the tract to the diameter of the anterior commissure was estimated as follows:

\[
B/A \times 100 \%
\]

A. Diameter of the anterior commissure measured at its midline in a coronal section.
B. Mean width of the optic tract measured on three successive coronal sections through 1 mm caudal to the posterior margin of the commissure (Fig. 1A).

Besides the unilateral microphthalmia, the optic tracts from 6 bilaterally microphthalmic rats\(^\text{40}\) without optic nerve on both sides were also examined. Ratio of the volume of the superficial tectal layers (the zonal, superficial gray and optic strata: SCS), of the deeper collicular layers (from the intermediate gray to the deep white laminae: SCD), and of PB to the volume of the central gray matter (SGC) beneath SC were determined as follows: Contours of SCS, SCD and SGC, and of PB in each coronal section were drawn with a camera lucida at a magnification of 44, and 100x, respectively, and then each area was measured with a digitizer system (Model G-6C, MUTOH Co., Tokyo). The measurement reliability was tested by comparing five subsequent analyses of a given figure by the same person (S.S.). Differences did not exceed 7%. Total volume of a given nucleus was estimated by multiplying the total area by the thickness of the section. Relative volume in percent was determined.
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Size of the eyeball

\[
\text{Total volume of a given nucleus} \times 100 \, (\%) \quad \frac{\text{Total volume of SGC}}{\text{Number of neurons in a given nucleus}}
\]

Number of neurons in a given nucleus was counted by means of an ocular grid \((25 \times 25 \, \mu m)\) which was superimposed on the microscopic image at randomly selected angles at a magnification of \(600 \times\). Then the cell density per unit area \((\text{cells}/50^2 \, \mu m^2)\) was determined.

Cell size was represented by the long and short axes of randomly selected neurons in a plane containing the cell nucleus. Statistical differences between the groups were examined by the Student's t-test.

Results

The unilaterally microphthalmic rats have one vestigial and one almost normally developed eyeball. The former eye completely lacks the optic nerve (Fig. 8A and B). The size of the latter eyeball was not significantly different from that of the normal (Fig. 1A). The bilaterally microphthalmic rat showed a thin but very apparent optic tract on both sides (about 40% of the width of the normal tract) (Figs. 1B and 8D). These non-retinal fibers of the optic tract form the commissure of Gudden. In the unilateral microphthalmia, the respective proportions between the optic tract contralateral (Fig. 8F) and ipsilateral (Fig. 8E) to the vestigial

Fig. 1. A: Size of the normally developed eyeball in unilaterally microphthalmic rats is not significantly different from that of the normal eyeball.

B: Relative thickness of the optic tract to the diameter of the anterior commissure; B/A \times 100 \, (\%). A: Diameter of the anterior commissure at its midline in a coronal section. B: Mean width of the optic tract measured on three successive coronal sections through 1 mm caudal to the posterior margin of the commissure.
eye were 49 and 76% of the width of the normal tract (Fig. 1B). This means that 9 and 36% of the width of the optic tract were occupied by the uncrossed and the crossed retinal fibers, respectively.

Superior colliculus:

In the unilaterally microphthalmic rats, conspicuous changes in the superficial layers of the superior colliculus (SCS) were found on the contralateral side to the vestigial eye, receiving uncrossed retinal fibers, but not on the ipsilateral SCS in which crossed retinal fibers terminate. Contralateral SCS showed a marked reduction in its size (Figs. 2 and 9B), and the midline between the two SCSs is displaced towards the contralateral side of the abnormal eye (Fig. 2). Small amounts of fibers were found in the medial 1/3 of the str. opticum (SO), with very few in the lateral 2/3.

The boundary between SCS and SCD (i.e., between SO and the intermediate gray layer) in the shrunken SC was determined by some medium-sized polygonal cells scattered in the ventral part of SO (Fig. 9B). The relative volume of SCS to SGC was decreased to about 50% of the normal on the contralateral side to the vestigial eye (p < 0.01), but not on the ipsilateral side (85% of the normal) (Fig. 3B). Cell density of contralateral SCS was markedly increased (133, 151 and 189% in the zonal, superficial gray and optic layers, respectively; p < 0.01), but not in the ipsilateral SCS (100, 102 and 108% in the respective three layers) (Fig. 4A). SCD showed no significant changes in not only the relative volume but also the cell density between the unilaterally microphthalmic and normal rats (Figs. 3B and 4B).

