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Spatiotemporal properties of intracellular calcium signaling in osteocytic and osteoblastic cell networks under fluid flow

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Mechanical stimuli can trigger intracellular calcium (Ca2+) responses in osteocytes and osteoblasts. Successful construction of bone cell networks necessitates more elaborate and systematic analysis for the spatiotemporal properties of Ca2+ signaling in the networks. In the present study, an unsupervised algorithm based on independent component analysis (ICA) was employed to extract the Ca2+ signals of bone cells in the network. We demonstrated that the ICA-based technology could yield higher signal fidelity than the manual region of interest (ROI) method. Second, the spatiotemporal properties of Ca2+ signaling in osteocyte-like MLO-Y4 and osteoblast-like MC3T3-E1 cell networks under laminar and steady fluid flow stimulation were systematically analyzed and compared. MLO-Y4 cells exhibited much more active Ca2+ transients than MC3T3-E1 cells, evidenced by more Ca2+ peaks, less time to the 1st peak and less time between the 1st and 2nd peaks. With respect to temporal properties, MLO-Y4 cells demonstrated higher spike rate and Ca2+ oscillating frequency. The spatial intercellular synchronous activities of Ca2+ signaling in MLO-Y4 cell networks were higher than those in MC3T3-E1 cell networks and also negatively correlated with the intercellular distance, revealing faster Ca2+ wave propagation in MLO-Y4 cell networks. Our findings show that the unsupervised ICA-based technique results in more sensitive and quantitative signal extraction than traditional ROI analysis, with the potential to be widely employed in Ca2+ signaling extraction in the cell networks. The present study also revealed a dramatic spatiotemporal difference in Ca2+ signaling for osteocytic and osteoblastic cell networks in processing the mechanical stimulus. The higher intracellular Ca2+ oscillatory behaviors and intercellular coordination of MLO-Y4 cells provided further evidences that osteocytes may behave as the major mechanical sensor in bone modeling and remodeling processes.

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Introduction

Bone, acting as an adaptive system, can remodel its structure in response to the external mechanical environment [1]. Understanding the mechanism of bone remodeling is of critical importance in osteoporosis, age-related fractures and bone loss in prolonged space flight. Osteocytes, accounting for almost 95% of the bone cell population, are encapsulated in a fluid-filled mineralized matrix and interconnected with each other through long dendritic processes to form extensive networks in the lacunae–canaliculi system. It has been long conjectured that this elaborate osteocytic network behaves as the central mechanosensory system in bone adaptation and bone homeostasis [2]. Osteocytic networks can transduce physical stimuli into biochemical signals to further regulate the behaviors of osteoblasts and osteoclasts, thus coordinating bone modeling and remodeling processes [3]. Several studies have shown that osteoblasts themselves could respond to mechanical stimuli with a cascade of cellular activities, such as calcium (Ca2+) signaling and release of nitric oxide (NO) and prostaglandin E2 (PGE2) [4–6]. Mechanical loading triggers Ca2+ release in both osteocytes and osteoblasts [7–11], which is a pivotal and ubiquitous second messenger regulating many downstream cellular activities, including cell proliferation, differentiation, and apoptosis [12]. Hence, a comprehensive understanding of Ca2+ signaling in osteocytes and osteoblasts under mechanical stimuli is of tremendous value in deciphering the mechanisms by which bone processes the biophysical stimulus.

