Two Synaptic Convergence Motifs Define Functional Roles for Inputs to Cochlear Nucleus Bushy Cells

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Abstract

Bushy neurons of the cochlear nucleus encode temporal fine structure and modulation of sound with high fidelity. However, the synaptic maps and electrotonic structures that underlie these properties are not specified in meaningful explanatory detail. We employed modern volume electron microscopy techniques to provide exact data on the numbers of synaptic inputs and their weights determined by the number of contained active zones, and the surface areas of all postsynaptic cellular compartments. Leveraging these high-resolution images, we discovered cabling of dendrite branches and new structures within dendrites, and identified non-innervated dendrites. We extend current nanoscale connectomic studies with methods to export cellular reconstructions into morphologically-constrained, biophysically-based predictive computational models. We reveal both coincidence detection and mixed supra/subthreshold modes of input convergence across the bushy cell population and show subthreshold inputs contribute to enhanced temporal encoding even in the presence of suprathreshold inputs. We demonstrate the variation of dendritic load and axon parameters and their importance in controlling excitability as potential homeostatic mechanisms, thereby defining heterogeneity in stimulus-evoked responses across the BC population.

Introduction

Bushy cells (BC) of the cochlear nucleus (CN), especially a subpopulation called globular bushy cells (GBC) located in the auditory nerve fiber (ANF) entry zone, encode temporal fine structure and modulation of sound with high fidelity (Bourk, 1976; Joris et al., 1994a,b; Wei et al., 2017; Spirou et al., 1990; Smith et al., 1991). They also play a central role in hearing as they are essential for binaural processing and are a key cell type defining and driving the early CNS stages of the lemniscal auditory pathway (Warr, 1966; Tolbert et al., 1982; Smith et al., 1991; Yin et al., 2019; Spirou et al., 1990; Smith et al., 1991).
We describe metrics for classifying the cells as GBCs, but use the generic terminology of BC through-
lineation of actual synaptic contacts to estimate synaptic weight. These parameters are essential
for prediction of neural activity and understanding the computational modes employed by BCs.

Although the preponderance of ANF inputs are somatically targeted, the dendrites of BCs ex-
hibit complex branching and multiple swellings that are difficult to resolve in light microscopic
(LM) reconstructions (Lorente de Nó, 1981). Consequently the dendritic contributions to the elec-
trical properties of BCs have not been explored. Innervation of dendrites and soma was revealed
from partial reconstruction from EM images (Ostapoff and Morest, 1991; Tolbert and Morest, 1982;
Smith and Rhode, 1987), but values are often estimated as percent coverage rather than absolute
areas. Sub-sampling using combined Golgi-EM histology has shown innervation of swellings and
dendritic shafts (Ostapoff and Morest, 1991), and immunohistochemistry has further indicated
the presence of at least a sparse dendritic input (Gómez-Nieto and Rubio, 2009). Nonetheless, a
complete map of synapse location across dendrite compartments, soma, and axon has not been
constructed.

To resolve these longstanding issues surrounding this key cell type, we employed volume elec-
tron microscopy in the auditory nerve entry zone of the mouse CN to provide exact data on num-
bers of synaptic inputs and their active zones along with surface areas of all cellular compartments.
We describe metrics for classifying the cells as GBCs, but use the generic terminology of BC through-
out the text, due to unresolved distinctions within the BC class in mice and when studied anew with
volume EM. We identify a new dendrite structure with a high branching order, which we term a hub,
describe cabling of dendrite branches not discernible by LM, and provide the first metrics for axon
initial segment (AIS) surface area and innervation by endbulbs. Nanoscale connectomic studies
typically provide neural connectivity maps at cell to cell resolution (Zheng et al., 2018; Scheffer
et al., 2020; Turner et al., 2022; Bae et al., 2021; Shapson-Coe et al., 2021; Cook et al., 2019). We
extend these studies and previous modeling studies of BCs, by using detailed reconstructions from
the EM images to generate and constrain compartmental models that, in turn, are used to explore
mechanisms for synaptic integration and responses to temporally modulated sounds. These pro-
cedures reveal that BCs fall into two populations, one that operates as a coincidence detector with
all subthreshold inputs, and one that operates in a mixed coincidence/first spike mode with com-
bined subthreshold and suprathreshold inputs. The presence of subthreshold inputs is essential
for enhanced temporal precision of spiking for amplitude modulated tones up to 300 Hz and for
transient sounds. We also provide the first complete map of synaptic inputs for a BC, and show
that entire branches of dendrites can lack synapses. The dendrite surface area and AIS length
emerge as key parameters, in addition to the membrane time constant, to adjust spike threshold,
weight synaptic inputs and provide homeostatic mechanisms by which GBCs can regulate their excitability.

Results

Cellular organization of the auditory nerve root region of the mouse cochlear nucleus

Despite many years of study, fundamental metrics on morphology of BC somata, dendrites and axons, and the synaptic map of innervation across these cellular compartments is far from complete. We chose volume electron microscopy (serial blockface electron microscopy (SBEM)) to systematically address these fundamental questions at high resolution and quantify structural metrics, such as membrane surface area and synaptic maps, in combination with compartmental modeling that is constrained by these measurements, to deepen our understanding of BC function. The image volume was taken from the auditory nerve entry zone of the mouse CN, which has a high concentration of BCs. The image volume was greater than 100 µm in each dimension and contained 26 complete BC somata and 5 complete somata of non-BCs that were likely multipolar cells (beige and rust colored, respectively, in Figure 1A). Fascicles of ANFs coursed perpendicular to other fascicles comprised, in part, of CN axons, including those of BCs, as they exit into the trapezoid body (ANF and BC (colored mauve) axons, respectively in Figure 1A).

Segmentation of neurons from the image volume revealed BC somata as having eccentrically located nuclei (25/26 BCs) with non-indented nuclear envelopes (25/26 BCs; the one indented nuclear envelope was eccentrically located), and stacks of endoplasmic reticulum only along the nuclear envelope facing the bulk of the cell cytoplasm (26/26 BCs; Figure 1B-C). Myelinated ANFs connected to large end-bulb terminals synapsing onto the BC somata. Reconstructions from volume EM permitted accurate measurement of the directly apposed surface area (ASA) between the endbulb terminal and postsynaptic membrane, and identification of synapses as clusters of vesicles along the presynaptic membrane (Figure 1B-D, D'). In a subset of terminals we counted the number of synapses. Because synapse density showed only a small decrease with increasing ASA (Figure 1F), we used the average density to estimate the number of synapses in each terminal and to set synaptic weights in computational models (Figure 1F, and see Methods).

An important goal of this project was to provide accurate measurements of membrane surface areas, in order to anchor compartmental models of BC function and facilitate comparison across species and with other cell types. We standardized a procedure based on a method to generate computational meshes (Lee et al., 2020), yet preserve small somatic processes (see Methods and Figure 1–Figure Supplement 1A-B). The population of BC surface areas was slightly skewed from a Gaussian distribution (1352 (SD 168.1) µm²), with one outlier (cell with indented nucleus) near 2000 µm² (Figure 1E). The MCs (red bars in Figure 1E) may represent two populations based on cells with smaller (<1700 µm²) and larger (>2000 µm²) somatic surface area.

The synaptic convergence motifs of bushy cells

With image segmentation parameters set, we next addressed a range of competing models for synaptic organization by which BCs can achieve higher temporal precision at the onset of sound and in phase locking to periodic stimuli than ANFs, and exhibit physiologically relevant values for spike regularity (Rothman et al., 1993; Joris et al., 1994b,a). These models are based on convergence of large, somatic endbulbs of Held (Rouiller et al., 1986; Liberman, 1991; Ryugo and Fekete, 1982) (Figure 2A,B). At one extreme, all convergent inputs, although harboring multiple release sites, are subthreshold for spike generation, and also of similar weight. With the functional attribute of a brief temporal integration window defined by the short membrane time constant, this convergence motif defines BC operation as a coincidence detector. At the other extreme, all so-
Figure 1. The imaged volume in the cochlear nucleus captures bushy (BC) and multipolar cells (MC), and reveals synaptic sites. (A) The VCN region that was imaged using SBEM is depicted within walls of the image volume. Twenty-six BCs (beige) and 5 MCs (orange) are shown with their axons (red). Left rear wall transects auditory nerve fiber (anf) fascicles, which run parallel to the right rear wall and floor. Non-anf axons exit into the right rear wall and floor as part of other fascicles that are cross-sectioned. The complete volume can be viewed at low resolution in Figure 1–video 1. (B) Example image, cropped from the full field of view, from the data set in panel A. Field of four BC (bc) cell bodies, myelinated axons in anf fiber fascicles, and capillaries (c). (C) Closeup of the cell body (cb) of lower right BC from panel B, illustrating the eccentrically located nucleus (n), short stacks of endoplasmic reticulum (er) aligned with the cytoplasm-facing side of the nuclear envelope, and contact by an endbulb (eb). (D) Closeup of the labeled endbulb contacting the cell in panel C (eb), revealing its initial expansion along the cell surface. Apposed pre- and postsynaptic surface area (ASA; green) are accurately determined by excluding regions with intercellular space (ASA is discontinuous), and synaptic sites (s1-4) are indicated as clusters of vesicles with some contacting the presynaptic membrane. (D') Inset in panel D is closeup of synapse at lower left in panel D. It shows defining features of synapses in these SBEM images, which include clustering of vesicles near the presynaptic membrane, convex shape of the postsynaptic membrane, and in many cases a narrow band of electron-dense material just under the membrane, as evident here between the “s1” symbol and the postsynaptic membrane. (E) Histogram of all somatic surface areas generated from computational meshes of the segmentation. BCs are denoted with grey bars and MCs with red bars. (F) Synapse density plotted against ASA shows a weak negative correlation. Marginal histogram of density values is plotted along the ordinate. Scale bars = 5 μm in B, 2 μm in C, 1 μm in D, 250 nm in D'.

Figure 1–Figure supplement 1. Steps in mesh generation and compartmental representation from EM volumes.

Figure 1–video 1. Exploration of the relation between an image volume and a bushy cell (BC) mesh derived from that volume. This video opens with a top-down view of the SBEM image volume from the ventral cochlear nucleus. The video zooms in as the volume is slowly cut away to reveal BC05, including its dendrites (red), cell body (beige), axon (pink), and all large somatic inputs (various colors). The perspective then shifts laterally to view several of the large terminals (various colors) contacting the cell body.
Figure 2. Two competing models for synaptic convergence. (A) Coincidence Detection model, all inputs are subthreshold (small circles), have similar weight, and at least two inputs are active in a short temporal window to drive a postsynaptic spike. Each vertical bar is a presynaptic spike and each row is a separate auditory nerve (AN) input. Bottom line is activity of postsynaptic bushy cell (BC). EPSPs are solid; action potentials indicated by vertical arrows. Dotted lines are inputs that occur during the refractory period (solid bar). Drawn after Joris et al. (1994a). (B) First-Arrival model, whereby all inputs are suprathreshold (large circles), have similar weight, and the shortest latency input drives a postsynaptic spike. For a periodic sound, both models yield improved phase-locking in the postsynaptic cell relative to their auditory nerve (AN) inputs. (C) Histogram of input sizes, measured by apposed surface area (ASA), onto BC somata. Minimum in histogram (vertical bar) used to define large somatic inputs (arrow). Inset: Top. Size distribution of somatic terminals traced to auditory nerve fibers within the image volume. Middle. Size distribution of somatic terminals with myelinated axons that were not traced to parent fibers within volume. Bottom. Size distribution of somatic terminals with unmyelinated axons. Some of these axons may become myelinated outside of the image volume. Small terminals (left of vertical dashed lines) form a subset of all small terminals across a population of 15 BCs. See Figure 2—Figure Supplement 1 for correlations between ASA and soma areas. (D) Plot of ASAs for all inputs to each cell, linked by lines and ranked from largest to smallest. Size of largest input onto each cell projected as a histogram onto the ordinate. Dotted line indicates a minimum separating two populations of BCs. Linked ASAs for BCs above this minimum are colored black; linked ASAs for BCs below this minimum are colored red. (E, F) All large inputs for two representative cells. View is from postsynaptic cell. (E) The largest input is below threshold defined in panel (D). All other inputs have similar range as the inputs in panel (E). See Figure 2—video 1 to view the somatic inputs on BC18.
matic ANF inputs are large and suprathreshold, also of similar weight. In this scenario, the BC operates as a latency detector, such that the shortest latency input on each stimulus cycle drives the cell. In both models, the BC refractory period suppresses delayed inputs.

In order to evaluate the predictions of these models, key metrics of the number of ANF terminal inputs and the weights of each are required. We first determined a size threshold to define endbulb terminals. All non-bouton (endbulb) and many bouton-sized somatic inputs onto 21 of 26 GBCs were reconstructed, including all somatic inputs onto 2 cells. We then compiled a histogram of input size based on ASA. A minimum in the distribution occurred at ~ 25 – 35 μm², so all inputs larger than 35μm µm² were defined as large terminals of the endbulb class (Figure 2C). We next investigated whether this threshold value captured those terminals originating from ANFs, by tracing retrogradely along the axons. Terminals traced to branch locations on ANFs within the volume matched the size range of large terminals estimated from the histogram (only two were smaller than the threshold value), and were all (except one branch) connected via myelinated axons (Figure 2C inset, top). Nearly all axons of the remaining large terminals were also myelinated (Figure 2C inset, middle). The remaining few unmyelinated axons associated with large terminals immediately exit the image volume, and may become myelinated outside of the field of view (Figure 2C inset, bottom, right of vertical dashed line). These data together lend confidence to the value of 35 μm² as the size threshold for our counts of endbulb terminals. We use the terminology "endbulb" or "large terminal" interchangeably throughout this report.

