Abstract

Physical activity shows a positive correlation with overall health, and vigorous intermittent lifestyle physical activity (VILPA) similarly offers advantages in reducing the risk of all-cause mortality. Might the short-time mechanical stimuli be discernible to cells, eliciting commensurate physiological responses? The study’s objective was to investigate the cellular response to short-time mechanical stimuli. Human umbilical cord-derived mesenchymal stem cells (hUCMSCs), isolated and thoroughly characterized, were subjected to various stimuli, including activation and mechanical stretching, with Ca\(^{2+}\) influx assessed through alterations in fluorescence intensity. Further validation of these findings was confirmed through short hairpin RNA (shRNA) and inhibitors. In addition, a comprehensive examination of PIEZO1 alterations was conducted through quantitative real-time polymerase chain reaction (qRT-PCR) and western blot (WB) techniques. The results show different frequencies of stretching stimulation and durations induced varying degrees of Ca\(^{2+}\) influx. The most substantial increase occurred within 2–3 minutes in the group subjected to 0.5 Hz stretching for 2 minutes (\(p < 0.05\)). Stretching at 0.5 Hz resulted in significant elevation in PIEZO1 mRNA expression at 15 minutes and 1 hour. Additionally, stretching cause a gradual rise in PIEZO1 protein levels, with a notable peak observed at 2 hours. In conclusion, cells primarily sense short-time mechanical stimuli through PIEZO1, predominantly mediated by regulated Ca\(^{2+}\) influx. This underscores PIEZO1’s crucial role in cellular responsiveness to transient mechanical cues, advancing our understanding of mechanosensory mechanisms in cellular physiology.

Keywords: Vigorous intermittent lifestyle physical activity, short-time mechanical stimuli, PIEZO1, Ca\(^{2+}\) influx, Yoda1.

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Introduction

Physical activity is intricately linked to human health, with the potential to substantially reduce mortality rates, encompassing all-cause mortality (Kraus et al., 2019), cardiovascular disease, diabetes, and certain tumors (Bull et al., 2020; Moore et al., 2016; Rezende et al., 2018). As physical activity guidelines have evolved, the concept of vigorous intermittent lifestyle physical activity (VILPA) has gained prominence. VILPA encompasses brief, sporadic episodes of vigorous physical activity, lasting up to one or two minutes, as an integral part of daily life (Stamatakis et al., 2021; Stamatakis et al., 2019). Recent data have shown a growing body of evidence suggesting that VILPA could decrease the risk of all-cause mortality, coronary artery disease, and tumor-related mortality (Ahmadi et al., 2023; Stamatakis et al., 2022).

Physical activity contributes significantly to human health, involving numerous physiological pathways, with diverse and complex molecular mechanisms at the cellular level, in which Ca\(^{2+}\) plays a pivotal role as a crucial second messenger in processes like muscle cell contraction, cell adhesion, cell cycle progression, cell growth, cell motility, and cell differentiation (McCarron et al., 2017). The irreplaceable role of Ca\(^{2+}\) in sustaining normal physiological function is integral to the overall promotion of health.

The initial process involves mechanical stimuli acting on various organ tissues. It is suggested that cells may directly perceive short-time mechanical stimuli, but the mechanisms behind this perception are not yet fully understood. This process may require the conversion of mechanical signals into biological signals, exerting widespread effects on downstream pathways. The involvement of mechanosensors in this process has been sparsely documented. The molecular landscape governing mechanotransduction is diverse, and we explored into several common and widely impactful mechanosensitive molecules, namely PIEZO1/2, TRPV4, KCN2, and TMEM63A/B. Our investigation revealed distinctive and regular alterations in PIEZO1 expression following short-time stretching stimuli. PIEZO1, primarily expressed in non-excitable cellular domains, assumes a pivotal mantle in the transduction of mechanical stimuli, both intracellular and extracellular in origin, thereby orchestrating a multifaceted role across a spectrum of physiological and pathological milieus (Bagriantsve et al., 2014; Martins et al., 2016; Murthy et al., 2017; Ranade et al., 2014). PIEZO1 functions as a non-selective cation channel, primarily facilitating the influx of Ca\(^{2+}\) (Lewis and Grandl, 2020; Wu et al., 2017; Zheng et al., 2019), which, in turn, participate as second messengers in the regulation of various downstream signaling pathways, profoundly influencing a spectrum of physiopathological processes. The activity of PIEZO1 is characterized by its transient and short-lived nature (Lewis et al., 2017; Nosyreva et al., 2021; Wijerathne et al., 2023; Yang et al., 2022; Zhao et al., 2018), coinciding with VILPA characteristics.

Building upon this foundation, this study delves into the dynamics of PIEZO1 response triggered by short-time stretch stimulation of hUCMSCs. It finds that cells primarily sense short-time mechanical stimuli through PIEZO1, predominantly mediated by regulated Ca\(^{2+}\) influx. This is accompanied by delayed, oscillatory changes in PIEZO1 mRNA and protein levels. This underscores PIEZO1’s crucial role in cellular responsiveness to transient mechanocues, advancing our understanding of mechanosensory mechanisms in cellular physiology.

Materials and Methods

Cell Isolation and Culture

In this study, hUCMSCs were successfully isolated from umbilical cords following established protocols, identified (Zhang et al., 2021), and then subjected to experiments at P3–4.

