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GSK-3β is essential for physiological electric field-directed Golgi polarization and optimal electrotaxis

Lin Cao · Jin Pu · Min Zhao

Abstract Endogenous electrical fields (EFs) at corneal and skin wounds send a powerful signal that directs cell migration during wound healing. This signal therefore may serve as a fundamental regulator directing cell polarization and migration. Very little is known of the intracellular and molecular mechanisms that mediate EF-induced cell polarization and migration. Here, we report that Chinese hamster ovary (CHO) cells show robust directional polarization and migration in a physiological EF (0.3–1 V/cm) in both dissociated cell culture and monolayer culture. An EF of 0.6 V/cm completely abolished cell migration into wounds in monolayer culture. An EF of higher strength (≥1 V/cm) is an overriding guidance cue for cell migration. Application of EF induced quick phosphorylation of glycogen synthase kinase 3β (GSK-3β) which reached a peak as early as 3 min in an EF. Inhibition of protein kinase C (PKC) significantly reduced EF-induced directedness of cell migration initially (in 1–2 h). Inhibition of GSK-3β completely abolished EF-induced GA polarization and significantly inhibited the directional cell migration, but at a later time (2–3 h in an EF). Those results suggest that GSK-3β is essential for physiological EF-induced Golgi apparatus (GA) polarization and optimal electrotactic cell migration.

Keywords Physiological electric field · Cell polarization · Golgi polarization · Directional cell migration · Wound healing

Abbreviations
EF(s) Electric field(s)
DC Direct current
GA Golgi apparatus
PI3K Phosphatidylinositol 3 kinases
GSK-3β Glycogen synthase kinase-3β

Introduction

It has long been proposed that electric fields (EFs) at the site of wounds may serve as a signal directing cells to migrate into the wounds [1–6]. Endogenous EFs were detected hundreds of years ago and have been confirmed in several laboratories with modern techniques [7–10]. At skin and corneal wounds in animals and in humans, several different techniques including micro-electrodes, micro-needle arrays, and EF-imagers measured the naturally occurring EFs about 0.42–1.7 V/cm in the superficial layers and at the surface of wounds [11, 12–14]. The field strength in the tissues, however, is likely to be smaller,
because of lower resistance in the tissues underneath the superficial epithelial layers and exponential decrease due to volume conductance of the deeper tissues. A reasonable estimation is a voltage value of about 40 mV/mm within the epidermis [14]. Most published data have used higher voltages and provided significant insights into cellular and molecular mechanisms of EF-induced biological effects. A key question remains as to how the applied EFs used are related to the physiological EFs.

Applied EFs induce cell polarization and directional cell migration [2, 3, 9, 10, 15–18]. We demonstrated that EFs of physiological strength are an overriding guidance cue that directs cell migration during wound healing of corneal epithelial wounds [19–21]. This signal therefore may serve as a fundamental regulator directing cell polarization and migration [7, 19, 20, 22, 23]. Very little is known of the molecular mechanisms that mediate EF-induced cell polarization and migration.

Golgi apparatus (GA) is important in cell polarization and directional cell migration [24]. In response to directional cues in wounds, cells extend membrane protrusion, elongate, and form a leading edge towards the direction of cues—the wound center [25–29]. Efficient migration requires coordination in many cellular activities, such as directed secretion to provide new membrane components to the leading edge [30, 31]. Primary polarization cues initiated at the cell’s leading edge are relayed to the GA complex, triggering a second set of signaling events that controls the subsequent remodeling of the secretary pathway [32]. Recent evidence suggests that the GA, in addition to its function in protein secretion, may be the site of signaling events important for diverse cellular processes [33–35]. GA-localized signaling pathway is required both for the clustering of the GA around the microtubule organizing center (MTOC) and for its polarization in migrating cells [36]. Disruption of GA by brefeldin A (BFA) inhibited lamellodipodia activity and significantly affected cell polarization [16, 37]. When cells in culture migrate to close a scratch wound, this directional extension is accompanied by GA reorientation, to face the leading wound edge, giving the motile cell inherent polarity aligned to the direction of cell migration at the wound edge [25, 26, 29, 38, 39].

