Charge injection based electrical stimulation on polypyrrole planar electrodes to regulate cellular osteogenic differentiation†

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In this study, polypyrrole (Ppy) electrodes were prepared to support an electrical stimulation to MC3T3-E1 cells for regulating their osteogenic differentiation. The charge injection capacity \( (C_{ij}) \) of the Ppy electrodes could be adjusted by the Ppy thickness, and a higher \( C_{ij} \) could make the electrode able to produce a higher charge injection quantity \( (Q_{ej}) \) at applied voltage. The \( Q_{ej} \) onto electrode could be considered as the intensity of the stimulation pulse to cells, and the pulse frequency means the number of electric stimulation with \( Q_{ej} \) at one second. Hence, we conducted the present work in the view of \( Q_{ej} \). When the cells were electrically stimulated for 1 hour per day, the electrodes with \( Q_{ej} \) ranged in 0.08–0.15 \( \mu Q \) had an obvious role in enhancing cellular osteogenic differentiation whereas \( Q_{ej} \) of lower than 0.03 \( \mu Q \) or more than 0.30 \( \mu Q \) gave the stimulations with no or negative effects. And the stimulation with 1 or 25 Hz showed to enhance the differentiation, whereas the stimulation with 50 Hz gave an inhibiting effect. We further found the osteogenic differentiation potential triggered by electrical stimulation was related to cell growth stage, and the stimulation carried out at early stage (day 2–5) during 8 days cell culture showed more contribution to enhancing osteogenic differentiation than that at later stage (day 6–8). It is proposed that the desired stimulation effects require that an appropriate voltage-gated calcium ion channel and efficient intracellular calcium ion oscillation are well activated. This work therefore reveals \( Q_{ej} \) as an important electrode parameter to decide effective stimulations and provides an insight into understanding of the role of electrode material characters in regulating cellular osteogenic differentiation during stimulation.

1 Introduction

Electric fields are believed to play a critical role in major biological processes such as embryogenesis, muscle contraction, wound healing and tissue regeneration.1−3 Inspired by the natural electrical properties of bone, an electric field stimulation generated by exogenous electric devices demonstrates a great potential to regulate osteogenic functions of stem and osteoblast-like cells and promote bone growth.4−10 To generate an effective electrical stimulation, the electric devices/electrodes and material selections are crucial. Compared with traditional parallel electrodes, planar interdigitated electrodes (IDE) integrate the pairing electrodes onto the same plane and generate stimulation under low voltage and in a direct, effective, highly reproducible and controlled manner.11−13 Various materials have been served as electrodes with different electric field intensities (or voltages) for electrical stimulation in osteogenic functions. For instance, McCullen et al. used a gold electrode with interdigitated shape to deliver stimulation under alternating electric field of 1 V cm\(^{-1}\), and the results showed an increase in cellular osteogenic differentiation potential of human adipose-derived stem cells (hASCs).4 An electrode material of conductive 3D silk foams produced stimulation with an interpenetrating network of poly (pyrrole-co-(2-hydroxy-5-sulfonic aniline)) bone tissue scaffolds with the electric field of 1 V cm\(^{-1}\) and had a potential to enhance the differentiation toward osteogenic outcomes of hMSCs.14 A Ti electrode covered by defined nanotubular TiO\(_2\) produced stimulation with constant electric field strength of 200 or 400 mV cm\(^{-1}\), which triggered the cellular osteogenic induction of mesenchymal stem cells (MSCs).5 An electrode material of conductive polypyrrole films supported stimulation with electric field of 350 mV cm\(^{-1}\), and showed its capability to promote osteogenesis of calcium deposition in the extracellular matrix of rat bone marrow mesenchymal stem cells (rBMSCs).15 Pelto
et al. showed an enhanced early osteogenic differentiation of adipose stem cell proliferation on an electrode material of chondroitin sulfate (CS)-doped polypyrrole-coated polylactide scaffolds, which provided stimulation with DC voltage of 200 mV cm\(^{-1}\).\(^{16}\) Hardy et al. developed electrode materials of bio-mineralized conducting polymers and conductive polycaprolactone (PCL) fiber-based bone tissue scaffolds to enhance the differentiation towards osteogenic outcomes of hMSCs with the electric field of 100 mV cm\(^{-1}\).\(^{17,18}\) A moderately conductive artificial extracellular matrix coated polyaniline substrates was employed by Thrivikraman to apply stimulation with intermittently electric field of 3.6 mV cm\(^{-1}\), and the data showed an enhancement in osteogenic differentiation potential of human mesenchymal stem cells (hMSCs).\(^6\) It is evident from this vast literatures that a large range (1 V to 3.6 mV cm\(^{-1}\)) of the applied voltage intensities were involved to stimulate to cells for osteogenic functions due to different electrode materials.