Ca2+ signaling in osteocytes and osteoblasts can propagate between neighboring cells and generate Ca2+ waves in the bone cell populations [10,13]. However, most studies in bone cell mechanotransduction have been performed on confluent or sub-confluent uncontrolled cell monolayers. In our previous studies, a two-dimensional patterned bone cell network was successfully constructed to mimic the elaborate in vivo bone cell network topology using microcontact printing and self-assembled monolayer (SAM) techniques [14]. Our recent findings demonstrated that the osteocytic network showed repetitive spike-like
Ca\textsuperscript{2+} peaks under fluid flow induced shear stress. These oscillations were dramatically different from those found in the osteoblastic network regardless of the magnitude of shear stress [15]. However, there are still two major obstacles in studying Ca\textsuperscript{2+} signaling in these cellular networks. First, bone cells patterned in the topologic network are spatially connected with their neighboring cells, so the time course of Ca\textsuperscript{2+} dynamics neglected the important spatial and temporal information embedded in the network responses. This information is critical to help provide essential insights into Ca\textsuperscript{2+} dynamics of individual cells and Ca\textsuperscript{2+} wave propagation in the cell network, which has attracted extensive attention in the signal analysis of neuronal cell types, such as astrocytes, glial cells and Purkinje cells [16–18]. Therefore, it necessitates more elaborate and systematic analysis of the spatiotemporal characteristics of Ca\textsuperscript{2+} signaling in bone cell networks. Second, most previous extraction methods for Ca\textsuperscript{2+} signaling in bone cells have been mainly based on a manual region of interest (ROI) analysis, which can be laborious and subjective, requiring users to select the target boundary manually according to the cell morphology. The manual ROI was subject to the constraints of image qualities, and the large number of cells in our bone cell network data also further increased the difficulty for manual extraction. Therefore, an unsupervised signal extraction technique is needed to reduce the workload and minimize the artificial errors.

Independent component analysis (ICA) is an unsupervised blind source separation procedure that transforms signal mixtures into a corresponding set of statistically independent source signals [19]. ICA has also been successfully applied for identifying and characterizing physiological signals in many research areas, such as electroencephalography (EEG), electrocardiography (ECG), magnetocardiography (MCG), and functional magnetic resonance imaging (fMRI) [20–23]. Isolating the individual Ca\textsuperscript{2+} signals in a bone cell network, sharing a number of similarities with separating the electrophysiological signals from the recorded mixtures, may also benefit from the unsupervised ICA technique. However, no study to date has employed this technique to extract the intracellular Ca\textsuperscript{2+} signaling of bone cells and systematically investigated the spatiotemporal properties of Ca\textsuperscript{2+} signaling in a cell network pattern.

In the present study, osteocyte-like MLO-Y4 and osteoblast-like MC3T3-E1 cell networks were respectively stimulated under physiological related fluid shear stress (0.5–4 Pa) and Ca\textsuperscript{2+} responses were extracted and analyzed using a set of novel unsupervised techniques. An ICA-based algorithm was used to separate the individual Ca\textsuperscript{2+} signals from the cell networks. Spike rate and power spectrum density (PSD) analysis were then employed to evaluate the temporal dynamics of Ca\textsuperscript{2+} signaling, and cell–cell spike synchronization and signal correlation were analyzed to reveal the spatial intercellular communications of Ca\textsuperscript{2+} signaling in the networks. This study represents the first effort to systematically study and compare the spatiotemporal characteristics of Ca\textsuperscript{2+} signaling in osteocytic and osteoblastic networks.

Materials and methods

Bone cell network

Osteocyte-like MLO-Y4 cells (a gift obtained from Lynda Bonewald, University of Missouri) were cultured on type I rat tail collagen (BD Biosciences, San Jose, CA, USA) coated Petri-dish in α-MEM containing 5% FBS, 5% CS and 1% P/S [24]. MC3T3-E1 cells were maintained in α-MEM supplemented with 10% FBS and 1% P/S. Microcontact printing and SAM surface chemistry technologies were employed to construct the in vitro bone cell networks, which have been described in detail in our previous studies [14,15]. The diameter of the round island for a cell to reside was 20 µm for MC3T3-E1 cells and 15 µm for MLO-Y4 cells, while the edge-to-edge distances between neighboring islands were 50 µm and 35 µm respectively. These geometric parameters were optimized based on cell morphology to obtain well-formed networks. A grid cell pattern was printed on a chromium mask, and then the pattern was replicated to a master made of positive photosist (Shipley 1818, MicroChem Corp, Newton, MA). Polydimethylsiloxane (PDMS, Sylgard 184, Dow Corning, Midland, MI) stamps were made using the master. The stamp was coated with an adhesive SAM (octadecanethiol; Sigma-Aldrich Co., St. Louis, MO), and pressed onto a gold-coated glass slide for 1 min. Ethylene glycol terminated SAM solution (HS-C11-EG3; Prochimia, Sopot, Poland) was then added onto the slide for 3 h, and the slide was then incubated in a fibronectin solution for 1 h to facilitate cell adhesion. Cells were then seeded onto the slide and cultured in α-MEM medium supplemented with 2% charcoal-stripped fetal bovine serum (CSFBS; Hyclone Laboratories Inc., Logan, UT) for 24 h before fluid flow stimulation. CSFBS was used here to control the growth of cells and to avoid double-cell-occupancy on a single circular spot on the pattern [10,15]. The fluorescence images of MC3T3-E1 and MLO-Y4 cell networks are shown in Figs. 1A & B.

