Temporal Coding of Voice Pitch Contours in Mandarin Tones

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Accurate perception of time-variant pitch is important for speech recognition, particularly for tonal languages with different lexical tones such as Mandarin, in which different tones convey different semantic information. Previous studies reported that the auditory nerve and cochlear nucleus can encode different pitches through phase-locked neural activities. However, little is known about how the inferior colliculus (IC) encodes the time-variant periodicity pitch of natural speech. In this study, the Mandarin syllable /ba/ pronounced with four lexical tones (flat, rising, falling then rising and falling) were used as stimuli. Local field potentials (LFPs) and single neuron activity were simultaneously recorded from 90 sites within contralateral IC of six urethane-anesthetized and decerebrate guinea pigs in response to the four stimuli. Analysis of the temporal information of LFPs showed that 93% of the LFPs exhibited robust encoding of periodicity pitch. Pitch strength of LFPs derived from the autocorrelogram was significantly (p < 0.001) stronger for rising tones than flat and falling tones. Pitch strength are also significantly increased (p < 0.05) with the characteristic frequency (CF). On the other hand, only 47% (42 or 90) of single neuron activities were significantly synchronized to the fundamental frequency of the stimulus suggesting that the temporal spiking pattern of single IC neuron could encode the time variant periodicity pitch of speech robustly. The difference between the number of LFPs and single neurons that encode the time-variant F0 voice pitch supports the notion of a transition at the level of IC from direct temporal coding in the spike trains of individual neurons to other form of neural representation.

Keywords: temporal coding, time-variant, voice pitch contours, natural speech, inferior colliculus, fundamental frequency

INTRODUCTION

Voice pitch is a perceptual quality of speech that varies with the fundamental frequency (F0), and is controlled by the rate of vibrations produced by the vocal folds. Natural sounds such as animal vocalizations and human speech are complex, quasi-periodic signals that can be partly characterized by their F0 and its harmonics. Periodicity in speech signals is related to the perception of low pitches (‘periodicity pitch’) (Plomp, 1967; Small, 1970; De Boer, 1976;
Studies have also shown that neural mechanisms coding periodicity have a temporal-to-rate coding transformation in thalamus and cortex (Bartlett and Wang, 2007; Wang et al., 2008; Bendor and Wang, 2010). The spike rate of pitch-selective neurons can also represent the periodicity pitch of temporally regular sounds in the low frequency region of auditory cortex in marmosets (Bendor and Wang, 2005, 2010). Together, these studies suggest that the temporal representation of periodicity pitch may be transformed to a rate-based representation (Bartlett and Wang, 2007; Wang et al., 2008; Bendor and Wang, 2010; Plack et al., 2014; Langner, 2015) or spatial-temporal representation (Loe et al., 1983; Oxenham et al., 2004; Bernstein and Oxenham, 2005) in the upper auditory pathway.

Scalp-recorded frequency following responses (FFRs) reflect sustained phase-locked activity of populations of neurons (Worden and Marsh, 1968), and are generated predominately from the IC (Smith et al., 1975; Chandrasekaran and Kraus, 2010). FFRs have been proposed to represent periodicity pitch of speech (Greenberg, 1981). For example, Krishnan et al. (2004) recorded FFRs to Mandarin speech sounds and found that the phase-locked interpeak intervals of the FFR waveform robustly matched the time-variant fundamental period of the stimuli. Gockel et al. (2011) also found that scalp-recorded FFR responses in humans were similar to modeled neural responses in the auditory periphery, and did not necessarily reflect any additional pitch-related processing. However, the periodicity strength of FFR was stronger for people with tonal language than English (Krishnan et al., 2005, 2010; Bidelman et al., 2011; Chandrasekaran et al., 2012), for people who are musicians compared to no musical experience (Wong et al., 2007; Bones et al., 2014), for adults compared to neonates (Jeng et al., 2011), and after musical training compared to before (Song et al., 2008). Because it is hard to control for these factors, human studies are limited in their ability to explore neural coding mechanisms, while studies using animal models can provide more detailed and comprehensive information to evaluate the neural mechanisms of coding pitch information. In the present study, neuronal activity in guinea pig IC was measured to investigate how it represent the time varying periodicity pitch of natural speech. To our knowledge, this is the first study to explore the correspondence between the temporal information of the single IC neurons with the time varying F0 contours in Mandarin speech.

Neural activity measured from an electrode in the IC includes two components: local field potentials (LFPs), which are dominated by the synaptic input information, and action potentials from single neurons or multiple neurons, which reveal axonal output information (Pettersen et al., 2012). LFPs can be recorded in IC with high spatial resolution and reflect neural ensembles with synchronized synaptic inputs in a small area around the electrode (Kraus et al., 1994; King et al., 1999). LFPs recorded in guinea pig IC have shown phase-locking to the acoustic characteristics of steady state vowels and formant transition periods of synthetic speech sounds such as the vowel-consonant-vowel token /ada/ (Cunningham et al., 2002). White-Schwoch et al. (2016) recently compared the waveforms of multi-unit activity recorded in
the central nucleus of guinea pig IC and scalp-recorded FFRs in humans in response to same speech sound /da/. The authors showed that the responses’ waveforms were morphologically similar and contained a periodic component corresponding to the periodicity of stimuli. In the light of these studies, we hypothesized that the LFPs of the IC could represent the time-variant periodicity pitch of speech.

Compared with LFPs recorded in the IC which are dominated by the input to the IC, spiking activity recorded from the IC reflects the output information of neurons processing pitch information (Pettersen et al., 2012). Studies have shown that the discharge pattern of IC neurons synchronize to the F0 of harmonic tones. For example, Sinex and Li (2007) measured IC single neuron responses to a single harmonic tone, and found that the response envelope of some neurons modulated at an F0 of 250 Hz. Shackleton et al. (2009) also found that the temporal patterns of most IC multi-neuron clusters significantly synchronized at a frequency equal to F0 (at and above 282.8 Hz).

