Mode-Locked Spike Trains in Responses of Ventral Cochlear Nucleus Chopper and Onset Neurons to Periodic Stimuli

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Laudanski J, Coombes S, Palmer AR, Sumner CJ. Mode-locked spike trains in responses of ventral cochlear nucleus chopper and onset neurons to periodic stimuli. J Neurophysiol 103: 1226–1237, 2010. First published December 30, 2009. doi:10.1152/jn.00070.2009. We report evidence of mode-locking to the envelope of a periodic stimulus in chopper units of the ventral cochlear nucleus (VCN). Mode-locking is a generalized description of how responses in periodically forced nonlinear systems can be closely linked to the input envelope, while showing temporal patterns of higher order than seen during pure phase-locking. Re-analyzing a previously unpublished dataset in response to amplitude modulated tones, we find that of 55% of cells (6/11) demonstrated stochastic mode-locking in response to sinusoidally amplitude modulated (SAM) pure tones at 50% modulation depth. At 100% modulation depth SAM, most units (3/4) showed mode-locking. We use interspike interval (ISI) scattergrams to unravel the temporal structure present in chopper mode-locked responses. These responses compared well to a leaky integrate-and-fire model (LIF) model of chopper units. Thus the timing of spikes in chopper unit responses to periodic stimuli can be understood in terms of the complex dynamics of periodically forced nonlinear systems. A larger set of onset (33) and chopper units (24) of the VCN also shows mode-locked responses to steady-state vowels and cosine-phase harmonic complexes. However, while 80% of chopper responses to complex stimuli meet our criterion for the presence of mode-locking, only 40% of onset cells show similar complex-modes of spike patterns. We found a correlation between a unit’s regularity and its tendency to display mode-locked spike trains as well as a correlation in the number of spikes per cycle and the presence of complex-modes of spike patterns. These spiking patterns are sensitive to the envelope as well as the fundamental frequency of complex sounds, suggesting that complex cell dynamics may play a role in encoding periodic stimuli and envelopes in the VCN.

INTRODUCTION

Spike timing is thought to play an important role in the neural representation of sound. It has long been studied in the context of phase-locking, in which a neuron tends to fire at a given phase of a periodic stimulus. This effect is robust in the auditory nerve (AN) and brain stem discharges to low-frequency tones (Rose et al. 1967) and complex harmonic sounds such as vowels (Palmer et al. 1986; Winter and Palmer 1990) and in responses to amplitude-modulated tones (for a review, see Joris et al. 2004). More recently, information theoretic studies have revealed how spike timing can be used to detect (Gai and Carney 2008) or distinguish categories of arbitrary spiking patterns to arbitrary stimuli (Huetz et al. 2006; Schnupp et al. 2006). However, this last approach does not consider the role of the dynamic properties of neurons in the encoding of stimulus information.

Stellate cells in the VCN are well characterized with respect to both their biophysical and coding properties. These cells exhibit highly regular spiking (chopping response) in response to constant stimulation (for both current steps and pure tones) (Manis and Marx 1991; Rhode et al. 1983). They also encode the frequency of AM using the temporal pattern of firing: they display a band-pass vector strength (VS) (Rhode and Greenberg 1994) and their interspike interval (ISI) statistics are related to the temporal periodicity of the stimuli presented (Wiegrebe and Winter 2001). These response properties are thought to result from summation of the inputs from auditory nerve fibers, temporal integration, and a simple spiking mechanism. Simulations of Hodgkin-Huxley (HH) type (Banks and Sachs 1991; Wang and Sachs 1995) or integrate-and-fire models (Hewitt et al. 1992), reproducing the linear subthreshold current-voltage (J-V) curve of stellate cells, have successfully replicated their responses to pure tone at characteristic frequency (CF), the VS in response to AM tones and the responses to other periodic stimuli.

In this article, we complement previous simulation studies using a mathematical theory explaining how nonlinear dynamical systems synchronize to time varying input. We observe higher-order temporal patterns, called “mode-locked” in response to several different classes of periodic stimuli: AM tones, steady-state vowels, and cosine-phase harmonic complexes. Mode-locking, of which phase-locking is a subset, results from the interaction between the dynamics of a nonlinear oscillator and a periodic stimulus. Such spiking patterns have been previously described in generic neuron models similar to models of stellate cells [for instance HH (Aihara et al. 1984; Lee and Kim 2006) or leaky integrate-and-fire (LIF) (Brette 2004; Coombes and Bressloff 1999; Keener et al. 1981)] and have been observed in vitro under DC injection (Aihara et al. 1984; Brumberg and Gutkin 2007; Guttman et al. 1980; Shreiber et al. 2004). Mode-locked responses are often stable across regions of parameter space called Arnold tongues (for a review, see Pikovsky et al. 2001). Our report is the first description of mode-locking in the auditory system and under sensory stimulation in vivo.

METHODS

Experimental data collection

The data were collected from 54 healthy adult pigmented guinea pigs (Cavia porcellus), weighing in the range of 300–400 g, which also provided extensive data for other studies. We analyzed the responses of 11 cells (not previously published) to SAM tones of 3-s duration presented 10 times with a range of modulation frequencies. Seven of these cells were classified as chopper units, two as onset choppers, and two as unusual. We also analyzed the responses of 24

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chopper units and 33 onset units in response to vowels and harmonic complexes (previously published in Winter et al. 2003).