The umbilical cords used in this study were approved by the Ethics Committee of the Seventh Affiliated Hospital of Sun Yat-sen University (Permit Number: 2019SYUSH-031), and informed consent was obtained from the participants and their families. Briefly, the umbilical cord was washed 3 times with phosphate-buffered saline (PBS) containing 2 % v/v penicillin/streptomycin within 2 hours after ex vivo and removed umbilical vessels from Wharton’s jelly. Then Wharton’s jelly was cut into pieces about 1 mm\(^3\) in size, evenly spread in the 6-well plate and placed upside down at 37 °C in a 5 % CO\(_2\) humified incubator for 30 min, and then added MSC special medium (Prim® hMSC SF-M, Premedical Laboratories Co., Ltd., Beijing, China) for culturing and changed the medium every 3 days. Digated when the confluence of the cells reaches about 80 %, subcultured at a density of 8000–10,000 cells per square centimeter (cm\(^2\)), and used Dulbecco’s Modified Eagle Medium-Nutrient Mixture F-12 (Cat NO. C11330500BT, ThermoFish, Waltham, MA, USA) medium for culture.

Cell Stretch

We employed a meticulously designed custom apparatus, which facilitates uniaxial cyclic stretching, offering adjustable parameters including frequency, amplitude, and duration. The stretching amplitude varies in a sinusoidal pattern over time while maintaining conditions akin to an incubator environment (5 % CO\(_2\), 37 °C). A concise overview is provided herein, and please refer to the literature for details (Zhu et al., 2023).

Post-sterilization of the culture chamber, a coating of collagen I (Cat NO. A1048301, ThermoFish, Waltham, MA, USA) was applied, and cells were seeded at a density of 1 × 10\(^4\) cells/cm\(^2\). Following a 24–48 h incubation, cells achieved 60–70 % confluence, underwent a 12-hour period of nutrient deprivation, and were subsequently replenished with fresh complete medium. Ensuring pre-stretched readi-
ness, the pulley system was engaged. Stretching parameters were meticulously calibrated, and upon achieving stability, stretching stimulation was conducted, adhering to the pertinent parameters aligned with the experimental plan (stretching amplitude 10%, stretching frequency 0.25/0.5/1.0 Hz, stretching time 0/0.5/1/2/4 minutes).

**Imaging of Intracellular Ca\(^{2+}\) and Data Analysis**

HUCMSCs were cultured to 60–70% confluence, followed by a simple wash with Hank’s Balanced Salt Solution (HBSS). Pre-treatments were conducted using GsMTx4 (5 µM (Gnanasambandam et al., 2017), Cat NO. HY-P1410, ThermoFisher, Waltham, MA, USA) according to the experimental design when required, following the instructions provided for the reagent. An appropriate concentration of Fluo-4 AM (Cat NO. F14201, ThermoFisher, Waltham, MA, USA), supplemented with F-127 (Cat NO. P3000MP, ThermoFisher, Waltham, MA, USA) at a concentration of 0.1% (v/v), was prepared according to reagent guidelines and incubated for 30 minutes. Subsequently, the liquid containing Fluo-4 AM was removed, and the cells were washed once with HBBS. Subsequently, the cells were incubated for an additional 10 minutes in HBSS.

The initial cell state was observed and captured using an inverted fluorescence microscope with a x10 objective lens. Subsequent experimental procedures, such as stretching or the addition of the PIEZO1 activator Yoda1 (Cat NO. SML1558-5MG, Merck & Co Inc, Shanghai, China), were carried out according to the predetermined plan. Immediate and sequential image capture was performed under consistent parameters, with a frame rate of 20 frames per minute for a duration of 3 to 5 minutes.

The images acquired were processed and analyzed using Image J software (version 1.54a, National Institutes of Health, Bethesda, MD, USA). Initially, the fluorescence values of images within the same group were quantified. Subsequently, the fluorescence values of images taken before treatment (such as stretching stimulation or Yoda1 activation) were used as a baseline (F₀) and images at different time points after treatment were normalized to this baseline (F/F₀). Finally, the relative increment changes in average fluorescence values at different time points were obtained to reflect the entry of Ca\(^{2+}\) into the cytoplasm. Each sample was analyzed once, and each experimental group underwent at least three repetitions.

**Virus Packaging and Cell Transfection**

HEK293T cells were plated at a density of 1.0 × 10⁵ cells/cm² and cultured in DMEM (Cat NO. C11995500BT, ThermoFisher, Waltham, MA, USA) supplemented with 10% FBS, while without penicillin-streptomycin. The day after plating, when the cells reached approximately 70% confluence, the medium was changed to fresh medium without FBS. Subsequently, the hU6-MCS-CBh-gcGFP-IRESpuromycin (Table 1) plasmid (8.4 µg) was added. Then, 37.8 µL of Lipofectamine 3000 (Cat NO. L3000015, ThermoFisher, Waltham, MA, USA) was used for co-transfection with the second-generation packaging plasmid (6.3 µg pSAX2, HedgehogBio Sci and Tec Ltd., Shanghai, China) and the envelope plasmid (4.2 µg pMD2.G, HedgehogBio Sci and Tec Ltd., Shanghai, China). After 6–8 hours of transfection, the medium was changed to complete medium, and lentiviruses were collected through conditioned medium 48 and 72 h post-transfection. The lentiviral supernatant was filtered through a 0.45 µm filter to remove residual cellular debris. MSCs were seeded in 100 mm dishes at a density of 1.0 × 10⁵ cells/cm². The next day, samples were infected with 1000 µL of lentivirus containing shRNA fragments or vectors. Transfected cells were screened with 1.5 µg/mL puromycin. Cells were cultured in DMEM/F12 for 4 days and then collected for RNA and protein extraction for subsequent experiments.