An electrode for electric stimulation to cells is desired to elicit effective cellular responses at applied voltages as low as possible in order to avoid electrolysis and accompanying pH- and ionic-gradients in long-term experiments.\(^{3,19,20}\) For these electrodes, electrical properties would be of greater concern.\(^{21-23}\) The interaction ability of the electrodes with external species is believed to be proportional to electrical property of charge injection capacity \((C_Q)\) of electrodes at a given applied voltage.\(^{24}\) Therefore, it is pursued for a material to support an electrode with high \(C_Q\).

In this study, conductive polymer polypyrrole (Ppy) was selected as electrode material due to its reversible-faradic reactivity which resulted in a high \(C_Q\). Ppy planar interdigitated electrodes were prepared by electro-polymerization, and the \(C_Q\) of electrodes was adjusted by changing Ppy polymerized conditions. An electrical stimulation with biphasic pulse mode was performed during culturing of MC3T3-E1. In the view of charge injection, the effect of electrical stimulation on the cellular osteogenic differentiation was studied at different applied voltages and stimulation times (the pulse frequency and the total stimulation time). Moreover, the possible mechanism of electrical stimulation to cells was also discussed.

## 2 Materials and methods

### 2.1 Preparation of Ppy electrodes

ITO glass (10 mm \( \times \) 10 mm) was patterned by laser etching to form 500 \( \mu \)m width ITO strip and spacing (Fig. 1A). The electro-polymerization of Ppy was conducted with a two-electrode electrochemical system under galvanostatic conditions. The patterned ITO substrate and platinum sheet were used as a working and counter electrodes, respectively. The current densities of 3 mA cm\(^{-2}\), 5 mA cm\(^{-2}\) and 8 mA cm\(^{-2}\) with 30 s were utilized to electro-polymerize for forming Ppy electrodes with different Ppy thickness. As-formed electrodes were washed with deionized water and followed by air-drying overnight. The electro-polymerization solution was consisted of 0.1 M pyrrole (Sigma-Aldrich, USA) and 0.1 M sodium \( p \)-toluene sulfonic acid (pTS) (Macklin, China).

### 2.2 Characterizations of Ppy electrodes

Raman spectra of the prepared electrodes on patterned ITO substrate were taken by OMNIC Dispersive Raman (Thermo Fisher Scientific, USA) with a DXR laser operating at 532 nm with incident power of 10 mW. The prepared electrode materials were peeled from patterned ITO substrate and mixed with KBr, and then FTIR spectrum was collected in the range of 4000–1000 cm\(^{-1}\). The thickness and morphologies of the prepared electrodes were examined by Field Emission Scanning Electron Microscopy (FESEM).

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**Fig. 1** Characterization of Ppy electrodes prepared by electrochemical polymerization. The interdigitated electrode (10 mm \( \times \) 10 mm) with 500 \( \mu \)m width and spacing (A). The photograph of patterned ITO substrate (left) and Ppy electrode (right) prepared by electrochemical polymerization (B). The Raman spectrum (C) and FTIR (D) of the Ppy electrodes; thickness of the patterned ITO substrate (E) and Ppy electrodes prepared at current density of 3 mA cm\(^{-2}\) (F), 5 mA cm\(^{-2}\) (G), and 8 mA cm\(^{-2}\) (H); surface morphologies of the patterned ITO substrate (I) and Ppy electrodes prepared at current density of 3 mA cm\(^{-2}\) (J), 5 mA cm\(^{-2}\) (K), and 8 mA cm\(^{-2}\) (L). (E–H) shares the same scale bar of 200 nm, (I–L) shares the same scale bar of 500 nm.
Mouse calvaria-derived pre-osteoblastic cells (MC3T3-E1) were cultured in a home-made culturing device made of polytetrafluoroethylene (PTFE) and the Ppy electrode was placed in a home-made culturing device made of PTFE. 2.4. Cell spreading, viability and proliferation