Glycogen synthase kinase-3 (GSK-3) was originally identified as one of several protein kinases capable of phosphorylating the rate-limiting enzyme of glycogen deposition, GS (glycogen synthase) [40]. Significantly, GSK is also a central molecule mediating cell polarization [25, 26]. Wounding results in a small GTPase Cdc42-dependent inhibition of GSK-3 through phosphorylation on Serine 9, this correlates with its dissociation from cell polarity molecule Par6. Inhibition of either Cdc42 or protein kinase C (PKC), or expression of a constitutively active version of GSK-3, blocks reorientation of the MTOC. Cdc42, Par6–PKC, and phosphorylated GSK-3 all orientated into the leading edge, where a polarizing signal should be active. Blocking GSK-3 or PKC, however, does not abolish the formation of membrane protrusions. Instead, they become randomly orientated, suggesting an overlap with other polarization mechanisms [41].

We previously reported that a strong EF of 3.0 V/cm induces pronounced GA polarization towards the cathode [16]. In this study, we aimed to answer the question whether physiological EFs induce directional GA polarization and cell migration. We found this to be the case. We further investigated the role of GSK-3/β and PKC in GA polarization and electrotaxis.

Materials and methods

Cell culture media and reagents were purchased from Invitrogen (USA). Human fibronectin was from Sigma–Aldrich (USA). GF109203X (inhibitor of PKC), SB216763 (GSK-3 inhibitor), Ly294002 and Wortmannin (PI3-kanase inhibitors) were purchased from EMD Chemical (USA). GSK-3/β inhibitor XII and GSK-3 inhibitor XVI were purchased from CALBIOCHEM (USA). CHO cells were purchased from ATCC (USA) and grown in NM-F12 medium, with 5% fetal bovine serum (FBS), 50 U/mL penicillin, and 50 μg/mL streptomycin. CO2-independent medium was used for experiments performed in room atmosphere. CHO cells were starved for 3 h in serum-free NM-F12 medium before wounding or treatment with inhibitors. For wound-healing assays, CHO cells were plated on the electrotactic chamber (as described previously [42]) and allowed to grow to confluence before wounding. A monolayer was manually wounded with a 10-μl tip. Wounded CHO monolayer were subject to an EF for 3 h and subsequently used for immunofluorescence analysis. All experiments were performed within passage five of CHO cells and triplicates.

EF stimulation

CHO cells were seeded at low density in electrotactic chambers on fibronectin-coated (10 μg/cm²) Falcon tissue culture dishes for 16–20 h, allowing them to settle and adhere to the base of the dish, before EF exposure. A roof consisting of a No. 1 coverglass was applied and sealed with high vacuum silicone grease (Dow Corning Corporation, USA) on top of the chamber as described previously [42]. The final dimensions of the chamber, through which the electric current was passed, were 40 mm × 10 mm × 0.3 mm. The different strengths of DC EFs were supplied
through agar–salt bridges connecting silver/silver chloride electrodes via beakers of Steinberg’s solution to pools of culture medium at either side of the chamber. This prevents diffusion of electrode products into the cultures. The dish was placed under a Zeiss Axiovert 100 (Germany) microscope with a stage incubator controlling temperature at 37°C.

Analysis of GA polarization

For the GA staining, CHO cells were subjected to different strengths of EFs for 3 h, then fixed in 4% paraformaldehyde (15 min), permeabilized (5 min in 0.2% Triton X-100) and blocked for 30 min with blocking solution (10% goat serum, 1% BSA and 0.02% NaN₃ in PBS). A monoclonal antibody against GM130 (1:100; BD, Franklin Lakes, USA) was used to label the GA (1 h at room temperature). After washing, the cells were incubated with Texas Red conjugated secondary antibodies (1:200; Jackson Immuno-Research Laboratories, USA) and phalloidin-FITC (1:100; Sigma–Aldrich, USA) for 1 h at room temperature. Nuclei were stained with DAPI. Images were obtained under ×40 fluor oil immersion lens of a Zeiss inverted fluorescence microscope (Zeiss Axiovert 100, Germany) with MetaMorph software. Cells in which the GA was within the quadrant facing to between 45 and 315° of the EF direction were scored as polarized in the EF direction (Fig. 1g) [16]. Cells with no EF supplied were

Fig. 1 EFs of physiological strength directed GA polarization in CHO cells. a–f Fluorescent images showed GA (red) in CHO cells cultured in the absence (a) or presence of an EF for 3 h. The cells were fixed and triple-labeled with GM130 antibody (GA marker, red), FITC-phalloidin (F-actin, green), and DAPI (blue). g GA polarization was quantified as the percentage of GA polarized into the quadrant between 45 and 315 degrees in the field direction as shown. h An EF of 0.3 V/cm induced significant GA polarization, which was more pronounced with increased field strength. *P < 0.01 compared to no EFs control. Scale bar 20 μm
regarded as control. The percentage of polarized cells ± SEM was calculated from three separate experiments.