The animal preparation has been described in detail previously (Winter and Palmer 1995) and will only be given very briefly. Animals were anesthetized with a single injection (1.3 g/kg ip) of 20% urethan (2 g urethan, Sigma U-2500, in 10 ml distilled water). Atropine sulfate (120 μg sc) was administered to reduce bronchial secretions (atropine sulfate injection BP 600 mg/ml). Depth of anesthesia was indirectly monitored using the pedal withdrawal reflex, and maintenance doses of phenoperidine (1 mg/kg im, intramuscular Operidine 1 mg/ml, Janssen Pharmaceuticals) were applied when required.

All procedures were carried out in a double-walled sound-attenuated chamber and in accordance with UK Home Office regulations. The animal was tracheostomized to allow artificial ventilation when necessary. The animal’s core body temperature was monitored rectally and maintained in the range 37–38°C by a heating blanket. The animal was placed in a stereotaxic frame, the left earbar of which had been replaced with a hollow Perspex (Plexiglas) specimen. A silver-wire electrode was placed on the round window of the cochlea via an opening in the bulla to monitor the compound action potential. A polythene tube was inserted into the opening to equalize middle ear pressure, and the hole was resealed with petroleum jelly. A posterior craniotomy exposed the cerebellum. The dura was removed and the exposed cerebellum was covered with 1.5% agar.

Single-unit extracellular recordings were made from the CN of the guinea pig using tungsten-in-glass microelectrodes (Bullock et al. 1988) inserted into the left VCN using stereotaxic co-ordinates. The microelectrode signal was passed via a high-impedance headstage (Axon Instruments HS-2) to an Axoprobe 1A microelectrode amplifier. The signal was filtered (300–2,000 Hz) before being further amplified. A spike discriminator produced an electrical pulse when the spike voltage exceeded a threshold level and a CED1401 laboratory interface time-stamped the electrical pulses with an accuracy of 10 μs.

Pure tone and SAM stimuli were produced by a Hewlett Packard HP3325A synthesizer/function generator. SAM tones were 3 s long with a 6.5-s repetition period and presented at 20 dB above unit threshold. The signals were attenuated before reaching the driver amplifier for a reverse-driven Bruel & Kjær 4134 12.7-mm condenser microphone, coupled to a damped, 4-mm inside-diameter probe tube. This earphone was fitted into the speculum at the left ear. The sound system was calibrated at the start of each experiment. An anti-distortion network and safe driving voltage level were used to ensure that harmonic distortion products from the condenser microphone were negligible.

Steady-state vowels were produced using the Klatt software synthesizer and sampled at 10 kHz. The vowel /a/ had a fundamental of 100 Hz and three formants at 730, 1,090, and 2440 Hz with band- phone were negligible.
The presence of ISI patterns can be displayed using ISI scattergrams, where each ISI is plotted against its following ISI. Mode-locked spike trains can be described through either I) their spike times sequence: \( T_n = (n/p) + \varphi, n \Delta \) where \( T_n \) denotes the nth spike time, \( n \) is the integer part of \( np \) and \( \varphi \), one of the \( p \) phases defining the spiking pattern, or 2) their sequence of ISI: \( I_{SI} = T_{n+1} - T_n \). A mode-locked sequence is thus defined by a set of \( p \) phases, or \( p \) ISIs, which repeats every \( q \Delta \) (where \( \Delta \) is the input period). The presence of ISI patterns can be displayed using ISI scattergrams, where each ISI is plotted against its following ISI. Because a periodic sequence of \( p \) ISIs is constituted of ordered pairs of ISIs, the sequence appears as a set of \( p \) points (see Fig. 1). ISI scattergrams have been used to reveal deterministic dynamic properties of datasets (Kantz and Schreiber 1997; Sauer 1994) and in electrophysiology, to show the presence of bursts in spike trains (Krahe and Gabbiani 2004).

**SURROGATE SPIKE TRAINS AND MODE-LOCKING CRITERION.** Mode-locked spike trains have by definition both specific ISI sequences and a set of preferred phases of spiking within an AM cycle. Displaying preferred ISI pairs with ISI scattergrams could have nothing to do with mode-locking. This would be the case if a spike train possessed the observed conditional ISI preference but was not locked to the AM cycle. Conversely, the observed conditional ISI structure could simply be inherited from a preference for spiking around certain phases. In this case, the observed ISI sequence would be much different from the obtained conditional structure (condition A) and if the stimulus locked structure was not enough to account for the observed conditional ISI structure (condition B). Hence for the spike train to be mode-locked, we specified that the timing of spikes needed to match both the phase and ISI description of mode-locked discharges.

