Behavioral/Systems/Cognitive

# Axonal Branching Patterns as Sources of Delay in the Mammalian Auditory Brainstem: A Re-Examination

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In models of temporal processing, time delays incurred by axonal propagation of action potentials play a prominent role. A pre-eminent model of temporal processing in audition is the binaural model of Jeffress (1948), which has dominated theories regarding our acute sensitivity to interaural time differences (ITDs). In Jeffress' model, a binaural cell is maximally active when the ITD is compensated by an internal delay, which brings the inputs from left and right ears in coincidence, and which would arise from axonal branching patterns of monaural input fibers. By arranging these patterns in systematic and opposite ways for the ipsilateral and contralateral inputs, a range of length differences, and thereby of internal delays, is created so that the ITD is transformed into a spatial activation pattern along the binaural nucleus. We reanalyze single, labeled, and physiologically characterized axons of spherical bushy cells of the cat anteroventral cochlear nucleus, which project to binaural coincidence detectors in the medial superior olive (MSO). The reconstructions largely confirm the observations of two previous reports, but several features are observed that are inconsistent with Jeffress' model. We found that ipsilateral projections can also form a caudally directed delay line pattern, which would counteract delays incurred by caudally directed contralateral projections. Comparisons of estimated axonal delays with binaural physiological data indicate that the suggestive anatomical patterns cannot account for the frequency-dependent distribution of best delays in the cat. Surprisingly, the tonotopic distribution of the afferent endings indicate that low characteristic frequencies are under-represented rather than over-represented in the MSO.

### Introduction

Coincidence detectors and time delays are common ingredients in models of temporal processing. In such models, neural equivalents of cross-correlation or autocorrelation are implemented by comparing spike trains with delayed versions of themselves or with other spike trains. A pre-eminent example is in binaural hearing, which affords the detection of minute time differences in the sound signals to the two ears [interaural time differences (ITDs)] that contain information about the spatial properties of sound sources. Neuronal ITD sensitivity in mammals originates in the medial superior olive (MSO). For >50 years, one model (Jeffress, 1948) has dominated the discussion of the neural mechanisms of ITD discrimination and its relationship to MSO activity. In this model, a binaural cell shows a "best delay": it is maximally active when the external acoustic delay (the ITD) is compensated by an internal delay, so that the inputs from left and right ear to the cell are coincident. Classic studies of ITD sensitivity (Rose et al., 1966; Goldberg and Brown, 1969; Yin and Chan, 1990) provided evidence for coincidence in accord with the Jeffress (1948) model (Joris et al., 1998; Joris, 2006). Jeffress hypothesized the source of internal delays as axonal length differences. He proposed that afferent axons are systematically arranged with length gradients in opposite directions for ipsilateral (Ipsi) and contralateral (Contra) inputs to an array of coincidence detectors. The result would be a systematic gradation in best delay along the binaural nucleus and the conversion of ITDs into a spatial activation pattern along the array.

In mammals, only two studies have examined the existence of delay lines anatomically. Smith et al. (1993) physiologically characterized and then labeled single afferent axons to the MSO from spherical bushy cells of the anteroventral cochlear nucleus (AVCN). They concluded that the projections to the contralateral MSO, but not the ipsilateral MSO, display a rostrocaudal (RC) delay line configuration. Beckius et al. (1999) labeled small populations of AVCN neurons with an extracellular deposit, and selected and reconstructed individual axons. Their results were similar to those of Smith et al. (1993) with the additional finding of a length gradient running in the opposite (caudorostral) direction in some ipsilateral projections. Physiologically, MSO neurons show a rostrocaudal gradient in ITD tuning consistent with the anatomical predictions (Yin and Chan, 1990).

Despite the anatomical and physiological support, the Jeffress (1948) model is no longer universally accepted for mammals (Palmer, 2004; McAlpine, 2005; Palmer and Kuwada, 2005; Joris

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DOI:10.1523/JNEUROSCI.5175-10.2011 Copyright © 2011 the authors 0270-6474/11/313016-16\$15.00/0 and Yin, 2007), while it remains unchallenged in the barn owl (Wagner et al., 2007). McAlpine et al. (1996, 2001) discovered a relationship between best delay and frequency tuning in binaural neurons, which is not necessarily contradictory to the Jeffress (1948) model but certainly is not predicted by it. Moreover, the anatomical evidence for delay lines in the mammal has been questioned (Grothe, 2003). Our purpose here is to re-examine the data of Smith et al. (1993) quantitatively and to examine the anatomical data in the light of the more recently discovered relationship between ITD tuning and frequency tuning.

#### Materials and Methods

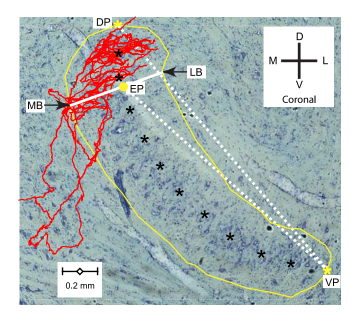
*Material.* The surgical procedure, stimulus delivery, and physiological and histological methods are described in the original study (Smith et al., 1993) and are only briefly outlined here. We will focus on the quantitative, computerized processing of the fibers labeled.

The ventral brainstem was exposed in cats of either sex, anesthetized with sodium pentobarbital. Sharp electrodes filled with either a 5% horseradish peroxidase (HRP) (type VI, Sigma) or 2% neurobiotin (Vector Laboratories) solution in 0.5 M KC1 were advanced into the trapezoid body while presenting calibrated search tones. When a fiber was isolated, the excitatory ear, the characteristic frequency (CF), spontaneous activity, and threshold at CF were determined by an automated threshold tracking program. Poststimulus time histograms for short (25 ms) tones at CF were then obtained at different sound pressure levels (in 10 or 20 dB steps) usually up to 50–60 dB above threshold. Intra-axonal impalement was achieved with current pulses or electrode advancement, followed by confirmation of the physiological responses. HRP or neurobiotin was then injected by iontophoresis. After 24–36 h, a lethal dose of sodium pentobarbital was given and the animal was perfused transcardially with phosphate-buffered glutaraldehyde/paraformaldehyde fixative.

For preparation of HRP-filled axons, frozen or vibratomed 60 or 70  $\mu$ m sections of the brainstem were processed using the DAB nickel–cobalt intensification method (Adams, 1981). For preparation of neurobiotin-filled cells, the sections were washed in 0.1 M phosphate buffer, incubated for 20 min in 0.5%  $\rm H_2O_2$ , rinsed in PBS, and left overnight in phosphate buffer containing avidin-biotin-HRP reagent (ABC kit, Vector Laboratories). The sections were then rinsed in buffer, treated with DAB (as above), mounted on glass slides, counterstained with cresyl violet, and coverslipped.

Reconstructions and analysis. Because the axons typically extended over many coronal sections, detailed analysis and three-dimensional reconstructions required computer reconstruction. We used a Neurolucida system (Microbrightfield) that included an Olympus BX61 light microscope, a video camera (MBF-CX9000, Microbrightfield), a motorized stage controller (MAC5000, Ludl Electronic Products), and a personal computer. Axons were traced and digitized directly on the microscope using a UPlanFLN 40× objective. The (x,y,z) positions accompanied with axon diameter were collected every 1.9 µm on average (538 points/ mm). Section thickness measured with the microscope z-axis ranged from 20 to 25  $\mu$ m. Corrections were made for shrinkage along this axis (i.e., the rostrocaudal direction) according to the section thickness specified during cutting and the thickness measured in the microscope. Estimates of axon diameter were made by adjusting the diameter of a circular cursor to the width of the labeled axon and do not include the myelin sheath. For several reasons, stated in Smith et al. (1993), these measurements should be regarded as coarse estimates.

The furthest distal points that could be traced were defined as endpoints (EPs; for ease of reading we use the full words in the text and the abbreviations in figures). These points are not identical to terminals: the darkness of the label typically decreased with distance from the injection site and toward the middle of the sections, so that the finest processes could not always be followed to unambiguous terminations. Moreover, all of the reconstructions analyzed here were done with the light microscope, so we cannot be sure that the afferent endpoints terminate at a synapse. However, we have analyzed a small number of the labeled axons with the electron microscope, and the afferent terminals were confirmed to be presynaptic (Smith et al., 1991, 1993). Also, comparison with



**Figure 1.** Illustration of measurements taken within sections. Coronal view of the contralateral projection of a traced fiber (CF, 1345 Hz) superimposed on a Nissl-stained single section, taken at the coronal level of the branch EP, indicated with the yellow dot. Yellow line, Contour of MSO on the section. Yellow asterisks, Most DP and most VP along the long axis of the MSO within the section, which is indicated with black asterisks. Drawing a line through the EP, which is approximately perpendicular to the long axis of the MSO, the arrows show the most medial and lateral points (MB and LB) of the MSO contour on this line. The two other lines (dotted white) indicate the Euclidian distances between EP and VP, and between VP and the DP of this section. Scale bar, 200 μm. Directions are indicated by the cross. D, Dorsal; V, ventral; M, medial; L, lateral.

