Possible involvement of calcium channels and plasma membrane receptors on Staurosporine-induced neurite outgrowth

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ABSTRACT

Staurosporine as a protein kinases inhibitor induced cell death or neurite outgrowth in PC12 cells. We investigated the involvement of calcium channel and plasma membrane receptors on staurosporine inducing neurite outgrowth in PC12 cells. PC12 cells were preincubated with NMDA receptor inhibitors (1.8 mM ketamine and 1μM MK801, treatment 1) or L-Type Calcium channels (100 μM nifedipine and 100 μM flavoxate hydrochloride, treatment 2) or calcium-calmoduline kinases (10 μM trifluoperazine, treatment 3) and nifedipine, MK801, flavoxate hydrochloride and ketamine (treatment 4) or without pretreatments (control). Then, the cells were cultured in RPMI culture medium containing 21μM staurosporine for induction of neurite outgrowth. The percentage of Cell cytotoxicity and apoptotic index was assessed. Total neurite length (TNL) and fraction of cell differentiation were assessed. After 24h, the percentage of cell cytotoxicity were increased in treatments 1, 2 and 4 compared with control (p<0.05). After 6h, apoptotic index was similar between all treatments. After 12h, apoptotic index were increased in treatment 3 compared with control (p<0.05). After 24h, apoptotic index were increased in treatments 1, 2 and 4 compared with control (p<0.05). TNL were decreased in treatments 1, 2 and 4 compared with control in different times of assessment (6, 12 and 24 h) (p<0.05). The fraction of cell differentiation were decreased in treatments 1, 2 and 4 compared with control (p<0.05). It can be concluded that the possible involvement of L-type calcium channel and the N-methyl D-aspartate receptor on staurosporine-induced neurite outgrowth process in PC12 cells.

INTRODUCTION

Staurosporine (STS) as a protein kinase inhibitor [1, 2] has dual effects on neuronal cells; induction of cell death and cell differentiation. STS induces apoptosis in high concentrations (μM) [3, 4] and neuronal differentiation in low concentrations (nM) by neurite extension in several types of cells [2, 3, 5, 6]. Although the detailed mechanism of STS action as a neurogenic morphogen remains unclear, it seems that it is associated with the inhibition of some protein kinases which may contribute to neurite outgrowth [7]. In previous studies it has been determined that STS can inhibit PKA, Ca2+/ calmodulin-dependent kinase II, cyclin dependent kinases, ion channels (Kv1.3, L-type Ca2+ channel, voltage-gated K+ channel) in myocyte [8-12]. Although, the role of STS in the inhibition of protein kinases during neurite outgrowth was clear but its function on plasma membrane calcium channels and receptors remains to be fully known [10,11]. Calcium plays an important role in the regulating a great variety of neuronal processes such as neuronal cell differentiation. In most neurons, multiple mechanisms exist whereby increases in intracellular calcium concentration may occur including for example in calcium entry through N-methyl-D-Aspartate (NMDA) glutamate receptors and various voltage-gated calcium channels such as L-type calcium channels (LTCC), as well as in the release of calcium from intracellular stores [13-15]. Calcium influx through LTCCs is particularly effective in neuronal migration, activation of transcription factors (e.g CREB), changes in gene expression that underlie plasticity and adaptive neuronal responses (e.g c-fos) [1, 16-22]. Although STS induced increasing of intracellular calcium in treated cells, its effect on plasma membrane and calcium channels and receptors located in the plasma membrane during neuronal differentiation and neurite outgrowth are not well known. In this study we aimed to determine whether plasma membrane calcium channels and receptors involves in staurosporine-induced neurite outgrowth.

MATERIALS AND METHODS

Cell culture

PC12 cells were cultured in complete culture medium containing RPMI1640 culture medium (Gibco), supplemented
with 0.2% bovine serum albumin (BSA, Gibco), 1% NEAA (Sigma), 2 mM L-glutamine (Sigma), 100 IU/ml penicillin and 100 μg/ml streptomycin (Sigma) in 10-cm tissue culture dishes. The cultures were incubated at 37°C in a humidified incubator containing 95% air and 5% CO₂. Culture medium was replaced every 2 days. When cell cultures reached to 80% confluency, they were trypsinated using trypsin-EDTA 0.25% (Sigma) and the cells were subcultured at a density of 104 cells/well in 24-well culture plates.