Cellular attachment and spreading on Ppy electrodes after 1 day of culture was determined by using immunofluorescence staining. In addition, typical morphology of cells on Ppy electrodes after 2 days of stimulation was also observed. The cells on the electrodes were fixed with paraformaldehyde (4% in PBS) for 15 min, and then permeabilized with 0.4% Triton X-100 for 15 min, blocked in solution contained BSA and FBS (2% in PBS) for 1 hour. After washing in PBS, cells were incubated with fluorescent dye of anti-vinculin (EPR8185, Abcam, UK), rhodamine phalloidin (Phalloidin-iFluor™ 594 Conjugate, AAT Bioquest, Inc. USA) and 4′,6-diamidino-2-phenylindole (DAPI, ENZ-594 Conjugate, AAT Bioquest, Inc. USA) for 1 hour. A Fluorescent staining. In addition, typical morphology of cells on Ppy electrodes was further collected using the miRNeasy Mini Kit (QIAGEN 217004, USA) via measuring the optical density at a wavelength of 405 nm. The ALP activities were obtained by normalizing the quantitative assay values to total protein contents tested in a BCA protein assay.

2.6. Alkaline phosphatase analysis

Alkaline phosphatase activity was measured after 8 days of culture. The cells on Ppy electrodes were rinsed with PBS for three times. The cells were lysed with CelLytic Buffer (Sigma, St. Louis), and the obtained cell lysate was centrifuged with the speed of 12 000 rpm at 4 °C for 15 min. The supernatants were assayed by LabAssay™ ALP (Wako Pure Chemical Industries, Ltd. Japan) using 40 cycles at 95 °C, 60 °C and 72 °C for 30 s, then 72 °C for 30 s and the total RNA was extracted using TRIzol reagent and the expression of osteogenesis-related genes (ALP, Runx2 and OCN) and Ca²⁺-calceinurin/NFAT signal genes (calmodulin (CaM), calcineurin (CaN), and NFAT) was analyzed by real-time (RT) polymerase chain reaction (PCR) assay. After 8 days of culture, the total RNA was extracted using TRIzol reagent and further collected using the miRNeasy Mini Kit (QIAGEN 217004, USA). DNA samples were reverse transcribed to cDNA in reactions using the PrimeScript™ RT reagent Kit with gDNA Eraser (Perfect Real Time) (Takara RR047A, Japan) according to manufacturer's protocol. The qPCR reactions were conducted on the Mastercycler® ep realplex system (Eppendorf, Germany) with a SYBR Green (PowerUp™ SYBR™ Green Master Mix (Applied Biosystems A25742, USA)) using 40 cycles at 95 °C for 2 min, 60 °C for 30 s, then 72 °C for 30 s and were performed in triplicate for each cDNA. The relative expression of genes was normalized to that of the reference gene β-actin. The following primer pairs related to the targeted RNA were used in this study: β-actin (5'-AATGGTGGCTAGGACTTTG-3' and 5'-GGGACCTCTGGTAAACCTATT3'); ALP (5'-CCAGAAGA-GACCCGTAAG-3' and 5'-TCTTGTCCGTCGCCTACAT-3'); OCN (5'-GCAAAGGTAGTGAACGACTC-3' and 5'-...
2.8. Measurement of intracellular Ca^{2+}

Intracellular Ca^{2+} levels were measured using Fluo4-AM calcium indicator (Dojindo Laboratories, Japan) after 2 days of stimulation or stimulation at later stage (day 4–5). After stimulation, the cells on Ppy electrodes were rinsed with HBSS and the 400 μL of Fluo4-AM dye loading solution (4 μM in HBSS) was added. The plates were incubated at 37 °C for 30 minutes for the complete reaction. Successively, the cells were washed with HBSS and collected by 0.5 mL trypsin. A flow cytometer (Cytoflex, Beckman Coulter, China) with the emission wavelength at 518 nm and excitation wavelength at 495 nm.