**RNA Extraction, Reverse Transcription and RT-qPCR Analysis**

Total RNA extraction was conducted using the RNA Isolation Kit (Cat NO. R0027, Beyotime, Shanghai, China) in strict accordance with the manufacturer’s protocols. The RNA isolation procedure comprised a temperature sequence of 37 °C for 15 minutes, followed by a 5-second exposure at 85 °C, and concluded with maintenance at 4 °C. Subsequently, 400 ng of RNA was subjected to reverse transcription along side 4 µL of the provided mix (Cat NO. RR036A, Takara, Beijing, China), and the volume was adjusted to 20 µL with nuclease-free water. The resultant cDNA was further diluted fivefold with 1× TE buffer.

Real-time quantitative PCR (RT-qPCR) analysis was performed using the PowerUp™ SYBR™ Green Master Mix (Cat NO. A25742, ThermoFisher, Waltham, MA, USA). A specific program was executed following established protocols (Zhu et al., 2023). Data analysis encompassed normalization against the GAPDH gene expression, employing the 2⁻ΔΔCt algorithm. Customized and purified primers, designed using Primer 6.0 software, were synthesized (Sangon Biotech Co., Ltd., Shanghai, China). The primer sequences employed in this research are presented in Table 2.

**Protein Extraction and Treatment**

Proteins were extracted with RIPA buffer (Cat NO. 89900, ThermoFisher, Waltham, MA, USA) following the manufacturer’s protocol. In brief, the culture medium was discarded, and the cells were washed twice with cold PBS. Cold RIPA buffer, cocktail inhibitors (Cat NO. 78430, ThermoFisher, Waltham, MA, USA), and phosphatase inhibitor (Cat NO. P1260-1ml, Solarbio, Beijing, China) were added to lyse the cells. The mixture was left to stand for 5 minutes, with occasional swirling to ensure even distribution. The lysate was collected using a pre-chilled scraper and transferred to microcentrifuge tubes. Centrifugation
Table 1. Sequence of shRNA (hU6-MCS-CBh-geGFP-IRES-puromycin).

<table>
<thead>
<tr>
<th>ID</th>
<th>5’ stem loop 3’</th>
<th>5’ stem loop 3’</th>
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<tbody>
<tr>
<td>Sh1-a</td>
<td>Cgg GCGTCTTCCTTAGCCATTACT CTCGAG AGTAATGGCTAAGGAAGACGC TTTTTg</td>
<td>Sh1-b</td>
</tr>
<tr>
<td>Sh2-a</td>
<td>Cgg GCTACGAGAACAAGCCCTACT CTCGAG AGTAGGGCTTGTTCTCGTAGC TTTTTg</td>
<td>Sh2-b</td>
</tr>
<tr>
<td>Sh3-a</td>
<td>Cgg GCGTCATCATCGTGTGTAAGA CTCGAG TCTTACACACGATGATGACGC TTTTTg</td>
<td>Sh3-b</td>
</tr>
</tbody>
</table>

Table 2. Primers used for qRT-PCR in our research.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer (5’-3’)</th>
<th>Primer sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Piezo1</td>
<td>Forward</td>
<td>ATGTTGCCTCTACACCCCTGACC</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>CCAGCACACACATAGATCCAGT</td>
</tr>
<tr>
<td>Piezo2</td>
<td>Forward</td>
<td>CACGTTGGTGAGCCTTGAAG</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>TCAAACTCAGGGTACTCTGT</td>
</tr>
<tr>
<td>KCNK2</td>
<td>Forward</td>
<td>TGGTGGTGAGCCTTGAAG</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>ATGGTGGTGACTCTGTGAG</td>
</tr>
<tr>
<td>TMEM63A</td>
<td>Forward</td>
<td>TGGTGGTGAGCCTTGAAG</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>ATGGTGGTGACTCTGTGAG</td>
</tr>
<tr>
<td>TMEM63B</td>
<td>Forward</td>
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<td></td>
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</tr>
<tr>
<td>GAPDH</td>
<td>Forward</td>
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</tr>
<tr>
<td></td>
<td>Reverse</td>
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</tr>
</tbody>
</table>

qRT-PCR, quantitative Real-time PCR.

was performed at 14,000 × g for 15 minutes to collect the total protein, followed by transfer to new centrifuge tubes. The entire process was conducted on ice. The protein concentration was determined using the BCA protein assay kit (Cat. No. P0012, Beyotime, Shanghai, China), and the concentration was adjusted based on the results obtained. Finally, the protein was boiled for 10 minutes and reserved for the following experiment.

Western Blot (WB)

In accordance with the established laboratory procedures, proteins underwent electrophoresis utilizing Pre-Cast 4–12 % gradient Gels (Cat NO. NP0335BOX, ThermoFisher, Waltham, MA, USA) to achieve optimal separation. Subsequently, the proteins were transferred onto a PVDF membrane using the iBlot™ 2 Dry Blotting System (ThermoFisher, Waltham, MA, USA), following the precise guidelines provided by the manufacturer. The membrane was then subjected to blocking for one hour to using 5 % non-fat powdered milk (Cat NO. D8340, Solarbio, Beijing, China) in 1× Tris Buffered Saline with Tween 20 (TBST). Following the blocking step, the membrane was incubated overnight at 4 °C with the specific primary antibody. Afterward, a suitable secondary antibody was applied, and the incubation continued for an additional hour at room temperature. To ensure proper antibody binding, the membrane was washed three for 10 minutes each time with 1xTBST. The protein expression was assessed using a gel imaging system in conjunction with an ECL chemiluminescence kit (Cat NO. SQ101, EpiZyme, Shanghai, China). Quantitative analysis of the bands was performed using Image J software to analyze the grayscale values. GAPDH was employed as an internal control throughout this research study. The details of all antibodies employed in this research are provided below: anti-PIEZO1 (Cat NO. MA5-32876, ThermoFisher, Waltham, MA, USA), anti-GAPDH (Cat NO. 2118S, Cell Signaling Technology, Danvers, MA, USA), second-antibody (Cat NO. A21010, ThermoFisher, Waltham, MA, USA).