To test condition A, we created surrogate ISI sequences which maintain the subsequent ISI dependencies using the following shuffling procedure (see Fig. 3A): 1) randomly pick from the data one ISI, say \( I_{SI,m} \), and assign ISI, to the surrogate sequence. 2) Find \( \{i | [I_{SI} = I_{SI,i}] \} \), i.e., all the ISIs equal to \( I_{SI,m} \), and build the list of ISI pairs: \( L_1 = \{I_{SI}=I_{SI,m}\} \). And 3) randomly pick one pair from \( L_1 \), say \( I_{SI,m}, I_{SI,m+1} \) and assign \( I_{SI,m+1} \) to the surrogate sequence. This maintains the ISIs pair dependencies: in the surrogate sequence an ISI of duration \( I_{SI} \) is followed by one of duration \( I_{SI,m+1} \) with a similar probability than the original ISI sequence. Then we repeat steps 1–3 until no more pairs remain. So we find in the data all ISIs that are equal to \( I_{SI,m+1} \) build \( L_2 = \{I_{SI}, I_{SI,m+1}\} \) choose a pair in \( L_2 \) and assign \( I_{SI,m+1} \) to surrogate sequence. To assess whether ISI dependencies are enough to produce phase-locking, we tested the surrogate spike train using a standard Rayleigh test. We denote by \( R_m \) the value of this test in the rest of this article.

To test condition B, a surrogate spike train is created by randomly shuffling the phases of the genuine spike train (see Fig. 3B). The dissimilarity of ISI sequences for surrogate versus genuine spike trains was assessed by computing the root mean squared error (RMSE)
between their ISI distributions. We compared this RMSe from the phase shuffling to the RMSe within data trials. The resulting statistic is expressed in the results as a z score \( Z_{ps} \):

\[
Z_{ps} = \frac{\text{RMSe}_{ps} - \text{RMSe}_{trials}}{\sigma_{\text{trials}}}
\]

We considered that the phase shuffling had destroyed some of the original ISI structure if the z score was >2 (i.e., the distance between the means was above 2 SD of the intertrial variability—see Fig. 3B). Because the trial length was different between the vowel and harmonic stimuli (120 ms) compared with AM tones (3 s), vowels and harmonic stimuli trials were grouped into sets of 30 randomly chosen trials to obtain a valid estimation of the phase distribution.

A neuron was considered to show stochastic mode-locking if conditions A and B were satisfied. For the AM stimulus, for which a few units were tested with many AM frequencies, we focus in the population statistic on the number of cells showing mode-locking. We consider that a neuron was mode-locking if conditions A and B were satisfied for >50% of modulation frequencies tested. For vowels and harmonic complex stimuli, for which the number of stimuli tested varied from unit to unit, we give the population statistic in terms of numbers of mode-locked responses. Hence we consider the presence of mode-locking for the response of each unit to each stimulus separately.

One possibility that concerned us was that the phase shuffling procedure introduced many very short intervals (<1 ms) that could not occur in the data simply due to refractory effects. To check that these short intervals were not affecting the z scores, we devised an alternative procedure to obtain surrogate spike trains with identical phase distributions but no ISIs <1 ms. To do so, we used a nonhomogeneous Poisson process with a 1-ms dead time (absolute refractoriness) the intensity of which was obtained from the phase distribution. We computed for all units the z score required to pass condition B using this nonhomogenous Poisson process with dead time. This z score is denoted by \( Z_{\text{poisson,B}} \) and is also shown in Figs. 4 and 5. The \( Z_{\text{poisson,B}} \) values are generally lower than the z scores, indicating that some of the measured difference was attributable to this refractory effect. However, these values remain well above two whenever the phase shuffle generated z score is high. Note that this is not quite the same as a Poisson process being driven by the input to the unit.

BOOTSTRAPPING TEST OF SIGNIFICANCE. Comparisons of some experimental measures across onset and chopper unit populations were made using bootstrap methods to predict the probability that two populations with the measured difference in means could arise if the two groups belonged to the same underlying population. The samples from both original populations were pooled, and then repeatedly randomly redistributed into two groups (500 times). For each resampling, the mean difference between the two groups was calculated. This generated a distribution of differences for the means, and allowed us to estimate the probability (our P value) that these two groups could have observed the difference in the mean if the samples were drawn from a single population.

Model

PERIODICALLY FORCED LIF MODEL. We used a LIF model throughout this article to exemplify mode-locked response patterns, to reproduce the mode-locked pattern of discharges of recorded cells, and finally to help understand the organization of the responses to AM tones for a whole range of stimulation parameters (Supplementary Fig. S4).1 The model has linear subthreshold properties, similar to a stellate cell, and generates action potentials by an ad hoc threshold mechanism. The dynamics of the membrane potential are given by the following equation

\[
\frac{dV}{dt} = -\frac{(V - V_o)}{\tau} + I_{\text{app}}(t)
\]

where \( V \) denotes the membrane potential, \( V_o \) the resting potential, \( \tau \) the membrane time constant and \( I_{\text{app}}(t) \) the input applied to the model. The input \( I_{\text{app}} \) was either deterministic of the form \( I_{\text{app}}(t) = I_o + e_o \cos(2\pi ft) \) or stochastic with \( I_{\text{app}}(t) = I_o + e_o \cos(2\pi ft) + \sqrt{2D}\delta(t) \) where \( \delta(t) \) is the additive Gaussian noise and D its intensity (see Fig. 1). When fitting the data, we used shot noise driven by a model of the auditory periphery (see details in next paragraph).