**Number of convergent auditory nerve endbulb inputs exceeds previous estimates**

After validating the size range for the endbulb class, we noted that the range of 5-12 convergent endings (Figure 2D and inset) exceeded estimates of 4-6 inputs from physiological measures in mouse (Cao and Oertel, 2010). We next inquired whether the range of input size was similar across all cells. Inspecting the largest input onto each cell revealed, however, two groups of BCs, which could be defined based on whether their largest input was greater than or less than 180 μm² (histogram along left ordinate in Figure 2D). Plotting endbulb size in rank order (largest to smallest) for each cell revealed that, excluding the largest input, the size distributions of the remaining inputs overlapped for both groups of BCs (black and red traces in Figure 2D). A catalogue of all inputs for the representative cells illustrates these two innervation patterns and reveals the heterogeneity of input shapes and sizes for each cell and across the cell population (Figure 2E,F; Figure 2–Figure Supplement 2 and Figure 2–Figure Supplement 3). We hypothesized from this structural analysis that one group of BCs follows the coincidence detection (CD) model depicted in Figure 2A where all inputs are subthreshold (12/21 cells; red lines in Figure 2D), and a second group of BCs follows a mixed coincidence-detection / first-arrival model (mixed-mode, MM) where one or two inputs are suprathreshold and the remainder are subthreshold (9/21 cells; black lines in Figure 2D). No cells strictly matched the latency detector model (all suprathreshold inputs) depicted in Figure 2B.

**Translating high-resolution neuron segmentations into compartmental models**

Ten of the BCs had their dendrites entirely or nearly entirely contained within the image volume, and were selected for compartmental modeling. The computational mesh structures (dendrites, cell body, axon hillock, axon initial segment, myelinated axon) were converted to a series of skeletonized nodes and radii (SWC file format (Cannon et al., 1998; Figure 3B, right and Figure 3–Figure Supplement 1 – mesh and SWC images of all 10 cells) by tracing in 3D virtual reality software (sy-Glass, IstoVisio, Inc.). The SWC files were in turn translated to the HOC file format for compartmental modeling using NEURON (Carnevale and Hines, 2006). The HOC versions of the cells were scaled to maintain the surface areas calculated from the meshes (see Methods). An efficient computational pipeline was constructed that imported cell geometry, populated cellular compartments
Figure 3. Steps in the generation of models from the mesh reconstructions of mouse VCN bushy neurons. (A) The mesh representation of the volume EM segmentation was traced using syGlass virtual reality software to generate an SWC file consisting of locations, radii, and the identity of cell parts (B). In (B), the myelinated axon is dark red, the axon initial segment is light blue, the axon hillock is red, the soma is black, the primary dendrite is purple, dendritic hubs are blue, the secondary dendrite is dark magenta, and the swellings are gold. The mesh reconstruction and SWC reconstructions are shown from different viewpoints. See Figure 3—Figure Supplement 1 for all reconstructions. See Figure 3—video 1 for a 3D view of the mesh and reconstructions for BC11. (C) The resulting SWC model is decorated with ion channels (see Figure 3—Figure Supplement 2 for approach), and receives inputs from multi-site synapses scaled by the apposed surface area of each ending. For simulations of auditory nerve input, sounds (blue) are converted to spike trains to drive synaptic release. (D). Comparison of responses to current pulses ranging from -1 to +2 nA for each dendrite decoration scheme. In the Passive scheme, the dendrites contain only leak channels; in the Active scheme, the dendrites are uniformly decorated with the same density of channels as in the soma. In the Half-active scheme, the dendritic channel density is one-half that of the soma. See Figure 3—autorefiggsupp:Fig3Supp3 for all current clamp responses and Figure 3—Figure supplement 3 for impedance plots. (E) Current voltage relationships for the 3 different decoration schemes shown in (D). Curves indicated with circles correspond to the peak voltage (exclusive of APs); curves indicated with squares correspond to the steady state voltage during the last 10 ms of the current step. Red dots indicate the AP threshold. (F) Example of voltage response to a tone pip in this cell. Action potentials are marked with red circles, and are defined by rate of depolarization and amplitude (see Methods). (G) Peri-stimulus time histogram (PSTH) for 50 trials of responses to a 4 kHz 100-ms duration tone burst at 30 dB SPL. The model shows a primary-like with notch response. See Figure 3—Figure Supplement 5 for all tone burst responses. (H) First spike latency (FSL; blue) and second spike latency (SSL; red) histograms for the responses to the tone pips in G. (F,G,H) The stimulus timing is indicated in blue, below the traces and histograms.

Figure 3—Figure supplement 1. Segmented neurons and their representations for compartmental modeling.

Figure 3—Figure supplement 2. Conductance scaling using voltage clamp simulations for different patterns of dendrite decoration.

Figure 3—Figure supplement 3. Summary of current-clamp responses for all 10 cells for each of the decoration conditions.

Figure 3—Figure supplement 4. Input impedance at rest.

Figure 3—Figure supplement 5. PSTH and spike latencies in response to tone bursts at CF.

Figure 3—video 1. Comparison of cell structures and regions between a 3D mesh and an SWC of a bushy cell (BC). This video opens with a full side-by-side view of the BC11 reconstruction (left), including its dendrites (red), cell body (beige), and axon (pink), and the SWC representation (right) of the same structures. The view then zooms into the cell body and axon, where the transition point from cell body to the axon (yellow arrow), the middle of the unmyelinated initial segment (green arrow), and the transition point where myelination of the axon begins (blue arrow) are indicated. Note that the diameter of the axon increases significantly where it is myelinated. Location of last paranodal loop of myelin is indicated by narrow, peach-colored band to right of the blue arrow on the SWC representation. The view then pans to a dorsolateral view of the cell, where a pink arrow indicates the proximal dendrite, and an orange arrow signifies the primary hub of the cell. The view then pans to a top-down location showing two secondary hubs (orange arrows). The view then shifts to reveal periodic dendrite swellings (three cyan arrows) separated by dendrite shafts (two red arrows). SWC color code: pink, cell body; axon hillock, orange; axon initial segment, light green; myelinated axon, light blue; proximal dendrite, maroon; dendrite hub, greenish-brown; dendrite swelling, cyan; dendrite shaft, gray.
with ionic conductances, assigned endbulb synaptic inputs accounting for synaptic weights, and simulated the activation of ANFs with arbitrary sounds (see Methods and Figure 3C).

Cell models were adjusted by mimicking in vitro measurements for $k_{il}$ to set channel densities (see Figure 3–Figure Supplement 2). The Half-active parameter set, in which dendrite conductances had values reduced by one-half relative to the soma, yielded BC-like phasic responses to current injection, a voltage sag in response to hyperpolarizing current and a non-linear IV plot (Figure 3D,E and Figure 3–Figure Supplement 3), with no further tuning. The Passive and Active parameter sets also resulted in phasic responses in most cells, although with passive dendrites, some cells showed trains of smaller spikes with stronger current injections, or 2-3 spikes with weaker currents (BCs 09, 10, 11 and 30). Similar patterns of repetitive spiking have been reported in BCs previously (Francis and Manis, 2000; Cao et al., 2007) Rebound spikes were larger and more frequent when the dendrites were passive, but were suppressed when dendrites were fully Active dendrites (BCs 05, 06 and 10).

Cell input impedance was assessed as a function of frequency for intact cells, cells without dendrites, and for the axon only (Figure 3–Figure Supplement 4). This analysis revealed a slightly peaked low-pass response for complete reconstructions. The peak was larger and narrower when the dendrites were removed, and was also larger when looking at the axon from the somatic end. This underscores the importance of including a full dendritic reconstruction when modeling BCs.

The number of synapses in each endbulb was based on ASA and average synapse density (Figure 1F). Terminal release was simulated with a stochastic multi-site release model in which each synapse in the terminal operated independently (Xie and Manis, 2013; Manis and Campagnola, 2018). Synaptic conductances were not tuned, but instead calculated based on experimental measurements (Manis and Campagnola, 2018). Action potentials (AP) (marked by red dots in Figure 3D,F) were detected based on amplitude, slope and width at half-height (Hight and Kalluri, 2016). For our model, ANFs were driven in response to arbitrary sounds via spike trains derived from a cochlear model (Zilany et al., 2014; Rudnicki et al., 2015) (Figure 3C, right). These spike trains generated primary-like (Pri) responses in ANFs and yielded Pri or primary-like with notch (Pri-N) responses in our BC models (Figure 3F-G; Figure 3–Figure Supplement 5). The standard deviation of the first spike latency, and the regularity were similar to values reported for mouse CN (Roos and May, 2012). These standardized implementations will underlie future computational studies of sound-encoding by cell populations.

Application of the model

Endbulb size does not strictly predict synaptic efficacy

The wide variation in size of the endbulb inputs (Figure 2C-F) suggests that inputs with a range of synaptic strengths converge onto the BCs. We then inquired whether individual cells followed the coincidence-detection or mixed-mode models hypothesized by input sizes shown in Figure 2D.

To address this question, we first modeled the responses by each of the 10 fully reconstructed BCs as their endbulb inputs were individually activated by spontaneous activity or 30 dB SPL, 16 kHz tones (responses at 30dB SPL for four representative cells (BC05, 30, 09, and 17 are shown in Figure 4A; the remaining BCs are shown in Figure 4–Figure Supplement 1). In Figure 4A, voltage responses to individual inputs are rank-ordered from largest (1) to smallest (7,8,or 9) for each cell. Without specific knowledge of the spontaneous rate or a justifiable morphological proxy measure, we modeled all ANFs as having high spontaneous rates since this group delivers the most contacts to GBCs in cat (Fig.9 in Liberman, 1991).

BC05 and BC30 (Figure 4A1, A2) fit the coincidence-detection model, in that none of their inputs individually drove postsynaptic APs except the largest input for BC30, which did so with very low efficacy (#postsynaptic APs/#presynaptic APs; see also BC10, BC06, BC02, BC13 in Figure 4 Supplement 1). BC09 and BC17 (Figure 4A3, A4) fit the mixed-model, in that the largest inputs (2
**Figure 4.** Efficacy of single ANF inputs is variable, depends on dendritic area, and participation of individual inputs in spike generation is rate dependent. (A1–A4) Simulations showing EPSPs and spikes in response to individual ANFs in 4 model BCs during a 30 dB SPL tone pip, arranged by efficacy of the largest input. Spikes indicated by red dots. Vertically, traces are ranked order by endbulb size. Responses for the other 6 cells are shown in Figure 4—Figure Supplement 1. (B1–B4) Reverse correlations between postsynaptic spikes and spikes from each input ANF during responses to 30 dB SPL tones (all inputs active). Trace colors correspond to the ASA of each input (color bar in (B1)). See Figure 4—Figure Supplement 1 for reverse correlations for the other 6 cells. (C1–C4): Voltage traces aligned on the spike peaks for each of the 4 cells in (B). Postsynaptic spikes without another spike within the preceding 5 ms were selected to show the subthreshold voltage trajectory more clearly. Zero time (0 ms; indicated by vertical red line) corresponds to the 0 time in (B1–B4). (D) BCs could be divided into two groups based on input efficacies. BCs 09, 11, 13, and 17 formed one group (“Group 1”), and BCs 02, 05, 06, 10, 18 and 30 formed a second group (“Group 2”). Red and light blue lines are best fits of logistic functions to the two groups. Stars indicate test ASA-efficacy points supporting group membership for cells 10, 17 and 30. (E) Comparison of the patterns of individual inputs that generate spikes. Ordinate: 1st+ indicates spikes driven by the largest input plus any smaller inputs, but not the 4 largest. 2nd+ indicates spikes driven by the second largest input plus any smaller inputs, excluding spikes in which the largest input was active. 3rd+ indicates spikes driven by the third largest input plus any smaller inputs, but not the first and second largest inputs. 4th+ indicates contributions from the fourth largest input plus any smaller inputs, but not the 3 largest. 5th+ indicates contributions from the fifth largest input plus any smaller inputs, but not the 4 largest. See Figure 4—Figure Supplement 3 for a additional summaries of spikes driven by different input patterns. (F) The participation of weaker inputs (smaller terminal area) is increased during driven activity at 30 dB SPL relative to participation during spontaneous activity. The dashed line indicates equal participation at the two levels. Each input is plotted separately. Colors and symbols are coded to individual cells as in (E). (G) Cumulative distribution of the number of inputs driving postsynaptic spikes during spontaneous activity and at 30 dB SPL. The color for each cell is the same as in the legend in (D). Symbols correspond to the stimulus condition. (H) Efficacy for a single 150 150 μm² input is inversely related to dendrite surface area. (I) Dendrite area and action potential threshold are highly correlated. Open circles (BC02 and BC05) were calculated using the average axon lengths, but are not included in the calculation. Colors and symbols as in (E). Figure 4—Figure Supplement 2 shows additional analyses of morphology, threshold and efficacy.

**Figure 4—Figure supplement 1.** Reverse Correlation plots for 6 additional modeled cells.

**Figure 4—Figure supplement 2.** Correlations between morphological parameters, efficacy, and action potential threshold.

**Figure 4—Figure supplement 3.** Contributions of different input patterns to postsynaptic spiking.
large inputs for BC17) individually drive APs with high efficacy (see also BC11, BC18 in Figure 4–
Figure Supplement 1). This result demonstrates two populations of BCs based on the absence or
presence of high efficacy suprathreshold inputs.

The second largest input for BC09 (132 μm²) had higher efficacy than the largest input for BC30
(172 μm²). The variation of efficacy for similar ASA was evident, especially between (125 – 175 μm²),
in a plot of all inputs across the ten BCs (Figure 4D). Since many cells lacked inputs in this range,
we created 3 different sizes of artificial synapses (150, 190 and 230 μm²) onto BCs 10, 17 and 30 to
predict the efficacy of a more complete range of input sizes. The addition of these inputs (stars
colored for each cell) reinforced the suggestion that there were two populations of BCs, of greater
(BCs 09, 11, 13, 17; red curve) or lesser excitability (BCs 02, 05, 06, 10, 18, 30; cyan curve). Therefore,
we combined all synapses (excluding the artificial synapses) from BCs 09, 11, 13 and 17 into
one group, and synapses from all the remaining cells, BCs 02, 05, 06, 10, 18 and 30, into a second
group. We then confirmed the efficacy data by fitting each group with logistic functions with
distinct parameters (Figure 4D). The group with the greater excitability had half-maximal size for
input ASAs of 148.6 (SD 1.1) μm² and a maximal efficacy of 0.72 (SD 0.01) μm², with a slope factor
of 14.3 (SD 1.1)/μm². The fit to the group with lesser excitability (Figure 4C, light blue line) yielded
a half-maximal size of 204.3 (SD 4.7) μm², and with a slope factor of 19.8 (SD 2.2)/μm². Cells with
lesser and greater excitability were found in both the coincidence-detection (lesser: BC02, 05, 06,
10 30; greater: BC13) and mixed-mode (lesser: BC18; greater BC09, 11, 17) categories described
above.