Statistical Analysis

Quantitative data were obtained from a minimum of three replicative experimental iterations, and expressed as mean ± standard deviation (SD) or mean ± standard error of mean (SEM) in the resultant formation dataset. Statistical analyses employed SPSS version 20. Significance was assessed using an unpaired Student’s t-test for two-group comparisons and a one-way ANOVA for three or more samples and multiple t-test with adjusted p-value by Bonferroni correction, with data conforming to standard distribution and aligned variance. Statistical significance was established at a threshold of p < 0.05 (ns not significant, * p < 0.05, ** p < 0.01, *** p < 0.001 and **** p < 0.0001).
Results

Yoda1 Stimulation: Ca\textsuperscript{2+} Influx Initial Peak and PIEZO1 Activation Lasting One Minute

The research involved the determination of optimal concentrations for two key components: the Ca\textsuperscript{2+} fluorescent probe, Fluo-4 AM 1 μM, and the PIEZO1 activator, Yoda1 30 μM. The line chart suggests that the most significant changes in relative fluorescence intensities for Fluo-4 AM and Yoda1 occurred at concentrations of 1 μM and 30 μM, respectively (Fig. 1A). It is noteworthy that, unless explicitly specified otherwise, these optimal concentrations were consistently employed in all subsequent experiments.

In contrast to the control group, stimulation of human hUCMSCs with Yoda1 elicited an immediate surge in Ca\textsuperscript{2+} influx, a phenomenon that persisted for approximately 60 seconds (Fig. 1B,C, \(p < 0.05\)). During this phase, a peak relative fluorescence was observed, surpassing baseline to 2× level, eventually converging to levels akin to the control group by the 90-second mark (Fig. 1B,C, Supplementary Fig. 1A). Upon transitioning to a Ca\textsuperscript{2+}-free D-Hanks extracellular medium, Yoda1 administration also appeared to induce an augmentation in Ca\textsuperscript{2+} influx (Fig. 1D). However, when rigorously compared with the control group (Supplementary Fig. 1B), no statistically significant difference was discerned. In cases where pretreatment with the PIEZO1 inhibitor GsMTx4 was implemented, Yoda1 stimulation similarly resulted in increased Ca\textsuperscript{2+} influx (Fig. 1E,F). Intriguingly, when the three experimental groups were subjected to a longitudinal comparative analysis in the absence of Yoda1 stimulation, the control group exhibited a noteworthy upsurge in Ca\textsuperscript{2+} influx at the 120-second mark compared to the GsMTx4 group (Fig. 1G,H). Conversely, with Yoda1 stimulation, the control group displayed an even more pronounced and rapidly peaking increase in Ca\textsuperscript{2+} influx within the initial 30 seconds (Fig. 1I,J). Notably, no significant distinctions were observed at other time points (Supplementary Fig. 1E). The time-dependent fluctuations in fluorescence intensity observed within the control and GsMTx4 groups provide indirect evidence of PIEZO1 activation and Ca\textsuperscript{2+} influx. Representative fluorescence images of different groups at characteristic time points were shown in Fig. 1K.

Optimal Stretching Stimulus Promotes PIEZO1 Activation and Ca\textsuperscript{2+} Influx

In this research, the dynamics of Ca\textsuperscript{2+} influx resulting from the mechanical stimulation of hUCMSCs through uniaxial cyclic stretching. Various frequencies (0.25/0.5/1.0 Hz) and stretch durations (0.5/1/2/4 minutes) at a 10 % stretch amplitude were employed as experimental parameters. First and foremost, when the stretch stimulation was administered at a frequency of 0.25 Hz for varying durations (0.5/1/2/4 minutes), a notable observation emerged: Ca\textsuperscript{2+} influx exhibited heightened levels across all stretch groups during the initial 30 seconds, gradually diminishing over time (Fig. 2A). Intriguingly, in some instances, Ca\textsuperscript{2+} influx even dipped below the peak observed in the static control group by the 150-second mark (Fig. 2D, Supplementary Fig. 2A). Comparable outcomes were evident with 1.0 Hz stretch stimulation, reflecting a gradual decline in Ca\textsuperscript{2+} influx across all stretch durations, albeit with a relatively extended period of elevated flow in the early phase (Fig. 2C). However, no significant deviation from the respective stretch groups was discerned at the peak when compared to the control group (Supplementary Fig. 2B). Subsequently, a resurgence in Ca\textsuperscript{2+} influx was noted in the stretch group after 210 seconds (Fig. 2G).