In contrast to synthesized harmonic tones with a fixed F0, animal vocalizations and human speech are amplitude and frequency modulated. For example, guinea pigs communicate using a characteristic ‘whistle’ – a vocalization with patterns of amplitude and frequency modulation (FM). Studies have demonstrated that neural responses in the IC can reflect the direction of amplitude and FMs (Suta et al., 2003; Woolley and Casseday, 2005). For instance, Suta et al. (2003) found that single IC neurons in guinea pig had a significantly stronger response to natural whistles compared to time-reversed whistles. Some neurons in the IC of bat are highly sensitive to the direction of amplitude and FMs (Suta et al., 2003; Woolley and Casseday, 2005). For example, Suta et al. (2003) found that single IC neurons in guinea pig had a significantly stronger response to natural whistles compared to time-reversed whistles. Some neurons in the IC of bat are highly sensitive to the direction of amplitude and FM. Studies have demonstrated that neural responses in the IC can reflect the direction of amplitude and FMs (Suta et al., 2003; Woolley and Casseday, 2005).

Studies have confirmed that spike activity patterns of IC neurons in animals are sufficiently abundant to explore the processing of speech (Perez et al., 2013; Steadman, 2015). For instance, Ranasinghe et al. (2013) found that IC neuronal activity patterns became more diverse when the speech added period information, and suggested that activity was determined by the F0 of speech.

In summary, previous research suggests that spike-timing information of single IC neurons could represent the periodicity of stimuli with a fixed F0, and the periodicity coding of IC neurons are affected by the stimulus type (i.e., click trains, sinusoidal amplitude modulated noise and iterated rippled noise) (Schnupp et al., 2015). However, how dynamic F0 patterns (such as are present in tonal languages) are encoded by IC neurons is currently unknown.

The aim of this study was to determine how the IC encodes time-variant F0 contours of natural speech. To address these aims, LFPs and single neuron activity were simultaneously recorded in the IC of anesthetized guinea pigs.

MATERIALS AND METHODS

Six healthy adult guinea pigs of either sex (weight 250–450 g) were used in this study. All guinea pigs were purchased from the Experimental Animal Centre of Chongqing Medical University. The procedures were performed in accordance with protocols of the Care and Use of Laboratory Animals approved by the Third Military Medical University.

Animal Anesthesia and Surgery

Surgical and neural data collection methods have been described previously (Peng et al., 2016). Healthy animals were maintained at a surgical level of anesthesia by an initial intraperitoneal injection of urethane (1.2 g/kg body weight in 20% sterile saline). Supplemental injections of urethane (0.16 g/kg) were administrated on indication of a paw withdraw reflex every 30–60 min. The body temperature of animals was maintained at 38°C by using a thermostatic bath (HSS-1B, Chengdu Instrument Factory, Chengdu, China).

The animal was placed inside a sound-attenuating room, which the ambient noise floor was 25 dB sound pressure level (SPL). Before surgery, auditory brainstem responses (ABRs) were recorded to assess the hearing sensitivity of animals. ABRs were evoked by a click stimulus (100 µs rectangular pulse) at a repetition rate of 4 Hz, and 600 clicks were presented at every intensity level (30 dB SPL to 80 dB SPL in 10 dB steps) in each animal. Stainless steel needle electrodes positioned at the vertex in the middle of ears, and the mastoid of the ipsilateral ear and nose served as active, reference and ground electrodes respectively. Normal hearing sensitivity was confirmed if the threshold of ABR was at or below 30 dB SPL.

After confirmation of normal hearing, a midline incision was made along the scalp surface, and the dorsal surface of the skull was exposed. The head was fixed in place anterior to the bregma by a custom head holder, which was held at its bottom by using 3 small screws and attached by dental acrylic. The right temporalis muscle was retracted and removed, and an approximate 5 × 5 mm opening was drilled in the skull dorsal to the temporoparietal suture and rostral to the tentorium (Snyder et al., 2004). The dura mater was then removed and the cortex was reflected and aspirated, allowing direct visualization of the surface of IC.

Acoustic Stimulation

Stimuli were four Mandarin words downloaded from the database of standard Mandarin speech (Mandarin, 2015) and the sample rate was 16 kHz. The four words were one syllable /ba/ pronounced by a male speaker with four lexical tones: bá ‘eight’, bá ‘pluck,’ bá ‘target,’ bá ‘father.’ The waveform and spectrogram of the speech used in this study are displayed in Figure 1; the duration of stimuli ranged from 400 to 600 ms. The F0 contours (Figure 1C) were extracted from the spectrogram by calculating the peak spectral energy in a sliding window (80 ms Hanning window shifted in 1 ms step). The F0 of the four tones varied with time (Figure 1C). The flat tone had a steady F0 of 170 Hz, the rising tone F0 increased from 120 to 210 Hz, the falling then
The waveform, spectrogram and F0 curve of the four Mandarin tones used in the current study. (A) From left to right, waveform of ‘bā’, ‘bá’, ‘bǎ’ and ‘bà’ stimulus, respectively. (B) Spectrogram of ‘bā’, ‘bá’, ‘bǎ’ and ‘bà’. (C) The low frequency (0–600 Hz) spectrogram with the F0 curve shown by a black line. The colors in the spectrogram represent spectral energy from blue (low) to red (high).

rising tone decreased from 120 to 90 then increased to 160 Hz, and the falling tone decreased from 220 to 80 Hz.

Sound stimuli were presented by a laptop using LabVIEW, converted by a 16-bit D/A converter (DAQ 9264; National Instruments) and delivered in free-field conditions from an earphone equipped with a custom-made box (Beyerdynamic DT 770, Germany). The earphone was located 1cm away from the animal’s ear canal contralateral to the exposed IC. The acoustic delivery system was calibrated (Tao et al., 2016) using a 1/4 microphone and amplifier (Piezotronics Inc., United States) to deliver a flat response (±50 dB) across the frequencies (0.5–8 kHz) used in this study.