STOCHASTIC INPUT USED WHEN FITTING THE LIF MODEL TO VCN CHOPPER UNITS. In our simulations of chord unit responses to acoustic stimuli, the LIF neuron model received its input from an AN model of the guinea pig cochlea (Sumner et al. 2003b). This first stage employed a dual resonance nonlinear (DRNL) filter-bank simulating the nonlinear mechanical filtering in the ear (Meddis et al. 2001), a neural transduction stage from a model of the inner hair cell (Sumner et al. 2002), and a nonhomogeneous Poisson process with a dead time modeling the AN discharges. These discharges were low-pass filtered (1st-order Butterworth, cut-off: 300 Hz) to simulate the effect of the dendrite on auditory nerve excitatory post synaptic potential (EPSPs) (following Hewitt and Meddis 1993). Parameters are given in Supplementary Table S1.

The LIF neuron model was fitted to the data by varying several model parameters (by hand). Characteristic frequencies of the input AN fibers were always set at the BF of the chopper unit. Bandwidths and gains of the nonlinear part of the DRNL filter were varied to match the frequency response area recorded from the data. For each cell from the data set, only one type of fiber (high medium or low spontaneous rate) was input to the LIF neuron model. The type was selected by varying two calcium parameters of the IHC synaptic release \( \left(G_{c_{\text{max}}, \left[C_{\text{calc}}\right]}\right) \). Four more parameters were varied in the LIF neuron model to replicate the ISI scattergram pattern: the membrane time constant, the threshold voltage, the synaptic strength of the AN inputs, and the numbers of AN fibers.

RESULTS

Spike timing and mode-locking in response to periodic stimulation

Models of neurons often show synchronized spike trains in response to a periodic input (Aihara et al. 1984; Keener et al. 1981). These patterns of synchronization can be described as \( q:p \) “mode-locked” spike trains when \( p \) spikes occur in an orderly manner during \( q \) input periods. Infinitely many periodic spiking patterns of this sort can be defined (2:3, 5:13, \( q:p \), etc.). Figure 1 shows two artificial examples of such firing patterns as well as their stochastic analogue. Figure 1A shows a 1:1 mode-locked spiking pattern in which there is one precisely timed spike on every cycle. The raster below it (Fig. 1B) shows a stochastic version of the 1:1 discharge pattern. Spikes no longer occur on every cycle but are still fairly precisely centered around a given phase. The VS quantifies the jitter around that single phase (Fig. 1E) and decreases when the noise increases. ISIs of the deterministic 1:1 mode are distributed at the period of stimulation (Fig. 1F). For the stochastic input, the ISI distribution has a multi-peaked structure centered at integer multiples of the stimulation period.

Figure 1C presents an example of a more complex spiking pattern: a 2:3 mode. The discharge pattern is defined by three

1 The online version of this article contains supplemental data.
precisely timed spikes distributed over two cycles. The phase distribution is multi-peaked leading to a VS = 0.5 although spikes are perfectly synchronized to the stimulus. A stochastic mode-locked response has even lower vector strength (Fig. 1D). Noise jitters the phases, widening their distribution. The ISI distribution for the deterministic pattern has three distinct intervals, but this structure is blurred for the stochastic 2:3 pattern (Fig. 1F).

To unravel mode-locked responses to a noisy periodic input, we use ISI scattergrams, which plot one interval against the previous. Figure 1J shows the deterministic 2:3 mode consists of three consecutive ISIs (9, 4.5, and 8.5 ms), which yields three points (black points at [9, 4.5], [4.5, 8.5] and [8.5, 9]) in the ISI scattergram. Note that the sum of ISIs adds ≤22 ms, twice the period of the stimulus. One can roughly consider two different actions of a noisy input (e.g., random arrivals of EPSPs from AN fibers) on the periodic response: a spike can be delayed or advanced (spike jitter), blurring the position of the points in the scattergram (Fig. 1I, I and J; gray points) and a spike creation or deletion can occur. Examples of spike deletion occur in Fig. 1H for the 1:1 mode. The absence of a spike will create an ISI twice as long as normal that appears on the ISI scattergrams as a regular grid of points.

The responses in Fig. 1 were obtained from a LIF model of neuron driven by a 90-Hz sinusoidal input. The mode-locking is the result of the interaction between the time needed for the LIF neuron model to integrate an input up to threshold and the periodicity of its input. LIF models are a type of nonlinear oscillator that often show mode-locking behavior when stimulated periodically and for which a considerable body of mathematics exists.

**Chopper VCN unit responses to AM tones**

Because LIF models like the example in Fig. 1 have successfully reproduced many aspects of VCN chopper unit responses, we reasoned that chopper cells should show these complex modes of firing. Examples of ISI scattergram representations are given for a VCN chopper unit (808009) in Fig. 2. This unit shows a regular (CV = 0.3 - Fig. 2B quadrant) “chopping” response in its peristimulus time histogram (PSTH, B) in response to pure tones at CF, which classified it as a sustained chopper (C). Adding Gaussian noise (D = 0.025 V/√Hz) to the previous LIF model broadens the distribution of phases decreasing the VS. E and F: period histogram and interspike interval (ISI) distribution of the 1:1 mode (black) and the stochastic 1:1 (gray). G and H: period histogram and ISI distribution of the 2:3 mode (black) and the stochastic 2:3 (gray). I and J: ISI scattergram representation of the modes. Black dots represent the sequence of intervals for the deterministic modes.