Mixed-mode cells operate in first-arrival and coincidence-detection modes when all in-
puts are active.

We next computed BC responses when all ANFs to a model cell were driven at 30dB SPL and active
at the same average rate of ~200 Hz, to provide the predictions about how interactions among
individual inputs contribute to initiation of postsynaptic APs. To accomplish this, we computed the
reverse correlation between the postsynaptic spikes and each individual input. These simulations
are summarized in Figure 4B–C, for the 4 cells shown in Figure 4A.

For BC05 and BC30, which had no suprathreshold inputs, all inputs had low coincidence rates.
However, not all inputs had equal contribution in that the largest input had a rate 3-4 times the rate
of the smallest input Figure 4B1, B2). In both cells the requirement to integrate multiple inputs was
evident in voltage traces exhibiting EPSPs preceding an AP (Figure 4C1, C2). BC09 and BC17 illustrate responses when cells have one or two secure suprathreshold inputs, respectively (Figure 4A3,
A4). The reverse correlation plots reveal the dominance of high probability suprathreshold inputs
in generating APs in BCs (yellow traces for BC09, 17). For BC09 but not BC17 (likely because it
has two suprathreshold inputs), all subthreshold inputs had appreciable coincidence rates. The
summation of inputs to generate many of the APs for BC09 is seen in the voltage traces preced-
ing spikes, but most APs for BC17 emerge rapidly without a clear preceding EPSP (Figure 4C3, C4,
respectively).

We further investigated the extent to which subthreshold inputs generated spikes, focused ini-
tially on mixed-mode cells when subthreshold inputs generated spikes in the absence of activity in
suprathreshold inputs (Figure 4E). We first calculated the fraction of postsynaptic spikes generated
by the largest input in any combination with other inputs. The preceding integration window for
inputs (-2.7 to -0.5 ms relative to the spike peak) was the same as that used for the reverse corre-
lation measures in Figure 4B). This fraction ranged from 40-60% in mixed-mode cells (hexagons,
1st+ in Figure 4E). The fraction of postsynaptic spikes generated by the second-largest input in
any combination with other smaller inputs was surprisingly large, ranging from 25-30% (excluding
BC17 which had 2 suprathreshold inputs; 2nd+ in Figure 4E). Notably, all combinations of inputs in-
cluding the 3rd largest and other smaller inputs accounted for about 25% of all postsynaptic spikes.
Thus, a significant fraction (about 50%) of postsynaptic spikes in mixed-mode cells were generated by combinations of subthreshold inputs operating in coincidence detection mode.

We next investigated the BCs operating in the coincidence-detection mode, expecting a more even distribution across the combinations of input sizes. Surprisingly, in several cells, the largest input in combination with all other input sizes also accounted for about 50% of all postsynaptic spikes (circles, 1st+ in Figure 4E). In the two cells with the largest inputs (BC02, BC30), this category accounted for a larger percentage of postsynaptic spikes than any of the mixed mode cells. These data illustrate the differential weights of subthreshold inputs, and that their size distribution factors into the activation profile for the postsynaptic target.

We then inquired whether the participation of weak inputs in AP generation varied across intensity (spontaneous activity at 0 dB SPL and driven activity at 30 dB SPL), or was normalized by the increase in postsynaptic firing rate. To address this question, we computed a participation metric for each endbulb as \#postsynaptic APs for which a presynaptic AP from a given input occurred in the integration window (-2.7 to -0.5 ms relative to the spike peak), divided by the total number of \#postsynaptic APs. Interestingly, smaller inputs have a higher relative participation at 30 dB SPL than larger inputs (Figure 4F), suggesting a rate-based increase in coincidence among weaker inputs. The level-dependent role of smaller inputs was also explored in cumulative probability plots of the number of inputs active prior to a spike between spontaneous and sound-driven ANFs. During spontaneous activity, often only one or two inputs were sufficient to generate an AP (Figure 4H, triangles). However, during tone-driven activity postsynaptic spikes were, on average, preceded by coincidence of more inputs (Figure 4H, filled circles). We conclude that mixed-mode cells can operate in either latency or coincidence-detection modes. Which mode predominates depends on the average afferent firing rates of the individual inputs (sound level dependent), and the distribution of input strengths. Furthermore, BCs operating in the coincidence-detection mode show similar participation bias toward their largest inputs.

Contribution of soma and dendrite area to bushy cell excitability

Although synapse ASA and size distributions play a critical role in innervation, the response to synaptic input also depends on postsynaptic electromorphology, which creates distinctions amongst cells by affecting their threshold to synaptic input. To further clarify these differences in excitability, we examined somatic and dendritic surface areas. The BC dendrite area spanned a broad range from 3000 - 4500 \(\mu m^2\). Interestingly, the BCs having the smallest dendrite surface area comprised the group with greater excitability as measured by current threshold and efficacy of a standard-ized 150 \(\mu m^2\) input (Figure 4H), revealing an important mechanism by which BCs can modulate their excitability. The large difference in excitability between BC17 and BC05 (Figure 4H), which have similar surface areas, indicates that other mechanisms are needed to explain these data fully. Somatic surface area was a lesser determinant of excitability, and was weakly correlated with dendrite surface area. Threshold was also only weakly correlated with soma area or the ration of dendrite to soma areas (Figure 4-Figure Supplement 2B,C).

Axon initial segment length modulates BC excitability

Another factor that can regulate excitability is the length of the AIS. Therefore, in the EM volume we also quantified the lengths of the axon hillock, defined as the taper of the cell body into the axon, and the axon initial segment (AIS), defined as the axon segment between the hillock and first myelin heminode. The axon hillock was short (2.3 (SD 0.9) \(\mu m\); measured in all 21 BCs with reconstructed endbulbs). The AIS length averaged 16.8 (SD 6.3) \(\mu m\) in length (range 14.2-21.4 \(\mu m\); \(n = 16\), the remaining five axons exited the volume before becoming myelinated) and was thinner than the myelinated axon. Because the conductance density of Na\(^+\) channels was modeled as constant across cells, the AIS length potentially emerges as a parameter affecting excitability. To characterize...
this relationship, in the 10 BCs used for compartmental modeling, we replaced the individual axons
with the population averaged axon hillock and initial myelinated axon, and parametrically varied
AIS length. Indeed, for each cell the threshold to a somatic current pulse decreased by nearly
40% with increasing AIS length across the measured range of values (Figure 4). However, using
each cell’s own AIS we found that although threshold varied by cell, the threshold and AIS length
were not significantly correlated (Figure 4—Figure Supplement 2E). This indicates that AIS length
and dendrite area together serve as mechanisms to tune excitability across the BC population,
although dendrite area has a greater effect.

In 20 of 21 cells for which all large inputs were reconstructed, at least one endbulb terminal
(range 1-4) extended onto the axon (hillock and/or the AIS), contacting an average of 18.5 (SD 10) \(\mu\)m²
of the axonal surface (range 0.7-35.2\(\mu\)m²). The combined hillock/initial segment of every cell was
also innervated by 11.8 (SD 5.6) smaller terminals (range 4 – 22; \(n = 16\)). These innervation fea-
tures will be further explored once the excitatory and inhibitory nature of the inputs, and the SR
of endbulb terminals are better understood.

**Innervation of BCs shows specificity for ANF fiber fascicles**

As indicated previously, most end bulbs could be traced along axon branches to parent ANFs con-
stituting fascicles within the image volume (98/158). Remaining branches exited the volume (2/6
and 3/8 branches (white arrowheads), respectively, for example cells in Figure 5 A, B). We then
asked if fascicle organization of the auditory nerve root related to innervation patterns, whereby
most inputs to a particular cell may associate in the same fascicle. We identified nine fascicles in
the image volume, containing in total 1,100 axons (section taken through middle of volume), which
is 7-15% of the total number of ANFs in mouse (7,300-16,600) (Burda et al., 1988; Anniko and Arnesen, 1988; Camarero et al., 2001). The largest five fascicles (containing between 115 and 260 axons/
fascicle) each split into as many as seven sub-fascicles along their trajectory (Figure 5A,B). Exclud-
ing 4 cells near the edge of the image volume (BCs 02, 24, 29, 14 plotted at left in right histogram
of Figure 5C), 2-9 endbulbs from individual cells were traced to ANFs in the same major fascicle (2
fascicles each contained 2 parent axons of inputs to each cell (fascicles 2, 3, and 2, 7, respectively)
in Figure 5A, B). None of the parent ANFs that were linked to endbulbs branched more than once
within the volume. The proportion of axons yielding endbulb terminals within the image volume
was low in some fascicles (#3, #4, #5, #6; #4 contributed no endbulbs), and high in others (#1, #7;
BC08 had 9 endbulbs traced to fascicle 7) indicating that fascicles preferentially innervate different
rostro-caudal regions of the same frequency region (Figure 5D).

The myelinated lengths of branches varied from 0 (endbulbs emerged en passant from parent
terminal in two cases) to 133 \(\mu\)m (Figure 5G). We then inquired whether the variance in length
might introduce sufficient conduction delay to decorrelate the somatic inputs. For a subset of 10
BCs with at least 4 branches traced to ANFs, we utilized the resolution and advantages of volume
EM to assay axon parameters. Branches were thinner than the parent ANF, (1.4 (SD 0.33) vs 2.7 (SD
0.30) \(\mu\)m), and both the parent ANF and branches had the same g-ratio of fiber (including myelin) to
axon diameter (Fig. 5F; ratio 0.76 across all axons). From these data we applied a conversion of 4.6
* fiber diameter in \(\mu\)m (Boyd and Kalu, 1979; Waxman and Bennett, 1972) to the distribution of fiber
lengths, yielding a conduction velocity range of 2.3 - 8.9 m/sec, and a delay range of 0 (en passant
terminal) - 15.9 \(\mu\)m. These values were then scaled by the \(L/d\) ratio, where \(L\) is the length between
the ANF node and the terminal heminode, and \(d\) is the axon diameter (Brill et al., 1977; Waxman,
1980). The \(L/d\) ratio slows conduction velocity to a greater extent in short branches, yielding a
latency range of 0-21 \(\mu\)s across the cell population, and a similar range among different branches
to individual cells (Figure 5G). Such small variations in delays could have a small effect on the timing
of spikes at sound onset, which can have a standard deviation of 0.39 ms in mouse (Roos and May,
2012), similar to values in cat (Young et al., 1988; Blackburn and Sachs, 1989; van Gisbergen et al.,
Figure 5. Large somatic terminals link to auditory nerve fibers (ANFs) through myelinated branch axons of varying length and fascicle organization. (A, B). ANFs and their branches leading to all large inputs for two representative cells. ANF, branch axon and large terminal have same color; each composite structure is a different color. Convergent inputs emerge from multiple fascicles (fascicles circled and named on back left EM wall), but at least two inputs emerge from the same fascicle for each cell (green, purple axons from fascicle 3 in panel (A); yellow, mauve axons from fascicle 7 and green, purple from fascicle 2 in panel (B)). Some branch axons leave image volume before parent ANF could be identified (white arrowheads). Bushy cell (BC) bodies colored beige, dendrites red, axons mauve and exit volume at back, right EM wall; axon in panel A is evident in this field of view. (C) Stacked histogram of branch axons traced to parent ANF (black), branch axons exiting volume without connection to parent ANF (open), small terminals linked to parent ANF (red; included to illustrate these were a minority of endings), arranged by increasing number of large terminals traced to a parent ANF per BC. BCs with fewest branch connections to parent ANF (BC02, 24, 29, 14) were at edge of image volume, so most branch axons could only be traced a short distance. Number of terminals per cell indicated in horizontal histogram at left. (D) Number of axons in each fascicle (left ordinate) and number of axons connected to endbulb terminals per fascicle (red symbols, right ordinate). (E) Example of en passant large terminal emerging directly from node of Ranvier in parent ANF. (F) Constant ratio of fiber diameter (axon + myelin) / axon diameter as demonstrated by linear fit to data. All branch fiber diameters were thinner than ANF parent axon diameters. (G) Selected cells for which most branch axons were traced to a parent ANF. Lines link the associated conduction delays from parent ANF branch location for each branch, computed using the individual fiber diameters (length / conduction velocity (leftmost circle, vertical dashed arrows) or values scaled by the axon length / axon diameter (rightmost circle, vertical solid arrow).

Figure 5–video 1. Exploration of all large somatic inputs onto a single bushy cell (BC), their branch axons, and parent auditory nerve fibers. This video opens with a full view of BC11, including its dendrites (red), cell body (beige), axon (pink), all of its large somatic inputs (various colors), and the auditory nerve fibers (ANFs) from which those inputs originate. Somatic inputs and their axon branches, including the parent ANF, share the same color. The view zooms into the cell body, where a large terminal (red) extends onto the axon hillock and initial segment (blue arrow). The view pans around the cell body to show all large terminals, and an axon (cyan) that exited the volume before likely linking to an ANF is indicated by a red arrow. All except two terminals are removed. The axons of these terminals are traced to branch points from their parent ANFs (magenta arrows). All inputs and their axons are replaced, and a second axon (green) that exits the volume before likely linking to an ANF is indicated (red arrow). The view pans to two ANFs (red and yellow) from the same nerve fascicle, and their branch points (magenta arrows). All other nerve terminals and axons are removed, and the two branch axons are traced to the cell body.
Although mice do not have a prominent population of low-frequency neurons that can encode auditory neurons can exhibit precisely-timed spikes in response to different features of sounds. (7 branches visible in Figure 6 µ (although one dendrite branched after 1.8 (total hubs/cell: range 4-13). One-half (11/22 BCs) of primary, and some secondary hubs contained additional summary metrics of dendrite structure. A novel observation was that nearly all dendritic trees exhibited regions where branches extended alongside one another and could exhibit braiding, whereby branches of the same or different parent branches intertwined. Dendrites were partitioned qualitatively into categories of little (n = 3), moderate (n = 4) and dense (n = 3) local branching and braiding (Figure 6A-C, respectively). EM images reveal the complexity of braided branches and frequent direct contact between them (Figure 6D-F, I).

