COLLAGEN VI PROTECTS AGAINST NEURONAL APOPTOSIS ELICITED BY ULTRAVIOLET IRRADIATION VIA AN AKT/PHOSPHATIDYLINOSITOL 3-KINASE SIGNALING PATHWAY

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Abstract—Collagen VI, one of the extracellular matrix proteins, has been implicated in regulating cell proliferation and reducing apoptosis in several different systems. However, the role of collagen VI in the central nervous system remains unclear. In this manuscript, we demonstrated that upon ultraviolet (UV) irradiation, mouse primary hippocampal neurons specifically up-regulate the expression of Col6α1, Col6α2, and Col6α3 mRNA and secreted collagen VI protein. Augmentation of collagen VI mRNA and protein after UV irradiation may have a neuroprotective role as suggested by the fact that extracellularly supplying soluble collagen VI protein, but not other collagen proteins, reduced UV-induced DNA damage, mitochondria dysfunction, and neurite shrinkage. We also tried to determine the signaling molecules that mediate the protective effect of collagen VI via Western blot and inhibitor analysis. After collagen VI treatment, UV-irradiated neurons increased phosphorylation of Akt and decreased phosphorylation of JNK. Inhibiting Akt/Phosphatidylinositol 3-kinases (PI3K) pathway diminished the protective effect of collagen VI. Our study suggested a potential protective mechanism by which neurons up-regulate collagen VI production under stress conditions to activate Akt/PI3K anti-apoptotic signaling pathway. © 2011 IBRO. Published by Elsevier Ltd. All rights reserved.

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Appearance of the proper type of extracellular matrix (ECM) proteins not only supports tissue structure, but also plays essential roles in the proliferation, differentiation, and survival for most types of cells (Sherman-Baust et al., 2003; Nelson and Bissell, 2006; Cheresh and Stupack, 2008). In the central nervous system, the ECM proteins derived from neurons and glia play a variety range of roles such as supporting cells, regulating cellular signaling, coordinating synaptogenesis, and synaptic activity (Dityatev and Schachner, 2003; Washbourne et al., 2004; Dityatev and Fellin, 2008; Lee et al., 2008). Collagens are part of ECM proteins that play active roles during the neurodevelopment and neurodegeneration (Fox, 2008; Hubert et al., 2009). However, few studies have dissected the consequent molecular mechanism of collagens in response to the stress conditions in the central nervous system.

Collagen VI is one of the 28 collagen protein families that composed of six subunits (Col6α1~6) encoded by separate genes (Col6α1~6). Col6α1, Col6α2, and Col6α3 subunits assemble in 1:1:1 ratio to form monomers and further secrete to extracellular space where they assemble into microfibrils (Brüns et al., 1986; Schreier et al., 1987; Kiely et al., 1992). Col6α4, Col6α5, and Col6α6 are newly identified as members in collagen VI family, but their structural and functional roles remain unclear ( Fitzgerald et al., 2008). Similar to many other molecules in ECM, collagen VI regulates cell survival and promotes tumor progression via binding to cell surface proteins and alternating downstream signaling cascades (Kuo et al., 1997; Howell and Doane, 1998; Ruehl et al., 1999; Milner and Campbell, 2002; Iyengar et al., 2005). The mutations on Col6 genes result in muscular diseases Bethlem myopathy and Ulrich congenital muscular dystrophy (Lampe and Bushby, 2005; Lampe et al., 2008). However, the role of collagen VI in the central nervous system is yet little understood.

Apoposis is the process of programmed cell death, which involves a series of morphological changes, including cell detachment, cell shrinkage, mitochondria leakage, chromatin condensation, and DNA fragmentation (Kannazawa, 2001). This process is controlled by the balance between multiple pro-apoptotic and anti-apoptotic signaling pathways, which may originate either extracellularly or intracellularly. Two of the key signaling molecules regulating this balance are Akt/phosphoinositide 3-kinase (PI3K) and Jun N-terminal kinase (JNK) cascades. Activation of Akt through phosphorylation has an anti-apoptotic role in a variety of tissue culture models against the withdrawal of growth factors, ultraviolet (UV) irradiation, matrix detachment, cell cycle disturbance, and DNA damages (Datta et al., 1997, 1999; Khwaja et al., 1997; Brazil et al., 2002; Burke, 2007; Parcellier et al., 2008). In contrast, activation of JNK through phosphorylation has a pro-apoptotic role under many different types of stresses, such as UV irradiation (Dénard et al., 1994; Caridad and Karin, 1996; Ro-
sette and Karin, 1996; Davis, 2000; Dhanasekaran and Reddy, 2008). Inhibition of JNK activity results in attenuation of apoptosis in UV irradiated cells (Zhang et al., 2008; Kim et al., 2009).