**FIG. 1.** Examples of different patterns of spike timing in response to a periodic signal, demonstrated using a leaky integrate-and-fire (LIF) model with a sinusoidally modulated input. From top to bottom: A, precise phase-locked pattern of discharge [LIF parameters: τ = 7 ms, Vth = -65 mV, Vrest = -50 mV, Vreset = -70 mV, g = 11.9*cos(2*π*90*Hz*t)], B: the influence of additive Gaussian noise added to the input (D = 0.025 V/√Hz). The noise creates the random skipping behavior and widens the period histogram. C: an example of a 2:3 mode-locked pattern [LIF parameters: τ = 7 ms, Vth = -65mV, Vrest = -50 mV, Vreset = -70 mV, g = 2.1 + 3.9*cos(2*π*90*Hz*t)], C: the sequence of discharges is defined by a set of phases and gives a low VS (0.51). D: adding Gaussian noise (D = 0.025 V/√Hz) to the previous LIF model broadens the distribution of phases decreasing the VS. E and F: period histogram and interspike interval (ISI) distribution of the 1:1 mode (black) and the stochastic 1:1 (gray). G and H: period histogram and ISI distribution of the 2:3 mode (black) and the stochastic 2:3 (gray). I and J: ISI scattergram representation of the modes. Black dots represent the sequence of intervals for the deterministic modes.
phases. We show in Supplementary Fig. S1 how a 2:3 mode can be sorted in cycles with one spike and cycles with two spikes (see supplementary methods and Fig. 1).

Figure 2E (see also red lines in B and C) shows that a LIF model can match the sequential temporal response of this chopper unit. We used a model of guinea pig auditory nerve fiber activity (Sumner et al. 2002) as input for a LIF model (see METHODS). The fit of the LIF parameters was realized for two of the frequencies of AM presented (1st and last quadrant of Fig. 2E). The LIF model gives ISI scattergrams very similar to the data at all AM frequencies (see supplementary movies). We interpret this match for all AM frequencies as a demonstration that the chopper unit and LIF model responses are constrained by the same underlying dynamic: their discharges are in both cases reminiscent of mode-locked states.

Mode-locking criteria from surrogate spike trains

The patterns of intervals seen in Fig. 2 were typical of those seen in other cells in our sample, indicating that the stimulus envelope was being encoded by mode-locked patterns rather than just phase-locked (see supplementary movie). To quantify this, we considered that mode-locking arose from an interaction of the time since the last spike and the phase within the AM cycle. Therefore mode-locked spiking patterns are characterized by a dependence on both the spiking phases and ISIs. We used two different shuffling schemes to rule out the possibility that the observed spike train temporal structure could arise solely from either the phase preference or the conditional dependence between successive ISIs (see METHODS). The first shuffled spike trains aimed at determining whether the ISI clusters observed were only due to a specific pair dependency of ISIs without any link to the periodicity of the stimulus. The second scheme was to clarify the difference between mode-locked discharge patterns and discharges showing solely phase preference, such as those observed at the level of the auditory nerve. Figure 3, A–C, shows the effect of each shuffling scheme on the period histogram and the ISI scattergram.
We applied a statistical criterion to each unit of our data set for both shuffling schemes. Specifically, we tested that after interval shuffling, there was no longer a significant phase preference at the modulation frequency ($R_{iw} < 13.8$) and that the mean squared error between ISI distributions before and after phase shuffling was at least twice the intertrial variability of the raw ISIs ($Z_{ps} > 2$; see Methods). Figure 3A shows how interval shuffling is performed. Figure 4A shows the results of ISI shuffling on the AM responses. Maintaining pair statistics is not enough to explain the locking observed. Figure 3B shows scattergrams for the same unit before and after the phase shuffling, and diagrams of the derivation of the $z$ score measure. Essentially this test compares how single trial ISI histograms differ before and after shuffling, relative to the intertrial variability of the unshuffled ISIs (see Methods). Figure 4B shows that nearly all chopper units (5/7) of our dataset had very different ISI statistics when only their phase preference was maintained. This in turns means that most units showed significant mode-locking behavior. Modulation depth was important: mode-locking was observed in half of the cells tested with 50% modulation depth (6/11 units including 5/7 choppers, 2/2 unusual, 0/2 onset), while in 75% of the cells tested at 100%
responses to a complex tone with same
so as to minimize the role of locking to individual harmonics.
sidered 24 chopper and 33 onset units, all with CF
also show stochastic mode-locked spiking patterns. We con-
complexes). We reasoned that responses to these stimuli might
spectrally extended stimuli (vowels and cosine phase harmonic

correlation, and VS) and the fundamental frequency (F0) of
ent measures of temporal coding (ISI histogram peak, auto-
(Winter et al. 2003) described the relationship between differ-
chopper and onset units. This previously published dataset
similar analyses to the responses of a larger population of
patterns occurred to other periodic stimuli, we then applied
modulation depth (3/4 choppers). Each chopper displaying a
mode-locking spiking pattern could be fitted with a LIF model
(see supplementary table). The root mean squared (RMS) error
between the VS of the data and of the LIF model over all seven
chopper cells and for all their AM frequencies, was 0.087 (the
Fig. 2 fit achieves an RMS error of 0.108).