2.9. Statistical analysis

All the quantitative data in this study were expressed as mean ± standard deviation (S.D.). Statistical analysis was performed using the software of Statistical Package for the Social Sciences (SPSS) version 19. The significant differences were performed by one-way analysis of variance (ANOVA) followed by Tukey’s test. In all of the statistical evaluations, the differences with p < 0.05 were considered as statistically significant.

3 Results

3.1. Characterizations of Ppy electrodes

As shown in Fig. 1A, the patterned electrode was interdigitated shape (10 mm × 10 mm) with 500 μm width and spacing. Fig. 1B shows the photograph of patterned ITO substrate (left) and the prepared electrode (right) that obtained by electro-chemical polymerization. Obviously, black materials with interdigitated shape were found after polymerization.

As shown in Fig. 1C, the obtained electrode exhibits typical Raman peaks at 1580 and 980 cm⁻¹, which are assigned to the symmetric C–C backbone stretching mode and C–H plane deformation. The FTIR spectrum (Fig. 1D) shows the broad band around 3000–3500 cm⁻¹, which could be ascribed to N–H and aromatic C–H stretching vibrations. The two absorption peaks at 2851 cm⁻¹ and 2922 cm⁻¹ are corresponded to symmetric and asymmetric stretching vibrations of C–H groups, respectively. The absorption peaks at 2360 cm⁻¹ is related to stretching vibrations of C–N. The bands at 1635 cm⁻¹ and 1540 cm⁻¹ are corresponded to C=C ring stretching vibration of pyrrole ring and benzene ring of dopant (pTS).

The Raman and FTIR spectra of the prepared electrode indicated that Ppy was formed on the patterned ITO substrate.

The SEM images of the electrode cross-section (Fig. 1F–H) show that the thickness of the Ppy electrodes on patterned ITO substrate (Fig. 1E) by electro-polymerization increased with current densities. The electrode thickness of 20, 50 ± 3 nm and 80 ± 5 nm corresponded to the electro-polymerization at current densities of 3 mA cm⁻², 5 mA cm⁻² and 8 mA cm⁻², respectively. Fig. 1J–L shows the morphologies of the obtained electrodes at different current densities. Ppy grains were observed to homogeneously cover the patterned ITO substrate (Fig. 1I) at current density of 3 mA cm⁻² (Fig. 1J), and the grains became large at current density of 5 mA cm⁻² (Fig. 1K). Further increasing the current density to 8 mA cm⁻², the large grains demonstrated a smoother and denser surface morphology (Fig. 1L). The Ppy electrodes prepared at current density of 3 mA cm⁻², 5 mA cm⁻² and 8 mA cm⁻² were named by Ppy-I, Ppy-II and Ppy-III, respectively.

3.2. The C_q and Q_{inj} of Ppy electrodes

The CV curves (Fig. 2A) demonstrate that three typical electrodes exhibit different loops within the voltage window ranging from 1 to −1 V in culture medium. The enclosed area in CV curve with the overall voltage range represents charge injection capacity (C_q) of the electrodes, and C_q increased with polymerized current densities. Obviously, the patterned ITO substrate demonstrated a much lower C_q compared to that of Ppy electrodes.

Fig. 2B shows the electric current responses of Ppy-I, Ppy-II and Ppy-III electrodes to the electric voltage pulses at 5 mV. The curve area of time integral of the current in charging period was considered to be the quantity of electric charge injected.

Fig. 2 Electrochemical analysis of Ppy electrodes in culture medium. (A) Cyclic voltammetric (CV) measurement; (B) current–time curve collected with a pulse signal; (C) the Q_{inj}–voltage curve of Ppy electrodes calculated from the time integral of the current in (B).
onto the electrodes, i.e. $Q_{inj}$ (Fig. S2†). Fig. 2C shows the $Q_{inj}$ obviously increased upon the increasing of electrode $C_Q$ and applied voltage.