The examination of Ca\textsuperscript{2+} influx over a 5-minute period following stretch stimulation, at a frequency of 0.5 Hz, revealed consistent elevation in Ca\textsuperscript{2+} levels within each stretch group in comparison to the control group. Additionally, each group exhibited distinct temporal trends (Fig. 2B). The fluorescence alterations at corresponding time points provided direct insights into the overall trajectory of each group’s response (Fig. 2F). Notably, the 0.5-minute group displayed a gradual increase in Ca\textsuperscript{2+} influx after 90 seconds, while the 4-minute group exhibited a slight and non-significant increase after 180 seconds (Fig. 2B,F). In contrast, the 1-minute group initially exhibited a higher Ca\textsuperscript{2+} influx level after stimulation, which gradually diminished over the course of 5 minutes, eventually approaching the baseline state (Fig. 2F). While, the 2-minute group displayed a gradual increase in Ca\textsuperscript{2+} influx beginning at the 40-second mark, reaching its zenith between 120 and 180 seconds (Fig. 2B,F, \(p < 0.05\)). During this peak period, Ca\textsuperscript{2+} influx exceeded that of the other groups, with F/F\textsubscript{0} measurements reaching approximately 1.5 or higher. Subsequently, after the peak, Ca\textsuperscript{2+} influx gradually declined (Fig. 2F, \(p < 0.05\)). Representative fluorescence images at specific time points after different duration of 0.5 Hz stretching were shown in Fig. 2E.

Inhibitor GsMTx4 Decreases Stretch-induced PIEZO1 Activation and Ca\textsuperscript{2+} Influx

Within the GsMTx4 pretreatment group, the Ca\textsuperscript{2+} influx exhibited distinct dynamics following stretch stimulation (0.5 Hz 2 minutes). Notably, the peak Ca\textsuperscript{2+} influx manifested earlier in this group post-stretch and was distinctly higher than that observed in the control group within the initial 60 seconds. However, the amplitude of this peak was comparatively smaller, and no significant disparity was discerned when compared to the stretch group in relation to the control group (Fig. 3A,B). As time elapsed, the intracellular Ca\textsuperscript{2+} influx within the GsMTx4 pretreatment group gradually declined, with no significant divergence from the control group and a significant reduction in comparison to the stretching group (Fig. 3A,B).
Fig. 1. Activation of PIEZO1 and Ca\(^{2+}\) influx for the first minute after Yoda1 stimulation. (A) Changes in relative fluorescence intensity over time after incubation with 1/2/4 µM fluo4-AM and treatment with 10 µM and 30 µM Yoda1, respectively, with each group repeated independently three times. (B,C) Curves of changes in relative fluorescence intensity and statistical graphs of the differences between the control group at static and after treatment with Yoda1. (D) Curves of changes in relative fluorescence intensity of the Ca\(^{2+}\)-free group. (E,F) Curves of changes in relative fluorescence intensity and statistical graphs of the differences in the GsMTx4 pretreatment group. (G,H) Curves of changes in relative fluorescence intensity and statistical graphs of the differences among the three groups at static. (I,J) Curves of changes in relative fluorescence intensity and statistical graphs of the differences among the three groups after Yoda1 treatment. (K) Representative fluorescent images of the control group and GsMTx4 pretreatment group at static and after Yoda1 treatment at different time. n, independent experiments, and the separate asterisks, relative to control group. Scale bars = 100 µm. All values were presented as mean ± SD. *: p < 0.05, **: p < 0.01; ***: p < 0.001.
Fig. 2. Changes in PIEZO1 activation and $\text{Ca}^{2+}$ influx regulated by different stretching stimuli. (A,D) Changes in relative fluorescence intensity for 0.25 Hz stretching for 0, 0.5, 1, 2, and 4 minutes. Continuous change curves in (A) and statistical graphs of the changes in relative fluorescence intensity for each group at 30 s, 120 s and 150 s in (D). (B,E,F) Changes in relative fluorescence intensity for 0.5 Hz stretching for 0, 0.5, 1, 2, and 4 minutes. Continuous change curves in (B), representative fluorescent images of each group in (E), and relative fluorescence intensity for each group at 30 s, 60 s, 90 s, 120 s, 150 s, 180 s, 210 s, and 240 s in (F). (C,G) Changes in relative fluorescence intensity for 1.0 Hz stretching for 0, 0.5, 1, 2, and 4 minutes. Continuous change curve in (C) and the change in relative fluorescence intensity of each group at 30 s, 60 s, 210 s, and 240 s in statistical graphs. n, independent experiments, and the separate asterisks, relative to control group. Scale bars = 100 $\mu$m. All values were presented as mean ± SD. *: $p < 0.05$, **: $p < 0.01$; ***: $p < 0.001$. 
Fig. 3. GsMTx4 decreases stretch-induced PIEZO1 activation and Ca$^{2+}$ influx. (A) Changes in relative fluorescence intensity of the static group, the stretching (2 min, 0.5 Hz) group, and the stretching (2 min, 0.5 Hz) group after pretreatment with GsMTx4. (B) Relative fluorescence intensity of each group at 30 s, 60 s, 90 s, 120 s, 150 s, 180 s, 210 s, and 240 s, showing variations and differences. n, independent experiments, and the separate asterisks, relative to control group. All values were presented as mean ± SD. *: p < 0.05, **: p < 0.01; ***: p < 0.001.

PIEZO1 Knockdown Diminishes Ca$^{2+}$ Influx Induced by Yoda1 and Stretching

Initially, we engineered lentiviruses for PIEZO1 knockdown and transfected them into hUCMSCs. Subsequently, we assessed the knockdown efficiency at the mRNA and protein levels, with the results indicating that the sh-1 and sh-3 achieved knockdown efficiencies of approximately 70% (Fig. 4A–C). Subsequently, we selected the sh-3 group for further experimentation.