Fluid flow stimulation and Ca\textsuperscript{2+} imaging

The patterned cells were loaded with 10 µM Fura-2AM (Molecular Probes, Eugene, OR) for 45 min. The glass slide was then mounted into a parallel plate flow chamber (Fig. 1C). The flow chamber was fixed on an inverted fluorescence microscope (Olympus IX71, Melville, NY) with a 10× objective. Steady, laminar fluid flow at different shear stress (0.5, 1, 2, and 4 Pa) on the cell surface was applied through the chamber by varying the power output of a magnetic gear pump connected to the flow chamber. α-MEM supplemented with 2% CSFBS was used as the flow medium for the experiment [15,31]. The Ca\textsuperscript{2+} responses of cells were recorded for a 10-min period: 1 min for baseline and 9 min after the onset of fluid flow. MetaFluor Imaging Software 7.0 (Molecular Devices, Downingtown, PA) was used to acquire and record the Fura-2 340 nm/380 nm ratio images every 3 s during the course of fluid flow stimulation.

ICA-based signal extraction algorithm

Flow chart for the unsupervised signal extraction algorithm and spatiotemporal analysis strategy for Ca\textsuperscript{2+} signaling in bone cell networks is shown in Fig. 2. All procedures were performed using Matlab (2010, The MathWorks Inc., Natick, MA). First, the pseudocolor fluorescence time-lapse images were transformed into grayscale images and resized from 1334 × 1024 pixels to 336 × 256 pixels for a total of 200 frames with the bilinear interpolation algorithm to reduce the computation load. Second, the signal in each pixel was normalized by the mean value over all time frames at that pixel. Then, principal
component analysis (PCA) was used to serve as a preprocessing procedure prior to ICA by whitening the original data matrix. Combining the PCA and ICA algorithms can simplify the separation of the underlying independent source signals and improve the robustness of ICA [25–27]. The major aim of PCA in the present study is to transform the original image sequence data \( X \) into a whitened matrix \( X_{\text{white}} \), in which components are uncorrelated with each other and the covariance matrix equals the identity matrix. As shown in Eq. (1), the original image sequence data \( X \) was decomposed into a series of spatial patterns \( U \) and their corresponding time series \( V \) using singular value decomposition.

\[
X \approx UDV^T.
\]  

(1)

\( R \) is the rank of \( X \), \( R \leq \min (M_{xy}, N_t) \), while \( D \) is the \( R \times R \) non-negative diagonal matrix of singular values in decreasing order. If \( R = N_t \), the approximation will be exact. \( V \) and \( D \) can be obtained by computing \( X^TX \). \( V \) are the eigenvectors of \( X^TX \), and \( D^2 \) are the eigenvalues of \( X^TX \). \( U \) can be computed as:

\[
U = XVD^{-1}.
\]

(2)

The smaller the eigenvalue, the less energy along the corresponding eigenvector there will be. Therefore, a small eigenvalue often corresponds to noise signals. In this study, the largest 40–60 singular values in matrix \( D \) were selected to ensure that the components encoding the random noise signals were excluded. After filtering the principal components, the whitened matrix \( X_{\text{white}} \) with reduced dimension was obtained:

\[
X_{\text{white}} = UV^T.
\]

(3)

ICA was applied to the whitened data \( X_{\text{white}} \) generated by PCA. Spatial ICA (sICA) was used to extract the signals to identify spatial components that maximized the independence of the spatial patterns, together with their corresponding time courses. Some well-developed ICA algorithms in the literature could be directly used [27,28]. We employed the FastICA algorithm, which is a fixed-point iteration algorithm by measuring the non-Gaussianity to seek the statistically independent sources from their mixtures [29]. The FastICA algorithm has been widely used in previous studies to separate physiological signals [18,20].