Western blot analysis

Cells were starved in serum-free culture medium overnight before to be applied to 2.0 V/cm EF in vitro. Protein levels of phosphorylated GSK-3β, total GSK-3β and phosphorylated Akt were determined at different time points after EF treatment. For inhibitor experiments, cells were preincubated with 20 μM GF109203X, 20 μM SB216763, 1 μM GSK-3β inhibitor XII, 1 μM GSK-3 inhibitor XVI, 30 μM Ly294002, and 200 nM Wortmannin, respectively, for 1 h and then were applied to EF for 5 min to display the phosphorylated GSK-3 or 3 h for detection of catenin. After various treatments, CHO cells were rinsed with cold PBS and lysed with lysis buffer (10 mM Tris–HCL, 50 mM NaCl, 5 mM EDTA, 50 mM sodium fluoride, 1% Triton X-100, 30 mM Na₃P₂O₇, 1 mM sodium orthovanadate and protease inhibitor cocktail; Boehringer–Ingelheim, Germany). Equal amounts of protein lysates were resolved by 4–12% SDS-PAGE, followed by electroblot analysis onto nitrocellulose membrane (Invitrogen, USA). The membranes were stained with Ponceau S for detection of transfer efficiency, then were blocked with 5% milk TBS (pH 7.4 with 0.1% (w/v) Tween 20) for 1 h. Incubate membrane with relevant primary antibodies respectively (1:1000, Anti-phospho-Akt, anti-β-catenin, anti-phospho-GSK-3β and total GSK-3β from Cell Signaling, USA, and anti-α-tubulin from Sigma–Aldrich) overnight at 4°C. Anti-rabbit/mouse secondary antibody with horseradish peroxidase (1:4,000; Sigma–Aldrich) was used, and the immunoblots were detected by an enhanced chemiluminescence (ECL) detection system (Amersham Pharmacia Biotech, UK).

Time-lapse imaging system and quantification of cell migration

Time-lapse images were recorded every 5 min and analyzed with a MetaMorph system (MDS Analytical Technologies, USA) [43]. Migration directedness (cosine θ) shows how directionally a cell migrated within the field, where θ is the angle between the EF vector and a straight line connecting the start and end position of a cell [42]. A cell moving perfectly toward the cathode would have a directedness of 1; a cell moving perfectly along the field lines toward the anode would have a directedness of −1. Therefore, the average of directedness values of a population of cells gives an objective quantification of how directionally the cells have moved. A group of cells migrating randomly would have an average directedness value of 0. Migration rate was analyzed with the following three parameters. Trajectory speed (T/t/T) is the total length of the migration trajectory of a cell (Tt) divided by the given period of time (T). Displacement speed (Td/T) is the straight-line distance between the start and end positions of a cell (Td) divided by the time (T). Displacement speed along the x-axis (Dx/T) is a cell’s displacement distance along the x-axis (Dx) divided by the time (T).

Cross flow experiment

To exclude the possibility that GA polarization might be an indirect effect mediated by chemical or ionic gradients, temperature, or pH variation, we conducted a more stringent control using a cross-current medium flow chamber modified from that described by Erickson and Nuccitelli [44, 45]. In this experimental setup, the electrotactic chamber was perfused constantly with the culture medium in a direction perpendicular the EF vector. A continuous flow of copious culture medium through the electrotactic chamber was maintained with a peristaltic pump during the entire experimental period (24 mL/h, through a 0.2-mm² channel). This eliminated possible build up of chemotactrant or ionic gradients and maintained stable temperature and pH across the chamber during experiments.

Results

Physiological EFs direct GA polarization

In cells not exposed to an EF, the GA was positioned randomly relative to the nucleus (Fig. 1a). In an EF of 0.3 V/cm, the GA polarization toward the cathode increased significantly from 24 to 41% (P < 0.01) (Fig. 1b, h). The percentage of GA polarized towards the cathode increased gradually when CHO cells were subjected to higher field strengths (Fig. 1b–f, h).