Tone bursts (duration 12 ms with 2 ms rise/fall ramps, frequency from 0.5 to 8 kHz steps 1/8 octave) were presented to estimate the characteristic frequency (CF) of the single neuron and LFP. Twelve levels were presented at each frequency (from 30 to 80 dB SPL in 5 dB steps). For each penetration into IC, neuronal responses to speech were obtained from 100 to 1000 ms relative to stimulus onset. All speech stimuli were presented at a level of 80 dB SPL, and delivered in pseudorandomized order once every 2 s. Each stimulus was presented either 9 (for 39 out of 90 recording sites) or 10 times.

Neuronal Activity Recording
Extracellular neuron activity and LFPs were simultaneously collected using a single channel tungsten electrode (1–3 MΩ, Global Biotech INC., China), and the ground electrode was placed at the temporal bone below the skin. The electrode was mounted and controlled by a microdrive (50-12-1C Hydraulic Probe Drive and 50-12-9 Manual Drum Unit, FHC, United States), and advanced in a dorso-ventral direction with 1 μm steps. Bursts of broadband white noise (duration 50 ms, 10 ms rise/fall ramps) presented at 4 Hz were used as the search stimulus and single neurons were identified by visual inspection of the response waveform morphology (Palmer et al., 2013).

Several parallel tracks were made in sagittal or coronal planes of the IC for each animal. Single neuron activity was amplified by a headstage connected with a Cerebus multichannel physiological signal acquisition system (Cerebus 6.01, Blackrock Microsystems, Salt Lake City, UT, United States). Single neuron spikes were band-pass filtered between 250 and 5 kHz and sampled at a rate of 30 kHz. For LFPs, the signal was band-pass filtered from 50 to 500 Hz and at a sampling rate of 2 kHz.

Data Analysis
All subsequent data analyses were processed offline in Matlab 2016b.

LFPs of IC Recordings
In order to verify whether the recorded LFPs in response to speech originated from IC, the latency between each LFP and a low pass filter version of a given stimulus was examined. The low-pass filter was an eighth order digital Butterworth filter with cut-off frequency at 500 Hz and was applied in both forward and reverse directions to minimize phase distortions. The latency of each LFP was defined as the time lag corresponding to the peak of the cross-correlation (LFPs and low pass filtered version of stimuli) function.

The biphasic waveform of LFPs in response to a pure tone have an upward-going shape with a first positive peak at ∼6.5 ms and duration of ∼30 ms (Orton et al., 2012). For each LFP, the response area was constructed by calculating the peak-to-peak amplitude in the response duration after the stimulus onset for each frequency and stimulus level combination (Fallon et al., 2016). The CF was defined as the frequency which evoked a detectable peak above the background noise at the lowest stimulus level.
To visualize the time-varying periodicity of LFPs corresponding to the F0 of each stimulus, an autocorrelogram was constructed using a periodicity detection short-term autocorrelation algorithm (Krishnan et al., 2004). The algorithm calculated the cross-correlation function within a rectangular window (40 ms window duration shifted in 1 ms steps) between the signal (LFPs and low pass filtered stimuli) and the copy signal shifted in 0.2 ms steps. In the resulting autocorrelogram, the vertical axis shows the time lag between the original signal and its copy signal, and the horizontal axis shows the response time relative to the onset. Colors represent the degree of correlation function between the signal and its copy at each response time and corresponding time lag.

To quantify the degree of periodicity in the neural activity underlying the LFPs, the pitch strength was calculated. The pitch strength at each time window (40 ms) was the maximum correlation magnitude (LFPs and copy LFPs) at the time lag corresponding to the stimulus F0 (converted to a period) in the same time window. The overall pitch strength of each LFP was calculated by averaging the pitch strength measures over all time windows. A one way analysis of variance (ANOVA) was constructed to examine whether the pitch strength measure was different for each of the four tones.

To examine whether the LFPs could represent the time-variant F0 and spectral components of a stimulus, narrow-band spectrograms of the stimuli and LFPs were compared. Narrow-band spectrograms were generated by calculating the short-term Fourier transform of 80 ms sliding hamming windowed LFPs and the stimulus waveforms, shifted in 1 ms steps. The windowed signal was zero-padded to 250 ms before the Fourier transform; as a result, the frequency resolution was 4 Hz.

In order to quantify the degree to which the LFPs represented the F0 dynamically, the stimulus-to-response correlation was calculated (Pearson’s correlation coefficient r). We calculated the cross-correlation between the F0 contours extracted from the narrow-band spectrogram of each response and its corresponding stimulus. This measure provides both the strength and direction of the relationship between two signals. The F0 curve was extracted from the spectrogram of the LFP responses by finding the maximum spectral magnitude in predefined frequency ranges (F0 ± 50 Hz).

To further quantify the accuracy of LFPs coding F0 across stimulus duration, the linear pitch tracking error was computed by finding the absolute difference between stimulus F0 and response F0 at each corresponding time bin then averaging them across all time bins (Song et al., 2008).

**Single Neuron Activity**

For each single neuron, a frequency response area (FRA) was created with a procedure that delivered tone bursts at different frequencies and intensities. The FRA of each single neuron was calculated as the driven spike rate (the recorded spike rate minus the spontaneous spike rate) in a 50 ms window following tone burst onset. Spontaneous spike rate was calculated in 50 ms window before the tone burst onset. CF was determined as the frequency of a tone burst that evoked a detectable response at the lowest stimulus level.

**RESULTS**

**Local Field Potentials**

Simultaneous recordings of LFPs and single neuron spike activity at 90 IC sites were carried out in six guinea pigs. The CF of LFPs ranged from 500 to 8000 Hz (Table 1), and they are all larger than the stimuli F0 (80–220 Hz).

To visually evaluate whether the most frequent interspike intervals of single neuron corresponded to the fundamental period, all-order intervals (time domain) were calculated using the intervals between both successive and non-successive spikes. The method was used by Cariani and Delgutte (1996a) to evaluate AN fibers in response to vowels with time-variant F0s. For each spike in a single stimulus presentation, all interspike intervals preceding the reference spike were calculated and listed. The running all-order intervals thus incorporate peristimulus time and interspike interval information.