Beckius et al. (1999) (see Results) suggests that our tracings must have covered the axons almost in their entirety in many cases. Importantly, the sheet-like mammalian MSO has limited thickness, and the delay line configuration is thought to span the rostrocaudal dimension: the main contributors to differential length are not the terminal axonal segments but rather the more proximal main stem of the axon, which gives off branches at different rostrocaudal positions (see Figs. 2*C*, 3). Assessment of collateral length differences along the rostrocaudal axis thus does not hinge on complete filling to the axon terminals.

We defined several markers to take measurements for numerical analysis. Figure 1 illustrates markers defined within single sections. The yellow line traces the MSO contour, and the black asterisks its long axis within the section. The yellow asterisks indicate the dorsal pole (DP) and ventral pole (VP). These are not necessarily literally the extreme dorsal and ventral points of the MSO outline in that section. For example, the ventral pole in Figure 1 is not the most ventral point of the MSO. Rather, dorsal pole and ventral pole are the extreme ends of the—often curved long axis of the MSO within the section. The yellow disk marks one endpoint (EP). To express the dorsoventral location of an endpoint within the MSO, we measured its Euclidian distance to the ventral pole, which we normalized to the Euclidian distance between dorsal pole and ventral pole (Fig. 1, dotted white lines). Normalization facilitates comparison across animals as it reduces variance resulting from differences in absolute dimensions of the MSO. Values close to 1 indicate an endpoint located close to the dorsal pole; values close to 0 indicate a position close to the ventral pole. We also measured the mediolateral location of endpoints based on their distance to the edge of the MSO outline, taken at the intersection of the outline with a line (Fig. 1, solid white line) that is approximately orthogonal to its long axis. These intersections give two measurements for each endpoint: the medial border (MB) and the lateral border (LB). These latter measurements are inherently coarse, particularly for endpoints close to the dorsal or ventral pole of the MSO. Again, we normalized the midline (ML) position as the (Euclidian distance between endpoint and medial border)/(Euclidian distance between lateral border and medial border). When this value is close to 1, the endpoint is located close to the lateral border.

Measurements taken across sections are illustrated in Figure 2 in the coronal (Fig. 2A), horizontal (Fig. 2B), and parasagittal (Fig. 2C) planes, for the same fiber used in Figure 1. First, the most rostral and most caudal sections where the MSO could be discerned are identified. The geometrical centers of the MSO outlines in these sections (Fig. 2A, green and magenta) were designated as the rostral pole (RP) and caudal pole (CP). The normalized RC distance of an endpoint relative to these poles was measured as (Euclidian distance between endpoint and rostral pole)/(Euclidian distance between caudal pole and rostral pole), illustrated with the dotted lines (Fig. 2B, C). An endpoint close to the caudal pole thus gives an RC value close to 1. For contralateral projections, we also marked the point where the axon crossed the ML. Finally, for both ipsilateral and contralateral projections we marked the first branch point (FB) (i.e., the last point of the axon that is common for all branches to the ipsilateral or contralateral MSO) (Fig. 2B, C). The axonal length from midline or first branch point to each endpoint was calculated as the sum of the distances between adjacent (x,y,z)coordinates in the course of the axon to that endpoint.

For the sake of brevity, we use the shorthand "delay line" to refer to a morphological pattern of systematic length differences in axon collaterals. We caution that the term inherently refers to a functional concept that can only be validated with physiological measurements—a point to which we will return in Discussion.

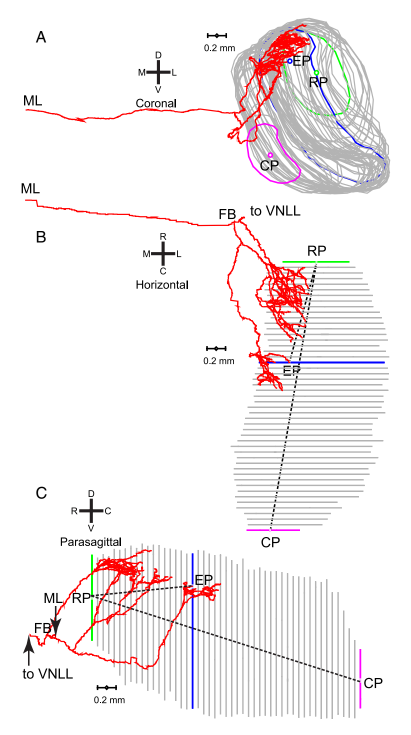
# Results

The material we analyzed here consists of 14 labeled fibers, obtained from 12 animals. We reconstructed the 16 projections (9 contralateral, 7 ipsilateral) reported in Smith et al. (1993). For only two fibers (with CFs of 1498 Hz and 2470 Hz) were both contralateral and ipsilateral projections sufficiently labeled for reconstruction. A qualitative description of the course and projections of the fibers is provided in Smith et al. (1993) and is summarily illustrated here for several fibers. We identify projections with the same number, symbol, and color scheme in all figures, where relevant.

#### Contralateral projections

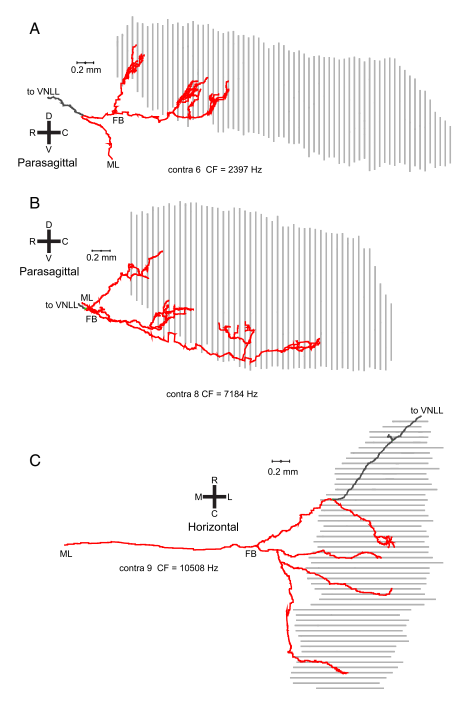
Nine contralaterally injected fibers were well labeled and used for the analyses. The axon in Figure 2 crossed the midline at a

level rostral to the MSO and projected multiple collateral branches caudally before heading rostrally toward the ventral nucleus of the lateral lemniscus (VNLL). The collateral projections span approximately half of the rostral-caudal extent of the MSO nucleus with a ladder-like branching pattern that is particularly clear in the parasagittal view (Fig. 2C). Similar inner-



**Figure 2.** Example of the contralateral MSO innervation by one axon (same axon as in Fig. 1). **A–C**, Coronal (**A**), horizontal (**B**), and parasagittal (**C**) views are shown. The RP and CP were defined as the geometrical center of the most rostral (green line) and most caudal (magenta line) section, respectively. The rostrocaudal position of endpoints was quantified by measuring the Euclidian distance (dashed line) between RP and EP, and was normalized to the distance between RP and CP. The endpoint illustrated is the same one as in Figure 1; the MSO contour outlined in Figure 1 is shown in blue here. ML indicates the point at which the axon crosses the midline. CF of this fiber was 1345 Hz. A computer reconstruction of this fiber was also shown in Smith et al. (1993, their Fig. 4).

vation patterns are shown for three fibers in Figure 3. The axons in Figure 3, *A* and *B*, also cross the midline in a rostral position and give caudally directed branches over a more restricted (Fig. 3*A*) or less restricted (Fig. 3*B*) rostrocaudal distance than in Figure 2. The axon in Figure 3*C*, shown in horizontal view, crosses the midline at a more caudal level. It shows three branches that are directed to the central and caudal MSO, as well as one branch



**Figure 3.** Delay line configuration in 3 contralaterally projecting fibers. **A**, **B**, Projections on a parasagittal plane. **C**, Projection on a horizontal plane. CFs are indicated. Branches that end within the MSO are indicated in red.

that is directed rostrally and does not seem to fit the caudally directed delay line patterns of the fibers discussed so far. A similar case is shown in parasagittal view in Figure 4A. The axon also crosses the midline at a rostrocaudal position that bisects the MSO in approximately equal halves. It shows a number of caudally directed collateral branches with increasing length, but also a branch that innervates the rostral half of the MSO and continues on toward the VNLL. Fibers that reach the MSO at an even more caudal position are shown in Figure 4, *B* and *C*. The fiber in Fig. 4*B* has a rostrocaudally restricted field. The axon in Fig. 4*C* shows no obvious bias toward increasing length for more caudally projecting collaterals.

