A Modified Chinese Herbal Decoction (Kai-Xin-San) Promotes NGF-Induced Neuronal Differentiation in PC12 Cells via Up-Regulating Trk A Signaling

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Kai-Xin-San (KXS), a Chinese herbal decoction, has been applied to medical care of depression for thousands of years. It is composed of two functional paired-herbs: Ginseng Radix et Rhizoma (GR)-Polygalae Radix (PR) and Acori Tatarinowii Rhizoma (ATR)-Poria (PO). The compatibility of the paired-herbs has been frequently changed to meet the criteria of syndrome differentiation and treatment variation. Currently, a modified KXS (namely KXS2012) was prepared by optimizing the combinations of GR-PR and ATR-PO: the new herbal formula was shown to be very effective in animal studies. However, the cellular mechanism of KXS2012 against depression has not been fully investigated. Here, the study on KXS2012-induced neuronal differentiation in cultured PC12 cells was analyzed. In PC12 cultures, single application of KXS2012 showed no effect on the neuronal differentiation, but which showed robust effects in potentiating nerve growth factor (NGF)-induced neurite outgrowth and neurofilament expression. The potentiating effect of KXS2012 was mediated through NGF receptor, tropomyosin receptor kinase (Trk) A: because the receptor expression and activity was markedly up-regulated in the presence of KXS2012, and the potentiating effect was blocked by k252a, an inhibitor of Trk A. Our current results in cell cultures fully support the therapeutic efficacy of KXS2012 against depression.

Keywords: Kai-Xin-San, depression, paired-herbs, neuronal differentiation, Trk A

INTRODUCTION

Depression, namely major depressive disorder, is a serious mental illness characterized by constant feeling of low mood, low self-esteem, and loss of interest or pleasure (Barlow and Durand, 2011; Ferrari and Villa, 2016). The major pathological alteration of depression includes neuron reduction, disorder in neurotransmitter system and loss of neurotrophic factors; these changes are more robust in hippocampus and cortex (Krishnan and Nestler, 2008; Masi and Brovedani, 2011). The dysfunction in neurotransmitter system has been widely recognized and considered as one of the most severe pathogenesis. However, anti-depressants targeted on regulating neurotransmitter levels are not able to exert an effect to all the patients, indicating the complex pathogenesis during...
depression (Pehrson and Sanchez, 2014; Duman and Aghajanian, 2015). In the brain of depressed patients, nerve reduction resulted from retarded neurogenesis deposits that depression is largely caused by an impairment of the brain’s ability to maintain neurogenesis. The disorder is reversible when neurogenesis is improved. During neurogenesis, neuronal differentiation having the neurite outgrowth is of great importance for brain function and low mood regulation. Thus, the induction of neuronal differentiation predicts a novel and effective therapeutic target for development of anti-depressants (Qin et al., 2008).

Kai-Xin-San (KXS) from Beiji Qianjin Yaofang (Thousand Formulae for Emergency) by Sun Simiao (581–685 A.D.) in China has been applied to medical care of depression for thousands of years (Sun, 1997; Yasunori, 2011). KXS comprises two functional paired-herbs, i.e., Ginseng Radix et Rhizoma (GR; root and rhizome of Panax ginseng C. A. Mey.; Araliaceae family)—Polygala Radix (PR; root of Polygala tenuifolia Wild.; Polygalaceae family) and Acori Tatarinowii Rhizoma (ATR; rhizome of Acorus tatarinowi Schott; Acoraceae family)—Poria [PO; sclerotium of Poria cocos (Schw.) Wolf; Polyporaceae family]. The make-up of the paired-herbs contributes to anti-depressant functions of KXS. The administration of KXS alleviated depressive symptoms in animal studies with the mechanism of restoring the levels of neurotransmitters and neurotrophic factors in the brain (Dang et al., 2009; Zhu et al., 2012), as well as in cultured astrocytes and neurons (Zhu et al., 2013; Hu et al., 2014). However, the combination of GR-PR and ATR-PO has been frequently changed to meet the criteria of syndrome differentiation and treatment variation, which hinders the development of anti-depressant. Thus, KXS2012 (GR-PR: ATR-PO = 1:5) was prepared by optimizing the compatibility of GR-PR and ATR-PO (Yan et al., 2015). This new formulation was able to significantly alleviate depression-like symptoms and to regulate neurotransmitter and neurotrophic factor levels in animal study (Yan et al., 2016). However, the cellular mechanism of KXS2012 against depression has not been fully illustrated.