To visualize whether the post-stimulus time histogram (PSTH) of single neurons could represent the F0 contours, a spectrogram of the PSTH was plotted. Spectrograms were constructed by calculating the short-term Fourier transform for 80 ms sliding hamming windowed PSTHs shifted in 1 ms steps. The signal was zero-padded to 250 ms, and the frequency resolution was 4 Hz.

To examine whether the single IC neuron discharge pattern was synchronized to the stimulus F0, the PSTH and the response spectrum were computed (Miller and Sachs, 1984). The PSTH and response spectrum were generated from spikes occurring at least 50 ms after stimulus onset to exclude the onset response. For each speech stimulus, the PSTH was divided into several successive time segments (256 bins per 50 ms). To examine the response frequency components, the discrete Fourier transform of the PSTH at each time segment was calculated. To increase frequency resolution, each 50 ms time segment was zero-padded to 200 ms before calculating the discrete Fourier transform; as a result, the frequency resolution was 5 Hz. The synchronization index (SI) was used to quantify the degree to which a single neuron’s spiking pattern represented all stimulus frequencies. The SI was computed as the magnitude of the Fourier component normalized by the average firing rate across each 50 ms window (Miller and Sachs, 1983; Sinex and Li, 2007; Sinex, 2008). The SI is an equivalent measurement to vector strength (Syka and Aitkin, 1981; Nakamoto et al., 2010). The value of SI can change between 0, representing no synchronization, and 1, for perfect synchronization. The statistical significance of synchronization was tested using Rayleigh values ($2n \pi^2$, where $n$ is the number of spikes in the 50 ms time segment, and $r$ is the SI) when Rayleigh values were >13.8 ($P < 0.001$) (Mardia, 1975).

| CF range (kHz) | 0.5–1 | 1–2 | 2–4 | 4–8 |
|---------------|-------|-----|-----|-----|
| No. of neurons | 14    | 24  | 28  | 24  |

**Table 1 | The distribution LFPs according to CF.**
To test the relationship between the LFPs and the stimulus waveform, we compared the waveform of the flat tone stimulus, a low-pass filtered version of the stimulus and a typical trace of the LFPs recorded in the IC (Figure 2A). The LFP waveforms were morphologically similar to the low pass filtered stimulus waveform, suggesting that the LFPs were phase-locked to the stimulus (Figure 2A right panel in dashed rectangular).

**Verification of LFPs Source**

To verify that the LFPs originated from the IC rather than far-field sources, the latency of all recorded LFPs was calculated. The latency of each LFP corresponded to the peak of the cross-correlation function between the LFP waveforms and low-pass filtered stimulus waveforms (Figure 2B). On average, the LFP latency [mean = 6.6 ms, standard deviation (SD) = 1.5 ms] was within the previously described for guinea pig IC (5–6 ms) (Harrison and Palmer, 1984; Orton et al., 2012).

**LFPs Represent Time-Variant Periodicity in the Stimulus**

Autocorrelograms were used to visualize the periodicity of signal. Autocorrelograms represent the degree of correlation between the signal and its copy at each response time and corresponding time lag. As shown in Figure 3A, bands with high correlation values can be seen, and reflect the fundamental period of the stimulus (or its multiples). Figure 3B shows the same analysis applied to each of the four LFP responses. There is a high
To quantify the degree to which periodicity in the LFP responses follows the fundamental period of the stimulus, the pitch strength of each LFP was computed (section 2.4.1.4). Pitch strengths for an example LFP were 0.76, 0.69, 0.68, and 0.59 for the four tones respectively (Figure 3). The pitch strength (mean ± SD) of all recorded LFPs for each of four tones was 0.5 ± 0.15, 0.53 ± 0.13, 0.61 ± 0.07, and 0.4 ± 0.09, respectively, and was significantly correlated with the CF ($r = 0.56$, 0.30, 0.32, 0.44, $p < 0.05$ respectively). The tone type had a significant effect on the pitch strength as determined by a one way ANOVA ($F(3,356) = 36.46$, $p < 0.0001$). A post hoc Tukey–Kramer test demonstrated that the pitch strengths of the rising tone and the falling then rising tone (both included a rising F0) were significantly larger than the flat tone and the falling tone ($p < 0.001$).

**Spectrogram of LFPs Represent Time-Variant Frequency Components**

To determine whether the LFPs represent the time-variant frequency components in the stimuli, the spectrograms of single LFP responses and the stimuli were compared (Figure 4). The spectrogram of single LFP responses (Figure 4B) show high spectral energy bands at the stimulus F0 and its harmonics (Figure 4A). For example, there are four time-variant spectral energy bands in the spectrogram of the falling then rising stimulus (Figure 4A, third column), and corresponding energy bands at the same frequencies are also evident in spectrogram of the response (Figure 4B, third column). When the spectrogram of all recorded LFPs was plotted (for all recording sites), some spectrograms only showed the spectral band at F0, but not at the harmonics. However, it is clear that the spectral bands in the LFPs follow the rising/falling dynamics of the changing F0 in the stimuli. The stimulus F0 (red line) and corresponding response F0 contours (black line) extracted from the spectrogram of an example LFPs are shown in Figure 4C. The correlation and linear pitch tracking error between the stimuli F0 contours and the corresponding response F0 contours were used to quantify the degree to which the LFP response tracked the direction and accuracy of the time-variant stimulus F0, respectively. The stimulus-to-response correlation for each tone of this example LFP was 0.86, 0.99, 0.99, and 0.99, respectively. The linear pitch tracking error of the representative LFPs for each tone was 0.96, 1.77, 1.34, and 3.4 Hz, respectively.

To quantify whether all recorded LFPs could match the time-variant F0 contours robustly, the stimulus-to-response correlation was computed individual for each LFP. The stimulus-to-response correlation (mean ± SD) for flat tone, rising tone, falling then rising tone, and falling tone stimulus was 0.66 ± 0.27, 0.93 ± 0.08, 0.95 ± 0.07, 0.98 ± 0.03 respectively, and 93% (84 of 90) of the cross-correlations between the stimulus and response F0 contours were significant ($p < 0.05$).