#### **Ipsilateral projections**

Seven ipsilaterally injected fibers were adequately labeled for our analyses. Because the pattern of the ipsilateral projections is more complicated than that of the contralateral projections described above, we show the projections in different views. The fiber illustrated in Figure 5 had several collaterals without an obvious delay line pattern. Two types of collateral projections can be distinguished: forward and backward. The coronal view (Fig. 5A) shows one branch coursing along the lateral aspect of the MSO (forward innervation), consistent with the known segregation of ipsilateral and contralateral inputs to the lateral and medial dendrites of MSO neurons, respectively (Stotler, 1953). Two more collaterals only branch off from the main axon after it has crossed underneath the MSO, to then loop back through the MSO to innervate the same re-(backward innervation). These branches have their origins medial to the ipsilateral MSO. All of the seven ipsilateral projections had "backward" components; four of the seven ipsilateral projections had "forward" components (Fig. 10B, summary).

Figure 6 shows another example of an ipsilateral projection. In this case, the forward and backward innervation contribute endpoints that are located at different rostrocaudal positions (Fig. 6B, C), where the backward branches innervate a more rostral part of the MSO than the forward branches. The horizontal and parasagittal views suggest that this fiber has greater length in its caudal than in its rostral innervation, but the complex course of the branches in the three dimensions precludes a simple visual assessment. In fact, length measurements reveal greater length in the rostral than in the caudal innervation (see Fig. 8D, green inverted triangles).

# Dendrograms

Because branches often extensively overlap in the projections of the computer reconstructions in Figures 1–6, these views do not give an adequate representation of

the complexity of branching. Figure 7 gives an overview, for the most completely filled fibers, of the sequence of branchings in contralateral (left column) and ipsilateral (right column) projections. The horizontal axis indicates the length of segments bordered by midline, branch points, or endpoints. The vertical axis only serves to offset the different segments after branching: it does not have the physical meaning of length, and there is no vertical sorting according to position in the MSO. The dendrograms therefore provide information about branch sequence and length (abscissa), but they contain no anatomical directional information (e.g., lateral vs medial, dorsal vs ventral, rostral vs caudal). There is obviously profuse branching in both contralat-

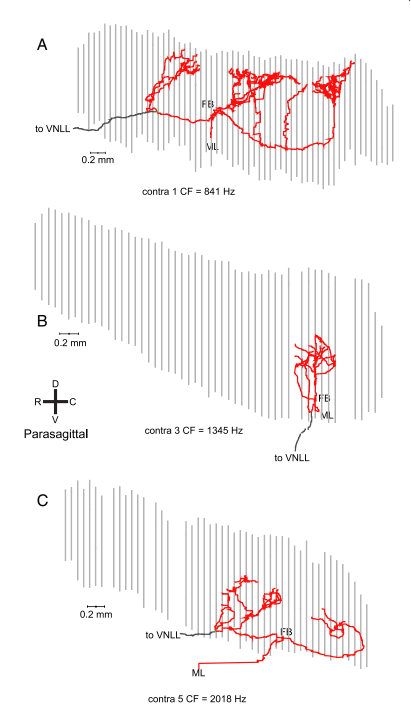
eral and ipsilateral projections. Considering the branching after the first branch point, the ipsilateral projections show longer (>1 mm; see scale bar in Fig. 7A) segments than the contralateral projections before further branching. These long segments are the stems of the forward and backward branches, for example, in Figures 5 and 6. As a result, the longest distances between first branch point and endpoint are found in ipsilateral rather than in contralateral projections. Comparison of the horizontal location of the endpoints within each dendrogram indicates the range of overall length differences within projections. It is of note that this range is a few millimeters for most projections (Fig. 7A-C,F), and that the largest ranges occur in ipsilateral projections (Fig. 7*D*,*E*).

# Distribution of endpoints in the parasagittal plane

Figure 8 shows the distributions of endpoints for contralateral (left column) and ipsilateral (right column) projections. Endpoints from one fiber are identified with the same colored symbol. The Figure 8 caption ranks the fibers by increasing CF. The two fibers for which both contralateral and ipsilateral projections were reconstructed are represented by the same symbols in left and right column: projections Contra 4 and Ipsi 2 (light blue inverted triangle, CF = 1498 Hz) and projections Contra 7 and Ipsi 5 (solid green triangle,  $CF = 2470 \,\text{Hz}$ ). In all panels, the abscissa is the normalized rostrocaudal position of the endpoints, measured as illustrated in Figure 2 (virtually identical results are obtained if the distance along the z-axis, i.e., the distance between coronal sections, is used). The ordinate of the top two panels is the normalized dorsoventral position of the endpoints, measured as illustrated in Figure 1. These two panels therefore represent a standardized side view of the MSO. As expected, the endpoint positions are orderly arranged with low-CF fibers most dorsal and high-CF fibers most ventral (this is further analyzed in Fig. 9). The endpoints of a given fiber tend to be distributed in an elongated rostrocau-

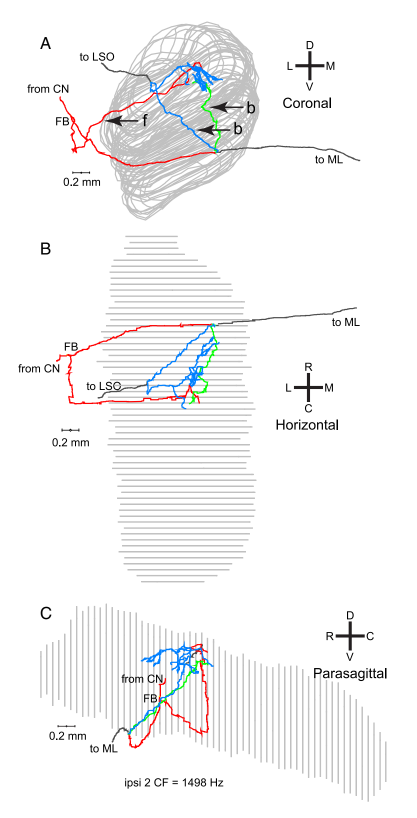
dal strip. This is particularly the case in contralateral projections (Fig. 8A) and less so in ipsilateral (Fig. 8B) projections, which can be elongated (projection 1), restricted (projection 7), or patchy (projection 6). In contralateral fibers, seven of the nine fibers showed a broad rostrocaudal extent, spanning 40–60% of the nucleus, but importantly none of the fibers spans the entire nucleus.

The bottom panels show the key measurement: the axonal length accumulated between the first branch point for the projection from that side, and the endpoints. For contralateral projections (Fig. 8*C*), this length tends to increase for endpoints positioned more caudally in the nucleus. The solid lines are linear



**Figure 4.** Parasagittal view of 3 more contralateral projections (*A–C*, with CFs indicated below each reconstruction) with a less clear-cut delay line configuration. In all cases, the FB is located near the center of the rostrocaudal range of EPs. The orientation shown in *B* applies to all panels. The color convention is as in Figure 3.

regressions; the dashed (black) line is the diagonal of equality. A "perfect" Jeffress delay line configuration would generate an endpoint distribution parallel to this line: the extra length incurred (ordinate) would equal the distance traveled rostrocaudally (abscissa). For six of nine contralateral fibers, the linear regression was significant (p < 0.05), with a correlation coefficient > 0.80 in four of these six cases. The mean slope for the six cases was 0.91 (range, 0.54–1.65). The slope of the regression line was negative in one fiber (#3), for which the endpoints were distributed over a narrow rostrocaudal range. In two fibers (#1 and #9), the relationship was nonmonotonic, with a rostral patch of endpoints of



**Figure 5.** *A–C*, Coronal (*A*), horizontal (*B*), and parasagittal (*C*) views of a reconstructed projection of one fiber to the ipsilateral MSO. The coronal (*A*) and horizontal (*B*) views clearly show the two kinds of ipsilateral branches. One branch originates lateral to the MSO and projects forward (*f*, red) to it. Two branches originate after the axon has crossed the plane of the MSO and looped back (b, blue and green) to innervate the same region.

larger length than the endpoints just caudal to it. These are fibers whose axon crosses the midline at a rather caudal position as illustrated in Figure 3C (fiber #9, CF = 10,508 Hz) and Figure 4A (fiber #1, CF = 841 Hz).

The topography of the length distributions of ipsilateral projections (Fig. 8D) is strikingly inhomogeneous and different from the contralateral ones. The length gradient is mostly negative (i.e., in the opposite direction than the contrateral gradient) (Fig. 8, note the difference in ordinate scale between C and D). For comparison, the negatively sloped diagonal (large dashes) is shown, which shows the slope of a perfect Jeffress delay line configuration in the direction opposite to the one dominant in contralateral fibers. Linear regressions were significant for all seven fibers (p < 0.01). For two fibers (#6 and #7), the number of endpoints is small. Surprisingly, the slope is positive in three fibers.