Under static conditions, the shPIEZO1 (PIEZO1 knocked down) group Ca$^{2+}$ current exhibited a continuous decline, whereas the negative control (NC) or empty vector (EV) group displayed an increase at 60 seconds, eventually reaching levels akin to the initial state. This disparity was statistically significant when compared to the shPIEZO1 group (Fig. 4D,E). Furthermore, in the NC group, Ca$^{2+}$ influx began to decrease gradually at 120 seconds and remained similar to the shPIEZO1 group until 240 seconds (Fig. 4D,E, Supplementary Fig. 3A).

As Yoda1 stimulated, the early peak occurred slightly earlier in the shPIEZO1 group, with a significantly reduced amplitude, leading to an observable difference compared to the NC group (Fig. 4F,G). While the NC group experienced a sharp decline after the peak, no significant divergence was observed between the two groups. Toward the end of the observation period, Ca$^{2+}$ influx in the NC group was lower than that in the shPIEZO1 group (Fig. 4F,G, Supplementary Fig. 3B).

As stretch stimulation groups, the peak Ca$^{2+}$ influx in the shPIEZO1 group occurred significantly earlier, albeit with a lower amplitude and shorter duration. Ca$^{2+}$ influx was higher in the shPIEZO1 group only at the 30-second mark compared to the NC group (Fig. 4H,L, p < 0.05). Subsequently, Ca$^{2+}$ influx in the shPIEZO1 group continued to decrease, with no significant difference observed between the two groups at 60 seconds (Fig. 4H). However, Ca$^{2+}$ flux in the shPIEZO1 group was significantly lower than that in the NC group at all other time points (Fig. 4I). Representative fluorescence images of the NC and shPIEZO1 groups at various time points following a 2-minute, 0.5 Hz stretching protocol were shown in Fig. 4J.
Fig. 4. ShPIEZO1 reduces Yoda1 and stretch-induced PIEZO1 activity and Ca^{2+} influx. (A–C) PIEZO1 was successfully knocked down. Cells were transfected with lentiviral vectors marked with green fluorescent protein (GFP) in (A). Western Blot (WB) results and statistics of gray values of the bands in the NC group and shPEIZO1 group in (B). mRNA expression in (C). n = 3. (D,E) Changes in relative fluorescence intensity at static conditions, with a continuous change curve in (D), and statistics graphs of relative fluorescence intensity in (E). (F,G) Changes in relative fluorescence intensity after Yoda1 stimulation, with a continuous change curve in (F), and statistics graphs of relative fluorescence intensity in (G). (H–J) Relative fluorescence intensity changes after stretching for 2 minutes, with a continuous change curve in (H) and statistics graphs of relative fluorescence intensity in (I), and representative fluorescent images of each group in (J). NC, negative control (or vector). n, independent experiments. Scale bars = 100 µm. Values, in B and C, were presented as mean ± SEM, and others as mean ± SD. *: \( p < 0.05 \), **: \( p < 0.01 \), ***: \( p < 0.001 \).
**PIEZO1 mRNA and Protein Dynamics in Response to Varied Stretching Time and Interval**

Beyond the evident influence of stretch stimulation on PIEZO1 activation and the consequential alterations in Ca\(^{2+}\) influx, we aimed to determine whether this mechanosensitive process was accompanied by corresponding shifts in gene and protein levels. To explore this, we subjected samples to 0.5 Hz (10 %) stretch stimulation for varying durations (2/5/10/15 minutes) and meticulously extracted total RNA at distinct time intervals (0/15/30/60 minutes). Our rigorous examination aimed to unveil fluctuations in PIEZO1 mRNA expresses, as well as potential shifts in mRNA levels pertaining to commonly observed mechanosensitive markers. The outcomes revealed a distinct pattern in PIEZO1 mRNA expression, characterized by dual peaks manifesting within the initial hour following stretch stimulation at various durations (Fig. 5A). Remarkably, the first of these peaks, occurring 15 minutes post-stretching, exhibited greater prominence, particularly evident in the 10-minute group (Fig. 5A, \(p < 0.05\)). However, it is worth noting that no clear, consistent pattern of change was observed across diverse stretch durations at each time point (Fig. 5B).

Diverse durations (5/10/15 minutes) of stretching stimuli led to a general reduction in Piezo2 mRNA levels (Supplementary Fig. 4A). Notably, Piezo2 mRNA exhibited relatively higher levels when subjected to 2 minutes of stretching stimulation at different time points (Supplementary Fig. 4B). Conversely, other commonly mechanosensitive molecules, including TRPV4 KCNK2 and TEME6A/B, displayed either no significant alterations or exhibited changes lacking a discernible pattern (Supplementary Fig. 4C,D). The findings from the aforementioned investigations have illuminated that a brief 10-minute bout of stretching at specific intervals can indeed elicit alterations in PIEZO1 mRNA expression. To comprehensively assess the corresponding modifications in protein levels, we established a temporal framework encompassing five distinct time points (0/0.5/1/2/4 hours). The results obtained through western blot analysis unveiled a discernible trend in PIEZO1 protein dynamics. Notably, an initial upsurge in PIEZO1 protein levels commenced at 1 hour post-stretching and had essentially reached its zenith by the 2-hour mark (Fig. 5C,D, \(p < 0.05\)).