To assess whether all recorded LFPs could code the stimulus F0 accurately, the linear pitch tracking error was calculated. Across all recordings, the linear pitch tracking error (mean ± SD) of flat tone, rising tone, falling then rising tone, and falling tone was 3.29 ± 4.33 Hz, 5.37 ± 3.25 Hz, 2.59 ± 1.79 Hz, and 6.64 ± 3.09 Hz, respectively. The maximum linear pitch tracking error was calculated for each LFP in response to the four tones.

**FIGURE 4 | Spectrograms of the stimulus (A) and single representative LFP recording (CF = 2.38 kHz) (B). (C) F0 curve of stimulus (red) and response (black). From left to right, each column corresponds to flat tone, rising tone, falling then rising tone and falling tone, respectively. In rows (A,B), the horizontal axis indicates the midpoint of each 80 ms hanning window, vertical axis indicates frequency, and the colors indicate spectral energy (red is highest). In row (C), the horizontal axis represents the midpoint of each time bin, the black line represents the response F0 curve and red line represents the stimulus F0 curve.**
The maximum linear pitch tracking error was lower than 5 Hz in 26 out of the 90 LFPs and lower than 10 Hz in 67 out of the 90 LFPs.

**Single Neuron Activity**

Single neuron activity in response to 10 presentations of each speech stimulus was recorded. The raster plots of five neurons with a sustained response to speech and the corresponding spike waveforms are displayed in Figure 5. The PSTHs of each neuron at one time segment are showed in Figure 5 (bottom panel), and the neurons only phased locked to the some of the stimulus periods. Responses were mostly stimulus-locked and the response duration was similar to the stimulus time. The CFs of single neurons was larger than the stimulus F0 (80–220 Hz) and ranged from 500 to 8000 Hz. The distribution of single neuron CFs at each frequency range was the same as the distribution of LFPs CF (Table 1).

**Running All-Order Intervals (Time Domain)**

To assess whether the temporal information of single neuron spikes could represent the F0 contours of Mandarin tone stimuli, running all-order interval distributions of all neurons were constructed. Running all-order intervals from two representative neurons in response to the four speech stimuli are displayed in Figure 6. Bands of most frequent interspike intervals for one neuron (Figure 6A) that match the time-variant fundamental period of the stimuli can be seen (red line in Figure 6A).

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**FIGURE 5** | The stimulus waveform (top row), five representative neuron raster plots (from top to bottom, CF was 2.59, 5.19, 6.73, 2.59, 5.19 kHz, respectively), and corresponding neuron spike waveforms (right column). The PSTH of each neuron in a short time segment (100–150 ms) is shown in the bottom panel. From (left) to (right), the stimulus waveforms were ‘bā’, ‘bá’, ‘bˇa’, ‘bà’, respectively. For the spike waveforms, the gray lines represent each single spike waveform and the black line represents the mean waveform. The short black bars below the horizontal axis in the stimulus waveform and raster plots represent the time segment used for calculating the PSTH.
Particularly, a band of most frequent interspike intervals for the rising tone decreased from 8 to 5 ms (Figure 6A, second column), and in the falling then rising tone increased from 8 to 11 ms then decreased to 6 ms (Figure 6A, third column). Specifically, two most frequent interspike interval bands in the flat tone (Figure 6A, first column) and the falling then rising tone (Figure 6A, third column) are evident, and the top interspike interval band corresponds to twice fundamental period. By visual inspection of the running all-order intervals for all neurons, only five neurons showed such clear bands of most frequent interspike intervals related to the fundamental period of the stimulus. The all-order intervals for most neurons did not clearly reflect the fundamental period (an example of another neuron with a similar CF is shown in Figure 6B).

PSTH, Spectrogram of PSTH and Response Spectrum (Frequency Domain)
Narrow-band spectrograms of the PSTHs were used to visualize whether the discharge pattern of single neurons could represent the F0 contours and harmonics of the stimuli. One example of a single neuron’s PSTHs (Figure 7A) and its corresponding spectrograms (Figure 7B) are displayed. Bands of high spectral energy can be clearly seen that correspond to the F0 contour (black dashed line) of the flat tone, the rising tone and the falling tone (Figure 7B, first, second, and fourth column).

![FIGURE 6](image1)

**FIGURE 6** | Example of running all-order intervals of two representative IC neurons’ responses to the four speech stimuli. (A) One neuron (CF = 2.83 kHz) in which the most frequent intervals followed the corresponding fundamental period of the stimulus (red line in each figure). (B) One neuron (CF = 3.08 kHz) with intervals that did not match the F0 period. From left to right, each figure corresponds to stimulus ‘bā’, ‘bá’, ‘bə’, ‘bà’, respectively. Each dot represents an interspike interval (vertical axis) at specific end time (horizontal axis) relative to stimulus onset.

![FIGURE 7](image2)

**FIGURE 7** | The PSTHs (A) and spectrograms of the corresponding PSTH (B) of a single neuron (CF = 8 kHz) in response to the four tones. The horizontal axis of the spectrogram indicates the midpoint of each 80 ms hanning window, the vertical axis indicates frequency, and the colors indicate the amplitude of the Fourier transform at each time bin (red is highest). The black dashed line in each figure corresponds to the stimulus F0 curve.
Harmonics can also be seen for the flat tone (Figure 7B, first column). However, the bands of high spectral energy in the falling then rising tone were not continuous throughout the response duration of this neuron (Figure 7B, third column).