The str. griseum superficiale (SGS) of the normal rat consisted mainly of small neurons (long and short axes: 8–12 and 5–8 μm respectively) with fusiform to oval cell bodies, orienting their long axis perpendicularly to the collicular surface (Figs. 5 and 10A). In the mutant rat, contralateral SCS was packed with small and roundish neurons (diameter of 5–8 μm) (Figs. 5, 9B and 10B). Although the size of the cell nucleus was almost identical to the normal, the SCS neurons were found to reduce their perikaryal size. In addition, the narrow perikaryal fringe, having a small amount of Nissl
Substantia grisea centralis beneath the superior colliculus. Relative volume of SCS to SGC: Total volume of SCS/total volume of SGC \times 100(\%). Relative volume of SCD: Total volume of SCD/total volume of SGC \times 100(\%). Number in parenthesis indicates number of animals examined.

B: Histogram of the relative volumes of SCS, and of SCD, to SGC. The relative volume of unilaterally microphthalmic SCS contralateral to the vestigial eye decreased to 50\% of the normal, while no significant differences were observed in ipsilateral SCS. There are no significant differences in SCD between the normal and the mutant rats.

Fig. 4: Cellular density per unit area in each collicular layer. Note that three superficial laminae of unilaterally microphthalmic SC on the contralateral side show very high neuronal densities (130–190\% of the normal), while there are no significant differences on the ipsilateral side to the abnormal eye. No noticeable changes are present in the deeper collicular layers.
Fig. 5. Cell size distribution of SCS neurons in normal and unilaterally microphthalmic rats. Most of the SCS neurons on the contralateral side in the mutant animals have shorter long axes as compared with those in normal SCS, so as to present a roundish appearance. However, no marked changes are found in ipsilateral SCS.
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There were no significant differences in the cell size of SCD neurons between the microphthalmia and the normal.

Parabigeminal nucleus:
The normal parabigeminal nucleus (PB) is subdivided into three subgroups: the dorsal (PBd), middle (Bm) and ventral parts (PBv) (Fig. 10A). In spite of its reduced size in the unilaterally microphthalmic rats, PB is still normally subdivisible into the three subgroups (Fig. 10B). Relative volumes of each PB subgroup on both sides to SGC were significantly decreased in the mutant animals (PBm and PBv, p<0.01, PBd, p<0.05) as compared to the normal (Fig. 6A). The relative volume of PBv to SGC was markedly decreased in the contralateral in comparison to the ipsilateral side to the vestigial eye (26 and 61% of the normal, respectively, p<0.01). However, the relative volumes of PBd and PBm to SGC showed no significant differences between the ipsilateral (65 and 60% of the normal, respectively) and the contralateral sides (58 and 57% of the normal, respectively).

Cell density was slightly lowered in the PBm on both sides as compared with the normal (ipsi- and contralateral sides; 86 and 84% of the normal, respectively, p<0.01), but not in the PBd (ipsi- and contralateral sides; 88 and 90% of the normal) (Fig. 6B).

In normal rats, PBm was mainly composed of neurons with round to polygonal cell body with a long axis of 10 to 17 \( \mu m \) (Figs. 7 and 10C). On the other hand, neurons in PBd and PBv were primarily fusiform in shape (long and short axes: 10—15 and 7—12 \( \mu m \), respectively) (Figs. 7 and 10D). In the unilaterally microphthalmic rats, the size of neurons in the three PB subgroups on both sides was definitely reduced and their cell body showed a fusiform to oval shape (long and short axes: 7—15 and 5—10\( \mu m \), respectively) (Fig. 7). Perikarya of the PB neurons contained a small amount of Nissl granules and had a pale appearance. The size of the cell nucleus was found to be nearly the same as that of the normal.

In all the mutant animals examined in this study, no other malformations were observed except for changes in relay nuclei in the central visual system associated with altered retinal axonal ingrowth.
Discussion

A study of the organogenesis of the microphthalmic eye revealed that growth inhibition of the inner layer of the optic cup at the 13th embryonic day resulted in complete loss of the optic nerve. The present observation disclosed that in the unilaterally microphthalmic rats uncrossed and crossed retinal fibers from the normally developed eye occupied 9 and 36% of the width of the optic tract, respectively. The ratio of uncrossed to crossed optic fibers is, therefore, estimated at 1 to 16. So it is assumed that in the mutant animals about 6% of retinal fibers entered into the ipsilateral optic tract as an uncrossed retinofugal projection. The proportion of uncrossed to crossed retinal fibers falls into the range of values that can be seen in rodents (5–10%).