Here, we evaluated the function of KXS2012 against depression via promoting neuronal differentiation, an important stage of neurogenesis. Among different cell models for neuronal differentiation, cultured PC12 cell is usually selected for detection of neuronal differentiation in responding to various stimuli, e.g., NGF (Lortie et al., 2005; Marin-Vicente et al., 2011). The status of differentiated PC12 cells was determined morphologically through measuring neurite outgrowth and biochemically through analyzing the expression of neurofilaments, i.e., neurofilament is the major structural components of differentiated neurons (Schimmelpfeng et al., 2004; Xu et al., 2012). In neurons, NGF activates its receptor, Trk A, to promote neuronal differentiation. Thus, the Trk A signaling was studied to predict the potential cellular mechanism against depression. Our results could accelerate the development of new therapy for anti-depression.

**MATERIALS AND METHODS**

**Preparation of Herbal Decoction**

The plant materials, purchased from Qining Market in Guangzhou China, were morphologically authenticated by one of the authors, Dr. Tina T. Dong. The corresponding voucher specimens were deposited in Center for Chinese Medicine of The Hong Kong University of Science and Technology. The herbs were tested to be qualified according to the requirements of Chinese Pharmacopoeia (2015 Edition) and Hong Kong Materia Medica Standards.

According to Yi Xin Fang (Yasunori, 2011), GR-PR and ATR-PO were prepared according to the weight ratio of 1:1 and 1:2, respectively. In preparing the herbal extracts, 20 g of herb mixture was boiled in 160 mL of water for 2 h and extracted twice. The extracts were combined, dried under vacuum and stored at −80°C. KXS2012 was obtained by mixing the water extracts of GR-PR and ATR-PO together in 1:5 weight ratio. The quality control of herbal extracts was described in previous study (Yan et al., 2015). The extracts were solubilized in dimethylsulfoxide (DMSO) to give stock solution at a series of concentration from 15 to 50 mg/mL and stored at −20°C.

**Cell Culture**

PC12 cells, originated from rat adrenal medulla, were purchased from American Type Culture Collection (ATCC, Manassas, VA). Cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM), supplemented with 6% fetal bovine serum (FBS) and horse serum (HS), 100 units/mL penicillin and 100 µg/mL streptomycin in a humidified CO₂ (7.5%) incubator at 37°C. Fresh medium was applied every other day. All culture reagents were purchased from Life Technologies (Grand Island, NY).

**Herbal Treatment**

Cultured PC12 cells, after serum starvation for 3 h in DMEM having 1% FBS, HS, penicillin, and streptomycin, were treated with KXS2012 for 48 h for western blotting, or 24 h for real-time quantitative PCR. The cell viability assay was performed to determine a safe concentration range (0–100 µg/mL) of each extract, at which all extracts did not induce cell proliferation or death (Yan et al., 2015). In the co-treatment, cultured PC12 cells were treated with NGF (1.5 ng/mL) and KXS2012 (15µg/mL) for 24 h for western blotting assay. In the pre-treatment, PC12 cells were treated with herbal extract for 48 h and washed by 1x PBS twice before NGF (1.5 ng/mL) application for 24 h. In phosphorylation study, the cultures were starved for 5 h in DMEM before NGF (1.5 ng/mL) or herbal application.