Mode-locked spike trains in response to complex harmonic stimuli

To further explore the extent to which complex spiking
patterns occurred to other periodic stimuli, we then applied
similar analyses to the responses of a large population of
chopper and onset units. This previously published dataset
(Winter et al. 2003) described the relationship between different-
measurets of temporal coding (ISI histogram peak, auto-
correlation, and VS) and the fundamental frequency (F0) of
spectrally extended stimuli (vowels and cosine phase harmonic
complexes). We reasoned that responses to these stimuli might
also show stochastic mode-locked spiking patterns. We con-
sidered 24 chopper and 33 onset units, all with CF > 1.5 kHz,
so as to minimize the role of locking to individual harmonics.

Figure 5 shows the responses of a sustained chopper (C, unit
293005) with high CF (3 kHz) to three different stimuli. In A
and B, we display the responses to two stimuli with similar fundamental frequency of a 100 Hz: a 120-ms steady-state vowel /l/ (A) and a cosine phase harmonic complex (B). Figure
5A shows responses close to a 1:2 mode (rate per cycle = 1.87)
with clusters in the ISI scattergram at (4 ms, 6 ms) and (6 ms,
4 ms), but almost no 10-ms intervals. Hence first-order ISIs
would not show $F_0$ encoding in this case. In Fig. 5B, this unit
responds to a complex tone with same $F_0$ but with a different
spiking pattern. The ISI scattergram shows in part similar clusters at (4, 6) and (6, 4) but shows also locking to the
fundamental period (10-ms intervals), which is clear even in
the spike raster (the 1st 20 repetitions of 100 are shown).
Therefore periodic stimuli with the same period but different
envelopes create patterns of spiking organized around a differ-
ent structure of intervals. In C, we show the responses of the
same chopper unit to the vowel /l/ with a fundamental of 125
Hz. The pattern of spikes is centered more closely around the
[2, 6]-ms ISI range but still considerable regularity is clear in
the raster. Two clusters are just about visible in the scattergram
at (5, 3) ms and (3, 5) ms. A bimodal structure is also clear in
the period and interval histograms for these stimuli. Indeed the
phase distribution resulted in low VS despite the precision of
firing within the phase.

For all three stimuli, the spiking patterns of this unit could
not be accounted for by their phase or paired ISI structure.
Applying the same phase and interval shuffling tests as for the
AM stimuli gave a nonsignificant Rayleigh test ($R_{is} < 13.8$,
shuffled phase is superimposed in black in Fig. 5) and large Z
score ($Z_{is} > 2$, shuffled ISI is shown in black in Fig. 5). The
CF of the unit is important here (3 kHz) as it falls above the
phase locking limit but is still well driven by the vowel stimuli.
This unit is unlikely to be responding to significant energy
from the low-numbered harmonics. Instead it is most likely
driven by envelope of the harmonics around 3 kHz. From this
example, it is clear that broadband harmonic stimuli produce
complex yet noisy periodic spiking patterns in chopper units,
and that the particular pattern of spiking is sensitive to the
stimulus envelope. Two further examples of chopper units are
given in the supplementary material.

We also observed complex responses in VCN onset units.
Figure 6 shows further examples of responses to the vowel /l/
(100-Hz fundamental) from three different onset units. All
three units have a high probability of firing at a particular
phase, but although all have a similar CF (~3 kHz), they
respond quite differently in other respects. The left-most panel
(an $O_v$ unit) is an example of a highly complex pattern of
spikes, and our criteria indicates that neither phase nor inter-
vals alone can account for this spiking pattern. In between the
spikes at the dominant phase, which form a large number of
10-ms intervals seen as in the scattergram at (10, 10) ms and
also in the ISIH, further spikes produce several sets of inter-
vals, forming a diagonal set of interval pairs in the ISI scat-
tergram. These diagonals tend to occur in our data whenever
the dominant interval corresponds to the fundamental, and the unit is firing in close to a 1:1 mode. This cell is firing almost once per cycle (1.26 spike/period, this is very clear in the raster) and thus is almost perfectly entrained to the \( F_0 \). However, the extra intervals markedly reduce the VS. The Oc unit in the middle panel is even more nearly perfectly entrained to the 100-Hz fundamental (VS 0.94, 1.05 spike/period) with an occasional extra spike occurring precisely 2 ms after the dominant phase. This unit also shows a high \( z \) score, but the Rayleigh indicates that the amount of phase structure that can be accounted for by the interval structure alone is significant.

In the right-most column of Fig. 6 is an example of an Oc unit in the middle panel is even more nearly perfectly entrained to the 100-Hz fundamental (VS 0.94, 1.05 spike/period) with an occasional extra spike occurring precisely 2 ms after the dominant phase. This unit also shows a high \( z \) score, but the Rayleigh indicates that the amount of phase structure that can be accounted for by the interval structure alone is significant. In the right-most column of Fig. 6 is an example of an Oc unit in the middle panel is even more nearly perfectly entrained to the 100-Hz fundamental (VS 0.94, 1.05 spike/period) with an occasional extra spike occurring precisely 2 ms after the dominant phase. This unit also shows a high \( z \) score, but the Rayleigh indicates that the amount of phase structure that can be accounted for by the interval structure alone is significant.