Particularly for cases where the distribution of endpoints is restricted rostrocaudally (Fig. 8 *D*, fiber #7, 8 *C*, fibers #3 and #4), it is questionable that the presence of a slope in the linear regressions is of functional significance. Nevertheless, overall the length gradients are in accordance with the scheme proposed by Jeffress (1948), with a majority of contralateral projections showing a rostrocaudal increase in collateral length, and a majority of ipsilateral projections showing an inverse gradient.

Interestingly, fibers tend to project to either the rostral or caudal half of the MSO, particularly on the contralateral side (Fig. 8*A*, *B*, with the exception of the fibers indicated by the empty red triangles). Moreover, the length gradients most consistent with a Jeffress delay configuration are found in the rostral half, both in the contralateral and in the ipsilateral branches.

Note that only the slope of the regressions is relevant to the issue of delay lines, not the vertical offset on the ordinate. This offset reflects our (arbitrary) choice of the first branch point as the point of reference for the length measurement and is affected by the absolute length of the collateral branches. A branch that is longer than the rostrocaudal MSO dimension has endpoints with ordinate values >1, as is the case for several contralateral and many ipsilateral projections. However, addition of an identical length of axon to all collateral branches (see below, Estimates of rostrocaudal axonal delay, on calculation of differences in path length between ipsilateral and contralateral projections) moves all data points of a projection up along the ordinate without affecting the presence or absence of a rostrocaudal delay.

# Tonotopic distribution along the dorsoventral axis

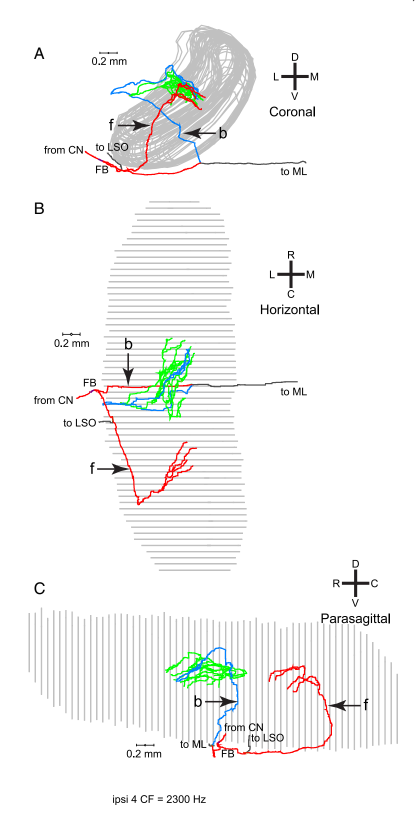
The rostrocaudal clustering of endpoints (Fig. 8 *A*, *B*) allows us to project a map of input CFs onto the MSO. Figure 9 shows the

relationship between fiber CF and the normalized dorsoventral position of all endpoints. The endpoints contributed by one fiber are shown as a column of circles in Figure 9A, color coded for laterality. The asterisks show data points taken from the physiological data of Guinan et al. (1972, their Fig. 17). The solid circles and line show the position of the borders drawn in the summary tonotopic map of Guinan et al. (1972, their Fig. 21), at CFs of 1000, 4000, 10,000, and 20,000 Hz. There is a reasonable agreement between our data and those of Guinan et al. (1972). Much to our surprise, however, there is an upward convexity in the CF-position relationship. If low CFs were to take up a disproportionate amount of space of the MSO, relative to a logarithmic CF distribution, the relationship should be concave rather than convex: if one were to progress from dorsal MSO in a ventral direction and plot CF, there would initially be "slow" progression in CF, which would accelerate as one comes closer to the ventral pole. This is clearly not the case.

Because the cochlear map deviates from a logarithmic relationship at low CFs, we incorporate the CF-position relationship for the cochlea in the abscissa of Figure 8B, using Greenwood's equation (Liberman, 1982; Greenwood, 1990). The means and SDs of endpoint positions and the data points of Guinan et al. (1972) are replotted directly on a cochlear distance scale in Figure 9B. The data points again show an upward convexity. Predictions of the MSO CF-position relationship are now straight lines (green). The solid green line shows the relationship expected if all cochlear CFs were represented proportionally in the MSO: at low CFs, the data points tend to lie above this line. Another factor to take into account is that not all cochlear CFs appear to be represented in the MSO: the highest CF reported by Guinan et al. (1972) was 22 kHz. The dashed green line in Figure 8 shows the CF-position relationship expected if the cochlea were proportionally represented in the MSO except for its base, so that CFs >22 kHz are absent (~5 mm of the extreme base). Clearly, consideration of a limited CF representation does not alleviate the upward convexity. The prediction based on a limited CF range follows the slope of the map Guinan et al. (1972) reasonably well, but again the representation of the cochlear apex is com-

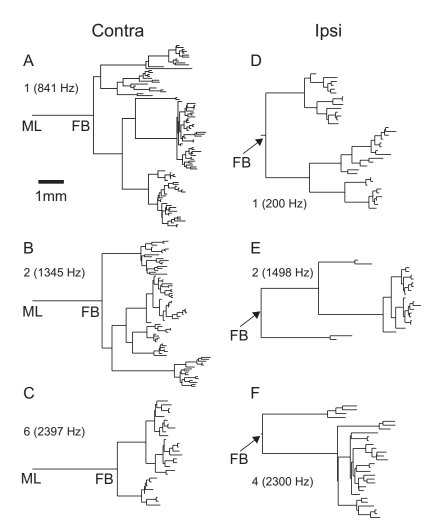
pressed for our data, with the majority of low-CF data points lying above the dashed line.

The MSO tends to show curvature in the coronal plane (Figs. 1, 2, 6), and this curvature can be more pronounced at the dorsal



**Figure 6.** Another example of an ipsilateral MSO projections of one fiber. **A**, This fiber ran underneath the MSO (coronal view) and also formed forward (f) and backward (b) projecting branches. **B**, **C**, The horizontal (**B**) and parasagittal (**C**) views reveal that the branches innervate different rostrocaudal portions of the MSO. The backward branches (green and blue) innervated a more rostral portion of the MSO and covered more length of axon from FB than the forward branches.

pole (Fig. 1). Because our measure of dorsoventral location does not take curvature into account, we recalculated the dorsoventral positions in Figures 8, 9, and 10 taking the Euclidean distance of the endpoint to the ventral pole (as before), but now normalized



**Figure 7.** *A, B,* Dendrograms showing the branching pattern and length of axonal segments for three contralateral (*A*) and three ipsilateral projections (*B*). The scale bar in *A* applies to *A*–*F*. The horizontal dimension represents the axonal length of branch segments; the vertical dimension is only used to offset these segments and has no meaning with regard to length or order. Fiber number and CF are indicated for each projection. Cross-linking to previous figures with computerized reconstructions is as follows: *A* = Figure 4 *A*; *B* = Figures 1 and 2; *C* = Figure 3*A*; *D* = Figure 5 of Smith et al. (1993); *E* = Figure 5; *F* = Figure 6.

to the sum of the distance of the endpoint to the ventral and dorsal poles (i.e., the normalization was now to the sum of the line segments EP–VP and EP–DP). This recalculation caused only minor shifts in ordinate position and did not remove the bias of low-CF data points above the green lines in Figure 9*A*, *B*.

#### Distribution of endpoints in the mediolateral dimension

Figure 10 shows the normalized mediolateral versus dorsoventral distribution of endpoints. As commented on in Materials, determination of the mediolateral endpoint position is inherently inaccurate. Relative to its small width, the medial and lateral border of the MSO cannot be determined with much precision (note that, on the square axes of Fig. 10, the mediolateral dimension is much magnified relative to the dorsoventral dimension, approximately by a factor 3.6). The uncertainty becomes particularly large for endpoints located close to the dorsal or ventral pole. This is visible in Figure 10: the mediolateral spread of points increases toward the dorsal pole. Nevertheless, as expected (Stotler, 1953), the contralateral endpoints occupy dominantly a medial position, and the ipsilateral endpoints a lateral position. Of interest is that the forward and backward branches of the ipsilateral projection seem to occupy the same dorsoventral (tonotopic) but not mediolateral position. For the four

fibers showing forward branches (Fig. 10 legend, + symbols), these branches give endpoints located more lateral than the backward branches (projections 2, 4, and 6).

#### Axon diameter and conduction velocity

The analysis so far has concerned branching patterns, endpoint position, and axon length. Other factors than axonal length contribute to the time delay in action potential propagation; most importantly myelination, axon diameter, and internodal distance. The only additional factor we can assess in our material is axon diameter, be it only crudely for the various reasons pointed out in Smith et al. (1993). Diameters were measured at each axonal (x,y,z) position entered. Using these measurements, we calculated the mean diameter of each "segment," which is the part of an axon between neighboring branch points; branch points and endpoints; or midline and branch point. Figure 11 shows the mean diameter as a function of distance from the midline (Fig. 11A) for contralateral projections, and from the first branch point (Fig. 11 B) for ipsilateral projections. These graphs are obtained by moving from left to right across the dendrograms (Fig. 7), and tallying and averaging all the segment diameters that are present for a given length from the starting point (Figs. 2–4, ML, 5, 6, FB). Close to the starting points, the mean diameter jumps in large steps because few branches are present, but the trace becomes smoother as more branching occurs. There is a general decrease in axon diameter toward the endpoints.