Physiological EFs direct cell migration

In the absence of an applied EF, CHO cells migrated in random directions. The cells have an average net cosine of 0.08 ± 0.06 (directedness) (Fig. 2a). When cultured in a physiological EF as low as 0.3 V/cm, CHO cells showed evident directional migration toward the cathode with a directedness of 0.3 (P < 0.01 compared to no EF control) (Fig. 2b). Cells extended lamellipodia in the direction of migration within 10–15 min after the onset of the EF. Electric stimulation increased the number of cells moving toward the cathode. Exposure to an EF of 0.6 V/cm for 3 h resulted in 91% of the cells moving toward the right quadrant (Fig. 2c). The directedness of CHO cells migrating to the cathode peaked at 3.0 V/cm (0.91 ± 0.02). CHO cells respond to EFs with a threshold at or less than 0.3 V/cm (Fig. 2f).
Exposure to EFs of physiological strength significantly increased the trajectory and translocation speed of cells (Fig. 2g). At 0.6 V/cm, the migration speed is twice and displacement speed is 2.5-fold of the value in control cells ($P < 0.01$). The fastest speed of cell migration is ~20.8 μm/h for trajectory speed ($Tt/T$) and 16.7 μm/h for displacement speed ($Td/T$) following application of the EF of 2.0 V/cm. The increases the migration speed showed a clear voltage dependency (Fig. 2g). The migration speeds reached a plateau EF at 2.0 V/cm and above.

Physiological EF directs GA polarization at the monolayer wound edges

Scratch wounding on a monolayer induces robust polarization of the cells towards the wound, as revealed by the reorientation of the GA in the direction of the movement in many cell types. We observed GA polarization of the wound-edge cells at 3 h after wounding with or without EF exposure. Directional cues such as mechanical stimulation, wound void, and chemical gradients are present in this model. Cells exhibited a polarized GA towards the wound in CHO cells at the wound edge (Fig. 3a). GA polarization at the first line of wound edge cells is 70 ± 2.6%, but just one-half of cells at the second and third line showed GA polarity toward the wound (second line 37 ± 4.2%, third line 39 ± 4.3%, data not shown). When an EF of 1.0 V/cm was applied with the field vector in the normal healing direction, GA polarized more directionally towards the wound (85 ± 3.9%, $P < 0.05$; Fig. 3b, e). Moreover, CHO cells showed obvious GA polarity not only at the first line of wound edges but also at the second and third line of cells (69 ± 11.9 and 54 ± 9.9%, respectively; $P < 0.001$, data not shown) compared to no field control. An additional

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**Fig. 2** EFs of physiological strength directed migration of CHO cells. a–e The migration trajectories of CHO cells for a period of 3 h in absence or presence of an EF showed significant directional migration in an EF of 0.3 V/cm or higher voltage. Each cell’s position at 0 h is positioned at the origin (0, 0). f Voltage dependence of migration directedness of CHO cells showed a threshold below 0.5 V/cm. g Trajectory speed and displacement speed along the x-axis all increased with the field strength of the applied EFs. All results were from three independent experiments. $^*P < 0.01$ compared to no field control. Directedness and migration speed values are shown as mean ± SEM, with total number of cells as n.
increase of EF strength to 2.0 and 3.0 V/cm did not further increase GA polarization ($P > 0.05$ compared to 1.0 V/cm; Fig. 3c–e). When an EF of 1.0 V/cm was applied opposite to the normal healing direction, GA still polarized toward the wound healing direction (Fig. 3f, g). When the field increased to 2.0 and 3.0 V/cm, GA polarized away from the wound toward the cathode (Fig. 3h, i, j).