**Polymerase Chain Reaction**

Total RNA was isolated from cell cultures by RNAzol RT reagent (Molecular Research Center, Cincinnati, OH) according to the manufacturer’s instruction. The amounts of RNAs were detected by UV absorbance at 260 nm. The total RNA was used to do the reverse transcription with moloney murine leukemia virus (MMLV) reverse transcriptase (Life Technologies), according to the protocol provided by the manufacturer. Real-time PCR was performed by using FastStart SYBR Green Master (Roche, Indianapolis, IN), according to the manufacturer's instruction. The SYBR green signal was detected by Mx3000P™ multiplex
quantitative PCR machine (BD Biosciences Clontech, San Jose, CA).

**SDS-Page and Immunoblotting**

After the treatment, the cells were solubilized in lysis buffer containing 0.125 M Tris-HCl, pH6.8, 4% SDS, 20% glycerol, 2% 2-mercaptoethanol, and analyzed immediately or stored frozen at −20°C. Proteins were separated on 8% SDS-polyacrylamide gels and transferred to a nitrocellulose. Transfer and equal loading of the samples was confirmed by staining the ponceau-S. The nitrocellulose was blocked with 5% fat-free milk in Tris-buffer saline/0.1% Tween 20 (TBS-T), and then incubated in the primary antibodies diluted in 2.5% fat-free milk in TBS-T over night at 4°C. The primary antibodies were: anti-NF200 (Sigma-Aldrich, St. Louis, MO, N4142, 1:1,000), anti-NF60 (Sigma-Aldrich, N2787, 1:5,000), anti-NF68 (Sigma-Aldrich, N5139, 1:2,000), anti-GAPDH (Abcam Ltd, Cambridge, UK, ab9485, 1:1,000,000), anti-phospho-Trk A (Cell Signaling, Danvers, MA, 9141, 1:1,000), anti-Trk A (Cell Signaling, 2505, 1:1,000), anti-phospho-Erk1/2 (Cell Signaling, 9101, 1:1,000), anti-Erk1/2 (Abcam, ab17942, 1:5,000), anti-phospho-CREB (Cell Signaling, 9198, 1:1,000), and anti-CREB (Cell Signaling, 9197, 1:5,000). After that, the nitrocellulose was rinsed with TBS-T and incubated for 2 h at room temperature in peroxidase (HRP)-conjugated anti-mouse secondary antibody, or peroxidase (HRP)-conjugated anti-rabbit secondary antibody (Life Technologies, 31430 or 31460, 1:5,000), diluted with 2.5% fat-free milk in TBS-T. After intensive washing with TBS-T, the immune complexes were visualized using the enhanced chemiluminescence (ECL) method (GE Healthcare, Piscataway, NJ).

**FIGURE 1** Standardization of herbal extracts. (A) The standardization of herbal extracts was described in our previous report (Zhu et al., 2010). The identification of 3, 6′-disnapoyl sucrose (330 nm) (1), α-asarone (258 nm) (8), and β-asarone (258 nm) (7) was made by a HPLC coupled with a DAD detector in paired-herb extracts. This HPLC profile served as fingerprint and determination of marker chemicals. (B) In chemical assessment, the identification of ginsenoside Rg1 (2), Rb1 (3), Rb1 (4), Rd (6), astragaloside IV (5) and pachymic acid (9) was made by a MS detector in standard markers and paired-herb extracts. The amounts of each chemical in mg/g of dried herbal extract were shown in (C). Mean ± SEM, n = 3.
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NJ). The intensities of bands in control and samples, run on the same gel and under strictly standardized ECL conditions, were compared on an image analyzer, using a calibration plot constructed from a parallel gel with serial dilutions of one of the sample.