**Discussion**

**VILPA could decrease the risk of related diseases and confer health benefits** (Ahmadi et al., 2023; Stamatakis et al., 2022), and in this study, cells discern short-time mechanical stimuli via the PIEZO1 protein. Our investigation shows that during resting state, PIEZO1 functionality maintains a state of low-level dynamic equilibrium. Upon exposure to brief mechanical stimuli, a notable dynamic shift occurs in PIEZO1’s functional state. The inhibitor GsMTx4 effectively attenuated the fluctuations but did not entirely suppress the Yoda1-induced calcium influx. GsMTx4 interacts independently of peptides and channels at membrane or membrane-protein interfaces and primarily reduces the mechanosensitivity of PIEZO1 by modulating membrane tension (Bae et al., 2011; Gnanasambandam et al., 2017; Suchyna et al., 2004). In contrast, Yoda1 predominantly stabilizes PIEZO1, thereby slowing down its inactivation phase (Syeda et al., 2015). Additionally, a knockdown of PIEZO1 expression substantially mitigates functional oscillation. Ongoing inquiries into PIEZO1 underscore its heightened mechanosensitivity, prompt functional alterations, and a subsequent refractory period post-activation (Lewis and Grandl, 2015; Wijeratne et al., 2022; Yang et al., 2022), aligning seamlessly with established findings.

However, in the process of responding to short-time mechanical stimuli via PIEZO1, it elicits a transient Ca\(^{2+}\) influx. Ca\(^{2+}\), serving as a second messenger, transmits diverse signals at the cellular level and impacts a broad spectrum of physiological functions. For instance, Ca\(^{2+}\) participates in regulating signaling pathways such as Notch (Wang et al., 2020; Wang et al., 2021), TGF-\(\beta\) (Jairaman et al., 2021), NFAT (Zhou et al., 2020), RhoA (Inaba et al., 2021; Tsuchiya et al., 2018), and Wnt (Wong et al., 2018), which govern cellular functions including proliferation, differentiation, migration, and metabolism. Consequently, these processes influence the physiological or pathological processes of the organism. In summary, transient cellular-level stimulation orchestrates the influx of Ca\(^{2+}\) through PIEZO1, eliciting a spectrum of effects.

While alternative mechanoreceptor and transducer molecules may contribute to brief mechanical stimuli, their primary role remains secondary. PIEZO2 is predominantly found in sensory neurons and the respiratory tract. Previous research has shown that PIEZO2 exhibits high sensitivity to cell membrane indentation (Ikeda and Gu, 2014; Moroni et al., 2018; Shin et al., 2019), but its sensitivity to stretching is relatively low (Coste et al., 2015; Moroni et al., 2018; Verkest et al., 2022). Our findings in hUCM-SCs also indicate that PIEZO2’s response to stretching is less pronounced compared to PIEZO1. The regulation of the transient receptor potential (TRP) superfamily remains debated. Some studies suggest TRP channels aren’t sensitive to cell membrane stretching (Nikolaev et al., 2019) but may be activated by cytoplasmic tethers (Yang et al., 2018). In contrast, TRPV4 responds to cell-substrate contact, unlike PIEZO1 activated by membrane stretching (Servin-Vences et al., 2017). The role of cytoskeletal stretching in TRP channel activation, however, remains unclear. In this investigation, TRPV4 displayed only marginal upregulation at the mRNA level without marked oscillations. Conversely, adhesion proteins and integrins predominantly perceive external mechanical stimuli through interactions with the extracellular matrix (Campbell and Humphries, 2011; Li et al., 2016; Valdivia et al., 2023). This intricate process involves the engagement of a larger array of protein...
molecules and exerts a more pronounced influence on prolonged mechanical stimuli.

However, with the escalating duration of stretching, the temporal activation of PIEZO1 and the consequent substantial Ca\(^{2+}\) influx manifest a non-linear correlation. Analogously, a akin pattern is discernible in the delayed oscillations of mRNA levels. The macroscopic ramifications of PIEZO1 channel opening predominantly derive from the stochastic activation of individual PIEZO1 molecules. The extension of stretching time does not propel each PIEZO1 molecule towards its activation threshold, as variables like the resurgence of PIEZO1 also factor into the equation (Lewis et al., 2017; Nosyreva et al., 2021; Wijerathne et al., 2023).

In studies involving uniaxial cyclic stretch-mechanical stimulation of cells, our findings indicate that appropriate stretch stimulation (0.5 Hz, 10 % for 2 minutes) can activate PIEZO1, resulting in a peak in channel opening between 90 to 180 seconds, subsequently leading to Ca\(^{2+}\) influx. As previously mentioned, existing research predominantly suggests that PIEZO1 detects changes in membrane tension by perceiving alterations in local membrane curvature, consequently regulating various functional states (Haselwandter et al., 2022; Jiang et al., 2021; Yang et al., 2022). Within a defined range, elevating membrane tension enhances the likelihood of channel opening through accelerating the rate of channel opening and decelerating the closing process, meanwhile by lengthening the opening interval and shortening the
closing interval (Wijerathne et al., 2023). It is noteworthy that stretch-induced stimulation exhibits notable variations among individual cells due to several factors. hUCMSCs, characterized by short shuttle-shaped morphology, exhibit disordered polarity and orientation on the substrate, resulting in distinct stimulation profiles. Furthermore, variations in PIEZO1 protein distribution on hUCMSCs may also contribute to differences in cellular response. Regions near the end, where membrane tension is most pronounced, exhibit relatively higher sensitivity of PIEZO1 activation. These findings further substantiate the notion that alterations in waveforms resulting from mechanical stretch stimuli primarily stem from changes in PIEZO1 function.

Furthermore, a precise correlation between the amplitude of stretching and activity intensity has not been definitively established. Prior investigations into stretch stimulation primarily employed amplitudes within the range of 5–15 %, with no observed significant cellular damage yet (Cao et al., 2018; Chen and Wu, 2019; Mathieu et al., 2020; Yang et al., 2021). We selected stretching magnitude (10 %) in this investigation emulates a level approaching the upper threshold of cellular endurance.