3.3. Cell adhesion, spreading, viability and proliferation on Ppy electrodes

Fig. 3A-C shows the cellular morphologies on the prepared electrodes indicated by immunofluorescent staining after 1 day of culture. F-actin and vinculin indicated the cytoskeleton-related proteins associated actin and integrins, respectively. The organization of actin and expression of vinculin throughout the spreading regions of cells grown on all Ppy electrodes confirmed the cytocompatibility of the electrodes. The results revealed that the prepared Ppy electrodes were suitable for cell adhesion and spreading.

Fig. 3D shows that cells on different electrodes presented the similar viability after 1 day of culture and cellular proliferation increased significantly after 5 days of culture. These results indicated the excellent adhesion and biocompatibility of the electrodes.

3.4. The ES effects of stimulation intensity on osteogenic differentiation

For electrical stimulation to cells, 5 mV and 25 mV for applied voltages were selected because the voltages could give the Ppy electrodes to have significant difference of $Q_{inj}$ in three orders of magnitude according to the $Q_{inj}$–voltage curve (Fig. 2C).

As shown in Fig. 4A, at the applied voltage of 5 mV, Ppy-I electrode with $Q_{inj}$ of 0.01 $\mu$Q showed no effect on osteogenic differentiation due to low $C_Q$. Ppy-II electrode with $Q_{inj}$ of 0.08 $\mu$Q expressed a promoted differentiation, while Ppy-III electrode with $Q_{inj}$ of 0.30 $\mu$Q inhibited the differentiation due to high $C_Q$.

At the applied voltage of 25 mV, Ppy-I electrode with $Q_{inj}$ of 0.03 $\mu$Q still showed no effect on osteogenic differentiation, whereas Ppy-II electrode with $Q_{inj}$ of 0.40 $\mu$Q and Ppy-III electrode with $Q_{inj}$ of 1.5 $\mu$Q inhibited differentiation. Those indicate the Ppy electrode with high $C_Q$ at low applied voltage is able to enhance differentiation.

Fig. 4B–D shows the osteogenesis-related gene expressions of the cells stimulated on Ppy-II electrode with $Q_{inj}$ of 0.08 $\mu$Q (5 mV) and 0.30 $\mu$Q (25 mV). The results showed that the gene expressions of ALP and Runx2 were up-regulated for $Q_{inj}$ of 0.08 $\mu$Q, whereas the stimulation with $Q_{inj}$ of 0.30 $\mu$Q showed no effect on ALP and Runx2 expression, and down-regulated the expression of OCN significantly.

The effect of $Q_{inj}$ on differentiation on Ppy-II electrode was shown in Fig. 5A. ALP activity increased gradually with $Q_{inj}$ increasing, and $Q_{inj}$ of 0.08 $\mu$Q showed the largest ALP activity. The $Q_{inj}$ of 0.4 $\mu$Q decreased the ALP activity significantly.

When $Q_{inj}$ of 0.08 $\mu$Q was set for other two Ppy electrodes using different applied voltages, Ppy-I (60 mV) and Ppy-III (1 mV) electrodes presented the same stimulation effect with Ppy-II electrode (5 mV) for enhancing osteogenic differentiation (Fig. 5B). Obviously, Ppy-II electrode with $Q_{inj}$ of 0.08 $\mu$Q (5 mV) showed the most significant upregulation of osteogenic differentiation compared with Ppy-I and Ppy-III electrodes (Fig. 5A), which was selected for further studies.

Fig. 3  Cellular morphologies on the Ppy electrodes were observed by immunofluorescent staining. (A) On Ppy-I; (B) on Ppy-II; (C) on Ppy-III. Cells were stained for the FA protein vinculin (green), cellular nuclei (blue) and actin cytoskeleton (red). (A–C) shares the same scale bar of 100 $\mu$m. Cellular viability and proliferation on the Ppy electrodes were evaluated by CCK-8 (D).