After performing sICA, the spatial patterns between each pair of components are as statistically independent as possible. In practice, however, one single IC sometimes contains several spatially separated cell regions, which have high correlation with each other. Therefore, a morphology-based segmentation algorithm was used to split each IC into spatially separated sub-ICs to ensure that each sub-IC corresponds to a target cell region. The spatial patterns were first transformed to a binary matrix. Then, the binary image segmentation procedure was implemented using the function ‘bwlabel’ in Matlab to distinguish the spatially disconnected regions [30]. After this procedure, any IC that

Fig. 2. Flow chart for the unsupervised signal extraction algorithm and spatiotemporal analysis strategy for Ca^{2+} signaling in bone cell networks.
contains multiple spatially separated cell regions can be split into the corresponding sub-ICs. Then, a threshold value of the minimal cell area (20 pixels in practice) was set to eliminate the ICs encoding the noise signals or background fluctuation. To identify the cell region occupied by multiple sub-ICs, the distance between the centroids of each two sub-IC regions was computed, and a threshold of minimal distance value (10 pixels in practice) was set to select the overlapped sub-ICs. The sub-IC with maximal area value among the overlapped sub-ICs was selected as the target spatial component. All these procedures ensured a one-to-one correspondence between each sub-IC and a cell region.

After signal extraction, a peak selection algorithm was developed to detect the peaks in each cellular signal. The signal was first normalized using the standardized z-score, and then the first derivative of curve was smoothed using the triangular smooth method. The downward zero-crossings in the smoothed first derivative were then detected. The points were selected as the peak points by setting the threshold values for the amplitude of the selected zero-crossings as:

\[
S_{\text{test}} - S_{\text{min}} > P_{\text{threshold}}
\]

where \(S_{\text{test}}\) is the value of the test point, \(S_{\text{min}}\) is the minimal value of the signal, \(S_{\text{max}}\) is the maximal value of the signal, and \(P_{\text{threshold}}\) is the threshold value. In this study, \(P_{\text{threshold}}\) was set to 1/4 [15].

Periodicity analysis

After signal extraction, information relating to spikes was automatically recorded, including the total responsive cell number, Ca\(^{2+}\) spike number in each cell, and time of spike occurrence in each cell. A color-code image with different color intensities corresponding to cells with different spike numbers was also generated automatically to facilitate describing the spatial distribution of Ca\(^{2+}\) signaling (Figs. 6A & B). The average Ca\(^{2+}\) spike number, occurrence time of the 1st spike and time between the 1st and 2nd spikes were compared between MC3T3-E1 and MLO-Y4 cells in various magnitudes of shear stress (0.5, 1, 2 and 4 Pa) [15,31]. Since the present ICA-based Ca\(^{2+}\) signaling extraction algorithm is only sensitive to the cells which exhibited Ca\(^{2+}\) response, the total cell number in each slide was manually counted to compare the responsive rate between MC3T3-E1 and MLO-Y4 cells. The spike rate was then systematically analyzed to elucidate the temporal distributions of spike activities of Ca\(^{2+}\) signals. In brief, a time bin (10 frames in practice) was set so that the time range was segmented into several sub-ranges with equal length (20 sub-ranges in practice). All the spikes of cells were mapped into the corresponding sub-range, and the spike number was determined in each sub-range. The spike rate in each sub-range was calculated as:

\[
R_{\text{spike}} = \frac{N_{\text{spike}}}{\Delta(t) N_{\text{cell}}}
\]

where \(R_{\text{spike}}\) is the spike rate, and \(N_{\text{spike}}\) is the total spike number in each sub-range, \(\Delta(t)\) is the bin size, and \(N_{\text{cell}}\) is the total cell number in each sub-range.