**EF-directed GA polarization in monolayer wound edge and local chemical gradients**

In our experimental system, the mechanical cues were present throughout. Were the chemical gradients formed at the wound affected by the applied EF? Is GA polarization in a wounded monolayer mainly due to the direct effects of the applied EF, or indirectly by chemical gradients caused by the applied EFs? To exclude possible chemotaxis effects, we did experiments by maintaining continuous flow of culture medium perpendicular to EF lines at different field strength 1.0, 2.0 and 3.0 V/cm. The continuous medium flow indeed had significant effects on the GA polarization either when the EF polarity was in the healing direction or against the default healing direction, especially at higher field strength (Fig. 3e, j). But for most experiments, the EF-induced GA polarization remained. These results suggest that the chemical gradients also play an

![Fig. 3](image)

*Fig. 3* A physiological EF directed GA polarization in the scratch wound model. a–d An EF applied in the default healing direction significantly increased GA polarization into the wound. e GA polarization in the first rows of cells at wound edge was shown. EFs of 1.0, 2.0 and 3.0 V/cm in the default healing direction significantly increased GA polarization into the wound to the same extent. GA polarization at 1.0, 2.0 and 3.0 V/cm was reduced by a continuous medium flow perpendicular to the EF lines with no significant difference. f–i An EF applied against the default healing direction polarized the GA away from the wound. j GA polarization in the first row at EF against wound edge with/without medium flow was shown. GA polarization at 3.0 V/cm was significantly reduced in medium flow. The medium flow was maintained during the experiments to exclude possible chemotaxis. The flow of copious culture medium through the electroatatic chamber was maintained with a peristaltic pump during the entire experimental period (24 mL/h, through 0.2 mm² channel). Data were grouped from three independent experiments. The percentage of cells with GA polarized in the EF direction is shown as mean ± SEM. Monolayer culture of CHO cells were wounded and allowed to heal for 3 h (a–i). The cells were triple-labeled with actin (green), GM130 (red), and DAPI (blue). *$P < 0.05$, **$P < 0.01$ when compared to control. Bar 20 μm
important role in the EF-induced GA polarization in this wounded monolayer model.

Physiological EF is a predominant cue directing cell migration at monolayer wounds

We analyzed the polarity and motility of the cells at the edge of a scratch wound made on the CHO cell monolayer. In a control condition without applied EFs, wounding induced migration of the remaining intact cell sheet into the gap (Fig. 4a, b). When a small EF of 0.3 V/cm was applied with a polarity requiring cells to migrate anodally to heal a wound, 40% of cells did not move into the wound and instead moved toward the cathode. The directedness changed from $-0.74$ (wound without EF) to $-0.37$ (Fig. 4c). When the field strength was increased to 0.6 V/cm, 56% of cells moved to the cathode with a directedness of 0.02 (Fig. 4d). The mean distance moving into the wound was $0.94 \pm 2.7 \mu m/h$ (10.8 $\pm$ 0.2 $\mu m/h$ for no field control). This result indicated that an EF of 0.6 V/cm (anode facing the default healing direction) stopped wound healing completely. With EFs larger than 0.6 V/cm, cell migration simply followed the direction of an applied EF and ignored all other cues generated by wounding a monolayer of CHO cells (Fig. 4e, f). We also analyzed the overriding effect on wound monolayer, which we found to be voltage dependent (Fig. 4g).

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**Fig. 4** A physiological EF was a predominant cue directing cell migration in monolayer wound healing. a Cells at one side of the wound edges migrated to the left. The following experiments were done on this side of the wound. b Migration trajectories of cells at the wound in the absence of an EF were shown. c–f An EF was applied with the field vector against the normal healing direction. Each cell’s position at $t = 0$ is positioned at the origin. Trajectories of cell migration were shown for a period 3 h in the EF. Migration of the cells into the wound was significantly reduced at 0.3 V/cm, and was stopped at 0.6 V/cm. The migration direction was completely reversed and cells migrated away from the wound in an EF of 1 and 2 V/cm. g Directedness of cell migration was voltage dependent with a threshold less than 1 V/cm. $Cos \theta$ is mean values of directedness $\pm$ SEM. $n$ the total number of cells at the given condition.
EFs induced phosphorylation of Akt, GSK-3β and down-regulation of β-catenin