To compare diameters of contralateral and ipsilateral branches, we first deter-

mined the number of segments of a given diameter (bin size, 0.1  $\mu$ m). Dividing by the number of contributing fibers (9 for contralateral and 7 for ipsilateral projections), we obtain the average numbers shown in Figure 12, A and B. The diameters are largely between 1 and 3  $\mu$ m; the mean is 1.8  $\mu$ m for both distributions (t test, p = 0.27). For the contralateral projections, the axonal segments include all segments of projections into the contralateral MSO arising after the midline crossing, while for the ipsilateral projections they include all segments projecting into the ipsilateral MSO after the first branch point. The reason that we included the "extra" segment between midline and first branch point in the contralateral projections is that the midline gives an "absolute" reference that can be identified in each animal, which allows an estimate of path length for the axon in the contralateral hemibrain. These proximal segments are shown in black in Figure 12, A and C. Removal of these segments did not affect the lack of a statistically significant difference between the diameter distributions of contralateral and ipsilateral projections (p = 0.13).

Numerically, the final segments, shown in gray in Figure 12, dominate the histograms, due to the profuse branching once the collaterals reach the tonotopic layers where they form their endpoints. Because the axons progressively taper (Fig. 11) and much

of the delay must accrue at the distal branches, it is of interest to compare contralateral and ipsilateral distributions for the final segments. Again, there is no difference in the distributions for contralateral and ipsilateral fibers (p = 0.26).

Next, to visualize the relationship between diameter and length of segments, we summed the lengths for all segments with a given diameter for all fibers (bin size  $0.1~\mu m$ ) and normalized to the number of contributing fibers (9 for contralateral and 7 for ipsilateral projections). Thus, Figure 12, C and D, shows average segment length of a given diameter for contralateral and ipsilateral projections, respectively. Statistical comparison shows that at none of the bins for which both contralateral and ipsilateral values are present is there a significant difference in segment length (t test, p > 0.05).

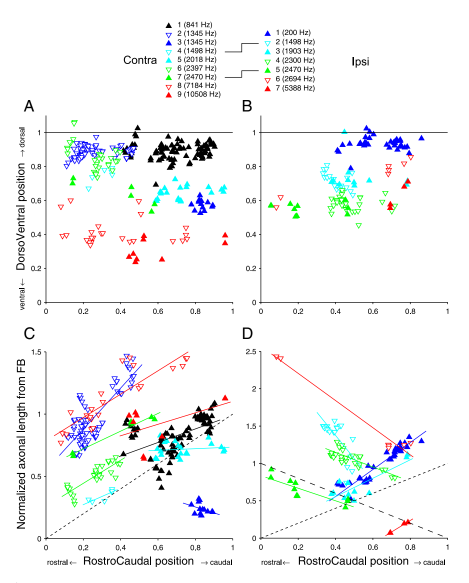
From the axon diameter and length, estimates for conduction speed and time can be calculated. We followed the method of Beckius et al. (1999) to estimate conduction time as follows:

$$L/(d^*RV),$$
 (1)

with L being the length (in millimeters) of axon between two branch points, d being the average diameter (in micrometers) of that segment, and RV being an experimentally derived constant (9.167 mm/ms/ $\mu$ m) (Waxman and Bennett, 1972). For each endpoint, the conduction times of all segments interposed between that endpoint and a fixed starting point were added.

In Figure 13 we show the estimated conduction times for all endpoints using as starting points the midline (Fig. 13*C*, contralateral projections) or the first branch point (Fig. 13*D*, ipsilateral projections) against the length of axon traversed to reach these endpoints. The relationship

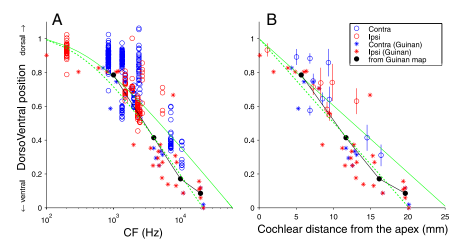
is very orderly; in all fibers, linear regression (data not shown) of length on conduction time was significant (p < 0.001; p < 0.01 in one fiber). This result is not surprising and may seem trivial because of the proportionality of estimated conduction time to length in Equation 1. However, if the average diameter to reach one endpoint were larger than that to reach another endpoint of the same tree, the conduction time would be shorter even if the average length were the same (Fig. 13A). Likewise, there could be a length difference that is compensated by a difference in axon diameter (Fig. 13B). The linear distributions of endpoints in Figure 13, C and D, show that d (Eq. 1) is "well behaved" in the sense that diameter decreases similarly with distance in all branches and that there is no systematic effect of diameter that offsets or enhances the "delay line effect" of the branching pattern. The slope of the linear regressions (data not shown) was on average 19.3 m/s for the contralateral and 17.9 m/s for the ipsilateral projections (ranges: 14–26 m/s for Contra; 13–21 m/s for Ipsi).



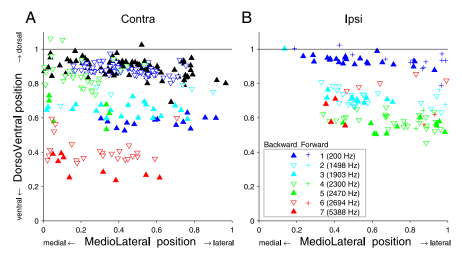
**Figure 8.** Length gradients along the rostrocaudal axis of the MSO. **A, B,** Distribution of endpoints of contralateral (**A**) and ipsilateral (**B**) projections on normalized MSO axes. **C, D,** Axonal length from FB to the different endpoints. Solid lines are linear regressions. The diagonals with short dashed lines indicate the diagonal of equality; the diagonal with the long dashed lines indicates the slope expected for an opposite length gradient. The caption shows the symbols and color used for the endpoints originating from the different fibers, numbered for increasing CF and reused in subsequent figures. The lines between left and right symbol captions indicate the two fibers for which both the contralateral and ipsilateral projections were reconstructed.

Note that these slopes do not give conduction speeds, but rather the speed at which excitation would "sweep" through the endpoints. If all endpoints were arranged in an orderly fashion as in Jeffress' model so that the endpoints with the shortest axonal length were located (e.g., most rostrally and those with the longest length most caudally), the range of conduction times spanned by the endpoints of one axon would equal the range of its x-values in Figure 13, C and D [e.g.,  $220-350~\mu$ s for the contralateral projection (#1), represented by the black triangles in C; i.e., a range of  $130~\mu$ s]. However, as already shown in Figure 8, C and D, the arrangement of endpoints is not that orderly. Again taking projection #1 as an example, it can be seen in Figure 8C that endpoints with similar axonal length occur over a significant rostrocaudal range. We return to this point in the section below, Estimates of rostrocaudal axonal delay (Fig. 14).

The lines in Figure 13, *C* and *D*, connect starting points, branch points, and endpoints of each projection. The slope of each line equals the speed of the corresponding segment. This



**Figure 9.** Tonotopic compression of low frequencies in MSO. *A, B,* The dorsoventral location, normalized to the dorsoventral extent of the MSO, is shown for all endpoints of all reconstructed fibers on a logarithmic CF abscissa (*A*) and for their mean and SD on a cochlear distance abscissa (*B*). Each circle in *A* indicates a single endpoint (blue, contralateral fiber; red, ipsilateral fiber). The asterisks and the black circles and line show extracellular and summary data from Guinan et al. (1972) (see main text). The green lines are predicted relationships based on the cochlear tonotopic map, based on Greenwood's formula (Greenwood, 1990). The solid green line is the prediction for a full representation of all CFs; the dashed line is for a representation limited to ≤22 kHz. The endpoints at low CFs cluster above the green lines. If low CFs were over-represented in MSO, those endpoints would be expected to cluster below rather than above the green lines.