The present morphological observation exhibited that the volume of SCS contralateral to the vestigial eye decreased to 50% of the normal, while its cell density increased up to 130–190%. On the other hand, ipsilateral SCS, receiving crossed optic fibers, and SCD on both sides of the mutant rats remained unchanged. It is well documented that the removal of one eye in newborn animals results in a marked reduction in the
volume of contralateral SCS\textsuperscript{6,7,9,15,27,29,37,41}. Tsang (\textsuperscript{37}\textsuperscript{41}) and Lund et al. (\textsuperscript{73}\textsuperscript{27}) reported respectively that in adult rats (4–19 months old) SC contralateral to the eye removed at birth shrunk by 45 and 55% of that of the unoperated rats. On the other hand, DeLong and Sidman (\textsuperscript{62}\textsuperscript{7}) observed a mild diminution of the collicular size (73–78% of the control SC) in 1.7 months-old mice, whose eyes had been unilaterally enucleated at birth. Tsang (\textsuperscript{37}\textsuperscript{41}) pointed out that the atrophy was proportional to the period after deaffrentation. It is, therefore, reasonable to consider that the survival period of 4–19 months (materials of Tsang\textsuperscript{44}, Lund et al.\textsuperscript{27} and the present study) induced a greater reduction of SC volume than that of 1.7 months (DeLong and Sidman\textsuperscript{7}). Although Godement et al. (\textsuperscript{80}\textsuperscript{9}) have described a marked reduction of size in both the lateral geniculate nucleus and SC of the unilaterally anophthalmic mice, quantitative morphological changes in the hereditary unilateral microphthalmia are hereby presented for the first time.

Many reports have been issued on the remarkable expansion of the terminal field in the uncrossed retinal projection following prenatal or neonatal removal of one eye \textsuperscript{5,9,10,17–19,21,22,27,28,23}. Godement et al. (\textsuperscript{80}\textsuperscript{9}) and Tokunaga et al. (\textsuperscript{85}\textsuperscript{39}) observed enlarged distribution of the uncrossed retinal pathway in the hereditary unilaterally anophthalmic animals, respectively. Rats with unilateral eye defects induced by tripan-blue administration\textsuperscript{23} also showed the expansion of the ipsilateral optic projection throughout the dorsal lateral geniculate nucleus (CGLd) and SCS. The relative volume of SCS to SGC in the bilaterally microphthalmic rats decreased to 35% of the normal\textsuperscript{40}. Therefore, it is reasonable to conclude that the expanded ipsilateral retinal projection caused a slight increase in the volume of SCS in the unilateral microphthalmia (about 15% of the normal). Cell size and morphological changes in neurons of SCS contralateral to the vestigial eye were similar to those found in the bilateral microphthalmia\textsuperscript{40}.

Sugita et al. (\textsuperscript{83}\textsuperscript{35}) demonstrated in their WGA–HRP study that neurons in str. griseum superficiale (SGC) projected their fibers to CGLd and those in the str. opticum (SO) to the lateroposterior thalamic nucleus (LP). This segregation of the tecto-thalamic projections was apparently preserved in the bilaterally microphthalmic rats\textsuperscript{36}. However, the amounts of the CGLd- and the LP-projection neurons decreased to 3 and 50% of the normal, respectively, in the mutant animals\textsuperscript{36}. As cell density was not so greatly changed between the unilateral and the bilateral microphthalmia\textsuperscript{40}, the slight increase of volume of SCS (15% of the normal) contralateral to the anomalous eye in the present materials suggests that the cell population itself had definitely increased in the shrunken SCS. Therefore, it is possible to assume that most of the SCS neurons are shared in forming the efferent projections to the thalamic nuclei.

Goodman et al. (\textsuperscript{73}\textsuperscript{8}) pointed out two routes of the ipsilateral tectal afferents: a rostral and a lateral route. The rostral route passes caudally through the nucleus of the optic tract and enters the rostral aspect of SC to distribute in its medial half. The lateral route, on the other hand, runs in the brachium of SC and enters the lateral aspect of SC to distribute in its lateral half. Unilateral microphthalmic SCS contralateral to the vestigial eye showed definite fiber bundles in the medial 1/3 of SO, but very few in the lateral 2/3. This finding suggests that in the unilaterally microphthalmic rats the expanded uncrossed retinotectal fibers arrive at SC mainly via the rostral route.