Proximal dendrites expanded into a novel structure, from which at least 3 and up to 14 branches extend (7.0 SD (3.8), n = 10). We name these structures hubs, due to their high node connectivity (7 branches visible in Figure 6G). Secondary hubs were positioned throughout the dendritic tree (total hubs/cell: range 4-13). One-half (11/22 BCs) of primary, and some secondary hubs contained a core of filaments that extended through the middle of the structure. This filamentous core was in contact with multiple mitochondria oriented along its axis (Figure 6; and Figure 6–Figure Supplement 1), and was also found in a thickened region of a second order dendrite of one of the two large MCs. Dendrites, as noted previously, have many swellings (Figure 6) along higher order branches. Swellings were more numerous than (range 51-126, mean = 74.9 (SD 26.8)), and did not correlate with the number of hubs (r² < 0.001; Fig. 6H). In rank order, dendrite surface area was comprised of dendritic shafts (58%), swellings (28%), hubs (10%) and the proximal dendrite (4%). (Figure 6K).

Temporal precision of BCs varies by patterns of ASA
Auditory neurons can exhibit precisely-timed spikes in response to different features of sounds. Although mice do not have a prominent population of low-frequency neurons that can encode temporal fine structure, they do have both behavioral (Cai and Dent, 2020) and physiological (Kopp-Scheinpflug et al., 2003; Walton et al., 2002) sensitivity to sinusoidal amplitude modulation (SAM) in the range from 10-1000 Hz on higher-frequency carriers. As amplitude modulation is an important temporal auditory cue in both communication and environmental sounds, we used SAM to assess the temporal precision of BC spiking which has been reported to exceed that of ANFs (Joris et al., 1994a; Louage et al., 2005; Frisina et al., 1990). Because temporal precision also exists for transient stimuli, we additionally used click trains. Given the variation of mixed-mode and coincidence-detection convergence motifs across BCs, we hypothesized that their temporal precision would differ across frequency and in relation to ANFs. The left columns of Figure 7 illustrate the flexibility of our modeling pipeline to generate and analyze responses to arbitrary complex sounds in order to test this hypothesis. SAM tones were presented with varying modulation frequency and a carrier frequency of 16 kHz at 15 and 30 dB SPL (see Figure 7–Figure Supplement 1 for comparison of SAM responses in ANFs and a simple BC model across stimulus intensities), and
Figure 6. Novel dendrite structure of BCs. (A-C). Dendrites vary in density of local branching and braiding of branches from the same cell, exhibiting (A) little, (B) medium or (C) dense branching and braiding. (D-E). Tangential view of dense braiding, showing (D) reconstruction of multiple branches in contact with one another and (E) a single EM cross section illustrating contact among the multitude of branches (individual branches identified with asterisks). (F, I). Two locations of cross-cut braided dendrites showing intertwining as change in location of branches (numbers) along the length of the braid. Images are lower resolution because viewing perspective is rotated 90° from image plane. (G). Reconstruction of dendrite hub (h) and its multiple branches (7 are visible and numbered in this image). (H). Swellings and hubs are prominent features of BC dendrites. Histograms of numbers of swellings and hubs plotted along abscissa and ordinate, respectively. (J). Core of many hubs is defined by a network of filaments (f); also see Figure 6–Figure Supplement 1. Many mitochondria are found in hubs and can be in apparent contact with the filament network. (K). Partitioning of dendrite surface area reveals that proximal dendrite (black), hub (light grey), swelling (dark grey) and shaft (medium grey) compartments, in increasing order, contribute to the total surface area for each cell. Averaged values indicated in stacked histogram, to right of vertical dashed line, as percent of total surface area (right ordinate), and aligned with mean sizes on left ordinate).

Scale bars: E, 2 microns; F, I, 0.5 microns; J, 1 micron.

Figure 6–Figure supplement 1. Dendritic Hubs
60Hz click trains were presented at 30 dB SPL. We implemented a standard measure of temporal fidelity (vector strength) for SAM stimuli. To analyze temporal precision of click trains, we used the less commonly employed shuffled autocorrelogram (SAC) metric, which removes potential contribution of the AP refractory period to temporal measures (Louage et al., 2004).

Here, we illustrate a representative range of cellular responses and analytics available in our pipeline, from intracellular voltage traces (Figure 7A, H) recorded in any cellular compartment (cell body depicted here), to event data with associated representations as raster plots and period histograms. BCs exhibited a more temporally-constrained distribution of BC spikes in response to SAM tones and click trains (Figure 7B-F, I-M, respectively, shown for BC17) relative to ANFs. Measures of temporal precision demonstrate an improvement between ANFs and BC responses to SAM tones (higher VS in Figure 7F). The responses to clicks consist of well-timed spikes, followed by a short refractory period before the ANF spontaneous activity recovers and drives the cell (Figure 7, L). The precision of responses to clicks is also better (narrower SAC half-width) in the BCs than in their ANF inputs (Figure 7M).

We then compared responses of BCs to ANFs across a range of modulation frequencies from 50 - 1000 Hz at 15 dB SPL, which revealed the tuning of BCs to SAM tones. BCs had higher VS at low modulation frequencies (< 300 Hz), and lower VS at higher modulation frequencies (> 300 Hz). Responses varied by convergence motif, whereby coincidence-detection BCs had enhanced VS relative to ANFs at 100 and 200 Hz (Figure 7O1-O2, BC02 and BC30), but mixed-mode BCs only at 200 Hz (Figure 7O3-O4, BC09 and BC17). To better understand how different ASA input patterns contributed to the VS, we ran additional simulations in which only the largest input was active or where we removed the largest input, leaving the remaining inputs active (and in BC17, removing the two largest inputs, since both were suprathreshold). For coincidence-detection BCs, activating only the largest input yielded VS values higher than ANFs at 50 and 100 Hz and much lower than ANFs for modulation frequencies greater than 00 Hz. BC02 and BC30 are illustrated here because their largest inputs could generate postsynaptic spikes but with low efficacy. For the mixed-mode BCs, activating only the largest input yielded responses closer to ANFs at all frequencies. For the coincidence-detection BCs, removing the largest input increased VS only at 50 Hz, likely because the cells are already operating as coincidence detectors. For the mixed-mode BCs, removing the largest input (or two largest for BC17) improved the VS at 100 Hz, resulting in similar frequency tuning to the coincidence detection BCs. Figure 7P1,P2 compare the SAM VS with activation of all inputs, the removal of the largest input (or two largest for BC17), or the largest input only, across all cells, at 15 and 30 dB SPL. The VS is higher when the subthreshold inputs are active, compared to when the largest input alone is active, for frequencies between 100 and 500 Hz at 15 dB SPL and between 200 and 500 Hz at 30 dB SPL. The VS is lower at 50 and 100 Hz at 30 dB SPL because of peak-splitting in ANF phase locking. Across the population, the results are mixed at 750 and 1000 Hz. These simulation results again highlight the contributions of subthreshold inputs in improving temporal precision compared to the suprathreshold inputs alone. Similarly, improvements in temporal precision were evident in response to click trains Figure 7-Figure Supplement 3. The half-widths of the SACs (when there were sufficient spikes for the computation) were consistently narrower and had higher correlation indices when all inputs, or all but the largest input were active than when only the largest input was active. The coincidence-detection BCs showed the highest correlation indices and slightly narrower half-width (Figure 7-Figure Supplement 3). Taken together, the different convergence motifs yielded a range of tuning (mixed-mode BCs more tuned) to the modulation frequency of SAM tones in comparison to ANFs. Notably, the mixed-mode BCs with the most pronounced tuning were those whose inputs most easily excited their postsynaptic BC (Figure 4), because their response at 100 Hz was similar to that of ANFs. Thus, the ANF convergence patterns play an important role in setting the temporal precision of individual BCs.
Figure 7. Temporal precision of spikes depends on the pattern of input ASA. Left column (A-G): Example of entrainment to 100% modulated SAM at 200 Hz, at 15 dB SPL. The sound level was chosen to be near the maximum for phase locking to the envelope in ANFs (see Figure 7–Figure Supplement 1). (A) Voltage showing spiking during a 150 ms window starting 300 ms into a 1-second long stimulus. (B) Spike raster for 100 trials shows precise firing. (C) PSTH for the spike raster in (B). (D) Spike raster for all ANF inputs across a subset of 5 trials. Inputs are color coded by ASA. (E) PSTH for the ANF. (F) Superimposition of the phase histograms for the BC (black) and all of its ANF inputs (red). (G) Stimulus waveform. Center column (H–N): responses to a 50 Hz click train at 30 dB SPL. (H) BC membrane potential. (I) Raster plot of spikes across 25 trials. (J) PSTH showing spike times from I. (K) ANF spike raster shows the ANFs responding to the clicks. (L) PSTH of AN firing. (M) The shuffled autocorrelation index shows that temporal precision is greater (smaller half-width) in the BC than in the ANS. See Figure 7–Figure Supplement 2 for SAC analysis of other cells. (N) Click stimulus waveform. Right column (O–P): Summary plots of vector strength. (O1–4) Vector strength as a function of modulation frequency at 15 dB SPL for 3 (4 for BC17) different input configurations. Vertical lines indicate the SD of the VS computed in (Figure Supplement 3 as described in the Methods. Red line: average ANF VS. See Figure 7–Figure Supplement 3 for the other cells. Legend in )O1 applies to all panels in (O) and (P). (P) Scatter plot across all cells showing VS as a function of modulation frequency for 3 (4 for BC17) different input configurations. (P1) VS at 15 dB SPL. (P2) VS at 30 dB SPL.

Figure 7–Figure supplement 1. Synchronization to stimulus envelope as a function of average stimulus intensity in ANF inputs.

Figure 7–Figure supplement 2. Vector strength of 6 other BCs in response to 100% SAM modulation at frequencies from 50 to 1000 Hz on a 16kHz carrier at 15 dB SPL.

Figure 7–Figure supplement 3. SACs in response to click trains show importance of weaker inputs in improving temporal precision.
Complete map of synaptic inputs onto a BC

We report here the first map for locations of all synaptic terminals onto soma, dendrites and axon of a BC (BC09; Figure 8A,B). In addition to 8 endbulb inputs from ANFs, 97 small terminals contacted the cell body. Together these inputs covered most 83% of its somatic surface (Figure 8C, D). This neuron had 224 inputs across all dendritic compartments (shaft, swelling, hub, proximal dendrite) (Figure 8). Dendritic and small somatic terminals were typically bouton-sized, contained one or two synaptic sites, and could be linked by small caliber axonal segments to other small terminals across the dendrite and/or soma (Figure 8A) (cyan arrowheads in Figure 8A’, C). Previous investigation suggested swellings as preferred sites for innervation (Ostapoff and Morest, 1991). However, in our reconstruction, innervation density was similar across most compartments (hubs, 10.4/100 μm²; swellings, 9.3/100 μm²; shafts 9.1/100 μm²), and greatest on the proximal dendrite (24/100 μm²; Figure 8A, E, G, H). At least one endbulb (typically 1 but up to 3) on nearly all BCs (20/21) extended onto the proximal dendrite (mean = 14.5% of endbulb ASA; black arrow in Figure 8A). Two endbulbs extended onto axonal compartments of BC09, indicating that this cell is not exceptional. Somatic endbulbs infrequently (8/159 terminals) innervated an adjacent dendrite of a different BC.

Notably, entire dendrite branches could be devoid of innervation (black arrows in Figure 8B), and instead were wrapped by glial cells, or extended into bundles of myelinated axons (Figure 8F). Even though they are not innervated, these branches will affect the passive electrical properties of the cell by adding surface area. We inquired whether these dendrites constitute sufficient surface area and are strategically located to affect excitability of the cell, by generating a model of BC09 with the non-innervated dendrites pruned. Pruning increased the input resistance from 20.2 to 25.1 MΩ (Figure 8I, J) and increased the time constant from 1.47 ms to 1.65 ms. The threshold for action potential generation for short current pulses decreased from 0.439 to 0.348 nA (Figure 8I), but the cell maintained its phasic firing pattern to current pulses (Figure 8I compared to Figure 3–Figure Supplement 3, "Half-active"). These seemingly subtle changes in biophysical parameters increased the efficacy for the 4 largest inputs (0.689 to 0.786 (14%); 0.136 to 0.431 (216%); 0.021 to 0.175 (733%); 0.00092 to 0.00893 (871%); Figure 8K, L). Note that the increase was fractionally larger for the 2nd and 3rd largest inputs compared to the first, reflecting a ceiling effect for the largest input. We also examined how pruning non-innervated dendrites affected phase locking to SAM tones (Figure 8M). Pruning decreased VS at 100 Hz, thereby sharpening tuning to 200 Hz relative to ANFs. From these simulations, we suggest that BCs can tune their excitability with functionally significant consequences by extension and retraction of dendritic branches, independent of changes in their synaptic map.
Non-innervated dendrites (d) can be embedded in bundles of myelinated (my) axons (ax), and also ensheathed by glial cells (gl) and their processes (lines). Both dendrite swellings (sw) and shafts (sh) can be innervated. (A) Cross section through primary hub (h), showing filamentous core (f), mitochondria (m), input terminals (asterisks), and contact with dendrite of another cell. (B) Top-down view of dendrites only, illustrating that some branches are not innervated (longest non-innervated branches indicated by arrows) and that other branches are innervated at varying density. (C) Bouton terminals innervate all regions of the cb surface. Some boutons linked by narrow connectors (cyan arrowheads), and extension of a somatic endbulb onto the basal dendrite (arrow). (D) Inside-out view of cb innervation by endbulbs (eb; each is numbered and a different color) reveals that they cover most of the cb surface. Cb removed to reveal synaptic face of eb's. (E) Proximal dendrites are innervated at highest density (number of inputs / surface area), and hubs, swellings and shafts are innervated at similar density. Scale bars: 1 μm in each panel. (I-M) Simulation results after pruning the non-innervated dendrites from this cell. (I) Voltage responses to current pulses, as in Figure 3–Figure Supplement 3, comparing the intact cell (black traces) with one in which non-innervated have been pruned (cyan traces). (J) IV relationship of data in (I) Cyan triangle indicates the spike threshold with the dendrites pruned compared to the intact cell (red circle). (K) Spikes elicited by the 4 largest individual inputs at 30 dB SPL with the dendrites pruned compared to data shown in Figure 4A3. (L) Comparison of the efficacy of individual inputs between intact and pruned as a function of ASA. The red and light blue lines (Group1 and Group2) are reproduced from Figure 4D. (M) Comparison of VS to SAM tones in the intact and pruned configuration. Minor ticks are at 200, 400 and 750 Hz.