Fig. 4  The ES effect of the applied voltages on osteogenic differentiation of the cells on Ppy electrodes. (A) ALP activity on Ppy electrodes with different voltages; (B–D) osteogenesis-related gene expressions on Ppy-II with applied voltage of 5 mV (0.08 $\mu$Q) and 25 mV (0.30 $\mu$Q). *$p < 0.05$, **$p < 0.01$, ***$p < 0.001$.

Fig. 5  The ES effect of the $Q_{inj}$ on osteogenic differentiation of the cells on Ppy electrodes. (A) ALP activity on Ppy-II electrodes with different $Q_{inj}$ (B) ALP activity on Ppy-I and Ppy-III electrodes with $Q_{inj}$ of 0.08 $\mu$Q. *$p < 0.05$, **$p < 0.01$, ***$p < 0.001$. 

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3.5. The ES effects on cell cytoskeleton

When cells were seeded onto the Ppy electrode, the cells grew to homogeneously distribute on whole electrode including conducting Ppy and spacing glass strips after 2 days of stimulation, this could result from that all cells on the electrode received the equal stimulation effect.

Fig. 6 shows the cellular cytoskeleton on the Ppy-II indicated by immunofluorescent staining after 2 days of stimulation under $Q_{\text{inj}}$ of 0.08 μQ (5 mV). The spreading area of actin and expression of vinculin in stimulation group was larger than that of non-stimulated group. The results indicated that electrical stimulation promoted the cellular spreading and migration on Ppy electrode.

3.6. The ES effect of stimulation time on cellular osteogenic differentiation

For electrical stimulation to cells at a biphasic electric pulse mode, the frequency of electric pulses means a stimulating pulse number in one second, and total stimulation time represents a stimulating day number in the whole cell culture period.

The effect of stimulating frequency on osteogenic differentiation for 1 hour stimulation per day was showed in Fig. 7A, 1 Hz or 25 Hz demonstrated to enhance the differentiation, whereas 50 Hz inhibited the differentiation. Fig. 7B–D shows the osteogenesis-related gene expressions under the stimulation with different frequency, the cellular gene expressions of ALP, Runx2 and OCN stimulated with 1 Hz or 25 Hz were up-regulated, especially OCN expression. However, the stimulation with 50 Hz showed no effect on ALP and Runx2 expressions, but down-regulated OCN expression.

The electrical stimulation carried out at early stage (day 2–5) showed more significant contribution to promotion of cellular osteogenic differentiation than that at early stage (day 6–8) in 8 day culture period (Fig. 8A), and a similar ALP level to that for 8 days.

The osteogenesis-related gene expressions of ALP (Fig. 8B), Runx2 (Fig. 8C) and OCN (Fig. 8D) that stimulated at early stage (day 2–5) were up-regulated. The stimulation carried out at later stage (day 6–8) showed no up-regulation of ALP gene expression, up-regulation of Runx2 and down-regulation of OCN.

3.7. The ES effect on $Ca^{2+}$-calcineurin/NFAT signal pathway

Fig. 9 shows the gene expressions of CaN/NFAT signal pathway of the cells stimulated on Ppy-II electrode with $Q_{\text{inj}}$ of 0.08 μQ (5 mV) or 0.30 μQ (25 mV) and 25 Hz or 50 Hz and stimulation at early or later stage.

For the stimulated cells, their CaM gene expression showed no significant change, and CaN gene expression was up-regulated. As the last downstream gene of $Ca^{2+}$ signal pathway, NFAT expression of the cells that stimulated with $Q_{\text{inj}}$ of 0.08 μQ and 25 Hz or at early stages showed a significant up-regulation.

3.8. The ES effect on intracellular $Ca^{2+}$

Fig. 10 shows the intracellular $Ca^{2+}$ concentration indicated by the mean fluorescence intensity (MFI) after 2 of stimulation or stimulation at later stage (day 4–5) on cell on Ppy-II with $Q_{\text{inj}}$ of 0.08 μQ at the applied voltage of 5 mV.
Fig. 9 The ES effects on the gene expressions of CaN/NFAT signal pathway of the cells on Ppy-II electrode during 8 days culture. (i) The effect of the applied voltage (1 Hz and the stimulation at day 2–8); (ii) the effect of the frequency (5 mV and the stimulation at day 2–8); (iii) the effect of the stimulating days (early stage at day 2–5 + later stage at day 6–8, 1 Hz). *p < 0.05, **p < 0.01.