In relation to PIEZO1 function, it is crucial to consider its dependence on the frequency of mechanical stimulation. Prior research has proposed a correlation between PIEZO1 opening and sinusoidal frequency, suggesting an augmented probability of PIEZO1 activation with increasing stimulus frequency within a specific range (Lewis et al., 2017). However, the scope of this study primarily encompasses a limited range of low-frequency variations, along with post-stimulus superimposed effects, which may yield distinct impacts on PIEZO1. The question arises as to whether the reduction in Ca\textsuperscript{2+} influx represents an adaptive response orchestrated by PIEZO1 to accommodate novel mechanical stimuli (Lewis et al., 2017). Existing data indicate that adaptation plays a relatively minor role in Ca\textsuperscript{2+} influx attenuation, with inactivation emerging as the predominant mechanism. Consequently, our study does not delve into this issue. Within our study, we subjected cells to stretch stimulation at a frequency of 0.25 Hz, revealing no significant disparities among groups. Notably, an initial decline in peak PIEZO1 opening was observed across all durations, with the middle period even exhibiting levels lower than the static control group. Contrastingly, at a frequency of 0.5 Hz, distinct variations emerged among groups during different time intervals. Specifically, for 0.5-minute and 4-minute stretches, PIEZO1 opening exhibited a gradual increase post-stretch, whereas a pronounced early peak followed by gradual decline was noted for the 1-minute stretch. Interestingly, the 2-minute stretch displayed a peak in the middle of the session. In Markov modeling, simulated studies have posited the existence of diverse states of PIEZO1 opening and closing, characterized by varying durations of open and closed states (Lewis et al., 2017; Nosyreva et al., 2021; Wijerathne et al., 2023). Lower frequencies, such as 0.25 Hz, may activate only a fraction of PIEZO1, leading to gradual inactivation toward the end of stretching. Conversely, elevated frequencies are closely associated with distinct states of opening and closing, hinting at a time-dependent stimulation influence on longer opening. The precise underlying mechanism is a focal point of our ongoing research efforts.

While our study explores PIEZO1’s role in cellular Ca\textsuperscript{2+} regulation in response to mechanical stimuli, we are keen to understand its relevance in living organisms. Given the challenges associated with direct animal modeling, we intend to utilize dynamic culture systems to develop in vitro organoid models. This approach will enable us to investigate cellular responses to short-term mechanical cues at a tissue level, identify key molecular pathways, and lay the groundwork for understanding the health implications of VILPA.

This study presents the following limitations. Firstly, it focused solely on a single cell type for validation and lacked validation at the tissue level or in vivo, which will be the focus of our next experimental steps. Additionally, there is attenuation in the fluorescence changes of Fluo4 AM, and the accuracy of Ca\textsuperscript{2+} influx in the later stage is slightly lower than in the early stage. This limitation cannot be avoided or overcome in the experiment.

Conclusions

In conclusion, our data evidenced that cells primarily sense short-time mechanical stimuli through PIEZO1, predominantly mediated by regulated Ca\textsuperscript{2+} influx. This underscores PIEZO1’s crucial role in cellular responsiveness to transient mechanical cues, advancing our understanding of mechanosensory mechanisms in cellular physiology.

List of Abbreviations

VILPA, vigorous intermittent lifestyle physical activity; hUCMSCs, human umbilical cord-derived mesenchymal stem cells; shRNA, short hairpin RNA; qRT-PCR, quantitative real-time polymerase chain reaction; WB, western blot; PBS, phosphate-buffered saline; HBSS, Hank’s Balanced Salt Solution; TBST, Tris Buffered Saline with Tween 20; SD, standard deviation; SEM, standard error of mean; NC, negative control; EV, empty vector; GFP, green fluorescent protein; TRP, transient receptor potential.

Declaration of AI and AI-assisted Technologies in the Writing Process

During the preparation of this work the authors used ChatGpt-3.5 in order to check spell and grammar. After using this tool, the authors reviewed and edited the content as needed and takes full responsibility for the content of the publication.
Availability of Data and Materials

The data supporting the conclusions of this study are available from the corresponding author, Z.Y. Zhou, in addition to those included in the published manuscript.

Author Contributions

FAW: administrative support, collection and assembly of data, data analysis and interpretation and manuscript writing. HKC: administrative support, collection and assembly of data and revise manuscript. ZYH: administrative support, collection and assembly of data and revise manuscript. JFL: collection and assembly of data and revise manuscript. ZYZhu: provision of study material and collection data. TT: provision of study material and collection data. XZL: financial support, assembly of data and revise manuscript. MMG: Conception and design, assembly of data and revise manuscript. SYL: financial support and final approval of the manuscript. All authors contributed to editorial changes in the manuscript, read and approved the final manuscript, and have participated sufficiently in the work to take public responsibility for appropriate portions of the content. All authors have agreed to be accountable for all aspects of the work.

Ethics Approval and Consent to Participate

This study was approved by the Ethics Committee at the Seventh Affiliated Hospital of Sun Yat-sen University, Shenzhen, China (NO. 2019SYSUSH-031), and informed consent was obtained from the participants and their families.

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Conflict of Interest

The authors declare no conflict of interest. MJS is serving the Editorial Board members of this journal. We declare that MJS had no involvement in the peer review of this article and has no access to information regarding its peer review. Full responsibility for the editorial process for this article was delegated to CE.

Supplementary Material

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**Editor’s note:** The Scientific Editor responsible for this paper was Chris Evans.