To determine the role of ECM proteins on neurons under stress conditions, we surveyed the level of known ECM proteins after UV irradiation. In this report, we described that increased levels of Col6a1-3 mRNA and collagen VI protein in neurons after UV irradiation. Soluble collagen VI may have a functional role in neuronal survival through the activating Akt/Pi3K and inhibiting JNK signaling pathway.

**EXPERIMENTAL PROCEDURES**

**Primary neuronal cultures**

The pregnant C57BL/6JNarl mice were purchased from National Laboratory Animal Center (NLAC, Taipei, Taiwan) with the approval of the Institutional Animal Care and Use Committee at National Yang Ming University. Primary hippocampal neuronal cultures, in which more than 95% of cells were immune reactive for microtubule-associated protein-2 (MAP-2, see Fig. 3), were generated from embryonic day 19 or post-natal day 0 mice as described (Chu et al., 2009). After growing 8 days in vitro, primary neurons were treated with collagen VI (354261, BD Biosciences, USA) or collagen IV (354245, BD Biosciences, Bedford, MA, USA) 2 h before UV irradiation. UV irradiation was carried out using UVC lamp emitting maximally at 253.7 nm (Philip, Eindhoven, Netherlands) for 20 min in collagen VI containing medium. UV irradiance was measured using a UVX radiometer UVX-25 (Ultra-violet Products, Rockford, IL, USA) before each use.

**Quantitative polymerase chain reaction (qPCR)**

Primary neuronal cultures were grown on six-well plates at 4 × 10^6 cells/well. After treatments, neurons were harvested with 0.5% trypsin and centrifuged for 20 s at 4700 × g, and their RNA was isolated using the Total RNA Mini Kit (RB050, Geneaid, Taiwan). RNA concentration was determined by NanoDrop Spectrophotometer (ND-1000, Thermo, Rockford, IL, USA). Total RNA was reverse transcribed with oligo(dT) primers and Moloney Murine leukemia virus reverse transcriptase (RT80110K, Epicentre, Madison, WI, USA). The expression level of different mRNA and collagen VI genes (Col6a1-3, forward, 5'-TACCCTACCCAGGACGAA-3', reverse, 5'-TCAGCTGGACTGGGCGTAGC-3') was determined by SYBR Green dye (04673484001, Roche, Indianapolis, IN, USA). The following primer sets (forward, 5'-ATGGTGTTGGAGCTTATTCTTCG-3', reverse, 5'-CAAATGCTTACAGCTTTTGGC-3') were used: mouse Col6a2 (forward, 5'-CATCTCACCCCAGGAGCAGGAA-3', reverse, 5'-TACACGTTGACTGGGCAGTCGG-3'); mouse Col6a3 (forward, 5'-AACCCTCCACATACTGCTAATTC-3', reverse, 5'-TGCCCTGTGGATC-TGTCTCTCACTAGTGTAATG-3').

**Immunoblotting**

Primary neuronal cultures were grown on six-well plates at 4 × 10^6 cells/well. To detect collagen VI protein level, culture media were collected at 0, 2, and 4 h after UV irradiation. For dot blot analysis, 20 μl of culture media was spotted on pre-soaked nitrocellulose membrane five times and probed with anti-collagen VI antibody (ab6588, Abcam, Cambridge, UK). To detect Akt and JNK levels, neurons were collected at 30 min after irradiation and lysed in ice-cold radioimmunoprecipitation assay (RIPA) buffer (50 mM Tris pH 7.5, 150 mM NaCl, 10 mM EDTA, 1% NP-40, 0.1% SDS) containing phosphatase inhibitor (04906845001, Roche, Indianapolis, IN, USA) and protease inhibitor (04693116001, Roche, USA). The cell lysates were agitated at 4 °C for 30 min and then centrifuged at 16100 × g at 4 °C for 10 min. The protein concentration was measured by Bradford assay (PAK500, Strong Biotech Corp, Taipei, Taiwan). Protein samples (66.5 μg) were heated at 95 °C for 10 min and separated in 10% SDS-PAGE. Western blotting was performed using anti-phospho-Akt at Ser-473 (9271, Cell signaling, Beverly, MA, USA), anti-total Akt (9272, Cell Signaling, Beverly, MA, USA), anti-phospho-JNK at Thr183/Tyr185 (9251, Cell signaling, USA), anti-total Akt (9252, Cell Signaling, Beverly, MA, USA), and anti-actin (MAB1501, Millipore, Billerica, MA, USA) antibodies. Immunoblot signals were developed by chemiluminescent HRP substrate (WBKLS0500, Millipore, Billerica, MA, USA) and detected with the Fujifilm Luminescence/Fluorescence Imaging System (LAS-4000, Fujifilm, Tokyo, Japan). Multi Gauge v3.0 was used for image acquisition and data analysis.