To examine whether the discharge pattern of each single IC neuron would significantly correlate with the time-variant stimulus spectral components, the response spectrum of successive 50 ms time segments of the single-unit's PSTH were calculated. The PSTHs and response spectrum for the four stimuli from one representative neuron (CF = 5.66 kHz) are plotted in Figure 8. Frequency components at which the Raleigh test indicated that the neural spiking pattern was significantly synchronous are indicated with triangle symbols. In this single representative neuron, we found that the significantly synchronized frequency in the response spectrum corresponded to the stimulus F0 (dashed red line), 2F0 (dashed blue line), or other harmonics [Figures 8B,C, TS (time segment) 1]. For the flat tone (Figure 8A), we found that the significantly synchronized frequency corresponded to F0 in four of the five time segments, and 2F0 in one of the five time segments (Figure 8A, TS4). However, for the rising tone (Figure 8B), only two of the seven time segments had significantly synchronized frequencies at F0 (Figure 8B TS1 and TS7), with activity in the remaining five time segments corresponding to 2F0. For the falling then rising tone (Figure 8C), the significantly synchronized frequencies correspond to both the F0 and 2F0 (Figure 8C TS1, TS4, TS5, TS6, TS7). For the falling tone (Figure 8D), there was no significantly synchronized frequency in the last time segment (Figure 8D TS5).

The results above were descriptive for a single neuron. To test whether the discharge pattern of all recorded single neurons could significantly synchronize to the stimuli F0s or harmonics, the significantly synchronized frequency components with the largest SI (47% of neurons) were computed and are shown in Figure 9. The stimuli F0s and their harmonics are shown with dashed lines. Twenty-four percent of 90 neurons had a significantly synchronized frequency in more than 5 time segments for all stimuli. The significantly synchronized frequency (open circles) was mostly at F0 (67%) or 2F0 (25%), and only a small number (7%) at other harmonics.

For each neuron, the significantly synchronized highest F0 was defined as the maximum F0 among the first four tones and all the time segments. To examine whether the CF of single neurons was correlated with the significantly synchronized highest F0, the relationship between them are plotted in Figure 10. Some neurons (53% of neurons) were not synchronized at any stimulus F0. In Figure 10 these neurons are plotted with highest F0 equal to 0. More than half of the remaining neurons (62% of 42 neurons) had significant synchronized highest F0 located as the range of unresolved harmonics (Figure 10, see ‘Discussion’), and the highest F0 was significantly positively correlated with the CF (r = 0.5, p < 0.001). However, the maximum SI at each neuron was not related to the CF (r = −0.17, p = 0.27). We suggest that periodicity of single IC neuron spike-timing patterns may be related to interactions of high-order frequency harmonics within the neuron’s pure tone response frequency area, such that the neuron may represent the F0 pitch associated with unresolved harmonics.

**DISCUSSION**

The aim of this study was to determine how the IC encodes time-variant F0 contours in Mandarin tones. The extent to which LFPs and single neuron activity represented the time-variant periodicity pitch of natural speech was examined in the IC of guinea pigs. In the time domain, LFPs reflected the time-variant fundamental period of speech, where there was consistency between interval bands and peak correlation bands in the autocorrelogram of response and stimulus. In the frequency domain, LFPs represented the F0 and harmonics, where the frequency bands with high spectral energy in the spectrogram of LFPs corresponded to the stimulus spectrogram. The F0 curves extracted from 93% of the responses significantly correlated with those extracted from the stimulus (p < 0.05), with 74% of the average linear pitch errors being lower than 10 Hz. These results suggested that LFPs recorded in IC code the F0 contours robustly.

For single neuron activity, phase-locking was also obvious both in the running all-order interspike interval distribution, where the most frequent interspike intervals corresponded to the time-variant fundamental period, and the Fourier transform of PSTHs, where the response significantly synchronized to F0 or 2F0 of the stimulus at corresponding time segments. While nearly all recorded LFPs could reflect F0 robustly, 47% (42/90) of single neurons significantly synchronized with the F0 or 2F0 of stimuli. These results are consistent with other studies suggesting LFPs reflect synchronized input activity (Denker et al., 2011), and that phase-locked temporal information is preserved in the IC (Gockel et al., 2011; Krishnan and Plack, 2011). Single neuron activity reflects the output of the IC processing of pitch information and, in our study we found that 47% of all recorded single neurons represented the F0 voice pitch robustly. To our knowledge, this is the first study to examine running all-order interval distributions of single IC neurons and to examine the correspondence between most frequent interspike intervals distribution and time-variant F0 contours in Mandarin speech.

**Phase-Locked LFPs of IC Represent the F0 Contours and Spectral Features**

In our study, the latency of LFPs and the FRA of single neurons were used to check the recording position in the IC. The shape of LFPs responding to pure tones was consistent with a first positive peak at ~6.5 ms across all the different frequencies (Orton et al., 2012), and the LFP morphology did not follow the stimulus waveform (as would be expected if the recording electrodes were directly picking up loudspeaker artifact). The latency (mean ± SD) of LFP responding to the speech stimuli was 6.6 ± 1.5 ms, and it was consistent with the previous studies (Harrison and Palmer, 1984). The FRA of single neuron responding to pure tones was similar with the study (Palmer et al., 2013).

In our study, we found that phase-locked LFPs recorded in the IC represented the time varying fundamental period in
FIGURE 8 | Example of the PSTHs at each time segment, and PSTH response spectra from a single neuron (CF = 5.66 kHz) response to four tones. (A–D) Correspond to flat tone, rising tone, falling then rising tone, and falling tone, respectively. For each PSTH, the horizontal axis represents time relative to onset of speech, the vertical axis indicates the spike rate per 0.195 ms bin, and the time segment (TS) is indicated in the right corner of each plot. The stimulus F0 (red line) and 2F0 (blue line) are plotted in each response spectrum. The triangle symbol in the response spectrum represents a significantly synchronized frequency.
natural speech. When examining the autocorrelogram in the time domain, we found correspondence between the peak correlation bands in the autocorrelogram of LFPs and the stimuli (Figure 3). This result reinforces the predominant interval hypothesis for pitch encoding in the IC. The LFPs coding the F0 contours both in strength and direction (stimulus-to-response correlation) suggest that neural coding pitch in the IC are dynamic. Similar results have also been observed in scalp-recorded FFRs in Mandarin-speaking participants (Krishnan et al., 2004), suggesting that the temporal patterns of phase-locked activity of neural ensembles in the IC play a role in coding the F0 of complex speech.