**Figure 10. A, B,** Distribution of endpoints in the coronal plane, for contralateral (**A**) and ipsilateral (**B**) projections. Symbol use for **A** is as in Figure 8. The "Backward" and "Forward" in **B** refer to backward and forward branches, as illustrated in Figures 5 and 6.

slope decreases with increasing distance from the starting point, due to the gradual decrease in axon diameter, and gives rise to the general curvature in the data. For each endpoint, the length traversed divided by the estimated conduction time gives an estimate of the average speed from starting point to each endpoint (this speed equals the slope of a line connecting the endpoint to the origin in Fig. 13C,D, illustrated for the diagrams in Fig. 13A,B with dashed lines). The ranges of estimated speeds are shown in Fig. 13E–G. They are similar across endpoints and CFs, and are also very similar for contralateral and ipsilateral fibers (means, 20.1 and 19.3 m/s, respectively), when taking midline (Fig. 13E) and first branch point (Fig. 13G), respectively, as a starting point. When the contralateral side is also referenced to the first branch point rather than to the midline (Fig. 13F), the estimated conduction speed drops to 18.4 ms/s on average. This reflects the long distance (several millimeters) of the segment from midline to first branch point in contralateral fibers (Figs. 7A–C, 11A) and the large diameter of that segment (Fig. 11A), while the ipsilateral projection tends to sustain long (Fig. 7E,F) and large-diameter (Fig. 11B) forward and backward segments after the first branch point. In summary, the overall distribution of axon diameter and its progression along the axon is similar in contralateral and ipsilateral projections.

#### Estimates of rostrocaudal axonal delay

The main quantitative check so far regarding Jeffress' (1948) hypothesized axonal delay lines was in Figure 8, C and D, which examined length as a function of rostrocaudal position. In a similar format, Figure 14, A and B, replots the data directly in dimensions of time and space: estimated conduction time, calculated from both diameter and length, versus rostrocaudal position in micrometers from the most rostral MSO section. The abscissa origin coincides with the section containing the rostral pole, while the location of the section of the caudal pole is indicated by an asterisk at the caudal end of a linear regression line drawn through all the endpoints of each projection. As expected from the general trend of a graded decrease in axon diameter (Figs. 11-13), the relationships of Figure 8, C and D, are preserved in Figure 14, A and B. Thus, it does not appear to be the case that afferent branch diameter is regulated to fine tune delay lines based on length, at least not at the level of assessment of diameter that is possible with our material.

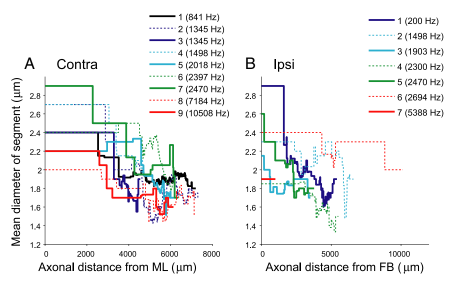
To summarize the regression data on estimated conduction times (Fig. 14A, B) and measured length gradients (Fig. 8C,D), Figure 14C shows the slopes of the linear regressions in these figures. Abscissa values of 1 and -1 indicate "perfect" slopes of delay, where the extra length of axon is identical to the extra dis-

tance in rostrocaudal (or caudorostral) direction (Fig. 8, compare diagonals in C and D). All significant (large symbols) contralateral projections, as well as three ipsilateral projections, contribute data to the upper right quadrant. The data points in the lower left quadrant consist only of ipsilateral projections, which form a length gradient in the caudorostral direction. Note that the most extreme positive and negative values are found in the ipsilateral projections. Also of note are the rather modest values of the estimated delays. To map the physiological range of delays (300-400  $\mu$ s in the cat) (Roth et al., 1980) on the length of MSO (3–4 mm) (Fig. 14A,B), the interaural delay accrued per millimeter should be  $\sim 100 \mu s$ . The ordinate values in Figure 14C show that the estimated "monaural" delays fall short for the contralateral projections. The ipsilateral delays come close but are in the wrong direction for the positive values, and for the negative values some of the projections are limited in spatial extent (Fig. 14B). In Jeffress' (1948) scheme, it is the combination of ipsilateral and contralateral axonal delays that establish the map of interaural delays. Thus, even if insufficient by themselves, a combination of the contralateral delays (triangles in upper right quadrant) and some of the ipsilateral delays (squares in the lower left quadrant) could sum to the needed physiological range of delays. The estimated delays are replotted versus CF in Figure 14D. For each fiber, the (extrapolated) delay at the rostral pole of the MSO is anchored to the horizontal line at 0 delay, and a vertical line is drawn to the extrapolated delay at the caudal pole. The line extends to positive delays for nearly all contralateral projections (triangles), and also for the four ipsilateral projections (squares) with negative slopes in Fig. 14B; these projections could all contribute to the pattern of interaural delays reported by Yin and Chan (1990). The individual data points show the estimated delays for individual end-

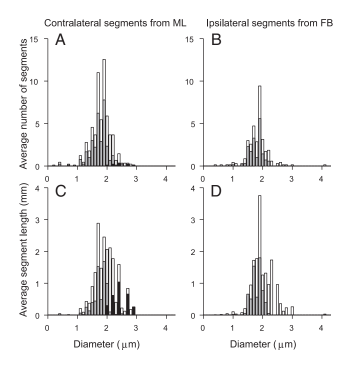
points, and the short horizontal bars delimit the range of delays of the linear regression over which endpoints are actually found. The gray rectangle approximates the range of physiological delays (400  $\mu$ s) that is presumed to be represented in the MSO of one side. Again, the range of delays subserved by contralateral projections is small, but could suffice when combined with the ipsilateral delays.

As was first reported in the guinea pig (McAlpine et al., 2001), the distribution of best delays in the inferior colliculus of the cat (Hancock and Delgutte, 2004; Joris et al., 2006) does not follow the shaded gray rectangle, but is largely restricted to the area defined by horizontal line at zero delay and the upper hyperbole at half a characteristic period (CF<sup>-1</sup>), and this is also the case for recordings of the MSO in rodents (Brand et al., 2002; Pecka et al., 2008). Binaural recordings have shown that best delays have a specific distribution pattern that is constrained by the upper hyperbole and the horizontal. Taken at face value, the axonal delay estimates of the present results are not consistent with the pattern of binaural delays thought to be represented in the MSO. There is no suggestion in the distribution of estimated delays toward larger delays at low than at high CFs.

Figure 14 focuses on the ranges of estimated delay, taking as anchor points the midline (Fig. 14A), the first branch point (Fig. 14B), or the rostral pole (Fig. 14D). The inherent longer path length of contralateral projections combined with the similarity of axon diameter in contralateral and ipsilateral projections (Fig. 12) implies a systematic additional delay for the contralateral inputs. Comparison of the entire contralateral and ipsilateral delay generated by an axon is difficult because that would require backfilling of the soma and labeling of the entire projection to both contralateral and ipsilateral MSO. However, we can make an estimate of the difference in delay between contralateral and ipsilateral inputs. Taking into account that individual fibers project both contralaterally and ipsilaterally, and, assuming mirror symmetry between left and right sides of the brainstem, we only need to increase the contralateral delay estimates of Figure 14A (made with respect to the midline) with the delay estimated for the axonal segment between the ipsilateral first branch point and the midline [Fig. 15B, distance from ipsilateral first branchpoint (FBi) to midline]. We could trace this segment in all ipsilaterally projecting fibers. The average segment length was 4994 µm

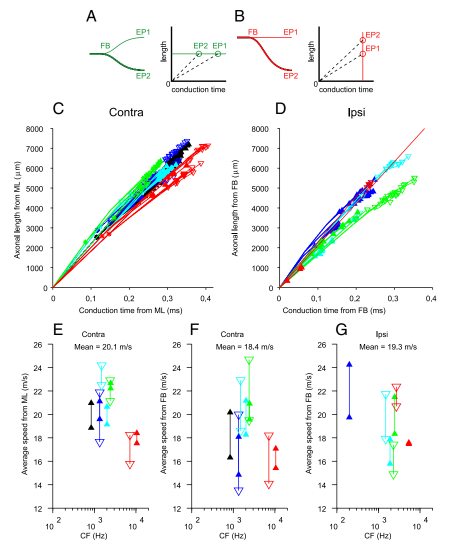


**Figure 11.** *A, B,* Average axon diameters decrease as a function of distance from the midline (*A*, Contra projections) or from the first branch point (*B*, lpsi projections).



**Figure 12.** *A–D,* Distribution of axon diameters does not differ between contralateral (*A, C*) and ipsilateral (*B, D*) projections. The top histograms show the average number of segments with a given diameter (0.1 μm bins); the bottom histograms show the corresponding average length (in mm) at that diameter. All histograms are averages calculated for the entire population of contralateral and ipsilateral fibers, normalized to the total number of fibers included. The starting segments were ML (contralateral fibers) and FB (ipsilateral fibers). White bars include all the segments from starting segment to endpoints. Gray bars show length of most distal segments, terminated by endpoints. Black bars show length of most proximal segments, which contain ML (contralateral fibers only).

(range,  $3748-6106~\mu m$ ), and the average diameter was  $2.6~\mu m$  (range,  $1.9-3.0~\mu m$ ), yielding an average extra delay (Eq. 1) of 221  $\mu s$  (range,  $139-351~\mu m$ ). Thus, on average the endpoint delay estimates in Figure 14A would need to be increased by 221  $\mu s$ . The value of 221  $\mu s$  may be an overestimate because the axon diameters of our small sample seem to be an underestimate (Eq. 1); larger axon diameters at the midline were reported by Smith et al. (1993) (2.5–5  $\mu m$ ) and Brownell (1975) (3–5  $\mu m$ ).