**Immunocytochemistry**

Primary neurons were grown on 12 mm coverslips in 24-well plates at 2.5 × 10^5 cells/well. Immediately after UV and collagen VI treatments, neurons were fixed in 4% paraformaldehyde for 60 min at room temperature. Fixed neurons were permeabilized with 0.1% Triton X-100 in phosphate buffered saline (PBS, 137 mM NaCl, 2.7 mM KCl, 10 mM Na_2PO_4, 2 mM KH_2PO_4, pH of 7.4) for 10 min, blocked with 10% fetal bovine serum (04001-1A, Biological Inc., Taipei, Taiwan), and incubated overnight at 4 °C with the primary antibodies, including monoclonal mouse anti-Collagen VI (MAB378, Millipore, Billerica, MA, USA), and polyclonal rabbit anti-glial fibrillary acidic protein (GFP, Z0334, DAKO, Glostrup, Denmark). Neurons were then rinsed with cold PBS and incubated with either Texas-Red-conjugated goat anti-mouse IgG (10007, AbD Serotec, Kidlington, UK) or FITC-conjugated goat anti-rabbit IgG (AP132F, Millipore, Billerica, MA, USA) for 1 h. The coverslips were mounted with VECTASHIELD® plus 4'-6-Diamidino-2-phenylindole (DAPI, H-1200, Vector Lab Inc., Burlingame, CA, USA). Omission of primary antibody served as negative controls. Fluorescent images of cells were captured on a CCD camera (DP50) mounted on an Olympus fluorescence microscope (BX-52, Olympus, Tokyo, Japan) equipped with a mercury arc lamp.

**Terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) assay**

TUNEL assay was performed using the DeadEnd™ Fluorometric TUNEL System (G3250, Promega, Madison, WI, USA). Primary neurons were cultured and treated in the same condition as immunocytochemistry experiments. The TUNEL staining was performed as described in Promega manual. The number of TUNEL-positive cells were counted in different 10× fields per sample using ImageJ software (free downloaded from NIH website: http://rsbweb.nih.gov/ij/).

**MTT assay**

Primary neuronal cultures were seeded in 96-well plates (10^5 cells/well). 2 h after all treatments, 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT, 298-93-1, Bio Basic Inc, Taipei, Taiwan) was added to the medium (0.5 mg/ml) and incubated at 37 °C. After 4 h, 100 μl of isyfer buffer containing 10% SDS and 20 mM HCl was applied to each well and incubated at 37 °C overnight to dissolve the formazan crystals. The absorbance at wavelength 570 nm in each well was measured on a
microenzyme-linked immunosorbent assay (ELISA) reader (TECAN, Männedorf, Switzerland).

**NeurotoxQ**

NeurotoxQ was implemented as an ImageJ macro. In order to perform all necessary operations, an additional ImageJ plugins “particle remover” (http://rsb.info.nih.gov/j/plugins/particle-remover.html) was incorporated. NeurotoxQ pre-processes the image using edge detection and background correction with user-defined parameters. To detect edges, the original image was subtracted from the image that had been smoothed by a Gaussian kernel. Uneven background was corrected using the “subtract background” function with a rolling ball radius of 50 pixels. Neuronal cell bodies were extracted from the pre-processed image using an open operation (minimum filter followed by maximum filter). The image with extracted neuronal cell bodies was binarized for quantification. The pre-processed image was also binarized, cell debris and small particles were removed via user-defined size criterion. The resulting “clean” image was skeletonized to reduce all objects into single pixel wide skeletons. Neuronal cell bodies were subtracted from the skeleton image to obtain the neurite length. In addition, neuronal cell bodies were subtracted from the “clean” image to obtain the neurite area. The color combined images were generated by combining the binarized neurite length image, the original image, and the binarized neuronal cell body image. NeurotoxQ was available for free download at http://life.nctu.edu.tw/~microtubule/macos.html (Username: neurotoxQ, Password: microtubule).