In our study, we also found pitch strength of LFP responses depending on the tone type. The tone with a rising F0 produced significantly greater pitch strength than the flat tone and the falling tone. Although similar trends were also observed in other studies (Krishnan et al., 2004), the values of pitch strength found in the current study were relatively low compared to those reported in human studies (Mandarin-speaking participants; Krishnan et al., 2005). This result is also consistent with studies of neural responses to guinea pig vocalizations. Suta et al. (2003) reported that IC neurons’ firing rates were higher for whistle than time-inversed whistle, and suggested that the IC neurons of the guinea pig may prefer rising frequencies compared to falling frequencies.

The pitch strength was also significantly correlated with CF – the pitch strength increased as the CF increased. This result may be related to the relatively broader spectral tunning curve at the speech level used in this study. The bandwidth of tuning curves increase with the CF, and the harmonics and periodicity information coded by the LFP with high CF were therefore more robust.

IC Single Neuron Activity Represent the F0 Voice Pitch Contours

In the current study, the most frequent intervals found in the running all-order intervals distribution of single IC neurons corresponded to the time-varying fundamental period of the stimuli. This finding was consistent with the studies of neural activity in the AN. Cariani and Delgutte (1996a,b) examined the AN fibers response to the time-variant F0 of synthesized vowel sounds. They were the first to pool interspike interval distributions of the population AN fibers and found the correspondence between most frequent interspike intervals and the human perceived pitch. They suggested that phase-locked neural activity is important for encoding the pitch of complex sounds. They also found that most single AN fibers could phase-lock to the pitch.

However, in the current study, we only found five single neurons in the IC with CFs higher than the F0 that clearly
reflected the fundamental period information. This may be because the IC receives phase-locked information not only from CN (Frisina et al., 1990), but also from medial and lateral divisions of the superior olive (Oliver et al., 2003) and lateral lemniscus (Schreiner and Winer, 2005), and subsequently, the accuracy of phase-locking preserved in the AN and CN is reduced at the level of the IC.

One possible explanation for finding few IC neurons in this study is that temporal information may be transformed into a rate-place code in the IC (Cariani, 1999). This idea has been examined in studies of IC neurons’ responses to the periodicity of amplitude modulation signals (Langner and Schreiner, 1988). Langner and Schreiner (1988) tested the best modulation function of single IC neurons responding to different modulation frequencies with a carrier frequency at the CF of the neuron. They found that only \( \sim 33\% \) of the 225 single IC neurons could tune to a best modulation frequency when it is tested by synchronization, while almost 75% when it is tested by firing rate (Langner and Schreiner, 1988).

Another possible reason for finding few IC neurons that the most frequency interspike intervals in the running all-order intervals showing stimulus periodicity is we only used 10 presentations of each stimulus in our study, while Cariani and Delgutte (1996a) presented stimuli 100 times, thereby increasing their ability to visualize the pattern of most frequent intervals. Miller and colleagues collected at least 600 spikes (Miller and Sachs, 1983, 1984) while we only collected 200 spikes on average.

In the current study, the spectra of the PSTHs (Figure 8) of some neurons (47%) at specific time segments showed discharge patterns that were significantly synchronized to the F0, 2F0 or other harmonics. The best-synchronized frequency was generally below 500 Hz (Figure 9), and all synchronized frequencies were below the best modulation frequency limit of IC neurons responding to amplitude-modulated signals (Langner and Schreiner, 1988; Langner et al., 2002). Previous research has also suggested that a periodicity map might exist in IC (Schreiner and Langner, 1988; Langner et al., 2002; Schnupp et al., 2015), and our synchronized neurons may have been located across that map. Krishna and Semple (2000) found that the spike firing rate tuned to the modulation frequency in IC and the rate modulation function was not dependent on the synchronization modulation function (Langner and Schreiner, 1988; Müller-Preuss et al., 1994). These results suggest that both rate and synchronization representation of periodicity might exist at the level of the IC. Henry et al. (2017) also showed that IC neurons in awake budgerigar encoded the vowel-like sounds by the neural synchrony and variations in average discharge rate related the envelope structure. Studies of the correlation between the behaviourally measured sensitivity to amplitude-modulated signals and neural responses in the IC showed that the synchronized temporal representation was more correlated with behavioral performance than rate representation for envelope detection in awake budgerigar (Henry et al., 2016). Similar studies in awake rabbits, however, showed the opposite effect, with rate representation was more correlated to behavioral performance (Nelson and Carney, 2007).

The limit of upper modulation frequency synchronization, known to decrease progressively from auditory periphery to the central auditory system (Palmer, 1982; Schreiner and Urban, 1988; Rees and Palmer, 1989; Joris and Yin, 1992, 1998; Rhode and Greenberg, 1994; Bieser and Müller-Preuss, 1996; Eggermont, 1998; Joris et al., 2004). Studies measuring from neurons in the auditory thalamus and auditory cortex have shown that the periodicity or repetition rate of temporally regular signals can be represented by the spiking timing (synchronized neurons) and/or average discharge-rate (non-synchronized neurons) (Lu et al., 2001; Lu and Wang, 2004; Bartlett and Wang, 2007). Bendor and Wang (2005) also found mixed-response neurons in the auditory cortex, and suggested that these neurons may play a role in temporal-to-rate transmission. The difference in the limits of neural synchronization between the IC (Batra et al., 1989), auditory thalamus (Bartlett and Wang, 2007) and auditory cortex (Lu et al., 2001) in awake animals suggests that the auditory thalamus may play the role of a transition stage for temporal-to-rate transformation (Wang et al., 2008), and that rate-place representation is more responsible than temporal representation for periodicity representation in auditory cortex (Bendor and Wang, 2005; Walker et al., 2011). It also has been suggested that the periodicity temporal code may transform to a rate-place code (Bartlett and Wang, 2007; Wang et al., 2008; Bendor and Wang, 2010; Plack et al., 2014; Langner, 2015) or spatial-temporal code (Loeb et al., 1983; Oxenham et al., 2004; Bernstein and Oxenham, 2005) in the upper auditory pathway. In the current study, 93% LFPs represented the time varying periodicity pitch, and only 47% single neurons represented the F0 pitch. The LFP reveal the synaptic input information, and single neurons reveal the axonal output information. The difference between the proportion of the LFPs and single neurons may suggest that the temporal coding transform to other form of neural coding.