The centroid of each cell was also automatically extracted in the signal extraction program for the spike synchrony and signal correlation analysis. All cells in the field of view from each movie were compared in a pair-wise manner. Since the center to center distance between neighboring round islands for cells to reside was 70 \(\mu\text{m}\) for osteoblasts and 50 \(\mu\text{m}\) for osteocytes, the cell pair was excluded for the spatial correlation analysis with intercellular distance lower than 60 \(\mu\text{m}\) for osteoblasts and 40 \(\mu\text{m}\) for osteocytes, given that the possible non-coincidence between the cell centroid and the island center. Because most cells’ 1st spikes occurred almost synchronously when fluid flow was applied, the spikes after the 1st main peak were analyzed to investigate the synchronization of multiple responses of Ca\(^{2+}\) signaling in bone cell networks. From our previous study, the 1st spike was resulted from Ca\(^{2+}\) entry into the cell while the subsequent spikes resulted from ATP-related signal propagation. Spikes between two cells were defined as synchronous ones if the time offset between the spike pair was less than 5 frames (15 s). In this study, the sliding-window cross correlation analysis (window width: 60 s; step size: 60 s) was employed to evaluate the signal correlation. This method can provide enhanced localization analysis in the temporal domain by splitting the signals with the sliding window, and thus quantitate the Ca\(^{2+}\) signal correlation more precisely compared with the normal correlation analysis [32–34]. We selected the maximum value over the lag range from −5 to 5 frames (−15 to 15 s) in each sub-range. The correlation value between two signals was calculated by averaging the values over all the sub-ranges.

To compare the periodicities and frequency distribution characteristics of Ca\(^{2+}\) signaling between MC3T3-E1 and MLO-Y4 cells, the normalized Ca\(^{2+}\) signals were analyzed using fast Fourier transform (FFT)-based power spectrum density (PSD) processing routines in Matlab. All the frequency curves of cellular signals in each group were averaged to identify the difference in frequency distribution between MC3T3-E1 and MLO-Y4 cells.

Statistical analysis

The number of cell networks (slides) and analyzed cells (responsive cells) in different experimental groups are shown in Table 1. All data were presented as the mean ± standard deviation (SD). Statistical analyses were performed using a SPSS version 10.0 for Microsoft Windows (SPSS, Chicago, IL, USA). At each fluid stimulation strength, differences of each parameter (the number of Ca\(^{2+}\) peaks, time to the 1st peak, time between the 1st and 2nd peaks, synchronous cell pair number, and sliding-window correlation coefficient) between MC3T3-E1 and MLO-Y4 cells were examined using a Student t-test. To determine the correlation between a parameter and the stimulation strength, linear regression analysis was performed. Linear regression analysis was also used to evaluate the correlation of the spike synchrony and signal correlation with the cell–cell spatial separation distance. \(P<0.05\) was considered statistically significant.

Results

The spatial patterns and temporal signals of the original simulated movies are shown in Fig. 3A (Left). Gaussian noise was artificially introduced to the movies (SNR ranged from 10 to 30 dB). The

<table>
<thead>
<tr>
<th>Shear stress (Pa)</th>
<th>0.5</th>
<th>1</th>
<th>2</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>MLO-Y4 Slides</td>
<td>3</td>
<td>5</td>
<td>4</td>
<td>5</td>
</tr>
<tr>
<td>Responsive cells</td>
<td>397</td>
<td>625</td>
<td>446</td>
<td>509</td>
</tr>
<tr>
<td>MC3T3-E1 Slides</td>
<td>7</td>
<td>6</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>Responsive cells</td>
<td>219</td>
<td>324</td>
<td>459</td>
<td>209</td>
</tr>
</tbody>
</table>
ICA-based and ROI-based signal extraction methods (B).

Fig. 3. Validation of the ICA-based Ca\textsuperscript{2+} signal extraction technique. Simulated time-lapse images were created to mimic typical cell shapes and curves of Ca\textsuperscript{2+} signals in bone cell networks (A, Left). Signals were artificially contaminated by generating Gaussian noise with different variances to create movies with signal-to-noise ratio (SNR) ranging from 10 to 30 dB, which covered the normal SNR range of our experimental fluorescence time-lapse images. (A, Middle) and (A, Right) showed the signal extraction in the noise-contaminated movie with 20 dB SNR using unsupervised ICA-based technique and manual ROI method, respectively. The signal fidelity, defined as the Pearson's correlation coefficient between true signal and extracted signal, was compared between ICA-based and ROI-based signal extraction methods (B).

unsupervised ICA-based technique was used to extract the signals in the noise-contaminated movies. To compare the extraction capacity between the ICA-based and ROI-based methods, signals in the movies were manually extracted by three analysts using the traditional ROI method. Fig. 3A (Middle & Right) shows the results of spatial separation and signal extraction by ICA-based and ROI-based methods when the SNR in the movie was 20 dB. These results demonstrated that the unsupervised ICA-based method achieved better performance than the manual ROI method in both spatial target and temporal signal extraction at 20 dB SNR. Moreover, the signal fidelity extracted by the ICA-based method was obviously superior to that identified by the manual ROI method as shown in Fig. 3B for SNR ranging from 10 to 20 dB. As SNR increased from 20 dB to 30 dB, the signal fidelity by the ROI method gradually approached that by the ICA-based method.