**Statistics**

All results were displayed as mean±SEM. Statistical analyses were carried out with SPSS (IBM SPSS, Chicago, IL, USA). Data were analyzed using one-way ANOVA and Student-Newman-Keuls post-test. P-values lower than 0.05 were considered significant.

**RESULTS**

UV irradiation up-regulates collagen VI gene expression after UV irradiation

The composition of ECM proteins could play a detrimental role in controlling pro- or anti-apoptotic signaling pathways under stress conditions. This study was aimed to identify the alteration of ECM genes expression after UV irradiation in neurons, and to determine the role of these genes in apoptosis. After 192 J/m² UV irradiation, mouse primary hippocampal neurons were harvested after 30 min for measuring mRNA levels by qPCR. One of the most up-regulated mRNA encoding ECM proteins was the Col6 gene family. Relative to no UV irradiated control group, expression level of mRNA encoding the α1 subunit (Col6a1) had 3.0-fold increase; the mRNA encoding α2 subunit (Col6a2) had 3.9-fold increase; the mRNA encoding α3 subunit (Col6a3) had 9.7-fold increase (P<0.05 for all three genes in UV vs. no UV groups (Fig. 1A)). However, the expression levels of mRNA encoding other collagen genes, such as Col1a1, Col4a1, did not have significant difference after UV irradiation.

Because the increase of mRNA level does not necessary lead to the increase in protein level, we therefore also measured the amount of collagen VI protein in culture medium at 0, 2, and 4 h after UV irradiation. Relative to 0 h group, the level of collagen VI protein increased 2.2-fold at 2 h and increased three-fold at 4 h after irradiation (both P<0.05, Fig. 1B). This expression analysis revealed that Col6 mRNA and collagen VI protein are quickly up-regulated after UV irradiation.

**Collagen VI protects neurons against UV irradiation**

The up-regulated collagen VI may serve as either an anti-apoptotic factor to protect neurons against UV, or a pro-apoptotic mediator to activate UV-induced cell death. To elucidate the role of collagen VI, we investigated if exogenous applying extra soluble collagen VI protein in the medium can make neurons more resistant or sensitive to UV irradiation. Primary neuronal cultures were pre-treated with media containing 0, 25 or 50 nM of purified human collagen VI protein for 2 h, and then exposed to 192 J/m² UV. The appearance of DNA strand fragmentation labeling by TUNEL assay is one of the hallmarks of apoptotic cells. We therefore labeled treated cells with TUNEL and calculated the ratio of TUNEL positive neurons as: (number of TUNEL positive cells (pink)/number of DAPI stained cells (blue))×100%. The ratio of TUNEL positive neurons were five-folds higher in UV irradiated culture than no UV control culture (P<0.05; Fig. 2A, B, F). However, under the same dose of UV, the ratio of TUNEL...
positive neurons decreased 70% in 25 nM collagen VI-treated cultures, and decreased 80% in 50 nM collagen VI-treated cultures (both $P<0.05$ vs. UV-irradiated group (Fig. 2C, D, F). Because pre-treating neurons with soluble collagen VI make neurons more resistant to UV induced apoptosis, collagen VI may have a protective role in neurons.

In addition to DNA breakage, we also investigated whether collagen VI treatment can rescue the mitochondria metabolic dysfunction induced by UV irradiation. We determined the mitochondria activity of primary neurons by MTT assay and compared the mean OD570 absorbance value to represent the mitochondria activity of live cells. Similar to the result from TUNEL assay, we found that mitochondria activities were impaired after UV irradiation ($P<0.01$ vs. no UV group) but could be rescued by collagen VI pre-treatment (Fig. 2G, $P<0.05$).