**Figure 13.** Axonal length is linearly related to estimated conduction time. Axonal lengths are measured as in Figures 7 and 11. Conduction time for each endpoint is the sum of conduction times of all segments leading up to that endpoint, calculated from segment length and diameter (see text). Diagrams in **A** and **B** illustrate configurations that would result in nonsloping relationships. **C**, Length and time for contralateral projections, using ML as reference point. **D**, Length and time for ipsilateral projections, using FB as reference point. One outlying endpoint is clipped off. **E**, **F**, Estimated average conduction speeds for all endpoints. This is the slope of the lines connecting the endpoints in **C** and **D** with the origin at (0,0), as illustrated with the dashed lines in **A** and **B**. **E**–**G**, Reference point at (0,0) was ML (**E**) or FB (**F**, **G**). Color code and symbols are identical to Figures 8 and 10. The symbols indicate the lowest and highest speed for the endpoints of a given fiber.

The values of Beckius et al. (1999) (slightly >2  $\mu$ m) are comparable to our own.

Note that inclusion of this extra delay does not alleviate the problem pointed out in Figure 14 D, since it adds a constant delay common to all endpoints of a given fiber and does not increase the range of available delays provided by a given fiber. Figure 15 A shows the estimated average (symbols) and range (lines) of path lengths from the last common axonal point (FBi) to all endpoints, as a function of CF, using the average of 4994  $\mu$ m for the distance from FBi to midline calculated above. On average, the contralateral pathway from FBi onward is 10.78 mm, and the ipsilateral pathway is 4.21 mm. Incorporating the measured diameters, the average delay from the first ipsilateral branch point to MSO is an estimated 511  $\mu$ s for the contralateral projections and 220  $\mu$ s for the ipsilateral projections. Thus, the difference in path length is on average 6.57 mm, which generates an extra delay for the contralateral projection of an estimated 290  $\mu$ s. Notwith-

standing the uncertainty of its exact value, this extra delay must contribute to the contralateral bias of best delays seen in all studies of the MSO and inferior colliculus. It is interesting to observe that this value is similar to the  $\sim 200~\mu s$  extra contralateral delay in the lateral superior olive, estimated with different physiological methods (Joris, 1996; Joris and Yin, 1998; Tollin and Yin, 2005).

# Axonal delay in the dorsoventral dimension

Differences in the CF of contralateral and ipsilateral inputs converging on MSO neurons have been proposed as a source of internal delays (Schroeder, 1977; Shamma, 1989; Bonham and Lewis, 1999). Crosscorrelation analysis of responses of auditory nerve fibers revealed that disparities of a fixed distance on the cochlear basilar membrane generate a wider range of internal delays at low than at high CFs, in accord with the pattern of internal delays inferred from recordings in the inferior colliculus (Joris et al., 2006). However, to explain not only the decrease in the range of best delays with CF, but also the decrease in mean best delay with CF, an additional mechanism is needed. Because most contralateral projections approach the MSO from a rostral and ventral position (Figs. 2C, 3A, B), it was hypothesized (Joris et al., 2006) that the longer distance toward the dorsal (low-CF) pole of the MSO compared with the ventral (high-CF) pole perhaps causes a systematically longer delay for low-CF than for high-CF contralateral inputs. This hypothesis is illustrated with the schematic in Figure 15C. In Figure 15D, we show the mean and SD of the axonal lengths from midline to endpoint for all contralateral projections as a function of fiber CF (for individual data points, see Fig. 13C). There is no hint of a CF dependency showing longer

lengths for the most dorsally projecting fibers (lowest CFs). The low-CF (Fig. 15*E*) and high-CF (Fig. 15*F*) projections illustrated in coronal view show that the tilting of the dorsal edge of the MSO toward the midline, combined with the angling of the contralateral low-CF fibers, is such that indeed no differential delay would be expected relative to the more ventral, high-CF projections. There is therefore no evidence in our data for a length gradient along the dorsoventral dimension in the excitatory inputs from the contralateral ear.

#### Discussion

Sensitivity to temporal differences in the sounds to the two ears provide a powerful model system to study neural temporal processing. Putative axonal delay lines have been an important component of models and of theorizing regarding such processing (Jeffress, 1948; Licklider, 1959; McFadden, 1973; Carr and Konishi, 1988; Cariani, 2004). There is general agreement that sensi-

tivity to ITDs involves internal delays causing a temporal shift of inputs from one ear relative to the other. What is increasingly questioned is the proposal (Jeffress, 1948) of a conversion of a temporal code to a place code by virtue of axonal delay lines. We quantitatively reanalyzed projection patterns of spherical bushy cells of the AVCN to MSO (Smith et al., 1993). The main findings of that study hold up and are consistent with a later study (Beckius et al., 1999); axonal configurations that are qualitatively of the nature surmised by Jeffress (1948) are found for contralateral projections, and less clearly for ipsilateral projections. Nonetheless, these configurations are not consistent in a simple form with the binaural data that have become available since.

#### Previous studies

The two previous reports on anatomical delay lines in the cat have complementary virtues. The power of the data of Smith et al. (1993) is that their 18 axons were physiologically characterized and that the labeling was unambiguously from single fibers, but filling was incomplete so that, typically, projections on only one side could be reconstructed. Their report showed one contralateral and one ipsilateral computer reconstruction. Beckius et al. (1999) labeled a large number of axons by injecting the rostral pole of the AVCN, of which 17 well filled axons (from 2 animals) were quantitatively reconstructed, with 7 reported in detail. Disadvantages of that material are the absence of physiology and the difficulty in connecting pieces across sections. Both studies have few data at very low CFs, where the largest best delays are observed. Best frequencies at the site of injection in Beckius et al. (1999) were estimated at  $\sim$ 1.5–1.75 kHz, but the tonotopic layering of the projections suggests that the actual range of CFs was much larger. In Smith et al. (1993), only two fibers had CF < 1 kHz.

Despite differences in technique, the two studies were consistent, finding caudally directed delay line patterns contralaterally and a complex, less systematic pattern ipsilaterally. Quantitative analysis allowed Beckius et al. (1999) to discern a weaker length gradient in ipsilateral branches, in a direction opposite to the contralateral pattern. The quantitative analysis presented here is largely consistent with that of Beckius et al. (1999). The main additional findings concern the ipsilateral projections. We found caudally directed delay lines in three (of seven)

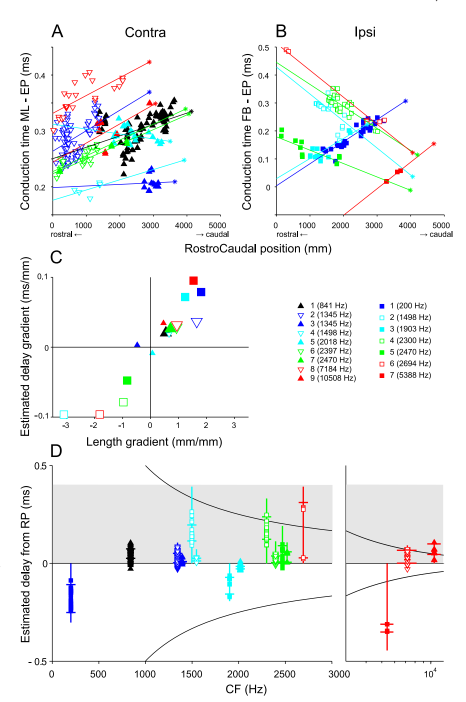
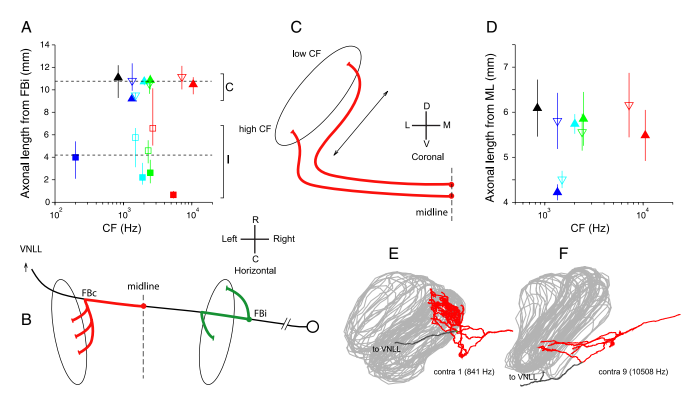