**Figure 8–video 1.** Exploration of a bushy cell (BC) and all of its synaptic inputs. This video opens with a full view of BC09, including its dendrites (red), cell body (beige), axon (pink), all somatic inputs (various colors), and all dendritic inputs (various colors). The cell undergoes a full rotation to display all of the inputs. The view zooms into the axon region, and rotates to illustrate all inputs onto the axon, including extensions of two large terminals (blue arrows pointing to purple and yellow terminals). The view zooms out to show the entire cell, the dendrites are removed, and the cell body is tilted. A cut plane passes from the edge to the middle of the cell, providing an inside-out view of the nearly complete synaptic coverage of the cell body. Large terminals are indicated by cyan arrows. All cellular elements are added, and the view shifts to reveal dense innervation of the proximal dendrite, including an extension of a large terminal (green terminal indicated by green arrow). The perspective shifts to a top-down view of the dendrites, indicating several dendritic branches (yellow arrows) that lack synaptic inputs.
Discussion

**Volume EM provides direct answers to longstanding questions**

Key questions about ANF projections onto BCs have persisted since the first descriptions of multiple large terminals contacting their cell bodies (Lorente de Nó, 1933; Cajal, 1971). Volume EM offers solutions to fundamental questions about network connectivity not accessible by LM, by revealing in unbiased sampling all cells and their intracellular structures, including sites of chemical synaptic transmission (for reviews, see (Briggman and Bock, 2012; Abbott et al., 2020)). By acquiring nearly 2,000 serial sections and visualizing a volume of over 100 μm in each dimension, we provided reconstruction of the largest number of BCs to date, permitting more detailed analysis than was possible with previous EM methods that subsampled tissue regions using serial sections (Nicol and Walmsley, 2002; Spirou et al., 2008; Ostapoff and Morest, 1991). Here, we report on a population of BCs in the auditory nerve root with eccentric, non-indented nuclei, ER partially encircling the nucleus, and somatic contact by a large number (5-12) of endbulbs of mostly smaller size. These features, except for ER patterns, define a subpopulation of BCs in mice more similar to globular (G)BCs than spherical (S)BCs as defined in larger mammals (Cant and Morest, 1979b,a; Tolbert et al., 1982; Osen, 1969; Hackney et al., 1990), also see (Lauer et al., 2013).

Nanoscale (EM-based) connectomic studies are providing increasingly large volumetric reconstruction of neurons and their connectivity (Bae et al., 2021; Scheffer et al., 2020; Witvliet et al., 2021). In this report, we add pipelines from neuron reconstruction to biophysically-inspired compartmental models of multiple cells. These models expand on previous BC models that used qualitative arguments, or single or double (soma, dendrite) compartments (Joris et al., 1994b,a; Rothman et al., 1993; Rothman and Manis, 2003c; Spirou et al., 2005; Koert and Kuenzel, 2021). By matching inputs to a cochlear model (Zilany et al., 2014; Rudnicki et al., 2015), we created a well-constrained data exploration framework that expands on previous work (Manis and Campagnola, 2018). We propose that generation of compartmental models, from high-resolution images, for multiple cells within a neuron class is an essential step to understand neural circuit function. This approach also reveals that there are additional critical parameters, such as ion channel densities in non-somatic cellular compartments, including non-innervated dendrites, that need to be measured. From these detailed models, more accurate reduced models that capture the natural biological variability within a cell-type can be generated for efficient exploration of large-scale population coding.

**Multiple cellular mechanisms to tune excitability**

The variability of responsiveness in cells and patterns of convergence in circuits are essential factors that help optimize the representation of sensory information (Ashida et al., 2019; Perez-Nieves et al., 2021). In this study, all model cells were assigned the same densities of channels on a per-compartment basis, and we chose to mimic driving auditory nerve synapses by high SR, low threshold ANFs because they are most numerous and tend to be associated with larger endbulbs in this region of the VCN (Liberman, 1991; Rouiller et al., 1986; Wang et al., 2021). This set of metrics represents an experimentally anchored starting point in a larger parameter space. Further measures, including collection of larger image volumes at higher resolution, will permit finer specification of ANF subtypes by assessing axon branching patterns and endbulb shape and size (Sento and Ryugo, 1989; Wang et al., 2021). This approach also will permit identification of non-ANF local sources of excitatory and inhibitory inputs (Campagnola and Manis, 2014; Cant and Morest, 1978; Ngodup et al., 2020) known to play important roles in spectral and temporal processing of BCs (Caspary et al., 1994; Gai and Carney, 2008; Keine and Rübsamen, 2015; Keine et al., 2016). Characterization of inputs by their putative excitatory or inhibitory function based on vesicle shape is one important next step in the evolution of these detailed models. The differences between the responses...
of simulated cells in our current models are governed by and provide insight into their detailed electromorphology and the cell-specific patterns of ANF input sizes.

We revealed that dendrite surface area varies sufficiently to adjust spike threshold across the BC population. Dendrite surface area defined two BC populations whereby smaller values increased excitability. These two populations did not respect BC grouping based on the profile of endbulb sizes (coincidence-detection or mixed-mode) or the density of local dendrite branching. Our demonstration of the lack of dendrite innervation along entire branches and increased excitability following their removal, offers an additional mechanism to tune excitability. Although BCs lack dendritic spines, they may grow or prune dendritic branches sufficiently rapidly to adjust to alterations in the acoustic environment or pathological changes in the inner ear.

We also found that the length of the AIS, which is the spike initiation zone for most neurons (Bender and Trussell, 2012), varied across BCs by 50% (14-21 μm). Changing AIS length, while assuming a constant density of Na⁺ channels, non-linearly changed rheobase by 50% (Figure 4). Interestingly, the AIS of each BC is contacted by multiple small inputs. Inhibitory inputs onto the AIS of other neuron types have been shown experimentally and computationally to modulate spike generation (Bae et al., 2021; Schneider-Mizell et al., 2021; Veres et al., 2014; Franken et al., 2021).

We reveal that in nearly all BCs one of the large somatic inputs extends onto the hillock and AIS. In our models, the proximal axon is electrotonically close to the somatic compartment, so further investigation is required to determine whether direct AIS innervation can increase synaptic efficacy for driving spikes. The AIS length and location of Na⁺ channels have been also shown to be sensitive to the history of neural activity (Kuba et al., 2010; Kuba, 2012; Grubb and Burrone, 2010), and we merit investigation in BCs.

Dendrite surface area and AIS geometry and innervation emerge as potential homeostatic mechanisms to regulate excitability. We expect that reconstructions of a larger population of BCs will better reveal the distribution of excitability, and may clarify additional regulatory mechanisms. Thus, the combination of high-resolution structural analysis and compartmental modeling specifies focused topics for further study.

Convergence of weak and strong inputs regulates temporal fidelity

We provide the first complete catalogue of numbers of ANF inputs and their sizes (38 – 270 μm²), revealing a broad range of subthreshold endbulb sizes and raising questions about the functions of smaller endbulbs. GBCs were proposed to achieve their highest temporal fidelity by acting as a coincidence detector for convergence of subthreshold endbulb inputs (Rothman et al., 1993; Rothman and Young, 1996). In the present simulations, we took advantage of the ability to selectively silence different sets of inputs, which allowed us to separately assess the contribution of suprathereshold and subthreshold inputs across a biologically relevant range of strengths. Our simulations predict that only about one-half of GBCs in mice operate strictly in the coincidence detection mode, whereas the remainder operate in a mixed integration mode. Furthermore, we find that by conventional measures of phase locking to an amplitude-modulated tone, the activity of the weaker inputs substantially improves temporal precision relative to individual ANFs for modulation frequencies up to 200 Hz. In contrast, the largest inputs alone provide better temporal precision than combined inputs only at high modulation frequencies, especially if they are supratreshold. Supporting the generality of these observations across stimuli, improved temporal precision in the coincidence and mixed modes is also mirrored when using a different measure, the shuffled correlation index, for transient stimuli. Our results are also consistent with simulations showing that small ANF synapses on dendrites can improve temporal precision in the presence of large somatic inputs (Koert and Kuenzel, 2021). The simulations may also help explain the experimental cross-correlation results between ANFs and BCs of Young and Sachs (2008). In that study, prepotentials, which represent arriving spikes in ANFs with larger endbulbs, were strongly correlated with
putative BC spiking at low stimulus intensities, but these correlations became weaker at higher intensities. Our simulations suggest that the decrease in the ANF-BC correlation is a consequence of the increased coincidences between smaller (non-prepotential generating) inputs at higher intensities, which then drive an increased fraction of postsynaptic spikes relative to the stronger inputs. The pattern of convergence of ANF inputs with a wide range of strengths thus provides a mechanism for improved temporal precision over part of the range of behaviorally relevant envelope modulation frequencies.

New dendrite structures
Our high-resolution images revealed a new dendrite structure, which we name a hub. The high branching order of hubs helps explain why GBC dendrites are contained locally to the cell body. We also revealed that dendrites branch and align adjacent to one another. This arrangement increases the surface area to volume ratio, which affects the excitability of the cell. Both of these features likely function in part to shorten the overall dendrite electrotonic length and increase the importance of the dendrites in the integration of somatic synaptic inputs. Inspection of published BC images based on Golgi or tract tracing techniques reveals cells with thickened proximal dendrites (Webster and Trune, 1982; Lorente de Nó, 1981; Brawer et al., 1974). We suggest that some of these represent unresolved dense local branching and hub structures that are better revealed by EM across many sections. We noted that swellings were a prevalent feature of the dendrites and, contrary to reports in cat based on subsampling (Ostapoff and Morest, 1991), swellings were innervated at similar densities to shafts. The partition of dendrite compartments into hubs, swellings and shafts may have functional significance if, for example, these structures have differential sources of innervation or are endowed with different densities of ion channels or pumps (Brownell and Manis, 2014). The latter may relate to filament bundles and concentrations of mitochondria inside of hubs. Although our SBEM volumes lacked resolution to assess vesicle shape, it is likely that some of the smaller dendritic inputs are inhibitory (Gómez-Nieto and Rubio, 2009). Hubs may also provide efficient sites to nullify excitatory inputs occurring along multiple distal branches through current shunting. Many of the dendritic inputs were linked by short branches. Thus, non-innervated dendrites also afford locations for adaptive strengthening of inputs via formation of short branches and new terminals. These functional hypotheses derive from unbiased sampling of neural structure at high resolution which, in combination with a computational pipeline, fuels cycles of new hypotheses and new experiments that increase understanding of neuron types and their circuits.

Methods and Materials

Ethics Approval
All procedures involving animals were approved by the West Virginia University (WVU) Institutional Animal Care and Use Committee and were in accordance with policies of the United States Public Health Service. No animal procedures in this study were performed at other institutions.

Serial Block-Face Scanning Electron Microscopy
All reagents for transcardial perfusion were purchased from Sigma-Aldrich, unless otherwise noted. An adult (P60) FVB/NJ mouse (NCI; Frederick, MD and Jackson Laboratory; Bar Harbor, ME) was anesthetized using Avertin (20 mg/kg) injection IP, and perfused transcardially with normal Ringer’s solution containing xylocaine (0.2 mg/ml) and heparin (20 U/ml) for 2 min at 35°C followed by 0.15 M cacodylate buffer containing 2.5% glutaraldehyde (Polysciences), 2% paraformaldehyde (Fisher Scientific) and 2 mM calcium chloride at 35°C for 5 min. The skull was placed on ice for 2 hours, then the brain was removed from the skull and post-fixed for an additional 18 h at 4°C in the same
solution. Brain tissue was cut into 200-μm-thick sections in the coronal plane using a vibratome (Ted Pella) in ice-cold 0.15 M cacodylate buffer containing 2 mM calcium chloride, then washed for 30 min in the same solution. The ventral cochlear nucleus (VCN) was identified in free-floating sections using a stereo-microscope, and sections were photographed before and after dissection of the CN from the surrounding tissue.

The tissue sections were prepared for Serial Block-Face Scanning Electron Microscopy Imaging (SBEM) using an established protocol in our group (Holcomb et al., 2013). All staining and embedding chemicals were purchased from EM Sciences unless otherwise indicated, and all water was nanopure filtered (Nanopure Diamond, Barnstead International). Initial staining was performed in a solution combining 3% potassium ferricyanide in 0.3 M cacodylate buffer with 4 mM calcium chloride with an equal volume of 4% aqueous osmium tetroxide, for 1 h at room temperature (RT). Tissue was processed sequentially through filtered 1% thiocarbohydrazide for 20 min at RT, 2% osmium for 30 min at RT, and 1% uranyl acetate overnight at 4°C. Tissue underwent triple rinses in H₂O for 5 min each between each step and was triple rinsed in H₂O at RT for 30 min after the final step. Sections were placed into filtered lead aspartate solution (0.066g lead nitrate dissolved in 10 ml of 0.003 M aspartic acid solution, pH adjusted to 5.5 with 1N KOH, warmed in a 60°C oven for 30 min). The tissue was rinsed five times (3 min each), photographed, then dehydrated through graded alcohols into acetone, and flat-embedded in Durcopan resin (Electron Microscopy Sciences) between mylar strips in a 60°C oven for 48 h. Tissue samples were again photographed and shipped to the National Center for Microscopy and Imaging Research (University of California San Diego) for imaging.