Neurite Outgrowth Assay
A light microscope (Diagnostic Instruments, Sterling Heights, MI) equipped with a phase-contrast condenser, 10x objective lens and a digital camera (Diagnostic Instruments) was used to capture the images with the manual setting. For analyzing the number and length of neurite, ∼100 cells were counted from at least 10 randomly chosen visual fields for each culture. Using the photoshop software, the cells were then analyzed for the number and length of neurite. The cells were scored as differentiated if one or more neuritis was longer than diameter of the cell body, and they were classified to different groups according to the length of neurite that it possessed, i.e., <15, 15–30, and >30 µm (Xu et al., 2012). The number of neurite per cell was counting only those neurite that was longer than cell body.

Statistical Analysis
All data were analyzed using one-way ANOVA followed by the Students t-test. Statistical significance were classed as *P < 0.05; **P < 0.01; ***P < 0.001.

RESULTS
Standardization of Herbal Extracts
According to ancient preparation method of herbal mixture, GR-PR and ATR-PO were prepared in a weight ratio of 1:1 and 1:2, respectively. The extraction efficiency of GR-PR and ATR-PO were about 25.79 ± 3.25 and 10.32 ± 2.78%, respectively (Mean ± SD, n = 3). Two approaches were selected for the quality control of paired-herb extracts: (i) chemical fingerprinting; and (ii) minimal-marker requirement. The identified chemicals were shown in fingerprints (Figures 1A,B), according to our published reports (Zhu et al., 2010; Yan et al., 2015). Eight chemical markers, i.e., GR-derived ginsenosides (Rb1, Rd, Re, Rg1), PR-derived 3,6′-disinapoyl sucrose (330 nm), ATR-derived α-asarone (258 nm), and β-asarone (258 nm) and PO-derived pachymic

FIGURE 2 | KXS2012 does not induce neuronal differentiation. (A) KXS2012 (5–25 µg/mL) were applied onto cultured PC12 cells for 48 h. The expressions of neurofilaments (NF68, NF160, and NF200) were determined by specific antibodies. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) served as a loading control. Quantification plot was shown in histograms. (B) Cultured PC12 cells were treated as in (A). The cells were fixed with ice-cold 4% paraformaldehyde. The % of differentiated cells (upper panel), the number of neurite per cell (middle panel), and the length of neurite (lower panel) were counted as described in the Materials and Methods section. Bar = 10 µm. NGF (50 ng/mL) served as a positive control. Representative images were shown. Values are expressed as % of total cells in 100 counted cells, Mean ± SEM, n = 4. *P < 0.05 and **P < 0.01 compared to the control.
acid, were selected for chemical quantification (Figure 1). In GR-PR extract, the amounts of Rb1, Rd, Re, Rg1, and 3,6′-disinapoyl sucrose were about 4.287 ± 0.065, 0.397 ± 0.046, 3.648 ± 0.079, 3.304 ± 0.084, and 9.678 ± 0.067, respectively, in mg/g of dried herbal extract. In ATR-PO extract, the amounts of α-asarone, β-asarone, and pachymic acid were 6.292 ± 0.017, 0.736 ± 0.014, and 0.021 ± 0.003 in mg/g of dried herbal extract (Mean ± SD, n = 3) (Figure 1C). The established chemical parameters served as the control for repeatability of below biochemical analyses. Thus, KXS2012 was referring to a mixture of GR-PR extract plus ATR-PO extract at 1:5 weight ratio.

**KXS2012 Promotes NGF-Induced Neuronal Differentiation**

The cellular study of KXS2012 on PC12 cells was employed to investigate the function of herbal extracts in neuronal differentiation. The neuronal differentiation of PC12 cells could be determined morphologically in measuring the length of neurite: NGF induced the neurite outgrowth in a dose-dependent manner (Supplementary Figure 1). Three mammalian neurofilament subunits, NF68 (~68 kDa), NF160 (~160 kDa), and NF200 (~200 kDa), form hetero-dimers in making the structural domain of neurite (Schimmelpfeng et al., 2004; Xiong et al., 2016). Application of NGF in cultured PC12 cells also induced the expressions of neurofilaments in dose-dependent manners (Supplementary Figure 2). In cultured PC 12 cells, application of KXS2012 caused up-regulation of neurofilaments, NF68, NF160, and NF200, in dose-dependent manners (Figure 2A). The treatment of KXS2012 at 25 µg/mL slightly induced neurite outgrowth in PC12 cells. As compared to the control group, the effect possessed no significant difference neither on the length of neurite nor the number of neurite (Figure 2B). Thus, the herbal extract was not able to promote neuronal differentiation as single treatment.