The flow cytometry assay showed that intracellular Ca\textsuperscript{2+} concentration increased with culture time from 2 days to 5 days without stimulation. And after 2 days of stimulation, intracellular Ca\textsuperscript{2+} concentration increased significantly, whereas stimulation at later stage did not affect the Ca\textsuperscript{2+} concentration.

4 Discussion

The Ppy electrodes showed that their thickness increased with electro-polymerization current densities (Fig. 1). As a result, higher thickness led the electrode have higher $C_Q$ (Fig. 1 and 2A) due to more spaces to accommodate electric charges derived from irreversible faradaic reaction.\textsuperscript{34}

For a given applied voltage, the electrode with higher $C_Q$ had a higher response to an applied voltage pulse (Fig. 2B) as well as a higher charge injection quantity ($Q_{ inj}$) (Fig. 2C) than those with lower $C_Q$. $Q_{ inj}$ of the electrodes could be also tuned by the applied voltage. The electrodes with higher $Q_{ inj}$ means to induce a stronger electrical simulation pulse intensity to cells (Fig. 2C), and could give a stronger simulation effect at lower applied voltage.

During electrical stimulations to cells, electric fields, currents and electrochemical factors derived from electrodes will contribute to the stimulation. In this study, we focused on the effects of electric field because there were no electrical current flow in Ppy stripes, the electrode under the present applied voltage and no redox peaks observed (Fig. 2A).

The Ppy film was testified to be very stable on the patterned ITO substrate for soaking in culture medium and showed to well support cellular adhesion, spreading and proliferation, good cytocompatibility (Fig. 3). The assay of cellular osteogenic differentiation (Fig. 4A) showed that the Ppy electrodes with higher $C_Q$ produced significantly upregulated effect at lower applied voltage as expected.

Since the stimulation pulse intensity is directly related to $Q_{ inj}$, the stimulation effects of the electrodes could be evaluated by their $Q_{ inj}$. The result (Fig. 4A) showed that no or negative simulation effect on cellular osteogenic differentiation was generated for the Ppy electrode with $Q_{ inj}$ of less than 0.03 $\mu$Q (Ppy-I at 5 mV) or more than 0.30 $\mu$Q (Ppy-III at 5 mV), while the Ppy electrode with $Q_{ inj}$ of 0.08 $\mu$Q (Ppy-II at 5 mV) produced the simulation to have an obvious role in enhancing osteogenic differentiation as well as the up-regulated the expression of osteogenesis-related genes (Fig. 4B–D). When the Ppy-I or Ppy-III electrode was applied by 60 mV or 1 mV voltage to have $Q_{ inj}$ of 0.08 $\mu$Q, the two electrodes also demonstrated the obvious stimulation effect to enhance the differentiation (Fig. 5B). Hence, the Ppy electrode with higher $C_Q$ and the stimulation pulse intensity of $Q_{ inj}$ of 0.08 $\mu$Q were appropriate to generate effective stimulations to cells for attaining the enhancement of osteogenic differentiation. In addition, this electrical stimulation promoted the cellular spreading and migration on electrode (Fig. 6).

For the electrical stimulation to cells, the stimulation time is another important parameter besides stimulation intensity of $Q_{ inj}$. The time could be divided into three parts, the time at one second with pulse number (the stimulation pulse frequency), the time at one day with stimulating hour (the daily stimulation time), and the time at the whole cell culture period with stimulating days (total stimulation time). The role of the daily stimulation time had been discussed in our previous work,\textsuperscript{11} and 1 hour per day showed appropriate in promoting cellular osteogenic differentiation, and was adopted in this study.