**Cell treatment**

One day after plating PC12 cells, cells were washed with phosphate buffer saline (PBS), pH 7.4. For inhibition of L-Type Calcium channel, NMDA receptor and CaM kinase, cells were preincubated with, adding 1.8 mM ketamine and 140 μM MK801, 15 min (treatment 1), 100 μM nifedipine and 100 μM flavoxate hydrochloride, 30 min (treatment 2) and 10 μM trifluoperazine, 30 min (treatment 3). In our experiment, we combined ketamine, MK801, flavoxate hydrochloride and nifedipine for inhibition of L-Type Calcium channels and NMDA receptors (treatment 4). Then, cells were cultured in differentiation medium containing complete culture medium supplemented with 214 nM staurosporine for 24h. PC12 cells were cultured in differentiation medium without inhibitor preincubation seems as control group. The cells were placed in the incubator at 37°C with 5% CO₂.

**Cell cytotoxicity measurement**

Cell cytotoxicity was quantified by measuring the release of lactate dehydrogenase (LDH) from damaged or destroyed cells into the medium. Cytotoxicity was measured with LDH Cytoxicity Detection Kit (Roche, Germany). This kit detects LDH release from dead cells. Therefore, increase of LDH activity in each treatment show that the treatment solution has further dead cells or cytotoxicity effects on PC12 cells. Cells were plated in 96 well culture plates with 104 cells/mL density for 12h. Then cells were pretreated in different treatments for certain time. Then, cells were cultured by differentiation medium for 24h. The percentage of cytotoxicity was measured by protocol of company; colorimetry of LDH activity measured by calculated the absorbance of samples at 490 or 492 nm using an ELISA Reader (EL800; USA). The references wavelength should be more than 600 nm. All experiments were replicated independently at least 3 times. Within each experiment, we replicated each condition 4 times.

**Quantification of cell death incidence**

Hoechst / PI nuclear staining was carried out as previously described [23]. Briefly, cells were plated in 24 well culture plates with 104 cells/mL density for 12h. Then cells were pre-treated in different treatment mediums for certain time. These were grown for a range of times in differentiation medium (6, 12 and 24h). Then cells were incubated for 15 min at 37°C with Hoechst 33342 dye (10 mg/ml in PBS), washed twice in PBS. PI (50 mg/ml in PBS) was added just before microscopy. Cells were visualized using an inverted-fluorescence microscope (Olympus IX-71, Japan). Nuclear morphology was scored as follows: 1, viable cells had blue-stained nuclei with smooth appearance; 2, viable apoptotic cells had blue-stained nuclei with multiple bright specks of condensed chromatin; 3, non-viable apoptotic cells had red-stained nuclei with either multiple bright specks of fragmented chromatin or one or more spheres of condensed chromatin (significantly more compact than normal nuclei); 4, non-viable necrotic cells had red-stained, smooth and homogeneous nuclei that were about the same size as normal (control) nuclei. The apoptotic index were calculated by the fraction of numbers of apoptotic cells on the total cell count in 100 (300 cells), respectively. All experiments were replicated independently at least 3 times. Within each experiment, we replicated each condition 4 times.

**Measurement of total neurite length**

Measurement of total neurite length was conducted as reported by previous study [24]. The assay is based on the measurement of total neurite length. Total neurite length (length of largest neurite on 100 cells) was assessed. Cells were plated in 24 well culture plates with 104 cells/well density for 12h. Then cells were pretreated in different treatments for certain time. These were then grown for a range of times at differentiation medium (6, 12 and 24h), fixed, and the morphology assessed by an inverted microscope (Olympus IX-71, Japan). Digital photos were taken of random fields of neurons derived from the treatments. Total neurite length was measured (Motic software; Ver. 2). All experiments were replicated independently at least 3 times. Within each experiment, we replicated each condition 4 times.

**The fraction of cell differentiation assessment (f (%))**

Fraction of cell differentiation was carried out as previous study [25]. PC12 cells were plated at a density of 2×104 cells/well on 24 well plates. Cells were pretreatment with different treatment mediums. These were then grown for a range of times at differentiation medium (6, 12 and 24 h), fixed, and the morphology microscopically assessed (Motic software; Ver. 2). The fraction of cell differentiation was evaluated under an inverted microscope by the fraction of neurite-bearing cells. The fraction of cell differentiation was the fraction of numbers of neurite-bearing cells with at last one neurite longer than the cell body diameter on the total cell count (300 cells). All experiments were replicated independently at least 3 times. Within each experiment, we replicated each condition 4 times.
Statistical analysis
Data were expressed as Mean ± SEM. All calculations were performed by SPSS (version 16; SPSS Inc.). The differences in the percentage of cytotoxicity, incidence of apoptotic index, total neurite length and fraction of cell differentiation, in PC12 cells between treatments were analyzed using t-test at significant level (p<0.05).