In depressed mammalian brain, the level of NGF is reduced, and NGF-induced neuronal differentiation is down-regulated. Thus, the effect of KXS2012 in the presence of low level of NGF in neuronal differentiation was determined here. KXS2012 at 15 µg/mL, a concentration just barely showed induction of neurofilament expressions, was applied together with a small amount of NGF onto cultured PC12 cells by the methods of pre- and co-treatments. A suitable concentration of NGF was selected: 1.5 ng/mL. Application of KXS2012 together with NGF caused up-regulation of NF68, NF160, and NF200 (Figure 3A). The treatment of KXS2012 at 50 ng/mL, a concentration that showed induction of neurofilament expressions, was applied together with a small amount of NGF onto cultured PC12 cells by the methods of pre- and co-treatments. A suitable concentration of NGF was selected: 50 ng/mL. Application of KXS2012 together with NGF caused up-regulation of NF68, NF160, and NF200 (Figure 3B). The treatment of KXS2012 at 100 ng/mL, a concentration that showed induction of neurofilament expressions, was applied together with a small amount of NGF onto cultured PC12 cells by the methods of pre- and co-treatments. A suitable concentration of NGF was selected: 100 ng/mL. Application of KXS2012 together with NGF caused up-regulation of NF68, NF160, and NF200 (Figure 3C). The treatment of KXS2012 at 150 ng/mL, a concentration that showed induction of neurofilament expressions, was applied together with a small amount of NGF onto cultured PC12 cells by the methods of pre- and co-treatments. A suitable concentration of NGF was selected: 150 ng/mL. Application of KXS2012 together with NGF caused up-regulation of NF68, NF160, and NF200 (Figure 3D). The treatment of KXS2012 at 200 ng/mL, a concentration that showed induction of neurofilament expressions, was applied together with a small amount of NGF onto cultured PC12 cells by the methods of pre- and co-treatments. A suitable concentration of NGF was selected: 200 ng/mL. Application of KXS2012 together with NGF caused up-regulation of NF68, NF160, and NF200 (Figure 3E).
this concentration should have no or little effect on induction of neurite outgrowth or neurofilament expressions. NGF at 1.5 ng/mL was selected in pre-treatment and co-treatment studies (see Supplementary Figures 1, 2): this concentration was to ensure the responses from a low amount of KXS2012 could be revealed. In both pre- and co-treatments, the inductions of NF68, NF160, and NF200, induced by KXS2012 plus NGF, were significantly higher at 3- to 4-folds (Figure 3). However, the effect triggered by pre-treatment was more robust than that of co-treatment. In neurite outgrowth assay, the number of differentiated cells by pre-treatment of KXS2012 and NGF was reached to ~30% (Figure 4A), and ~20% increase was revealed in the co-treatment (Figure 4B). By counting the number of neurite per cell, the pre- and co-treatment of KXS2012 plus low dose of NGF caused an increase, significantly, by about 10% of cells having neurite (Figure 4). In addition, the differentiated cells, induced by KXS2012 plus low amount of NGF, contained longer neurite, e.g., >15 and >30 µm (Figure 4). Compared to