To identify signaling elements involved in EF-directed GA polarization, we examined the phosphorylation state of GSK-3β and total GSK-3β, a critical protein in the establishment of cell polarity [26, 46, 47]. The level of phosphorylation at serine 9 of GSK-3β reaches a maximum level at 3 min after EF application and then gradually decreases to the basal level at 1 h (Fig. 5b). This is one of the earliest molecules we found to be activated by an applied EF. The phosphorylation level of Akt also reached a peak 3 min after exposure to an EF, which persists for 1 h (Fig. 5a). The total GSK-3β and Akt did not change at EF-stimulated samples (Fig. 5a, b, lower channels). When PI3K were inhibited by 30 μM Ly294002 and 200 nM Wortmannin, the phosphorylation of GSK-3β was significantly reduced (Fig. 5e). It indicates that the EF can induce the phosphorylation of GSK-3β mediated by PI3K/Akt. In addition, we found that the level of β-catenin as a marker for function of GSK-3 is reduced by EF application (Fig. 5f). Inhibition of GSK-3/GSK-3β (1 μM GSK-3 inhibitor XVI or 1 μM GSK-3β inhibitor XII) can significant increased the level of β-catenin (Fig. 5f). This result suggests that EF can affected the function of total GSK-3β protein.

Inhibition of PKC abolished phosphorylation of GSK-3β

Atypical protein kinase (PKC) is required to establish and control cell polarity [26, 47]. To test whether GSK-3β phosphorylation induced by an EF is dependent on PKC signaling, cells were pretreated with PKC inhibitor GF109203X (GF) prior to EF application. Inhibition of PKC completely prevented GSK-3β phosphorylation (Fig. 5c). SB216763 (SB, inhibitor of GSK-3β) did not inhibit the phosphorylation of GSK-3β (Fig. 5d). Phosphorylation of GSK-3β could be inhibited by inhibitors of PI3K (30 μM Ly294002 and 200 nM Wortmannin) and inhibitor of PKC (20 μM GF109203X). GF109203X (GF, PKC inhibitor) completely inhibited the phosphorylation of GSK-3β (Fig. 5e). β-catenin as a marker of GSK-3 function was down-regulated by EF. Inhibition of GSK-3 with 1 μM GSK-3 inhibitor XVI (GSK-3in XVI) or GSK-3β with 1 μM GSK-3β inhibitor XII (GSK-3βin XII) could increase the expression of β-catenin. α-Tubulin or GAPDH was used as protein loading control. Histograms show the relative intensity of phospho-GSK3β and phospho-Akt over the tubulin control. EF = 2.0 V/cm. All results were from three independent experiments.

Fig. 5 An EF induced the phosphorylation of GSK-3β.

a, b (upper channels) Time courses of phosphorylation of Akt and GSK-3β were analyzed with western blotting. a, b (lower channels) The expression of total GSK-3β and Akt were no different between the EF-supplied and no EF control. c GF109203X (GF, PKC inhibitor) completely inhibited the phosphorylation of GSK-3β. d SB216763 (SB, inhibitor of GSK-3β) did not inhibit the phosphorylation of GSK-3β. e Phosphorylation of GSK-3β could be inhibited by inhibitors of PI3K (30 μM Ly294002 and 200 nM Wortmannin) and inhibitor of PKC (20 μM GF109203X). GF109203X (GF, PKC inhibitor) completely inhibited the phosphorylation of GSK-3β. f β-catenin as a marker of GSK-3 function was down-regulated by EF. Inhibition of GSK-3 with 1 μM GSK-3 inhibitor XVI (GSK-3in XVI) or GSK-3β with 1 μM GSK-3β inhibitor XII (GSK-3βin XII) could increase the expression of β-catenin. α-Tubulin or GAPDH was used as protein loading control. Histograms show the relative intensity of phospho-GSK3β and phospho-Akt over the tubulin control. EF = 2.0 V/cm. All results were from three independent experiments.
Fig. 5c), but only partly inhibited the GA polarization (from 64 to 52%) in an EF (Fig. 6a, b). It suggests that GA polarization induced by EF is not entirely dependent on PKC and other pathway may play an important role.

Inhibition of GSK-3β abolished EF-induced GA polarization

To investigate the effect of GSK-3β on directed migration and GA polarization, cells were pretreated for 1 h with SB216763 (SB, inhibitor of GSK-3β), 1 μM GSK-3β inhibitor XII and 1 μM GSK-3 inhibitor XVI. The results showed that the inhibition of GSK-3β by SB cannot inhibit the phosphorylation of GSK-3β induced by EF (Fig. 5d), but it significant reduced GA polarization induced by EF (from 64 to ~30%) (Fig. 6a, b). This suggests that the GSK-3β is required in GA polarization induced with EF.