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The relationship between $z$ score and CV partly reflects differences between choppers and onset units. In our sample of cells that passed both our criteria, we found that onset cells have on average a significantly lower $z$ score than chopper units (chopper units: 19.4 ± 9.3; onset units: 15.1 ± 10.6, means ± SD; $P < 0.05$ nonparametric bootstrap test), and onset cells also tend to have a higher average CV than chopper units (due to the tendency of onset units to show less sustained discharge to a pure tone) (Winter and Palmer 1995). However, this relationship is not simply a difference between different cell types. Noticeably the relationship still holds when considering chopper responses alone ($R = 0.5$, $P < 0.001$). This negative relationship between the regularity (CV) of the response and the $z$ score implies that more regular units tend to show more complex ISI structure in response to sounds with envelope modulations.

The second relationship, between $z$ score and spikes per cycle, again partly reflects differences between unit types. In addition to having slightly higher $z$ scores, chopper units have a larger number of spikes per period (1.6 ± 0.7) than the onset population (1.1 ± 0.6, different at $P < 0.01$ for a bootstrap test). However, this relationship is also strong in the chopper and onset populations considered separately (chopper units: $R = 0.62$, $P < 0.001$; onset units: $R = 0.72$, $P < 0.001$). This suggests that the more spikes per period, the less of the interval structure could be accounted for by the phase structure. It is important to note that this is not what would be expected of a Poisson process, where spike rate increases without any increase in $z$ score (there was no correlation in a simulation of modulated Poisson processes—not shown). There was also a relationship between the number of spikes per cycle and VS (Fig. 7C, $R = 0.48$, $P < 0.0001$). This of course fits with the example models shown in Fig. 1, which makes it clear that VS does not well characterize the precision of periodic coding when there is more than one spike per period. We observed no correlation between VS and $z$ score. All these relationships held if we derived the $z$ score from a modulated Poisson-process with a 1-ms dead time (see Methods).

The points on the left of Fig. 7D indicate the proportion of each population that failed the Rayleigh test. In other words, after interval shuffling the phase locking to the fundamental was still significant. Clearly, a much larger proportion of onset responses (56%) failed this first criteria compared with choppers (20%). These proportions correspond almost exactly to the proportions that passed both tests, indicating that almost all failed to make the criteria for mode-locking because they failed the Rayleigh test. We observed that the failure of the Rayleigh test only happens when VS is high. We interpreted this in the following way: very reliably spiking near the same phase was resulting in very small phase jitter buildup during interval shuffling and hence in a significant phase locking after ISI shuffling. This is true for both the chopper (0.81 ± 0.11) and
the onset ($0.91 \pm 0.11$, $P < 0.01$ for a bootstrap test of significance) populations. Thus we see that nearly all cells fail to pass our test for mode-locking because they are phase locking almost perfectly to the fundamental. However, VS and the Rayleigh test do not distinguish between Poisson like stochastic firing at a particular phase in the period, and when a cell is reliably firing every cycle, as in a 1:1 ratio mode-locked state, or entrainment. The number of spikes per cycle for the chopper units that failed was close to 1 ($0.96 \pm 0.28$), suggesting that these cells were well entrained to the stimulus period. For onset cells the number of spikes per cycle was slightly lower and more variable ($0.77 \pm 0.46$, the difference of means was not significant for a bootstrap test). Our interpretation here is that some cells are behaving in a roughly Poisson like manner, while others are responding in a 1:1 mode.

From these analyses, it is clear that both chopper and onset units show complex spiking patterns in response to periodic stimuli of different types. Consistent with our starting hypothesis these are reminiscent of mode-locking, poorly described by the vector strength and strongest in chopper units. Furthermore, these spiking patterns are sensitive to both the fundamental and the spectrum of the stimulus.

**DISCUSSION**

We have shown here the existence of mode-locked spiking patterns in VCN chopper units in response to SAM tones, steady-state vowels, and harmonic complexes. Mode-locking behavior is observed in chopper and onset units (all of which are presumed to be multipolar cells) but is strongest in chopper units. Our insights originate from a LIF model that reproduces the sequential pattern of ISIs. The mode-locked behavior in VCN neurons is stochastic, and although clearly not-random, it is difficult to ascribe a particular mode to a particular spike train.

We specifically hypothesized that chopper unit responses should show mode-locked responses because of the previous success of LIF models in modeling them. The model best fits were obtained for few ANF inputs (4–14, see supplementary table), close to estimations obtained in mice VCN slices using minimal ANF electrical stimulation (Ferragamo et al. 1998). Although we did not attempt to fit models to the responses to vowels, the responses seen to these stimuli were nevertheless qualitatively similar to the mode-locking seen in response to AM, and it is likely that a large portion of these responses could be reproduced with fairly simple LIF based models (Hewitt and Meddis 1993).