Resin-embedded tissue was mounted on an aluminum specimen pin (Gatan) using cyanoacrylic glue and precision trimmed with a glass knife to a rectangle ≈0.5 × 0.75 mm so that tissue was exposed on all four sides. Silver paint (Ted Pella) was applied to electrically ground the edges of the tissue block to the aluminum pin. The entire specimen was then sputter coated with a thin layer of gold/palladium to enhance conductivity. After the block was faced with a 3View ultramicrotome unit (Gatan) to remove the top layer of gold/palladium, the tissue morphology became visible by back-scattered electron detector imaging using a Merlin scanning electron microscope (Carl Zeiss, Inc.). A low-magnification image (≈500X) was collected to identify the proper location in the VCN (caudal and in the auditory nerve root) for serial image collection. This region was selected because it has a high concentration of globular bushy cells (GBC, (Harrison and Irving, 1966; Osen, 1969; Brawer et al., 1974)). The imaged volume was located at approximately the mid dorsal-ventral location of the VCN. Imaging was performed using a pixel dwell time of 0.5 μs, tissue was sectioned at a thickness of 60 nm, and the imaging run required 7.5 days. Accuracy of section thickness was estimated by assuming circularity of mitochondria and comparing the diameter of longitudinally oriented organelles with diameters measured in the image plane (Wilke et al., 2013).

A volume of 148 μm x 158 μm x 111 μm was imaged with an in-plane pixel resolution of 5.5 nm. The image volume contained 31 complete cell bodies, including 26 GBCs. Due to the large size of the volume (1.4 TB) and the goal of reducing noise in the image, most of the analysis was performed by down-sampling in the image plane. Voxel averaging at 2 x 2 binning increased the dimensions of each voxel to 11.0 nm x 11.0 nm x 60.0 nm. With these imaging parameters, synaptic vesicles can be identified and, in many cases, a post-synaptic density, which appears as darkening on the post-synaptic membrane. Synapses were defined by collections of vesicles near the presynaptic membrane across at least 3 sections and with at least one vesicle in contact with the membrane (Jackson et al., 2021). Images were assessed to be of high quality for segmentation due to well preserved membranes, as evidenced also by uniform preservation of tightly wrapped myelin, and the absence of degenerating profiles.
Segmentation

Seg3D (https://www.sci.utah.edu/cibc-software/seg3d.html, University of Utah, Scientific Computing and Imaging Institute) was used to manually segment the structures of interest from the raw data volume. These structures (somata, nuclei, dendrites, axons, nerve terminals) were identified and segmented according to accepted morphological criteria for the mammalian CNS (Peters et al., 1997). The tracing tool was used to paint all pixels interior to the membrane. This strategy permitted the creation of 3D meshes for adjacent structures that did not overlap. Student segmenters were organized into small teams of trained workers supervised by an expert segmenter (who completed a course called Connectomics taught by Dr. Spirou). Expert segmenters reviewed all work by their team of trained segmenters. The 3D meshes of all dendrites were reviewed by expert segmenters and Dr. Spirou in VR (syGlass software; IstoVisio, Inc.), overlaid onto the EM image volume so that anomalous branches and structures could be identified, and enclosed ultrastructure and membranes could be incorporated into the evaluation. Tracing the dendrites of all 31 cells provided an internal self reference preventing incorrect assignment of branches to a particular cell. Tracing of dendrites for import into the modeling environment provided additional rigorous review for the subset of 10 cells with complete or near-complete dendritic trees, Endbulb terminals were traced by the same segmenting teams with the same review procedures. Tracing all large inputs and several smaller inputs onto the 21 BCs reported here also provided an internal check that branches of inputs were not missed or assigned to the incorrect terminal. Testing methods for calculation of the ASA followed by performing the calculation for all large inputs onto all cells provided additional rigorous review of the large terminal segmentations.

Fascicles of nerve fibers traverse the volume in the coronal and sagittal planes. ANFs formed the fascicles in the coronal plane. These fascicles were outlined in every 100th section so they could be tracked to determine their extent of splitting and merging. Branches from axons within the fascicles that led to endbulb terminals were also segmented and tabulated, to determine whether axons in particular fascicles gave rise to endbulb terminals within the volume or tended to converge onto the same cellular targets. Terminal size was quantified by measuring the apposed surface area with the postsynaptic membrane, omitting regions where the membranes were separated by intervening glia or extracellular space. We reconstructed the terminals onto each cell that appeared larger than bouton terminals. On two cells we reconstructed all terminals, and from these data we created a histogram of terminal sizes and a definition of minimum size for the large terminal class. We then verified that terminals larger than this threshold were indeed branches of ANFs (see Results). All endbulb axons were traced visually from the terminal retrogradely to their parent ANF or to the location where they exited the image volume. The axon and fiber diameters were calculated from a subset of fibers that had a segment with a straight trajectory either parallel or perpendicular to the image plane, in order to calculate their axon and fiber diameters. A similar procedure was applied to a subset of ANFs (see Fig. 5F). To visualize the spatial relationship of endbulbs and ANF branches to ANF fascicles, all of these structural elements for all endbulb inputs to four cells were segmented using the tracing tool in syGlass.

Three-Dimensional Reconstruction

3D models of the structure of interest were exported from Seg3D as a VTK file and converted to OBJ format using a custom Python script or, in newer versions of the software, exported directly as OBJ files. The meshes in OBJ format were imported into Blender (https://www.blender.org) for processing. Meshes were first decimated by using the decimate modifier tool in collapse mode to merge neighboring vertices progressively while considering the shape of the mesh Low (1997). The meshes are then smoothed using the smooth modifier tool. While these mesh processing steps are suitable for visualization, they do not produce sufficiently accurate surface area or volume measurements. Thus, we evaluated more consistent mesh processing algorithms.
We implemented accurate mesh processing by applying the GAMer2 algorithms and procedures systematically to all meshes in order to create so-called computational meshes (Lee et al., 2020). Surface meshes of segmented objects were generated by performing marching cubes, and produced structures having greater than 1 million vertices due to the high-resolution images and anisotropic sampling during imaging (resolution in x-y plane was ten times resolution in z direction). Anisotropic sampling generates a stair-step effect in the rendering (Figure Supplement 1A). Initial vertex decimation was designed to generate meshes containing 100,000 – 300,000 vertices and reduced time to perform subsequent processing. Experimentation revealed this size range to be the minimum that preserved geometry upon visual inspection. Next, twenty iterations of angle-weighted smoothing (AWS) were applied, which generated nearly equilateral triangles for the mesh faces (Figure Supplement 1B). This geometry is a characteristic of a well-conditioned mesh, which maintains complete surfaces through subsequent processing (Shewchuk, 2002). Two iterations of normal smoothing (NS) were then applied which, in combination with AWS, resulted in a reduction of surface area. The surface area reached an asymptote after the second NS step, confirmed by running three cell bodies through a second round of AWS and NS, indicating that the stair-step effect was minimized after the first round of AWS and NS (Figure Supplement 1C). We visually inspected the meshes during mesh processing and confirmed that all features of the mesh were well-preserved and stair step features were removed after one round of AWS and NS (Figure Supplement 1B). Therefore, we determined this stage of mesh processing to be an accurate stopping point.

### Assignment of Synaptic Weights

We assigned synaptic weights as a density of synapses per square micron of directly apposed presynaptic and postsynaptic membrane, the latter of which we term the apposed surface area (ASA). EM affords the opportunity to measure accurately the membrane apposition, and account for features such as extended extracellular space (Cant and Morest, 1979a; Rowland et al., 2000), where the membranes separate, and interposition of glial processes. We generated an algorithm and custom Python script to identify only the ASA and calculate its summed value for each nerve terminal [https://github.com/MCKersting12/nrrd_tools](https://github.com/MCKersting12/nrrd_tools). This script reads the original segmented image volumes of the two objects contacting one another, which may have been traced in different subvolumes of the original volume (subvolumes were created to permit multiple segmenters to work in parallel), and transforms them to have the same origin (pixel-spacing, height, width, and length). If the segmented terminal and postsynaptic cell have overlapping voxels, the overlap is removed from the soma because the terminal segmentations were typically more accurate. Next, the terminal is dilated by 3 voxels in the x-y plane and then, because the volume is anisotropic, another 3 voxels in all directions. The dilation in z was tested and this value was chosen based on visual inspection to provide overlap selectively of the ASA. The overlapping region between the dilated terminal and the soma volume is extracted as a separate volume, and the marching cubes algorithm is performed on this separated volume. The surface area of the resultant mesh, which appears as a flattened volume, is divided by two because we are only interested in the contact area to generate the ASA.

Synapses can be identified in our SBEM volume by clustering of synaptic vesicles along the presynaptic membrane in at least 3 serial sections, direct contact of at least one vesicle with the presynaptic membrane, and a concavity in the postsynaptic membrane, the latter of which is typical of endbulb terminals in the cochlear nucleus in aldehyde fixed tissue (Spirou et al., 2008; Cant and Morest, 1979a; Ryugo et al., 1997). A postsynaptic density is typically found but is not present in all cases, so was not used as an explicit criterion. Each large input contains multiple synapses, so the number of synapses was quantified for 23 terminals of varying sizes, and density (#synapses/μm²) was calculated using the ASA for each terminal. The average synapse density was applied to termi-
nals for which the ASA was determined but synapses were not counted, to achieve an estimate of the number of synapses in each terminal reconstructed in this study.

**Model Generation**

Biophysically-based models were generated for each reconstructed cell, using the ASA data for individual auditory nerve inputs, and the compartmental reconstructions. The modeling was performed as a predictive exercise, using previously measured biophysical parameters for synapse release dynamics, postsynaptic receptors, and ion channels, along with a standard model of auditory nerve responses to sound. The principal free parameters were the densities of channels in different cell compartments. The channel densities were calculated based on the ratios of densities for somatic models in a previous study (Rothman and Manis, 2003c), measured densities in voltage clamp from mouse BCs for the low-threshold potassium conductance, and relative densities in the axon initial segment and hillock from other central neurons. Because ion channel densities in the dendrites of bushy cells have not been measured, we bracketed the likely range by testing models with passive dendrites, fully active dendrites (densities were the same as in the soma) and half-active dendrites. Thus, the models are predictive given the constraints of unmeasured channel densities. To accomplish this, the models were built up in a series of steps: morphological reconstruction, surface area adjustments, base channel density adjustment, and overall channel density assignment. Synaptic conductances were constrained by previous measurements (Raman and Trussell, 1992; Xie and Manis, 2013), and the only free variable was the number of sites for each multi-site synapse, which was set according to the ASA measurements and release site counts from the SBEM material.

**Translating Reconstructions to NEURON models**

We rendered the SBEM mesh into a modified version of the SWC file format (Cannon et al., 1998) using the tracing tool in syGlass. Each reconstructed part of the cell is represented as a series of conical frustums with starting and ending radii. We also annotated groups of points with a named morphological feature of the section. Identified morphological features were given new tags in the SWC file, and included the myelinated axon, axon initial segment, axon hillock, soma, proximal dendrites, dendritic hubs, distal dendrites, and dendritic swellings. Next, the SWC files were translated to HOC files using a Python script. The script added groups of SWC points in a 3D shape format (pt3d) to create short sections composed of at least three and up to 50 segments. This translation retained the detailed geometry of the cells. Comment fields in the HOC files referenced the original SWC point for each 3D point in Neuron, which facilitated mapping voltages in processes back to the original mesh representation, and confirming that the translation proceeded correctly. This annotation also allowed us to perform manipulations that removed specific parts of the original reconstruction.

We then compared the original SBEM mesh files’ surface area representations with those of the 3D geometry HOC files. The mesh represented the cell surface at a high resolution that captured membrane crenelations, even after reducing the mesh density with GAMer2 (Lee et al., 2020) and subsequent smoothing. In contrast, the SWC and HOC representations capture the mesh structure using simple frustrated cones, which have smooth surfaces. Consequently, the mesh surface area was always significantly greater than the surface area computed from the HOC representation. The surface area determines the capacitance and plays a fundamental role in establishing ion channel densities and the transmembrane leak resistance in the model cells. We therefore compensated for these surface area differences by inflating the compartment diameters in the HOC file by the ratio between the mesh and HOC areas, while not changing the lengths. Separate inflation factors were calculated for the soma and for the entirety of the dendritic tree, because the mesh’s ratio to HOC surface areas for these regions was different. NEURON instantiates compartments (as
Axons: Axons were reconstructed from the soma to the first internodal (myelinated) region for 8 of the 10 reconstructed bushy cells. Data from mouse bushy cells from Yang et al. (2016) indicates that the Na⁺ channel density is lower in the soma than in the axon hillock and that the action potential initiation begins distally, likely in the AIS. Lacking direct measurements in bushy cells, we used the experimental and model data from Kole et al. (2008) from layer V cortical neurons to guide the relative channel densities. The axon hillock channel density for Na⁺ channels was set to five times that of the soma, and the initial segment was 100 times that of the soma. The hillock and AIS compartments were each decorated uniformly, to approximate the uniform distribution reported for immunostaining of Na⁺ channels (Kuba et al., 2015), although there is some data suggesting that channel density and composition vary with distance from the soma (Lorincz and Nusser, 2008; Hu et al., 2009). The assignment of spatially uniform conductance densities to the AIS represents a first-order assumption, as we lack experimental data with appropriate resolution to justify other distributions in BCs. With this decoration, the total AIS Na⁺ conductance in the model is a function of AIS length, and therefore also affects action potential threshold and amplitude. Variations in AIS length have been correlated with neuronal excitability (Grubb and Burrone, 2010; Kuba et al., 2019; Kaphzan et al., 2011), and tonotopic position in nucleus laminaris (Kuba et al., 2006). Na⁺, K⁺ channel and Iₜ channel densities are shown in Table 1.