RESULTS

Cell cytotoxicity
The percentage of cytotoxicity of inhibitors in PC12 cells cultured in culture medium containing 214 nM staurosporine was assessed by evaluation of the lactate dehydrogenase activity. In PC12 cells the percentage of cytotoxicity were increased in treatments 1, 2 and 4 (36% ± 2%, 32% ± 2% and 46% ± 3%, respectively) compared with control (20% ± 2%), (p<0.05). The percentage of cytotoxicity in treatment 3 (21% ± 3%) were decreased compared with treatments 1, 2 and 4 (p<0.05) and was similar to control (Figure 1).

Effects of inhibitors on apoptosis index
The evaluation of apoptotic index of inhibitors for PC12 cells cultured in culture medium containing 214 nM staurosporine was assessed by PI/Hoechst fluorescence staining. After 6h, the apoptotic index were increased in treatments 1 and 2 (18% ± 3% and 19% ± 2%); respectively and were similar in treatment 3 (15% ± 3%) compared with control cells (16% ± 2%) but these differences were not significant. The apoptosis index in treatment 4 (26% ± 4%) was increased compared with control and treatments 1-3 (p<0.05). After 12 h, the apoptosis index were increased in treatments 1 (31% ± 2%), 2 (28% ± 2%) and 4 (43% ± 4%) compared with control (21% ± 3%) (p<0.05). The apoptotic index in treatment 3 (22% ± 4%) was decreased compared with treatments 1, 2 and 4 (p<0.05) and was similar to control. After 24 h, the apoptosis index were increased in treatments 1, 2 and 4 (42% ± 4%, 38% ± 3% and 55% ± 5%, respectively) compared with control (23% ± 4%) (p<0.05). The apoptosis index in treatment 3 was increased (25% ± 3%) compared with control but this difference was not significant (Figures 2; A and B).

Neurite outgrowth measurement
The average of total neurite length for PC12 cells was assessed. The total neurite length (TNL) was calculated. Af-
After inhibitors preincubation, TNL significantly were decreased compared with control cells. After 5h, TNL were decreased in treatments 1, 2 and 4 (123 ± 0.67, 118 ± 0.72 and 98 ± 0.83, respectively) compared with control cells (142 ± 0.89) (p<0.05). TNL in treatment 3 (141 ± 0.64) was similar to control. After 12h, TNL were decreased in treatments 1, 2 and 4 (102 ± 0.92, 98 ± 0.87 and 83 ± 0.93, respectively) compared with control cells (128 ± 0.94) (p<0.05). TNL in treatment 3 (125 ± 0.85) was similar to control. After 12h, TNL were decreased in treatments 1, 2 and 4 (76 ± 0.85, 71 ± 0.88 and 65 ± 0.82, respectively) compared with control cells (108 ± 0.81) (p<0.05). TNL in treatment 3 (102 ± 0.92) was similar to control (Figures 3 and 4).

**Fraction of cell differentiation assessment**

The evaluation of the fraction of cell differentiation of inhibitors for PC12 cells cultured in culture medium containing 214nM staurosporine was assessed. After 6 h, f (%) were not significantly decreased in treatments 1, 2 and 4 (98% ± 1%, 98% ± 0.7% and 96% ± 1%, respectively) compared with control (100%). f (%) in treatment 3 (100%) was similar to control. After 12 h, The fraction of cell differentiation f (%) was decreased in treatment 4 (92% ± 1.2%) (p<0.05). f (%) were not significantly decreased in treatments 1 and 2 (95% ± 2% and 94% ± 2%) compared with control (100%) (p<0.05). f (%) in treatment 3 (p<0.05) was similar to control cells. After 24h, f (%) were decreased in treatments 1, 2 and 4 (87% ± 3%, 78% ± 3% and 63% ± 5%, respectively) compared with control cells (98% ± 2%), (p<0.05). f (%) in treatment 3 was similar to control (Figure 5).