Signals extracted by the ICA-based technique from experimental data of typical MC3T3-E1 and MLO-Y4 cell networks under 2 Pa shear stress are shown in Fig. 4. Most MC3T3-E1 cells and MLO-Y4 cells released 1st spikes at the onset of flow. After the 1st main spike, most MC3T3-E1 cells exhibited a few lower spikes during the 9-min fluid flow stimulation period (Fig. 4B). MLO-Y4 cells showed significantly more frequent Ca\textsuperscript{2+} oscillatory activities than MC3T3-E1 cells, and some of them exhibited repetitive and robust multiple peak responses in addition to the 1st peak (Fig. 4E). Each individual dot in Figs. 4C & F represents an identified spike. These two plots clearly showed that MLO-Y4 cells exhibited many more Ca\textsuperscript{2+} peaks under fluid flow than MC3T3-E1 cells.

With an increase in fluid shear strength, the average number of Ca\textsuperscript{2+} peaks increased in both cell types. MLO-Y4 cells exhibited significantly more Ca\textsuperscript{2+} peaks than MC3T3-E1 cells at all levels of shear stress (P<0.01, Fig. 5B). The percentage of responsive MC3T3-E1 cells increased with the shear stress levels, and the responsive percentage in MLO-Y4 cells was higher than that in MC3T3-E1 cells at 0.5, 1 and 2 Pa shear stress (Fig. 5A). MLO-Y4 cells also took significantly less time than MC3T3-E1 cells to release the 1st spike in response to the fluid flow under 0.5, 1 and 2 Pa shear stress, and recovered faster from previous response to release the next Ca\textsuperscript{2+} peak under 0.5 and 2 Pa shear stress (Figs. 5C & D). Linear regression indicated that Ca\textsuperscript{2+} peak number was positively correlated, and the time to the 1st peak and time between the 1st and 2nd peaks were negatively correlated for both cell types with fluid flow levels. These results are consistent with our previous findings using the manual extraction method [15].

As shown in Fig. 5E, the spike rate of both MC3T3-E1 and MLO-Y4 cells reached a peak at the onset of flow. MLO-Y4 cells also showed relatively higher spike rate in the whole flow experimental period under all loading levels, which further suggested that MLO-Y4 cells had more frequent spike activities and more obvious features of multiple spikes than MC3T3-E1 cells (Fig. 5E).

As shown in Figs. 6C & E, MLO-Y4 cell networks had significantly higher spike synchrony and signal correlation than MC3T3-E1 cell networks in all sub-groups with different cell–cell separation distance at all fluid flow levels. Linear regression showed that the spike synchrony and signal correlation decreased with the increase of cell–cell separation distance at 1, 2 and 4 Pa shear stress for MLO-Y4 cell networks, indicating that cells with shorter separation distance exhibited higher cross-talk than those with farther separation distance. However, the cross-talk between cells in MC3T3-E1 cell networks only exhibited minor negative correlation with the cell–cell separation distance in the 4 Pa shear stress group. Figs. 6D & F revealed significantly higher spike synchrony in MLO-Y4 cell networks under 1, 2 and 4 Pa, and also higher signal correlation in MLO-Y4 cell networks under all fluid flow levels (P<0.01). Linear regression further showed an increasing trend for both spike synchrony and signal correlation in the two types of cells with the increase of the fluid flow levels.

In the frequency spectral analysis (Fig. 7), both MC3T3-E1 and MLO-Y4 cells had a peak in the lower frequency band (0.001–0.002 Hz) at all loading levels, and the energy distribution of MC3T3-E1 cells was much more prominent than MLO-Y4 cells in this frequency range. However, MLO-Y4 cells exhibited striking 2nd peak in the higher frequency band and the frequency corresponding to the peak was increased with the fluid flow rate (0.0047 Hz at 0.5 Pa, 0.0068 Hz at 1 Pa, 0.0106 Hz at 2 Pa, and 0.0124 Hz at 4 Pa). This indicated that the Ca\textsuperscript{2+} oscillations in MLO-Y4 cells were more dramatic than MC3T3-E1 cells, and also positively correlated with the stimulation strength.