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\text{FIGURE 4} \ | \ \text{KXS}_{2012} \text{promotes NGF-induced neurite outgrowth. (A) KXS}_{2012} \text{ (15 g/mL) was applied onto cultured PC12 cells for 48h before NGF application (1.5 ng/mL) for 24h. (B) KXS}_{2012} \text{ (15 g/mL) and NGF (1.5 ng/mL) were applied onto cultured PC12 cells for 24h. NGF (50 ng/mL) was applied as a positive control. Cells were fixed with ice-cold 4% paraformaldehyde, and the extension of neurite was revealed. Bar = 10 µm. The % of differentiated cells (upper panel), the number of neurite per cell (middle panel), and the length of neurite (lower panel) were counted as described in the Materials and Methods section. Values are expressed as % of total cells in 100 counted cells, Mean ± SEM, } n = 4, \ *p < 0.05 \text{ and } **p < 0.01 \text{ compared to the control.}
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pre-treatment study, the inductive role of KXS$_{2012}$ showed less robust effect in promoting NGF-induced neurite outgrowth by co-treatment (Figure 4).

**The Effect of KXS$_{2012}$ on Neuronal Differentiation via Up-Regulating Trk A Signaling**

NGF achieves its function by binding to and activating Trk A in PC12 cells. The NGF-activated Trk A stimulates downstream signaling pathways leading to neuronal differentiation and promoting cell survival (Zhang et al., 2000; Vaudry et al., 2002). The role of KXS$_{2012}$ in phosphorylating Trk A was tested here. NGF induced Trk A phosphorylation in a dose-dependent manner, and NGF at 1.5 ng/mL was selected in phosphorylation study, which barely showed phosphorylation capability (Supplementary Figure 3). KXS$_{2012}$ or co-treatment procedure did not effectively phosphorylate Trk A; however, the pre-treatment of KXS$_{2012}$ together with low level of NGF showed much better responsiveness (Figure 5A).

In addition to Trk A, the downstream effectors of CREB and Erk1/2 were also determined here by pre- and co-treatment of KXS$_{2012}$ with low amount of NGF. In cultured PC12 cells, NGF induced CREB phosphorylation (∼40 kDa) at ∼5-folds; however, co-treatment with KXS$_{2012}$ did not show significant difference, as compared to NGF single treatment (Figure 5B). In contrast, the phosphorylation of CREB was significantly induced by ∼7-folds in the scenario of pre-treatment of KXS$_{2012}$ plus NGF (1.5 ng/mL; Figure 5B). Similarly, the Erk1/2 phosphorylation (∼42/44 kDa) was also being markedly up-regulated with pre-treatment of KXS$_{2012}$, while no significant difference, compared to NGF group, was observed in the co-treatment (Figure 5C).

To explore the role of KXS$_{2012}$ in NGF signaling, the expression of Trk A in cultured PC12 cells were analyzed. The expression of Trk A at ∼140 kDa, recognized by its specific antibody, was increased by 160% in KXS$_{2012}$ (15 µg/mL)-treated cultures (Figure 6A). NF68 served as a positive control. The mRNA encoding Trk A was analyzed by real-time PCR. Application of KXS$_{2012}$ in PC12 cultures induced Trk A mRNA in a dose-dependent manner: the maximal induction of over 150% increase was revealed at ∼25 µg/mL (Figure 6B). This was in consistent with the phosphorylation study of KXS$_{2012}$ that the up-regulation of Trk A expression could be an outcome of increasing sensitivity to low level of NGF during differentiation in PC12 cells.

Moreover, K252a, an inhibitor of tyrosine phosphorylation of Trk A, was applied onto cultured PC12 cells for 3 h before the pre- and co-treatment procedures. In the presence of K252a (0.1 µM), the protein expression of NF68 was totally decreased to that of control in KXS$_{2012}$-treated cultures (Figure 7), which suggested