**DISCUSSION**

The current study investigated the involvement of calcium channel and plasma membrane receptors on staurosporine...
inducing neurite outgrowth in PC12 cells. In this work, we used PC12 cells as the best cell model for study of effect of materials on neurite outgrowth [25]. PC12, a neuron-like cell line, expresses voltage-dependent Ca channels appear to dihydropyridine-sensitive voltage-dependent Ca channels demonstrable by different techniques [26, 27]. Stauroporine was employed as a strong inducer of neurite outgrowth with inhibition of protein kinases in vitro model. The results obtained in this study showed that nifedipine and ketamine could effectively inhibit neurite outgrowth induced by staurosporine and increase cell death incidence in PC12 cells. We observed that when cells were preincubated with nifedipine and flavoxate hydrochloride or ketamine and MK801, they dramatically suppressed the neurite outgrowth and increased cell death and cytotoxicity in PC12 cells. Meanwhile, preincubation with ketamine and MK801 together with nifedipine and flavoxate hydrochloride result in powerful inhibition of neurite outgrowth and induce cell death in PC12 cells. It could be suggested that the possible involvement of voltage dependent calcium channels and NMDA receptors on staurosporine-calcium dependent signal transduction. Meanwhile, PC12 application of trifluoperazine does not the same age dependent calcium channels and NMDA receptors result in regulation of neurite outgrowth through the Ca²⁺ influx via extracellular Ca²⁺ -sensing receptor localized to neurons or signals derived from intermediate or final target tissues [28]. Previously, it has been shown that external Ca²⁺ evoke the signal transduction through the Ca²⁺ influx via extracellular Ca²⁺-sensing receptor localized to neurons and their nerve terminals [29]. It demonstrated that neurite outgrowth of PC12 is induced via the Ca²⁺-signal transduction pathway by the Ca²⁺ influxes through channels [30]. On the other hand, recent study showed that staurosporine leads to intracellular calcium overload, which induce apoptosis in PC12 cells [31]. In the recent study, showed that staurosporine caused a large increase in [Ca²⁺]c even after the depletion of Ca²⁺ from the ER, the IP₃-sensitive Ca²⁺ store, in the absence of perfusate Ca²⁺. This result indicates that IP₃-insensitive, non-ER compartments are responsible for the staurosporine-induced [Ca²⁺]c increase in rat submandibular acinar cells [32]. We reported previously that Stauroporine use extracellular calcium stores tend to increase intracellular calcium concentration [33]. In addition, previously, it is known that cytosolic Ca²⁺ increase caused by staurosporine that mobilize Ca²⁺ from different sources might cause apoptosis in astrocytes [34]. Ca²⁺ in DDTIMF-2 smooth muscle cells by influx but also by intracellular mobilization from thapsigargin-sensitive and -insensitive Ca²⁺ stores. Furthermore, the high local Ca²⁺ gradient just under the plasma membrane, which can be preserved over long periods of time in Ca²⁺-free medium despite the presence of EGTA, indicates that the efflux mechanism is also affected [35]. The stores of Ca²⁺ ion entry from extracellular into intracellular during staurosporine-induced neurite outgrowth is still not completely understood. Many studies in different cells showed that staurosporine result in an increase cytosolic calcium concentration and induction of apoptosis in NGF-differentiated cells [36, 37]. In another study showed that the rate of apoptotic cells is greater in differentiated cells than undifferentiated cells [28]. Different study showed that neurotrophins factors like NGF result in increase of mRNA encoding of calcium channels like voltage-dependent calcium channels and glutamate-sensitive ion channels like NMDA [38-42]. It has shown that compared with undifferentiated cells maybe activation of calcium channels and plasma membrane receptors by staurosporine lead to increase of staurosporine-induced apoptosis in differentiated cells. If true, these receptors and channels play important role in increasing intracellular calcium concentration during staurosporine-induced cell differentiation in PC12 cells. Meanwhile, We suggest it possible that staurosporine by a protein kinase-independent mechanism (PKC, PKA and CaMKs) by activation of plasma membrane Ca²⁺ channels lead to enhance of neurite outgrowth and increases cell viability and fraction of cell differentiation in PC12 cells.

CONCLUSION

According to the results of present study, application of stauroporine with activation of calcium channels may lead to enhance of neurite outgrowth and have effects on neuronal cell differentiation in PC12 cells. However, more key receptors and enzymes need to be investigated in these effects.

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DECLARATION OF INTEREST

There is no conflict of interest.
REFERENCES