Discussion

In the present study on Ca\textsuperscript{2+} signaling in bone cell networks, a signal processing technique allowing for a cell’s spatial target separation, temporal signal extraction and spike detection was implemented based on PCA followed by sICA algorithm. The PCA followed by ICA algorithm has been extensively applied in ECG, MCG and fMRI studies for separating target signals from the background, which can increase the robustness and simplify the implementation complexity than the single ICA method [21,22,26,27]. Previous optical imaging studies in neuroscience fields have used a combined PCA–ICA method to identify the distinct...
functional region and to explore the spatiotemporal relationship between vessels and surrounding tissue in the cerebral cortex \[35,36\]. Mukamel et al. employed the PCA followed by spatiotemporal ICA method to identify Purkinje cells or Bergmann glial signals in two-photon images \[18\]. In our bone cell fluorescence imaging data, one IC sometimes contained multiple spatially separated target regions after sICA. ICA tends to sort these disconnected regions to be one IC, since they may have strong high correlations with each other. Therefore, morphology-based image processing techniques were necessary to further segment each IC into spatially separated sub-ICs and ensure the one-to-one correspondence between each sub-IC and cellular signal. Validation studies proved that the ICA-based technique was more accurate than the manual ROI method for our image data captured with a fluorescence microscope, which had high image quality with the SNR ranging from 10 to 30 dB. Moreover, the large cell number in our bone cell network data (100–200 cells for each set of data) also greatly increased the difficulty and artificial errors during manual ROI selection. In our studies of \(\text{Ca}^{2+}\) signaling in bone cell networks, it took an experienced analyst 1–2 h to analyze one set of data using commercial analysis software, including manually selecting the cell regions, extracting the signals, and identifying the spikes of each signals. The current ICA-based signal extraction technique greatly reduced the workload and also facilitated the analysis of \(\text{Ca}^{2+}\) spatiotemporal characteristics. Therefore, this study suggests that the unsupervised ICA-based algorithm has great potential in analyzing \(\text{Ca}^{2+}\) signals of bone cell networks.

Under fluid flow stimulation with the same levels, more frequent \(\text{Ca}^{2+}\) spike activities and vigorous temporal \(\text{Ca}^{2+}\) oscillations were demonstrated in MLO-Y4 cell networks. The osteocyte-like MLO-Y4 cells, exhibiting unique repetitive \(\text{Ca}^{2+}\) firing activities stimulated by fluid flow, act similar with the action potentials of a neural network. This further supported our previous hypothesis that the osteocyte network holds the key to cellular memory of mechanical loading history in bone \[14,15,37\]. The prominent difference in temporal \(\text{Ca}^{2+}\) oscillatory activities implied different mechanisms of \(\text{Ca}^{2+}\) responses between the two types of cells. Previous studies indicated that the \([\text{Ca}^{2+}]_i\) elevation under fluid flow stimulation was mainly from two sources: extracellular calcium from the environment and \(\text{Ca}^{2+}\) stores (e.g. endoplasmic reticulum, ER) \[38,39\]. The influx of extracellular calcium was the major source for the rapid initial \(\text{Ca}^{2+}\) spike, and the ER \(\text{Ca}^{2+}\) stores and ATP pathways played a critical role in the subsequent multiple \(\text{Ca}^{2+}\) peaks in osteoblasts \[15,31\]. T-type voltage gated calcium channels (VGCC), which regulate the pacemaking and repetitive firing activities in cardiac and neuronal cells \[40,41\], have also proven to be specifically expressed in osteocytes, but not in osteoblasts \[42\]. Therefore, the mechanism mediating more dramatic temporal oscillatory activities in osteocytic networks might be attributed to multiple channels and their relevant downstream signal pathways, including ATP-gated P2R ion channels and T-type VGCC, etc. The exact cellular and molecular mechanisms are being studied by our group.