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**FIGURE 5 | KXS$_{2012}$ promotes NGF-induced phosphorylation.** In pre-treatment, KXS$_{2012}$ (15 µg/mL) was applied onto cultured PC12 cells for 48 h before serum starvation for 5 h, with or without application of NGF (1.5 ng/mL), for different time. In co-treatment, cultured PC12 cells, serum starvation for 5 h, were co-treated with NGF (1.5 ng/mL), KXS$_{2012}$ (15 µg/mL) and KXS$_{2012}$ (15 µg/mL) + NGF (1.5 ng/mL) for different time. (A) Total Trk A and phosphorylated Trk A were revealed by using specific antibodies. (B) Total CREB and phosphorylated CREB were revealed by using specific antibodies. (C) Total Erk1/2 and phosphorylated Erk1/2 were revealed by using specific antibodies. Quantification plot of the phosphorylation level in treatment of 5 (A) and 10 (B, C) min was shown. Values are expressed x Basal, where control value is set as 1. Mean ± SEM, n = 4. *p < 0.05, **p < 0.01, and ***p < 0.001 compared to the control.
that KXS$_{2012}$-promoted neuronal differentiation was mediated by Trk A. These findings suggested that KXS$_{2012}$ promoted neuronal differentiation by up-regulating Trk A expression, which could increase the cellular response to low level of NGF.

**DISCUSSION AND CONCLUSION**

Depression is a common mental disorder, which shows severe neuron loss and dysfunction, mood disorder and behavioral disturbance (Barlow and Durand, 2011; Masi and Brovedani, 2011). In addition to neurodegenerative symptoms, depression is more likely to progress to suicide and other concurrent metabolic disorders, e.g., diabetes and cardiovascular diseases (Compare et al., 2014; Koponen et al., 2015). However, current anti-depressants are not able to deal with all the depression-related injuries, which make efficient drug development for anti-depression a must. KXS, a classical Chinese herbal formula is expected to develop as a new anti-depressant. KXS is composed of two functional paired-herbs, i.e., GR-PR and ATR-PO. According to the theory of syndrome differentiation and treatment variation, the varied combination of paired-herbs in KXS has been recorded in Chinese medical books, which directs possible development of anti-depressants (Sun, 1997; Yasunori, 2011). Thus, a re-formulated KXS$_{2012}$ was prepared by optimizing the pairing of GR-PR and ATR-PO: the new formula presented robust anti-depressant action in animal study. It could also induce the expressions of neurotrophic factors as well as their receptors in rat cortical and hippocampal neurons (Yan et al., 2016). However, the underlying mechanism of KXS$_{2012}$ against depression has not been fully investigated. In the study, we investigated the cellular mechanism of KXS$_{2012}$ against depression by using PC12 cell as an *in vitro* model.

It is suggested that depression is largely caused by an impairment of neurogenesis in the brain. During neurogenesis, neuronal differentiation, in presence of neurite outgrowth, plays a critical role in synapse formation, which predicts a promising target of anti-depression (Hatiangady and Shetty, 2010; Snyder et al., 2011). Here, the potential drug targets and cellular mechanism on neuronal differentiation were addressed. The cultured PC12 cells are probably the most valid cell model for the detection of neuronal differentiation. The status of differentiated PC12 cells was determined morphologically through measuring neurite outgrowth and biochemically through analyzing the expression of neurofilaments (Schimmelpfeng et al., 2004; Xu et al., 2012). Our results indicated that KXS$_{2012}$ significantly potentiated NGF-induced neuronal differentiation. The potentiating effect was mediated through NGF receptor, Trk A. KXS$_{2012}$ greatly up-regulated Trk A signaling in the cultures, and the potentiating effect was blocked by k252a, an inhibitor of Trk A. Interestingly, the promoting mechanism of KXS$_{2012}$ in Trk A phosphorylation did not show great effect on the co-treatment situation. This discrepancy could be accounted by KXS$_{2012}$-induced Trk A expression in cultured PC12 cells, which required longer time for Trk A activation. Moreover, the phosphorylations of Erk 1/2 and CREB, the downstream activators of Trk A, were explored here. The co-treatment of KXS$_{2012}$ and NGF did not show marked induction on Erk 1/2 and CREB phosphorylations, as compared to NGF group, which was in accord with our hypothesis that the promoting effect of KXS$_{2012}$ might require longer time of activation. By up-regulating Trk A expression and activation, KXS$_{2012}$ could increase the cellular response to low level of NGF; as to improve neuron adaptation and depression (Welberg, 2008; Peña et al., 2014). In line to this hypothesis, the study on siRNA knockdown of Trk A expression in cultured PC12 cells will be carried out as to further illustrate the function of KXS$_{2012}$ via Trk A signaling.