One of the possible explanations for the protective effect of collagen VI is that the presence of extra 50 nM of protein in the culture medium may block UV irradiation. To test this possibility, 50 nM of soluble human collagen IV protein was also added into medium as control. In contrast to collagen VI treatment, pre-treating neurons with collagen IV failed to res-
cule DNA damage and mitochondria dysfunction induced by UV irradiation (Fig. 2F, G). To further confirm the essential role of collagen VI, we also added 250 nM of anti-collagen VI antibody to naturalize the effect of both exogenous and endogenous collagen VI. The protective effect of 50 nM exogenous collagen VI was diminished in the presence of specific antibody in both TUNEL and MTT assays (Fig. 2F, G). After naturalizing endogenous collagen VI, these cells had a significantly worse mitochondria dysfunction in MTT assay (Fig. 2G, \( P < 0.05 \) vs. UV group), but only a trend toward more DNA damage in TUNEL assay (Fig. 2E, F, \( P = 0.07 \) vs. UV group). These results suggested that this protective effect is collagen VI specific.

During the apoptosis process, shrinkage of neurites and somas are commonly seen and easy to detect (Kanazawa, 2001). To quantify these morphological alterations, nevertheless, manual measurement is extremely time consuming and biased. We therefore developed an ImageJ-based morphology quantification algorithm, NeurotoxQ, which provides automatic measurements of the following neuronal morphological parameters: (1) the average length of the neurite segments of each neuron, and (2) the average size of each neuron cell body (soma), (3) the number of neurons.

To examine whether collagen VI treatments can rescue the alteration of cell number and morphology after UV irradiation, primary neuronal cultures were pre-treated with 50 nM of collagen VI for 2 h before UV irradiation. These cells were immunolabeled with neuron specific marker MAP-2 (red), and their nuclei were labeled with a nuclei dye DAPI (blue) (Fig. 3A–D). After UV irradiation, the average neurite length per neuron was >50% shorter than no UV control (\( P < 0.01 \), Fig. 3A, B, E). However, under the same dosage of UV, the average neurite length was 1.5- and 1.7-fold longer in 25 and 50 nM collagen VI-treated cultures (both \( P < 0.05 \) vs. after UV-irradiated group, Fig. 3C, D, E). Nevertheless, there was no significant difference in the average size of soma with or without UV irradiation and collagen VI treatment (Fig. 3F). In addition, the number of MAP positive neurons per image field did not have significant difference (Fig. 3G). Therefore, the average length of neurites, but not the size of soma, was consistent with the data from TUNEL and MTT assays indicating that pre-treating cultures with collagen VI rescues neurons from UV-induced cell damage.

To evaluate our algorithm, we compared neurite length determined manually using NeuronJ with that obtained using NeurotoxQ. Sixteen images of mouse hippocampal neurons were used to evaluate NeurotoxQ (Fig. 4A). NeurotoxQ-produced neurite tracings were highly correlated (Pearson’s correlation coefficients \( R = 0.95 \)) (Fig. 4B) and statistically indistinguishable (\( P = 0.2275 \) from the two-tailed Student’s \( t \)-test) from manual tracings.

**Collagen VI alters apoptotic signaling pathways**

Because soluble collagen VI made neurons more resistant to UV irradiation, we speculated that collagen VI may bind to unidentified cell surface protein and alter survival-related signaling pathways. In most cell types, the balance of pro-apoptotic and anti-apoptotic signals determines cell fate under stress conditions. To decipher the molecular paths that mediate the protective effect of collagen VI, we used Western blot to monitor phosphorylation states of Akt and JNK, whose activities are critical for regulating cell survival/death upon UV irradiation. After UV irradiation, the ratio of phosphorylated Akt protein was not significantly altered in primary neurons. However, under the same dosage of UV, the ratio of phosphorylated Akt increased 1.6-folds with 25 nM collagen VI treatment, and increased 1.8-folds with 50 nM collagen VI treatment (both \( P < 0.05 \) vs. UV irradiated group, Fig. 5A, B). In contrast, the ratio of phosphorylated JNK protein increased 2.6-fold after UV irradiation (\( P < 0.05 \) vs. no UV control, Fig. 5A, C). Under the same dosage of UV, ratio of phosphorylated JNK decreased 49% in 25 nM collagen VI treated cells, and decreased 60% in 50 nM collagen VI treated cells (both \( P < 0.05 \) vs. UV-irradiated group, Fig. 5A, C). In summary, soluble collagen VI could stimulate Akt and inhibit JNK phosphorylation in UV-irradiated neurons.