Two of the 11 units stimulated with AM tones were onset-choppers, and these also showed clustered interspike interval scattergrams, which could be accounted for by the phase distribution alone. This tendency was also evident in the population of onset units stimulated with broad band signals (complex tones and vowel sounds) where around 50% of the spiking patterns observed can be explained by their phase preference. However, the other half of onset units showed more complex spiking patterns. In many instances, onset cells locked almost perfectly to the fundamental. This behavior, often called
entainment, corresponds to a 1:1 mode with a single preferred phase and ISI. Onset-choppers are stellate cells, they show regular spiking in response to injected current (type-I spiking) (Oertel et al. 1990) and considerable temporal integration (Palmer and Winter 1996), but they differ from other stellate cells in their intrinsic currents (Oertel et al. 1990). They also receive convergent inputs from many different ANFs. These factors may conspire to ensure that firing in these cells remains close to the 1:1 mode.

In VCN, the regular spiking of stellate cells contrasts with the phasic spiking of bushy cells and octopus cells (Bal and Oertel 2001), known to come from \( I_{KL} \) (Rothman and Manis 2003), a low-threshold potassium current, present in bushy and octopus cells. This current limits integration and fosters coincidence detection. For spherical bushy cells responses are similar to those of the auditory nerve because they are connected by only a few large auditory nerve synapses (endbulbs). Thus we can expect both types of bushy cells to show a stochastic type of behavior more similar to the ANFs. We did not have any data from primary-like units with which to test this hypothesis. However, it is possible that some of the onset responders in our sample are octopus cells.

We have undertaken here a description of the observed spiking pattern from the point of view of forced dynamical systems. An alternative approach is to use a purely statistical description of spike times. Johnston and colleagues (1986) modeled chopper responses to pure tones in the lateral superior olive using stochastic point processes. In this case, the point process needs to be more regular than a Poisson process, i.e., the model has a preferred ISI. Rather than obtaining regularity through the hazard function of a renewal process, we set it from biophysical parameters in the LIF model, one advantage being that the description remains valid for time varying stimuli, such as AM tones. It also shows that chopper intrinsic dynamics is a main feature of the response to periodic stimuli. The intrinsic frequency of chopper units was already crucial in some pitch processing models (Wiegrebe and Meddis 2004) and has been reported by Kim et al. (1990) to correlate with the best temporal modulation frequency.

The relation between intrinsic chopping frequency and best temporal modulation frequency is clearly addressed by the theoretical framework of mode-locked synchronization. Indeed there is a clear organization of mode-locked responses in chopper units across frequency of AM as is displayed in Fig. 2 (see also the supplementary movies). For the fitted model, reducing the noise shows how mode-locked discharges are organized in an orderly manner in parameter space (see Supplementary Fig. S4). Progressively increasing stimulation frequency, one goes from \( p > q \) modes, through the 1:1 mode and toward \( q > p \) modes for higher frequencies. The observation of Kim et al. is thus in accordance with our observation that the 1:1 mode displays the strongest VS and that in models with sufficiently low noise, the 1:1 mode converges to the chopping frequency at 0% modulation depth (Supplementary Fig. S4). Previous studies on spike timing reliability (Hunter et al. 1998; Schreiber et al. 2009) also point at maximum reliability at a modulation frequency equal to the intrinsic frequency.

In conclusion, this study demonstrates mode-locked spiking responses to AM tones, vowels, and harmonic complexes in VCN chopper and onset units. Early work (Knight et al. 1972) on 1:1 locking pointed at the deleterious effect that synchronization represents for stimulus encoding in LIF populations when compared with asynchronous firing. However, recent work has shown how LIFs complex spiking patterns can provide specific coding advantages (e.g., nonuniform sampling stimulus reconstruction) (Lazar 2005). The ISI dependencies displayed by mode-locked spike trains are intriguing as it is known that subsequent ISI dependencies can increase information transfer by spike trains (Chacron et al. 2004). Moreover, knowledge of both the phases and \( p,q \) value are important to characterize precisely the stimulus parameters. For example, although a 1:2 mode exists at 44 Hz with 100 to 30% of modulation depth in Supplementary Fig. S4C, the phases at 100 to 30% are different (as seen by the change in VS) and allow disambiguation. Likewise AM depth, frequency, and envelope shape all influence the mode-locked spike train pattern. For example, it is clear in Fig. 5 (and the supplementary figures) that the patterns of spikes are sensitive to whether the stimulus is a vowel or a harmonic complex. Hence the exact sequence of ISIs might carry additional information about the stimulus. For instance, ramped and damped AM tones, which are known to affect neural responses in CN (Pressnitzer et al. 2000) and change roughness perception (Pressnitzer and McAdams 1999), will produce different sets of mode-locked patterns. Mode-locking could thus play a role in encoding multiple features of complex stimuli, possibly more sensitive to stimulus parameters than a pure Poisson based coding. Further work, describing the responses to a larger range of stimuli (such as IRNs or periodic signals in noise) is needed to fully relate mode-locking to auditory temporal processing. A linear analysis of spatiotemporal interactions along the frequency axis has already proven to be advantageous for temporal representation (e.g., Shamma 1985). Thus a full characterization of spatiotemporal mode-locked encoding constitutes an attractive problem from both the experimental and theoretical perspective.

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