<table>
<thead>
<tr>
<th>Decoration Type</th>
<th>Channel</th>
<th>Myelinated axon</th>
<th>AIS</th>
<th>AH</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Na⁺</td>
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<td>100.0</td>
<td>5.0</td>
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<tr>
<td></td>
<td>K⁺HT</td>
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<tr>
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<td>K⁺LT</td>
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<td>1.0</td>
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<td></td>
<td>Iₜ</td>
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<td>0.5</td>
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<tr>
<td></td>
<td>Leak</td>
<td>0.00025</td>
<td>1.0</td>
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</tbody>
</table>

Table 1. Densities of channels used to decorate the axon compartments of bushy cells. Values are given as ratios relative to the standard decoration of the somatic conductances.

"segments") from the 3D reconstructions. However, there is no analytical solution to the reverse problem of recalculating the total area's diameters. Therefore, we computed the inflation factor iteratively until the reconstructed area, as computed from NEURON, matched the mesh area. For the bushy cells, the soma's inflation factor averaged 1.486 (SD 0.227), and the factor for the dendritic tree averaged 1.506 (SD 0.145). The ratio of the soma inflation factor to the dendrite inflation factor for different reconstructions varied from 0.78-1.38 (mean 0.995, SD 0.195). The last step in establishing the geometry for simulations was determining the number of segments necessary to maintain an appropriate spatial discretization. The number of segments for each section was recomputed using the d-λ rule (Carnevale and Hines, 2006), at 1000 Hz. Because many of the reconstructions already had short section lengths, this step affected only a fraction of the sections for any given cell. All current clamp simulations were run with a time step of 25 μs.

Ion Channels and Receptors

Cells were "decorated" with Hodgkin-Huxley style ion channels based on biophysical measurements from previous studies. The kinetic measurements for K⁺ channels were obtained from acutely isolated bushy neurons that lacked dendritic trees (Rothman and Manis, 2003a), scaled to 37°C (Rothman and Manis, 2003b). We drew K⁺ channel density estimates from measurements made from cells in mouse brain slices (Cao et al., 2007), scaled as described below. Sodium channels were represented by a modified model (Xie and Manis, 2013), which incorporated cooperative interactions between channels (Huang et al., 2012; Ilin et al., 2013; Manis and Campagnola, 2018). Actual conductance densities for the dendrites, axon hillock, axon initial segment, and nodes of Ranvier are not known. To address these uncertainties, we decorated the cell compartments using density distributions that have been estimated for other neurons, as described next.

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Table 1. Densities of channels used to decorate the axon compartments of bushy cells. Values are given as ratios relative to the standard decoration of the somatic conductances.
For BC02 and BC05, the axon left the tissue block before becoming myelinated. To compensate, we replaced the axon hillock, initial segment and first myelinated region with a “standard axon” based on the average axon lengths and diameters from the other 8 cells for simulations of these cells. These cells were not used in evaluating the effects of AIS length on excitability, although their data is plotted alongside the other cells for comparison.

Dendrites: Based on the SBEM measurements, the surface area of bushy cell dendrites ranged from 2.43-3.23 (mean 2.76 SD 0.24) times the cell body area. Although bushy cell dendrites are short, they have a large diameter and consequently represent a substantial capacitance and conductive electrical load to the soma. The distribution of ion channels on BC dendrites is not known. Qualitative immunostaining studies hint at the presence of HCN and low-voltage activated K⁺ channels in at least the proximal BC dendrites (Koch et al., 2004; Oertel et al., 2008; Pál et al., 2005; Wang et al., 1993) (but see (Perney and Kaczmarek, 1997) where dendritic staining for the high-voltage activated channel K₃.1 is visible in stellate cell dendrites but not clearly visible in bushy cell dendrites in rat). However, with relatively few synaptic inputs and a limited role for active dendritic integration, it seems likely that voltage-gated ion channels may not be present at high densities in the dendrites. To account for the potential roles of dendritic channels, we therefore bracketed the conductance density range with three models. In each of these models, we decorated all types of dendritic compartments (proximal and distal dendrites, dendritic hubs, and dendritic swellings) with the same conductance densities. First, we used a model in which the densities of the channels in the dendrites were half of those in the soma (“Half-active”). The other two models addressed the extremes of possible channel densities. In the “Passive dendrite” model, the dendrites were uniformly decorated only with leak channels. In the “Active dendrite” model, the dendritic channel density was set uniformly to the somatic channel density for all channels. We refer to these models below as the “dendritic decoration configurations”.

Conductance Scaling: To properly scale the conductances into the somatic and dendritic compartments, we began with the low-voltage activated channel, gKLT, which was measured under voltage clamp to be 80.9 (SE 16.7) nS in CBA mice (Cao et al., 2007). Next, to set a baseline value for the conductances, we first computed the mean somatic surface area from the SBEM reconstructions (1352.1 (SD 164.9) μm², N=26 bushy cells), and for dendrites from the ten complete reconstructions (3799.5 (SD 435.8) μm², N=10 bushy cells). We then chose one cell whose somatic and dendritic areas were closest to the mean of these distributions (BC17: somatic surface area = 1357.6 μm², dendritic 3707.7 μm²) to adjust gKLT. The use of the “average” cell for this step was chosen to be consistent with the use of the mean value from Cao et al. (2007). We then adjusted gKLT by computing the measured gKLT from a voltage clamp protocol that mimicked experimental measurements (steady-state currents with 100 ms pulses) with only gKLT and a leak conductance inserted into the soma and dendrites for each of the three dendritic distribution assumptions. The soma was initially decorated with gKLT channels at a fixed density of 2.769 mS/cm² based on a maximum conductance measured in vitro of 80 nS and a measured cell capacitance of 26 pF (Cao et al., 2007). However, this capacitance corresponds to a surface area of 2889 μm², which is more than twice the area of the measured somas, and is also significantly larger than other previously reported values (12 pF in acutely isolated neurons from guinea pig Rothman and Manis (2003a), 9-12 pF in rat pup bushy cells in slices (Xu-Friedman and Regehr, 2008), 9-22 pF in adult CBA mouse bushy cells, Xie and Manis, unpublished). To investigate this discrepancy, we measured the input capacitance (as seen by a somatic electrode) using voltage clamp simulations of the reconstructed cells. The voltage-clamp simulations were stopped at 5 μs, with 1 MΩ of uncompensated series resistance (Rₛ), to approximate the experimental situation that used 90% compensation of ~ 11 MΩRₛ (Cao et al., 2007). Voltage steps from -80 to -90 mV were applied to models with only gKLT and gL channels in the membrane, which yielded values of 13 pF, based on the fastest membrane charging time constant of ~ 15 μsec, consistent with the studies cited above. This corresponds to a membrane area of...
1460 µm², close to 1358 µm² measured for the soma area of this cell. We then ran additional voltage clamp simulations with steps from -80 to +20 mV to measure $g_{KLT}$. Total $g_{KLT}$ was measured from V-I relationship by fitting a Boltzmann function to the steady-state portion of the simulated currents (Figure 3–Figure Supplement 3), after correcting the membrane voltage for the drop across the series resistance, $R_s$. We iteratively made a linear prediction after each adjustment, by calculating the ratio between the measured conductance and the target value of 80 nS, and applied this to rescale $g_{KLT}$. Three to five iterations were adequate to arrive within 1% of the target value for $g_{KLT}$ for each of the three dendritic decoration models for the test cell. Once $g_{KLT}$ was determined, the ratio of $g_{KLT}$ to the original model density was then calculated, and applied to all of the other channels at the soma, relative to their total cell conductances in the original models (based on the measurements and models of Xie and Manis (2013) and measurements of Cao et al. (2007) ($g_{KLT}$: 80 nS; $g_{KL}$: 500 nS, $g_{KHT}$: 58 nS, $g_{H}$: 30 nS). The resulting densities, expressed in mS/cm² and listed in Table 2, were used to decorate all reconstructed cells. Thus, with this approach, we anchored the model ion channel densities according to our morphological measurements to experimental measurements of $g_{KLT}$ in the same species.

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Table 2. Densities of channels in dendrites for 3 models spanning the likely range. All values are in mS/cm².

Auditory Nerve Inputs

Auditory nerve spike trains were computed using the cochlea package (Rudnicki et al., 2015), which is a Python wrapper around the widely-used model of Zilany et al. (2014). These simulations were incorporated into, and controlled by, cnmodel (Manis and Campagnola, 2018). Although the spike trains generated by these simulators were based on data from cat ANFs, the responses for mouse auditory nerve are quite similar, including irregular interspike intervals and the thresholds are similar in the central range of mouse hearing (Taberner and Liberman, 2005). Tonal acoustic stimuli were generated at 100 kHz with rise-fall times of 2.5 ms, and durations from 100 to 1000 ms. Clicks were generated as 100 µs pulses. The intensity was expressed in dB re 2x10⁻²Pa (dB SPL). For tonal stimuli, the frequency was set to 16 kHz to avoid low-frequency phase locking.

For some simulations, single-frequency tones at 16 kHz were amplitude modulated with a sinusoidal envelope (100% modulation) at frequencies between 50 and 1000 Hz. The depth of response modulation in ANFs is critically dependent on the average stimulus intensity as well as ANF SR (Smith and Brachman, 1980; Joris and Yin, 1992; Joris et al., 2004; Wang and Sachs, 1993) and this sensitivity continues to be evident in cochlear nucleus neurons (Moller, 1972; Frisina et al., 1990; Wang and Sachs, 1994). We tested responses of the BC models to SAM tones at an intensity that produces the highest synchronization in the high-spontaneous ANFs, 15 dB SPL, as well as at 30 dB SPL (see Figure 7–Figure Supplement 1 for the VS as a function of level in the ANF model). Testing was performed with only high-SR ANFs as inputs, consistent with observations in cats that GBCs are principally innervated by high-SR inputs (Liberman, 1991). Testing by including other SR groups would be expected to show higher synchronization at high sound levels (Wang and Sachs, 1994) as the medium and low SR fibers continue to synchronize to the envelope. However this would require making specific assumptions about the relationship between ASA and SR in order to appropriately assign SR groups. While recent data (Wang et al., 2021) suggests that some mouse...
GBCs may receive a greater proportion of medium and low-SR inputs than previously suggested for cat, we considered exploration of this dimension in the context of our simulations beyond the goals of the current study.

Endbulb Synapses

The endbulb synapses were modeled using a stochastic multisite release model, as described previously (Xie and Manis, 2013; Manis and Campagnola, 2018) and incorporated into cmodel. Briefly, the release at each endbulb terminal is initiated when an action potential is generated by the auditory nerve model. Each synapse in the terminal then can release transmitter with a release probability, $P_r$ in the range $[0,1]$. In the present simulations, the release probability was held fixed over time (it was not a function of the history of release event times). Whether a synapse will release or not is determined by drawing a random number from a uniform distribution, and if the number is less than $P_r$, then a release event is initiated. Transmitter time course was computed by convolution of a Dirac pulse with a bi-exponential function to mimic diffusion across the synaptic cleft, and the concentration time course at the postsynaptic receptors is computed by summing each release event with an ongoing cleft concentration. This glutamate transient then drives postsynaptic receptors. The postsynaptic receptors are based on fast AMPA receptors at the endbulbs in the nucleus magnocellularis of chicken (Raman and Trussell, 1992), with kinetics adjusted to match recorded currents at the mouse endbulb (Xie and Manis, 2013). The AMPA receptor model conductances were also adjusted to match measurements of mEPSCs at mouse bushy cells. The receptor model includes desensitization, and the current through the receptor channels includes rectification of the current-voltage relationship by internal channel block from charged polyamines (Woodhull, 1973; Donevan and Rogawski, 1995). The cleft glutamate also interacts with NMDA receptors in the synapse, based on the model of Kampa et al. (2004). NMDA receptor conductances were scaled to match the to the voltage-clamp measurements in Cao and Oertel (2010). Each release site of the terminal is treated independently, ignoring the possible consequences of transmitter “spillover”. A time-dependent increase in release latency is observed experimentally (see Manis and Campagnola (2018)), but was disabled in the simulations reported here because it has not been fully characterized. The number of synapses at each endbulb is calculated using the ASA and average synapse density as determined from the SBEM data. For all simulations here, the density was $0.7686 \text{synapses/\mu m}^2$.

Spike Detection

Spikes in bushy neurons are often small and of variable amplitude, and the EPSPs can be large (10’s of mV). Simple approaches using a fixed voltage or slope threshold are not reliable for discerning spikes from EPSPs with somatic recordings. We, therefore, used the method of Hight and Kalluri (2016) to detect spikes based on the width of the peak and the rising and falling slopes. Spike detection parameters were set exactly as in Hight and Kalluri (2016).

Spike Timing Analysis

Vector strength was computed using the standard equations (Goldberg and Brown, 1969), using spikes taken from the last 800 ms of 100 repetitions of 1-s long SAM stimuli. To estimate the error of the vector strength calculation, vector strength was calculated for 10 groups of 10 consecutive repetitions, and the mean and SD computed. Responses with fewer than 50 spikes were not calculated (this appeared only for BC10 for the configuration with only the largest input active). Vector strength for ANFs was calculated across all spikes of all ANFs connected to the postsynaptic cell. We also calculated shuffled autocorrelations using the method of Louage et al. (2004) for both SAM stimuli and click stimuli. These calculations were verified to reproduce Fig 2. of Louage et al. (2004).
Action Potential Current Threshold Measurement

The minimum current required to elicit an action potential (rheobase) was measured in response to a brief current pulse (20 ms) of variable amplitude. An iterative binary search procedure was used to identify the threshold, with a terminal step size of 1 pA. Ten to twenty iterations were sufficient to resolve threshold to this precision.