Inhibition of PKC or GSK-3β inhibited EF-induced directional cell migration

We further investigated electrotrans when PKC or GSK-3β was inhibited. In cells pretreated with SB216763, GSK-3β inhibitor XII, GSK-3 inhibitor XVI or GF109203X (PKC inhibitor), ~50% of the directedness of CHO cell migration in an EF was inhibited respectively ($P < 0.01$, Fig 7b). However, the effects of inhibition of PKC and GSK-3β had different time courses on electrotrans. The effect of PKC inhibition on the migration directedness happened within 1 h in an EF. The effect of SB (inhibition of GSK-3β), on the other hand, was not significant until 3 h in an EF (Fig. 7a). Migration speed ($Tt/T$ and $Dx/T$) was significantly affected by SB216763 but not by GF109203X (Fig. 7c). These data suggest that inhibition of GSK-3β affect EF-induced GA polarization more significantly than electrotrans.

Discussion

Proper cell polarization is a prerequisite for directional cell migration. We reported previously that very strong EFs (3.0 V/cm) induce GA polarization and better electrotrans towards the cathode in CHO cells caused by the parallel alignment of cell body along the EF vector [16]. CHO cells might provide an excellent system in which to investigate...
microtubule-directed GA localization due to sustain a variety of their vesicular transport events [48]. In this study we show that physiological EFs induce GA polarization and directional cell migration: (1) an EF of 0.3 V/cm that is well within the physiological range induces GA polarization and directional cell migration; (2) a physiological EF is a predominant cue directing cell’s migration in the monolayer scratch wound; (3) an EF induces rapid phosphorylation (3–30 min) of GSK-3β and Akt; (4) EF-induced GA polarization is mediated by GSK-3β but not phosphorylation of GSK-3β; and (5) EF-induced GA polarization appears to contribute to electrotaxis at a much later stage.

The physiological EFs induced GA polarization and directed migration

Physiological EFs exist in vivo and have profound influences on cell migration, cell division, cell proliferation, and nerve regeneration [17, 49–55]. We showed that a strong EF of 3.0 V/cm directed GA polarization, which in turn reinforces and maintains optimal electrotaxis in CHO cells [16]. Our present data demonstrate that an EF as low as 0.3 V/cm induces GA polarization and directs cell migration towards the cathode, which is the equivalent of a small voltage drop of 0.75 mV extracellularly across a cell 25–30 μm in diameter. GA polarization and directed migration by EF are voltage dependent (Figs. 1–3). The migration directedness and rate increase when the strength of the EFs increases from 0.3 to 3 V/cm. The directedness of cell migration had a similar voltage dependence and threshold as the GA polarization (Fig. 2).

GA might act as a signaling platform, integrating extracellular signal cues and thereby participating in the regulation of downstream events of directional cell migration [33]. Inhibition of traffic from the GA to plasma membrane blocks cell migration [56]. Disintegration of GA significantly inhibited cell migration [16]. EF-induced GA polarization therefore may set up a directed supply of membrane components to the front of the cells, thus facilitating and maintaining optimal directional cell migration. Our present results show that indeed the GA polarization and directional cell migration and migration rate are consistent (Figs. 1, 2). When GA polarization was abolished, the directedness of cell migration was partly reduced at the third hour after EF was applied (Figs. 6 and 7a, b). This suggests GA polarization is a facilitating and maintaining factor in directional cell migration in an EF, rather than an initial factor.

A physiological EF is a predominant cue directing cells migration

Wounding the monolayer generates unequivocal directional movement of the surrounding cells into the wound [57–62]. In this model, multiple candidate cues are generated for cell polarization and migration. These include the initial mechanical stimulation, loss of contact inhibition at the wound edge, and chemical gradients formed upon wounding [63–65]. An additional important signal may be the wound-induced EF. Endogenous EFs at the wounds in vivo are about 1.5 V/cm [5]. Our data demonstrate that EFs as low as 1 V/cm are able to direct cell migration in a predominant manner in the monolayer wound healing
model (Fig. 4). The responses observed at lower fields of 0.6 and 1.0 V/cm indicate that even voltage drops as low as 1.5–2.5 mV per responding cell diameters are able to affect directional cell migration. An EF of 0.3 V/cm significantly inhibited wound edge cells’ migration into the wound (the directedness decreased from ~0.74 to ~0.37, P < 0.001; Fig. 4). Cells at the wound edge stopped moving into the wound completely when subjected to an EF of 0.6 V/cm applied opposite to the normal healing direction. With EFs larger than 0.6 V/cm with the anode facing the default healing direction, CHO cells restored their movement but now followed the direction of electric signal regardless the direction of wound (Fig. 4e, f). Therefore, a threshold for an applied EF to predominantly affect the direction of migration of wound edge cells lies between 0.6 and 1.0 V/cm. The EF-induced opening of the wound has a profound implication. Because several guidance cues collectively drive healing in this model, the complete reversal of healing must mean that an applied physiological EF has an overriding influence on the combined efforts of the other guidance cues (Fig. 4f, g). Thus, we conclude that a physiological EF is a predominant guidance cue directing cell migration in this scrape wound model.