Based on the Western blot result and well-established anti-apoptotic role of Akt, we hypothesized that collagen VI may stimulate Akt/PI3K signaling pathway to protect neurons against UV irradiation. To further confirm the role of Akt/PI3K, primary neuronal cultures were co-treated with 50 nM of collagen VI together with 50 \( \mu \)M Akt/PI3K inhibitor LY294002 for 2 h before UV irradiation. Treating neurons with collagen VI alone increased the ratio of phosphorylated Akt, but treating neurons with collagen VI and LY294002 together lowered the ratio of phosphorylated Akt to untreated level (Fig. 6F). We determined the degree of damage under these treatments by neurite length alteration and cell counts. Similar to our finding in Fig. 3, we found that UV irradiation reduced the average length of neurites, but collagen VI treatment rescued this damage effect (Fig. 6A–C, G). Under the same dosage of UV and collagen VI, Akt/PI3K inhibitor treated neurons had shorter neurite length than no inhibitor treated neurons (\( P < 0.05 \) vs. collagen VI only, Fig. 6C, D, G). Without UV irradiation, treating neurons with collagen VI and Akt/PI3K inhibitor did not affect neurite length. Nevertheless, there was no significant difference in the total number of neurons under all treatments (Fig. 6H).

Furthermore, we investigated the effect of collagen VI, Akt/PI3K inhibitor and the combination treatments on the UV-induced DNA damage and mitochondria dysfunction. After UV irradiation, neuronal culture co-treated with Akt/PI3K inhibitor and collagen VI had higher ratio of damaged cells than collagen VI only culture (both \( P < 0.05 \), Fig. 7). In the absence of collagen VI, neurons with or without Akt/PI3K inhibitor treatment did not have significant difference in cell survival under the same does of UV irradiation. In the absence of UV irradiation, treating neurons with Akt/PI3K inhibitor alone or in combination with collagen VI did not affect the cell survival (Fig. 7). These results support the notion that Akt/PI3K signaling plays an anti-apoptotic role in the collagen VI-dependent protection of neurons against UV-induced apoptosis.

**DISCUSSION**

In this study, we found that neurons have increased Col6a1, Col6a2, and Col6a3 mRNA levels along with se-
creted collagen VI protein level after UV irradiation. UV usually causes multiple cellular damages and induces apoptosis process in many types of cells (Lu and Lane, 1993). To determine whether the up-regulation of collagen VI plays a protective or detrimental role after UV irradiation, we exogenously applied soluble collagen VI into the

![Image of neurons with untreated and treated conditions](image)

**Fig. 3.** Collagen VI rescued UV-induced neurite shrinkage. Primary neurons with or without 2 h collagen VI pre-treatment were irradiated with 192 J/m² UV. (A–D) Representative images of MAP-2 positive neurons with following treatments: control (A), UV (B), UV+25 nM collagen VI (C), UV+50 nM collagen VI (D). Blue, DAPI staining for nuclei; green, GFAP staining for astrocytes; red, MAP-2 staining for neurons. Scale bar represents 200 µm. (E–G) Quantification of average neurite length (E), average soma area (F), and total cell count (G) of MAP-2 positive neurons determined by NeurotoxQ. The results were averaged from four independent experiments. Three pictures were taken from each well, and three to six repeated wells were performed in each condition per experiment. * P<0.05 versus no UV control.
primary neuronal culture medium. Assays for DNA damage, mitochondria dysfunction, and cell morphology indicated that the higher concentration of soluble collagen VI protein made neurons more resistant to UV irradiation. In addition, we demonstrated that collagen VI-treated neurons have higher Akt activity and lower JNK activity after UV irradiation. Inhibiting Akt/PI3K activity diminished the protective effect of collagen VI. Our findings suggested that collagen VI may enhance neuronal survival after UV irradiation through inducing Akt/PI3K anti-apoptotic pathway.