When $Q_{ inj}$ of Ppy-II electrode was set at 0.08 $\mu$Q (5 mV), the stimulation pulse frequency and total stimulation time were changed, the simulation pulse with 1 Hz or 25 Hz showed to enhance the differentiation, whereas that with 50 Hz inhibited the differentiation and osteogenesis-related gene expressions (Fig. 7). During cell culture for 8 days, the electrical stimulation carried out at early stage (day 2–5) demonstrated more obvious role in promoting cellular osteogenic differentiation than that
at later stage (day 6–8) (Fig. 8). This suggests that the early stimulation could make more significant contribution in enhancing the differentiation in 8 day cell growth.

The assay of CaN and NFAT (Fig. 9) showed that their changing tendency was almost same with one of the electrical stimulation dependent osteogenic differentiation activity (Fig. 4, 7 and 8), revealing that the electrical simulation derived up-regulation in osteogenic differentiation could be attributed to that the calcium ion signaling pathway was activated to trigger osteogenic differentiation.

Based on the mechanism of electrical stimulation on cellular function, the electric field of stimulation manipulates transmembrane potentials of cells and activates the voltage-gated calcium ion channels (VGICs). As a result, the fluxes of calcium ions through the activated VGICs induce the intracellular calcium ion oscillation, and the osteogenic differentiation is triggered.35–38 The VGICs have been proved to play an important role in enhancement of AD-MSCs’ and osteoblasts’ functions of osteogenic differentiation during electrical stimulation.36 VGICs include high voltage-activated (HVA) channels that sub-classified into L-, N-, P-, Q- and R-types and low voltage-activated (LVA) channels (or T-type). VGICs to produce the biological function usually require specific stimulation voltage correspond to specific cells LVA channels are activated by lower membrane potentials, and HVA channels are opened in response to large membrane depolarizations and are activated at higher membrane potentials.39,40

In this study, the electrical simulation generated by \( Q_{\text{ inj }} \) of 0.08 \( \mu \text{mC} \) showed the optimum effects in osteogenic differentiation, which might be attributed to that the studied \( Q_{\text{ inj }} \) range from 0.03 \( \mu \text{mC} \) to 0.30 \( \mu \text{mC} \) for Ppy electrodes involves activation of different VGICs, and the stimulation with \( Q_{\text{ inj }} \) of 0.08 \( \mu \text{mC} \) just activates a VGIC which can trigger well the osteogenic differentiation. The simulation to cells with high frequency of 50 Hz showed no role in promoting osteogenic differentiation (Fig. 6). Since cell growth experiences adhesion, skeletal spreading, proliferation and differentiation, etc., we suggest that the cells at different growth stage could have specific stimulation sensitivity to VGIC activation and intracellular calcium ion oscillation. The obvious enhancement in intracellular \( \text{Ca}^{2+} \) concentration at early stage may activate calcium ion pathway, whereas no effect on \( \text{Ca}^{2+} \) concentration with stimulation at later stage (Fig. 10). The cells at early growth stage could have good sensitivity and result in the early stimulation with more contributions to promoting cellular osteogenic differentiation.

5 Conclusion

Charge injection capacity (\( C_{\text{ inj }} \)) of polypyrrole (Ppy) electrodes could be tuned by electro-polymerization controlled Ppy thickness. The charge injection quantity (\( Q_{\text{ inj }} \)) onto the electrodes was a vital parameter and considered as stimulation pulse intensity to stimulate cells. The electrodes with \( Q_{\text{ inj }} \) of 0.08 \( \mu \text{mC} \) resulted in the stimulation with a good effect in enhancing cellular osteogenic differentiation. The stimulation with 1 or 25 Hz showed an obvious role in improving the differentiation, and the early stimulation (day 2–5) in cell culture period of 8 days had more significant contribution to promote cellular osteogenic differentiation. These are attributed to activating of an appropriate VGIC and intracellular calcium ion oscillation, which induces up-regulation of the expression of osteogenesis-related gene and subsequently triggers osteogenic differentiation. Therefore, this study provides a way to regulate the osteogenic differentiation through electrical stimulation in the view of charge injection, a guide to design electrode in considering materials with \( C_{\text{ inj }} \) and an alternative insight into understanding of the mechanism of electrical stimulation to cellular responses.

Conflicts of interest

There are no conflicts to declare.

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