It is well established in many experimental animals that PB is reciprocally and
almost restrictively connected with SC\textsuperscript{11,12,16,25,34,42}. The present study revealed that PB in the unilaterally microphthalmic rats was subdivided normally into the 3 subgroups. The relative volumes of each subgroup on both sides to SGC decreased to 57–65\% of the normal, except for that of PBv on the contralateral side to the abnormal eye (26\%). On the other hand, no subdivisions were observed in the bilaterally microphthalmic PB\textsuperscript{40). The reduction of the relative volume and of the cell density were much more severe in the bilateral (30 and 75\%, respectively)\textsuperscript{40) than in the unilateral microphthalmia.  

Altman and Bayer (81)\textsuperscript{1) studied the cytogenesis of the rat brain stem structures with the \textsuperscript{3}H-thymidine autoradiographic method. Neurons of PB were produced between 13- and 15-day embryonic stages with a peak on the 14th day\textsuperscript{11}. The generation of SCS neurons begins on the 11th or 12th embryonic day and ends by the 18th embryonic day\textsuperscript{1,3,7,39). The crossed and uncrossed retinal fibers of the rat are reported to reach SCS by the 16th and 17th embryonic day, respectively. Furthermore, Stevenson and Lund (82)\textsuperscript{34) demonstrated a corresponding intra-SCS distribution pattern of projection fibers from contralateral PBm and from the ipsilateral retina of rats with unilateral eye removal at birth. These findings indicate not only developmental but also some functional independence between SCS and PB. Therefore, the mild changes in the unilateral microphthalmic PB can be attributed to the existence of expanded distribution of the uncrossed retinotectal fibers in SCS.  

The present finding showing severe reduction in the volume of PBv on the same side as the collapsed SCS is difficult to explain. Tokunaga et al. (85)\textsuperscript{40) pointed out that the uncrossed retinal fibers in the unilaterally microphthalmic rats were densely distributed to the entire depth of the medial 2/3 of SCS, while scarcely to the lateral 1/3. Their result may suggest a fiber connection between PBv and the lateral 1/3 of SCS. On the other hand, a topographical study on the tecto-PB projection in rats\textsuperscript{25) seems to eliminate such a possibility. Further research needs to be done on the connectivity of the parabigeminal nucleus.

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References

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PLATES
Plate I

Explanation of Figures

Plate I

Fig. 8. Ventral view of a normal (A), and a unilaterally microphthalmic rat brain (B). Arrow in B indicates the optic nerve from a normally developed eye.

C-D: The optic tract of a normal (C), a bilateral (D) and a unilateral microphthalmia (E and F). Very thin optic tract (arrows in D) is definitely present in the bilaterally microphthalmic rat, in spite of its complete loss of the optic nerve. E: Ipsilateral side to the vestigial eye. The tract contains crossed retinal fibers from a normally developed eye. F: Contra lateral optic tract to the anomalous eyeball contains uncrossed retinal fibers. *: nucleus supraopticus. Klüver-Barrera staining. Bar indicates 500 μm.
Plate II

Fig. 9. Cytoarchitecture of the unilaterally microphthalmic superior colliculus (SC) on the ipsilateral (A) and contralateral (B) sides to the vestigial eye. No remarkable changes were found between the ipsilateral side of mutant rats and the normal. On the other hand, the superficial layers (I to III) of the contralateral SC in unilateral microphthalmia are definitely collapsed and packed densely with small roundish neurons. The deeper tectal layers (IV to VII) of the mutant rats show no marked changes on both sides. Klüver-Barreta staining. Bar indicates 500 μm.

Lateral microphthalmia are definitely collapsed and packed densely with small roundish neurons. The deeper tectal layers (IV to VII) of the mutant rats show no marked changes on both sides. Klüver-Barrera staining. Bar indicates 500 μm.

Plate III

Fig.10  A and B. Neurons of the zonal (Z) and the superficial gray layers (GS) on the ipsilateral (A) and contralateral (B) sides of a unilaterally microphthalmic rat. Many neurons in the ipsilateral superficial gray lamina (SGC) and characterized by a fusiform cell body with its long axis oriented perpendicularly to the tectal surface, as seen in the normal. However, the contralateral SGC is densely packed with small round cells and the intralaminar arrangement of the SGS neurons is highly disordered. Klüver-Barrera staining. Bar indicates 100 μm.

C and D: The parabigeminal nucleus (PB) of a normal (C) and a unilaterally microphthalmic rat (D). Three subdivisions of PB, as shown in the normal (C), are also identifiable in the mutant rat. D: Dorsal subdivision of PB, M: Middle subdivision of PB, V: Ventral subdivision of PB. Klüver-Barrera staining. Bar indicates 100 μm.