Figure 14. Rostrocaudal gradients of conduction time. A, B, Relationship between estimated conduction time and location of EP in rostrocaudal dimension in contralateral (A) and ipsilateral (B) fibers. The abscissa is zeroed to the position of the most rostral section in which MSO could be identified, and abscissa values are the distance of endpoints to that most rostral section. Ordinate values are the estimated conduction times from the midline ( $\mathbf{A}$ , contralateral projections) or first branch point (B, ipsilateral projections). Solid lines are linear regressions. The asterisks at the end of each line indicate the most caudal MSO section. Note that the ordinate in **B** has a wider range than in **A**. **C**, Summary of regression slopes of Figure 8, A and B (abscissa), and of A and B (ordinate). Large symbols indicate values that are significant for both abscissa and ordinate. Contra 4 showed significance ( p < 0.05) for length but not for delay; and vice versa for Contra 9.  $\boldsymbol{D}$ , Relationship between estimated delay and CF. The anchor point of each colored vertical line at 0 delay represents the RP of the MSO. The opposite end shows the extrapolated delay at the CP (corresponding to the delay accumulated between RP and the asterisk in A and B). The small horizontal bars show the range of delays of the linear regression over which endpoints are present. Symbols and lines at positive delays, in the shaded region, are for fibers with a pattern consistent with the trend observed by Yin and Chan (1990); these are the fibers with positive slope in  $\boldsymbol{A}$  or negative slope in  $\boldsymbol{B}$ . Symbols and lines at negative delays are for fibers with an opposite branching pattern (negative slope in  $\boldsymbol{A}$  or positive slope in  $\boldsymbol{B}$ ). Hyperbolic curves indicate the  $\pi$  limit (i.e., the extent of one period equaling CF  $^{-1}$ ). The scale of the abscissa is linear in the left panel and logarithmic in the right panel.



**Figure 15. A, B,** Differences in path length between ipsilateral and contralateral inputs. **A,** Estimated average and range of lengths for contralateral and ipsilateral projections between the FB of the ipsilateral MSO projection and the endpoints. Dashed lines show overall contralateral (C) and ipsilateral (I) mean. **B,** Schematic of the measurement. The length of the trajectory between midline and contralateral endpoints (green), was measured (same values as in Fig. 13C,D). To estimate the difference in length of contralateral and ipsilateral inputs, the red segment needs to be augmented by the distance FBi to midline: we use the average measured on ipsilateral projections (4994 μm, see main text). **C–F,** Absence of length differences in the innervation from the contralateral ear along the dorsoventral dimension of MSO. **C,** Hypothetical scheme of innervation by contralateral fibers, suggesting extra length for the most dorsally projecting (low-CF) fibers. If input fibers approach the MSO from ventral, low-CF fibers may have longer axonal length (arrow) to innervate the dorsal part of MSO than high-CF fibers for the ventral part. **D,** relationship between axonal length from ML to endpoints, and CF. Symbols in **A** and **D** indicate average axonal length; lines are range (**A**) and SD (**D**) for each projection (symbol color and shape as in Fig. 8). **E, F,** Coronal view of contralateral projections of fibers with lowest and highest CF (840 and 10,508 Hz, respectively).

ipsilateral reconstructions (i.e., where the length gradient was in the same direction as that of contralateral projections). This pattern was particularly convincing in the fiber with the lowest CF (200 Hz). Also, the rostrally directed delay line in the remaining four ipsilateral reconstructions was at least as steep (though of opposite sign) as that of the contralateral fibers (Figs. 8 *B*, 14*C*), while in Beckius et al. (1999) the gradients on the ipsilateral side were shallow and constrained in rostrocaudal extent.

#### Anatomical patterns and best delays

Ultimately, "delay" is a functional concept requiring functional measurements. Particularly appealing in the morphological observations is their apparent consistency with the relationship between binaural tuning and cell location observed by Yin and Chan (1990), both in sign and in size. In that study, small best delays were found rostrally and large delays caudally, consistent with a caudally directed delay line contralaterally and/or a rostrally directed delay line ipsilaterally. Moreover, the range of physiological delays is reasonably in accord with estimated length differences and conduction speeds: a range of  ${\sim}400~\mu s$  could be covered by combining "monaural" contralateral and ipsilateral delays.

Nevertheless, our quantitative analysis brings out several problems. Most importantly, there is nothing in the pattern of estimated delays observed that would suggest a CF-dependent range of delays (Fig.  $14\,D$ ). The anatomical branching patterns, as illustrated in Figures 2 and 3, are equally present at high and low CFs. In contrast, the physiological distribution of best delays measured in the MSO and inferior colliculus shows a clear depen-

dence on CF, in cat and in other mammals studied (Brand et al., 2002; McAlpine and Grothe, 2003; Hancock and Delgutte, 2004; Joris et al., 2006; Pecka et al., 2008).

A second problem, of relevance beyond the issue of axonal delay lines, is that of correct concatenation. None of the afferents illustrated here or in Beckius et al. (1999) innervates the entire rostrocaudal extent of the MSO. Thus, each tonotopic strip of MSO neurons must be supplied by a patchwork of afferents. This complicates the structural basis for a systematic rostrocaudal gradient, since it requires correct temporal stacking of collaterals. For example, contralateral projections 2 and 3 have the same CF but innervate different portions of the MSO: projection 2 is to the rostral half and shows a positive slope, while projection 3 is restricted toward the caudal pole (Figs. 8C, 14A). Obviously, if these two fibers were representative for a single animal, the delay pattern generated by projection 2 would not be complemented by the inputs from projection 3. In other cases, fibers of similar CF appear to concatenate in an orderly manner (e.g., contralateral projections 5, 6, and 7). The fibers labeled here were derived from different animals, and offsets in the ordinate values of Figures 8, C and D, and 14, A and B, could reflect differences in anatomy between animals (e.g., in MSO size and location). Within-animal data are reasonably consistent in one animal studied by Beckius et al. (1999, their Fig. 11) but not in the other, which showed large offsets among fibers. However, the latter case also showed large variation in dorsoventral location of the projections, so that the labeled population must reflect a wide range of CFs. Together, the available material does not allow a prediction of the degree of

temporal dispersion that may be present across afferents of equal CF.

A third problem may be in the range of delays. The estimated contralateral delays are small relative to the values of best delay observed, particularly at low CFs where best delays can be >1 ms (Fig. 14, upper hyperbole) (Hancock and Delgutte, 2004; Joris et al., 2006). Ipsilateral delays tend to be larger than contralateral delays but are also more erratic in sign and rostrocaudal coverage. Combination of contralateral and ipsilateral delays can enlarge the net interaural delay (when the slopes are of opposite sign for the contralateral and ipsilateral projections), but sources of delay in addition to the delays provided by the within-fiber branching pattern are needed to account for the full range of best delays observed at low CFs. We caution that the diameter measured on our material is necessarily a rough estimate, as is the relationship between fiber diameter and conduction speed (Waxman and Bennett, 1972), and is suited for observing trends rather than absolute values.

Our reanalysis casts doubts on the relevance of the rostrocaudal axonal branching patterns for interaural delay, but does not by itself contradict the evidence for a spatial map of delay in MSO (Yin and Chan, 1990). Nevertheless, taking into account the noise in the relationship observed by Yin and Chan (1990), the absence of a clear relationship in multiunit recordings (Oliver et al., 2003), and the problems in tying the distribution of best delays to axonal branching patterns (Fig. 14*D*), it appears increasingly doubtful that these patterns are an essential component of the binaural circuit.

A final qualification is that other features of the branching pattern—different from the simple within-axon delay line configuration—may contribute to internal delay. For example, contralateral/ipsilateral asymmetries in temporal dispersion, in convergence across CFs, and in convergence ratio could all contribute to the pattern of best delays observed. Also, based on our material we cannot exclude that patches of contralateral and ipsilateral fibers would provide the range of best delays observed (for comparison, see Goldberg and Brown, 1969). It is interesting in this regard that there is more consistency in the pattern of projections to the rostral than to the caudal half of MSO (Fig. 8). Finally, there are other determinants of conduction speed, internodal distance in particular, which may show systematic variations in mammalian binaural circuits, as in birds (Carr and Konishi, 1990; Seidl et al., 2010).

#### Tonotopy: over- or under-representation of low frequencies?

A surprising finding is the tonotopic distribution of afferents. Their layering suggests that low frequencies are underrepresented rather than over-represented in the MSO. This is puzzling—functionally and anatomically—and is inconsistent with the "duplex" organization of the superior olivary complex (Irvine, 1986), comprising a low-frequency circuit computing ITD and a high-frequency circuit computing interaural level differences (ILDs). Taking the limited sample sizes of our data and those of Guinan et al. (1972) into account, a conservative statement is that MSO is tonotopically mapped isomorphic to the cochlea and has a low-frequency bias to the extent that very high CFs are not represented in MSO. The main duplex asymmetry is in the ILD (not the ITD) circuit: the lateral superior olive and medial nucleus of the trapezoid body show an apparent over-representation of high frequencies (Guinan et al., 1972).

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