In the present study, MLO-Y4 cell networks possessed more robust intercellular spike synchronization and signal correlation than MC3T3-E1 cells under all fluid flow levels. The synchronous activities in MLO-Y4 cells also negatively correlated with the cell–cell separation distance, but tended to be flat in MC3T3-E1 cells. These observations share similarities with many previous in vivo \(\text{Ca}^{2+}\) dynamics studies on cells of nervous system. Hirase et al. found that the coordination of

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**Fig. 4.** Signal extraction from the ICA-based technique for the experimental data of typical MC3T3-E1 and MLO-Y4 cell networks under 2 Pa shear stress. The spatial patterns and temporal signals from MC3T3-E1 (A & B) and MLO-Y4 (D & E) cell networks are shown. Each spike in one cellular signal is depicted as a dot (C for MC3T3-E1 cells and F for MLO-Y4 cells).
Ca\textsuperscript{2+} events was low and similar between nearby and distant astrocytes in the cerebral cortex in the absence of provoking conditions. After bicuculline treatment, a significantly negative correlation of synchronization with the cell–cell distance was found, which revealed more obvious Ca\textsuperscript{2+} wave propagation between cells in the astrocytic networks [16]. Studies also found that the Purkinje cells in cerebellar microzones also exhibited faster Ca\textsuperscript{2+} wave propagation when mice arose from rest to locomotion, evidenced by the increase of synchronous activities and negative correlation of cell crosstalk with the cell–cell distance [18]. Therefore, the results in our present study, which were similar with these findings in neural cells, indicated that the Ca\textsuperscript{2+} wave propagation might be much more obvious and faster in MLO-Y4 cell networks than that in MC3T3-E1 cell networks. Our previous study demonstrated that ATP played a dominant role in mediating the intercellular Ca\textsuperscript{2+} wave propagation from the single nanoindented bone cell to its neighboring cells over gap junctions in the network pattern [10], and inhibition for the ATP-related signal pathway can also reduce Ca\textsuperscript{2+} oscillation to one single spike in both osteoblasts and osteocytes under fluid
Therefore, a possible faster extracellular ATP diffusion between neighboring cells in osteocytic network may act as a critical factor to dominate the dramatic difference of spatial intercellular coordination of Ca\(^{2+}\) events between these two kinds of cells. Extracellular ATP can activate the membrane-bound P2 nucleotide receptors, leading to sequential activation of phospholipase C (PLC), generation of IP3, and...
thus triggering the release from the IP3-sensitive ER calcium stores [43,44]. Our previous studies have shown that depletion of the ER calcium stores could severely hamper both the intracellular calcium response and intercellular calcium wave propagation in bone cells [10,15]. The exact cellular and molecular mechanisms underlying the spatiotemporal differences in Ca\(^{2+}\) signaling between osteocyte and osteoblasts are still unknown. We believe that the governing mechanisms depend on unique fast kinetics of membrane Ca\(^{2+}\) channels and the refilling of the ER, as well as faster extracellular ATP diffusion between neighboring cells in osteocytes to generate much more frequent and synchronous Ca\(^{2+}\) events, and also promote many more nearby cells to generate intracellular Ca\(^{2+}\) spikes more synchronously than the distant cells. Ongoing studies on bioluminescence trace for extracellular ATP, and also simultaneous visualization for the cytoplasmic and ER Ca\(^{2+}\) dynamics using FRET-based bioluminescence technique exhibited more accurate signal extraction capacity and molecular mechanisms.

**Conclusion**

The present study clearly demonstrated that the ICA-based signal extraction technique exhibited more accurate signal extraction capacity than traditional ROI analysis for Ca\(^{2+}\) signaling in bone cell networks. This unsupervised technique can dramatically reduce our effort to extract the Ca\(^{2+}\) signals and also facilitate more elaborate spatiotemporal analysis for Ca\(^{2+}\) signaling in bone cell networks. Our results showed that MLO-Y4 cell networks exhibited more frequent and synchronous Ca\(^{2+}\) events, and also promote many more nearby cells to generate intracellular Ca\(^{2+}\) spikes more synchronously than the distant cells. Ongoing studies on bioluminescence trace for extracellular ATP, and also simultaneous visualization for the cytoplasmic and ER Ca\(^{2+}\) dynamics using FRET-based bioluminescence technique exhibited more accurate signal extraction capacity and molecular mechanisms.

**Disclosure**

All authors state that they have no conflicts of interest.

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