The relationship between the CF of single neurons in guinea pig with the equivalent rectangular bandwidth (ERB) (ERB = 0.3 CF \( 0.56 \)). Shackleton et al. (2009) could be used to estimate whether any given stimulus contained resolved or unresolved components (Figure 10). In general, a sound can be said to contain only resolved harmonics when the ERB includes fewer than 2 harmonics, and unresolved when the ERB includes more than 3.25 harmonics (Shackleton and Carlyon, 1994). In the current study, the CF of most neurons corresponded to the unresolved harmonics. The results suggest that most neurons represent F0 voice pitch (Figure 10), and that this is due to the interaction of harmonics in the FRA of pure tones (Langner and Schreiner, 1988). In 47% of all recording neurons synchronized to the F0, 26 of 42 neurons (62%) had CFs located in the unresolved range. The unresolved range was decided by the stimulus F0. For a stimulus F0 of 100 Hz, 34% (26 of 76) of the recorded neurons were synchronized; and for a stimulus F0 of 150 Hz, 50% (23 of 46) were synchronized. The largest SI of each neuron was used to assess the ability of single neuron coding periodicity, and we found that there is no relationship between the CF of single neuron and the largest SI \( (r = -0.17, \ p = 0.27) \). The results suggest that there is no relationship between the proportion of
synchronized neurons in the resolved and unresolved ranges in current study, and the results was consistent with the study (Shackleton et al., 2009). One possible reason is that the sound level was 80 dB SPL, and the FRA at this level may have included more stimulus harmonics even though the CF of the neuron were located in the range between the resolved and unresolved. Although the amplitude of the stimuli was changing with the time, the largest SI of all synchronized neurons at each time segment was not correlated with the root mean square of the stimulus amplitude \( r = -0.28, p = 0.17 \).

Three of the neurons we recorded did have CFs corresponding to resolved harmonics. The maximum SI of these three neurons in the resolved harmonics was not significant different in the unresolved harmonics. Other researchers have found similar results in studies of IC neurons in response to harmonic complex tones (Shackleton et al., 2009). However, there are some discrepancies between our results and others. Sinex and Li (2007) examined IC neurons’ responses to single harmonic tones with an F0 of 250 Hz and found that the discharge pattern of single neurons was modulated at F0, but only in some cases. Shackleton et al. (2009) tested multi-unit activity from the IC with different F0s (50, 100, 200, 400 Hz) of harmonic tones and found that most clusters were synchronized to F0 above 280 Hz. Compared with their studies, differences include the stimuli and the response analysis time windows used. In the study by Shackleton et al. (2009), synthesized harmonic tones with fixed F0 were used. In our study, complex speech sounds with time-variant F0 from 80 to 220 Hz were used (Figure 1C). Another difference is the method used in the two studies: Shackleton et al. (2009) recorded multiunit responses to stimuli of 100 ms duration, and the entire response waveform was analyzed. In Sinex and Li (2007), the stimulus was 500 ms and the response after 100 ms was analyzed. They found that the discharge pattern was generally not related to the F0, except in some cases in which the discharge pattern was modulated by an F0 of 250 Hz. However, in our study the stimulus duration ranged from 400 to 600 ms, and the response was analyzed in 50 ms successive time segments. As a result, the difference of synchronized frequency components may be due to the time windows of analysis.

Method Limitations

Although the results of LFPs are similar to FFRs in human studies, an important limitation is that the Mandarin tone is not a natural or behaviourally relevant sound to the guinea pig. As previously discussed, spike rate is affected by a sound’s relevance to behavior (Suta et al., 2003). Urethane is a common anesthetic for acute recordings of neural responses in the auditory midbrain; however, studies have demonstrated that it inhibits spontaneous and evoked spike rate (Albrecht and Davidowa, 1989; Capsius and Leppelsack, 1996). Other studies suggested that there is no significant effect on spectral tuning properties in the anesthetic state (Schumacher et al., 2011; Palmer et al., 2013). The phase-locking to the F0 was recorded in the IC of urethane anesthetic guinea pig in our study, so as the study by Shackleton et al. (2009). The CF of most neurons was higher than 1 kHz and corresponded to the unresolved harmonics of stimulus, so our results only showed that single neurons represent the F0 pitch of stimulus.

AUTHOR CONTRIBUTIONS

FP, WH, and YZ designed the experiments. FP, XW, and NH performed the experiments and collected the data. FP, HI-B, CM, and JF analyzed the data and interpreted results of experiments. FP, HI-B, and JF drafted the manuscript. All authors edited and revised manuscript critically and approved final version of manuscript.

FUNDING

This work was supported by the Nature Science Foundation of China (NSFC31771069 and 31700856) and the Natural Science Foundation of Chongqing in China (cstc2015jcyjB0538). HI-B was supported by an Australian National Health and Medical Research Council Early-Career Fellowship.

ACKNOWLEDGMENTS

We would like to thank Nan Xia and Manqing Wang for providing assistance of collecting data. We thank Dr. Ying Xiong from the Neurobiology Department at the Third Military Medical University for their guidance of animal surgery. We also thank Dr. Janine Clarey from the Bionics Institute for editing the manuscript. The Bionics Institute acknowledges the support it receives from the Victorian Government through its Operational Infrastructure Support Program.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Title:
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Date:
2018-07-24

Citation:
Peng, F., Innes-Brown, H., McKay, C. M., Fallon, J. B., Zhou, Y., Wang, X., Hu, N. & Hou, W. (2018). Temporal Coding of Voice Pitch Contours in Mandarin Tones. FRONTIERS IN NEURAL CIRCUITS, 12, https://doi.org/10.3389/fncir.2018.00055.

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