Modeling Software Environment

The entire set of simulations were controlled and analyzed by additional Python (V3.7.8, 3.8.6, 3.9.1, 3.10.0) scripts (VCNModel). VCNModel controlled simulations and organized simulation result files, read cell morphology files into NEURON Carnevale and Hines (2006), and decorated the cells with channels using tools from cnmodel (www.github.com/cnmodel). Parametric simulations were managed by shell scripts (bash, zsh) that called the Python scripts. Simulations reported here were run with NEURON 7.7, 7.8.1, 8.0 and 8.1 on an 8-core MacPro (2013), a MacBook Pro (2017), and a 20-core MacStudio (2022); there was no difference in the results of the underlying auditory nerve, bushy cell, or synapse models as determined by the unit tests in cnmodel for any versions of NEURON, Python, or hardware. The anatomical structure of the reconstructions was defined by the NEURON HOC files, and the channel densities were set from text (human readable) tables managed in cnmodel. The VCNModel scripts computed scaling of cell areas (inflation of the SWC/HOC files to match the mesh areas), control of “experiments” (for example, only activating selected AN terminals), data management, plotting, and analysis. Analysis of current voltage relationships and spike detection was handled by the ephys package (www.github.com/pbmanis/ephys). Plots were generated using matplotlib (versions 3.2.0-3.5.2) and seaborn (version 0.11.2).

Data and Code Availability

Data (Excel worksheets) and code (Matlab R2022a) for graphs in Figures 1, 2, 5, 6 and 8H are available at www.github.com/gaspirou/pub_file_share. Simulation source code, documentation, and a shell script to set up a working environment is available at www.github.com/pbmanis/VCNModel (“release” branch). Simulation result files used to generate figures 3, 4, 7 and 8I-M and their associated supplemental figures have been uploaded to Dryad, and can be accessed at www.dryad.org for review. Code and data for Figure 2, Supplement 1 is included in VCNModel. Simulation figures and figure panels can be generated using the DataTables script in the VCNModel package after downloading the simulation result files.

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Additional Information

Funding
The funding agencies had no role in study design, data collection and interpretation, or the decision to submit this work for publication.

Ethics
George A. Spirou is the Chief Scientific Officer of IstoVisio, Inc., which makes the syGlass software used to visualize the EM volume and reconstruct the neurons. Carolyna Y. Alves-Pinto, Sean Carr, Mariah Dawson, Mark Ellisman, Matthew Kersting, Paul B. Manis, and Bayan Razzaq have no declarations or affiliations to report.

Author Contributions
George Spirou: Conceptualization, Formal analysis, Software, Investigation, Visualization, Methodology, Funding acquisition, Project administration, Writing - reviewing and editing. Matthew Kersting: Formal analysis, Software, Investigation, Visualization, Writing (portions of original draft). Carolyna Y. Alves-Pinto, Sean Carr, Mariah Dawson, and Bayan Razzaq contributed Formal analysis, Investigation, Visualization. Mark Ellisman: Investigation, Methodology, Visualization. Paul Manis: Conceptualization, Formal analysis, Simulations, Software, Investigation, Visualization, Methodology, Funding acquisition, Writing - reviewing and editing. Portions of this work are derived from Matthew Kersting's Master's thesis in Medical Engineering (University of South Florida).

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Bourk TR. Electrical responses of neural units in the anteroventral cochlear nucleus of the cat. Ph.D., Massachusetts Institute of Technology; 1976.


## Key Resources

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Figure 1—Figure supplement 1. Steps in mesh generation and compartmental representation from EM volumes. (A) Cell membranes of objects (cell body of a bushy cell in this case) were traced and assembled into stacks. Each tissue section appears as a slab, seen clearly in expanded view at right. The slab thickness is the 60nm section thickness used during imaging. (B) The GAMer2 algorithm (Lee et al., 2020) was used to create a mesh surface enclosing the volume, comprised of isosceles triangles, that preserved real surface irregularities such as small protrusions. (C) Decimation and smoothing algorithms were successively applied until the meshed surface reached an asymptotic change surface area (vertical arrow). The change in area for all reconstructed somas are shown. After testing multiple cycles on three cells, values at NS-2 stage (see Methods) were used for all cells.
**Figure 2—Figure supplement 1.** Morphological correlations for synapse and somatic area. (A) There was no relationship between cell body surface area (SA) and the apposed surface area (ASA) of the largest input. (B) There was a weak correlation between the somatic coverage by large inputs and the largest input area. (C) There was no correlation between cell body area and the number of large inputs. (D) There was no correlation between the area coverage by large inputs and the cell body area.
Figure 2–Figure supplement 2. Large somatic terminals onto each bushy cell (BC) that fits the Coincidence Detection model. Terminal groups are arranged from top left to bottom right in order of decreasing size of their largest input. The number of terminals ranges from 5 (BC12) to 12 (BC19). Apposed surface area values are indicated nearby each terminal. Six of these BCs (label enclosed in blue box) had dendrites fully reconstructed and were used for compartmental modeling. The extent of their local dendrite branching is indicated by L, low; M, moderate; D, dense. Scale bar omitted because these are 3D structures and most of the terminal would be out of the plane of the scale bar.
Figure 2—Figure supplement 3. Large somatic terminals onto each bushy cell that fit the Mixed Coincidence Detection/First Arrival model. Terminal groups are arranged from top left to bottom right in order of decreasing size of their largest input. The number of terminals ranges from 5 (BC14, BC16) to 12 (BC08). Four of these BCs (label enclosed in blue box) had dendrites fully reconstructed and were used for compartmental modeling. The extent of their local dendrite branching is indicated by L, low; M, moderate; D, dense. Scale bar omitted because these are 3D structures and most of the terminal would be out of the plane of the scale bar.
**Figure 3—Figure supplement 1.** Panels are labeled by cell number. For each bushy cell (BC), the segmentation as computational meshes is depicted at left. Cell bodies are light grey, dendrites are red, and axons are pink. Cells are aligned to permit the most encompassing views of their dendrites. Meshes were converted to SWC format, which represents structures as a series of linked skeleton points with associated radius, using 3D virtual reality software (syGlass). The SWC representations are shown at right for each pair of images. Axon and dendrite subcompartments were annotated during SWC creation, which permitted quantification of their surface areas. Cells are clustered into three groups by the extent of local dendrite branching and braiding: low (BCs 02, 06, 18), moderate (BCs 09, 11, 13, 30), and dense (BCs 05, 10, 17). Cellular compartments are color coded in the SWC files, as depicted in the legend. Scale bar is not included because it does not apply throughout the depth of the 3D structure. Images scaled to fit within figure panels so cells are pictured at different scales.
Figure 3—Figure supplement 2. Conductance scaling using voltage clamp simulations for different patterns of dendrite decoration. Voltage-clamp simulations were used to compute total channel densities to match experimental data for different dendritic decoration configurations. (A) Passive dendrites. (A1): voltage clamp protocol with steps as in panel (A2). The transients at the start and end of the steps represent the charging of the cell membrane capacitance, which was not compensated in the model cell. (A3) Steady-state conductance calculated from the data in A1. The inset text indicates the fits to a Boltzmann function of the form

\[ g(V) = \frac{1}{1 + e^{(V-V_{0.5})/k}} \]

where \( V_{0.5} \) is the half-activation voltage and \( k \) is the slope factor. C_m is the cell capacitance, calculated from an exponential fit to the initial charging curve for small negative voltage steps, and \( t_0 \) is the clamp charging time constant (fastest time constant, representing somatic capacitance). (B) and (C) 1-3 are in the same format as (A1-3), with different channel decoration of the dendrites. (D) Input resistance measures from the 10 completely reconstructed cells for each of the decoration conditions from current-clamp simulations (see Figure 3—Figure Supplement 2). Boxplots: Shaded area indicates interquartile distances, whiskers indicate 5-95% confidence limits. (E) Time constant measurements from the 10 completely reconstructed cells for each of the decoration conditions from current clamp simulations. Boxplots are formatted as in (D). Scale bars in (A) apply to (B) and (C).
Figure 3—Figure supplement 3. Summary of current-clamp responses for all 10 cells for each of the decoration conditions. All current-voltage relationships show traces from -1 to + 2 nA, in 0.2 nA steps. Action potentials are indicated by red dots. The resting potential is indicated to the left of each group of traces, in mV. The calibration bar in the top row applies to all traces. The right most columns hows the steady-state current-voltage relationships (squares) and peak (circles, for hyperpolarization only) for each of the decoration conditions (Passive: cyan; Half-active: black, Active: magenta). Note that some model cells (9, 11, 30) show small repetitive spikes with stronger depolarization with the passive dendrites. Most cells show spikes at anodal break, but these are attenuated or absent when the dendrites are fully active.
Figure 3—Figure supplement 4. Input impedance at rest. Input impedance was calculated as seen from the soma for the Half-active condition, for intact cells (real part: (A), phase angle: (D)), cells with no dendrites (real part: (B), phase angle: (E)) and for the axon only as seen from somatic end (real part: (C)). The impedance is highly frequency dependent, with a peaked-low pass characteristic. The dendrite presents a substantial load to the cell body, as evidenced by the much smaller real part in (A) compared to (B), and phase shift (D,E). (F) The conductance load of the dendrite and the soma as seen from the axon was also calculated as in Eyal et al. (2014), Eq. 3, as a function of frequency. The values below 250 Hz vary between cells by a factor of 2, which will affect the ability of the cells to phase lock at higher frequencies. (G, H, I) Nyquist plots for intact, no dendrite and axon conditions (plotted imaginary vs real impedance components with parametric frequency) further reveal the differences between the cells. The frequency progression runs counterclockwise as indicated in panel (H).
Figure 3—Figure supplement 5. PTH and spike latencies in response to tone bursts at CF. Each row shows the responses for one cell. BC02 and BC05 whose reconstructed axon left the volume prior to myelination were simulated with a substitute axon ("Sub. axon") that had the average hillock, initial segment and myelinated axon lengths and diameters from the other 8 cells. Otherwise, each cell was simulated using its own reconstructed axon. The first column shows the soma voltage for one trial for a stimulus at CF at 30 dB SPL. The stimulus starts after 50 ms and is 100 ms in duration. The PTH shows the spike rate as a function of time averaged over 100 repetitions of the tone pip, with 0.5 ms bins. The FSL/SSL column shows the first (FSL, blue) and second (SSL, red) spike latency distributions; text shows the mean and SD of the FSL and SSL. The rightmost column plots the coefficient of variation corrected for a 0.7 ms refractory period (CV') of interspike intervals through the stimulus, ending approximately 25 ms before the end of the stimulus to minimize end effects. The CV value is indicated to the right of each plot. All CV values fall in the range of 0.3-0.7 reported for mouse primary-like neurons (Roos and May, 2012). The blue plots in the bottom row show the stimulus waveform timing for each column.
Figure 4–Figure supplement 1. Reverse Correlation plots for 6 additional modeled cells. Each cell is plotted in the same format as in Figure 4A, D and E. Summary information is presented in panels Figure 4(D-J).
Figure 4–Figure supplement 2. Correlations amongst morphological parameters and action potential threshold. In all panels, filled points are colored according to the cell number, and symbols correspond to Coincidence Detectors (circles) and Mixed-Mode (diamonds), as in Figure 4E. (A) Dendrite area is weakly related to soma area. (B) Current threshold for spike generation is weakly correlated with soma area. (C) Current threshold is correlated with the ratio of dendrite to soma area. (D) Current threshold is weakly negatively correlated with AIS length. (E) The efficacy of the largest input to each cell is not correlated with AIS length. Regression calculations in panels (B) and (C) excluded cells BC02 and BC05 (un/filled circles), but their thresholds are plotted assuming an axon (hillock, initial segment, myelinated section) that is the population average length of all of the other cells. Linear regression statistics are printed on each panel.
Figure 4—Figure supplement 3. Contributions of different input patterns to postsynaptic spiking. For each panel, the contributions are measured in cells 2, 5, 6, 10, 13, 30 ("Coincidence" group) and contributions measured in the mixed-mode group of cells 9, 11, 17, 18 ("Mixed-mode" group). The panels are titled according to the particular patterns of input, and plot the percent of postsynaptic spikes that were generated by each pattern. Cell colors are the same as in other figures. The bars show the median, interquartile distances, and 5-95% whiskers. Points that fall outside of the expected distribution are indicated with diamonds. Results are shown for spontaneous activity. Individual cells are noted by the color in the legend.
Figure 6—Figure supplement 1. Filament core (white arrows) in primary BC hubs and dendrite of one MC. A-L. MC04, BCs 05, 14, 27, 08, 10, 09, 24, 17, 30, 29, 16. Filaments appear in close apposition to mitochondrion outer membranes and can fill narrow spaces defined by those membranes (white arrowheads). Scale bar = 1 micron in panel L applies to all panels except panel I scale bar = 2 microns.
Figure 7—Figure supplement 1. Synchronization to stimulus envelope as a function of average stimulus intensity in ANF inputs. Vector strength of response to 100% SAM across frequency for the ANF model. Carrier frequency was 16 kHz. Color bars indicate 15 dB SPL (gold) used in Figure 7, O1-4 and P1, and 30 dB SPL (gray), as used in Figure 7, P2.
Figure 7–Figure supplement 2. Vector strength of BCs in response to 100% SAM modulation at frequencies from 50 to 1000 Hz on a 16kHz carrier at 15 dB SPL. These simulation results are also included in Figure 7P1. Each plot shows vector strength for 3 different synaptic input configurations. The vertical lines indicate the SD of the vector strength (VS) computed as described in the Methods. (A) Cell BC05, (B) Cell BC06, (C) Cell BC10, (D) Cell BC11, (E) Cell BC13, (F) Cell BC18. BC10 had too few spikes with only the largest input to compute VS above 100 Hz.
**Figure 7—Figure supplement 3.** SACs in response to click trains show importance of weaker inputs in improving temporal precision. SACs were computed for click evoked spike trains as shown in Figure 7. (A-J) SACs computed for each of the BCs for 3 different input configurations (colors are indicated in panel (K)). For BC05 (B) and BC10 (E), there were insufficient spikes in the largest input only condition for the SAC calculation. (K) Half-width of the SAC for each cell and configuration computed from Gaussian fits to the SACs in panels (A-J). (L) SAC correlation index (CI) at 0 time for each cell and configuration.