Expression of Col6 genes are increased after UV damage

After neuronal damage by UV irradiation, we found that both collagen VI mRNA and protein levels were significantly up-regulated. Although collagen VI protein is undetectable inside the neurons, it has significantly higher level in culture medium after UV irradiation, which is consistent with its role as an extracellular protein. Similarly, when neurons expose to amyloid-β peptide (Aβ), the primary toxin causing Alzheimer’s disease, Col6a1 gene expression is up-regulated through the TGF-β/Smad3 pathway to make neurons more resistant to Aβ (Cheng et al., 2009). The dynamic expression of Col6 gene family has been founded in multiple types of cells under normal and pathogenic conditions (Hatamochi et al., 1989; Adriaenssens et al., 2009). For example, during tumor progression, Col6 genes are up-regulated to promote survival of tumor cells through the β-catenin/TCF signaling pathway (Sherman-Baust et al., 2003; Iyengar et al., 2005). Therefore, our result is consistent with others suggesting that up-regulating Col6 gene expression could be a general response under stress condition in multiple types of cells including neurons.

Soluble collagen VI protects neurons against UV damage

Collagen VI protein has important roles in the control of DNA synthesis, cell survival, proliferation, and metabolism of normal and transformed cells (Ruehl et al., 1999; Ruhl et al., 1999; Irwin et al., 2003; Sherman-Baust et al., 2003; Iyengar et al., 2005; Cheng et al., 2009; Khan et al., 2009). In contrast, Col6a1 knockout mouse and zebrafish have mitochondrial dysfunction, ultrastructural defects, and increased incidence of apoptosis in myofibers (Palma et al., 2009; Telfer et al., 2010). Furthermore, primary neurons generated from Col6a1 knockout mice is more sensitive to Aβ (Cheng et al., 2009). Our result agrees with others suggesting that collagen VI could have a pro-survival role in neurons.
Collagen VI α1–3 subunits are believed to be stable as a heterotrimERIC chain complex (Bruns et al., 1986; Kielty et al., 1992), but it remains possible that each subunit has specific functions. The collagen VI used in this study is
well on collagen VI coated surface. Therefore, this protective effect is less likely due to the enhancement of neuronal adhesion but more likely due to the activation of the anti-apoptotic signaling.

Collagen VI alters Akt/PI3K signaling pathway

We identified that the activation of Akt/PI3K pathway may involve in collagen VI dependent protective effect in primary neurons. Phosphorylation of Akt was induced in neurons after collagen VI treatment, and inhibition of Akt/PI3K signaling pathway diminished the protective effect of collagen VI against UV irradiation. Akt/PI3K is highly expressed in the brain and many other tissues, and is known as one of the major signaling molecules for cell proliferation and survival mediated by extracellular stimuli (Brunet et al., 2001; Brazil et al., 2002; Parcellier et al., 2008). Akt/PI3K activation is known to increase the cell resistance to UV irradiation via inhibiting of JNK activation in several types of cells (Sunayama et al., 2005; Kim et al., 2009). Similar to our findings, collagen VI can also increase phosphorylation of Akt/PI3K to promote its growth-stimulatory and pro-survival effects of tumor cells (Iyengar et al., 2005).

NeurotoxQ, a free, automatic morphological quantification program

Improvement of tools to study cellular and molecular biology is playing an increasingly important role in assisting researchers to study the mechanisms of diseases or screening novel drugs. Neuronal morphological changes after neuronal damages are typically quantified by manual examination. This kind of quantification is time consuming, inconsistent, and highly subjective. Programs for high-throughput screening are not always easy to adopt for small scale laboratory use (Zhang et al., 2007; Daub et al., 2009). The utilization of NeurotoxQ allows rapid, consistent, and objective quantification of neuronal cell body, neurite area, neurite length, and neurite thickness using image processing principles. We benchmarked NeurotoxQ and found that a typical 1 × 10⁶-pixel image acquired using a 10× objective takes roughly 15 s to complete. This is a significant improvement over another ImageJ-based automatic tracing algorithm which takes 2–3 min (Narro et al., 2005) or a computer-aided manual tracing algorithm which can take hours (Meijering et al., 2004). However, it is important to note that the result from NeurotoxQ quantification is on a per image field basis. Due to the complexity of the neuronal network, it is extremely difficult, if not impossible, to distinguish neurites originated from different neurons; this prevents us from developing an algorithm that can generate quantitative data on a per cell basis. As a result, it is crucial to normalize the neurite length or neurite area using neuronal cell body if one wishes to compare results on a per cell basis. In addition, NeurotoxQ cannot operate on images acquired using high magnification objectives (higher than 20×). This is due to the skeletonize operation utilized. When high magnification images are processed, skeletonize operation produces tree-like branches within thick neu-
rites and hence results in an over-estimation of the neurite length.

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