Scratch-wounding monolayer generates reproducible and consistent cues to induce GA polarization in fibroblasts, astrocytes, and endothelial cells [25, 26, 29, 38, 39]. A physiological EF with the polarity in the healing direction induces GA to polarize directionally towards the wound (Fig. 3a–e). When the EF was applied with the field vector against the normal healing direction, the field strength of 2 V/cm was required to induce GA polarization in the field direction and away from the wound.

Chemical gradients may form at the wound site. We used cross-current flow experiments in which a continuous medium flow is maintained perpendicular to the EF vector, which would exclude chemical gradient. The result showed that fewer wound edge cells displayed GA polarization than those in non-flow EF control experiments. This indicates that chemical gradients may contribute to GA polarization.

GSK-3β mediates EF-directed GA polarization and migration

GSK-3β is a multi-task kinase that plays important roles in many signaling pathways, such as Wnt and Hedgehog signaling pathways. GSK-3β has been indicated in the regulation of cell division, stem cell renewal and differentiation, apoptosis, circadian rhythm, transcription, and insulin action [66–73].

Applied EFs transiently induce phosphorylation of GSK-3β as well as Akt (Fig. 5a, b). Applied EFs induce tyrosine phosphorylation of acetylcholine receptors, epidermal growth factor receptors (EGFR) [74, 75]. Those membrane receptors are upstream of PI3 kinase and PKC. Mitogen-activated protein kinase (MAP kinase), PI3K/Akt and PKC are upstream signaling molecules that may control phosphorylation of GSK-3β [76–79]. Indeed, inhibition of PI3 kinase with either LY294002 or wortmannin significantly inhibited phosphorylation of GSK-3β (Fig. 5e). Cross-talk amongst MAP kinase, PI3 kinase and PKC may happen, for example, PKC can be activated by PI-3,4,5-(PO4)3 (PIP3), a product of PI3 kinase activation [80–83]. Our results suggest a requirement for PI3 kinase and PKC in phosphorylation of GSK-3β, thus mediating signaling from applied EFs.

A striking role for GSK-3β is in the regulation of cell polarization in a monolayer wound healing model [26, 84]. In the presence of GSK-3 inhibitors (SB216763 and LiCl), centrosome polarization was blocked at the wound edge. Phosphorylation of GSK-3β, however, appears not to be essential for centrosome/GA reorientation in an astrocyte monolayer culture [26, 85].

We used three different inhibitors for GSK-3β, SB216763, GSK-3β inhibitor XII and GSK-3 inhibitor XVI. All those inhibitors completely blocked EF-induced GA polarization, but did not block the phosphorylation of GSK-3β induced by EF (Figs. 5d and 6). Inhibition of PKC with GF109203X prevented GSK-3β phosphorylation induced by EF, but only partially blocked the EF-induced GA polarization (Figs. 5c and 6). Those results suggest that the EF-induced GA polarization was mediated by GSK-3β, but not through phosphorylation of GSK-3β. When GSK-3β was inhibited with SB216763, GSK-3β inhibitor XII, GSK-3 inhibitor XVI or PKC was inhibited with GF109203X, the EF-induced directedness of cell migration was reduced to ~50%, respectively (Fig. 7b).

In conclusion, an EF well within the physiological range (~0.3 V/cm) directs GA polarization and cell migration. Most importantly, GSK-3β and PKC are two important molecules for EF-induced establishment of cell polarization and directedness of cells migration. PKC may play a role to initiate electrotaxis, while GSK-3β in optimizing the electrotaxis via polarizing GA. Our data suggest that the GSK-3β may play an essential signaling role in physiological EF-induced GA polarization and optimal cell migration.

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