Considering the major ingredients in KXS$_{2012}$ responsible for anti-depression, different chemicals abundant in the herbal extract have been studied. GR-derived ginsenosides were reported to attenuate depression-like symptom by balancing neuronal cell proliferation and apoptosis and up-regulating neurotransmitter system (Xue et al., 2006; Li et al., 2014; Zhang et al., 2017). PR-derived 3,6′-disinapoyl sucrose reversed mental disorder by improving hippocampal neuron plasticity and neurotrophic signaling pathway in stressed rats (Hu et al.,
FIGURE 7 | K252a inhibits KXS2012-promoted neurofilament expression. (A) K252a (0.1 µM) was applied onto cultured PC12 cells, 3 h before KXS2012 (15 µg/mL) treatment. After 48 h, PC12 cells were washed by PBS twice and treated with NGF (1.5 ng/mL) for 24 h. (B) K252a (0.1 µM) was applied onto cultured PC12 cells 3 h before KXS2012 (15 µg/mL) treatment together with NGF (1.5 ng/mL) for 24 h. The cell lysates were collected to determine the expression of NF68. GAPDH served as a loading control. Quantification plot was shown in low panel. Values are expressed as × Basal, where control value is set as 1, Mean ± SEM, n = 4. *p < 0.05 and **p < 0.01 compared to the control.

2010). Asarones, derived from ATR, were more likely to pass the blood brain barrier, which may help improve distributions of other ingredients in the brain (Lu et al., 2014). The finding was consistent with our preliminary results that GR-derived ginsenoside Rb1 and Rg1, PR-derived 3,6′-disinapoyl sucrose, ATR-derived α-asarone and β-asarone were able to promote neuronal differentiation in neuronal cells. Therefore, it is reasonable to use KXS2012 as a new regimen for anti-depression for its robust effect on neuronal differentiation and Trk A signaling.

AUTHOR CONTRIBUTIONS

LY and KT: conceived and designed the experiments; LY: performed the experiments; LY, MW, SX, and PS: analyzed the data; CB, JL, AX, and TD: contributed reagents; LY, AG, and KT: contributed to the writing of the manuscript.

ACKNOWLEDGMENTS

Supported by Hong Kong Research Grants Council Theme-based Research Scheme (T13-607/12R), GRF (663012, 662713, M-HKUST604/13), TUYF15SC01, The Hong Kong Jockey Club Charities Trust (HKJCCT12SC01), Foundation of The Awareness of Nature (TAON12SC01); Shenzhen Science and Technology Innovation (JCYJ20160229205726699, JCYJ20160229205812004, JCYJ20160229210027564 and 20170326); National Natural Science Foundation of China (81403087); the program of Jiangsu Key laboratory for the Research and Utilization of Plant Resources (JSPKLB201607).
SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcell.2017.00118/full#supplementary-material

Supplementary Figure 1 | NGF induces neurite outgrowth. Cultured PC12 cells were treated with NGF (0.5–50 ng/mL) for 24 h (A) or 48 h (B). Cells were fixed with ice-cold 4% paraformaldehyde. Bar = 10 μm. The % of differentiated cell and length of neurite were counted as described in the Materials and Methods section. Values are expressed as % of total cells in 100 counted cells, Mean ± SEM, n = 4. *p < 0.05 and **p < 0.01 compared to the control.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.