Synergistic Effects of Combined Nurr1 Overexpression and Natural Inducers on the More Efficient Production of Dopaminergic Neuron-Like Cells From Stem Cells

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The development of dopaminergic (DA) neurons is a very complex process, and a combination of extrinsic and intrinsic factors involves their differentiation. Transcription factor, Nurr1 plays an essential role in the differentiation and maintenance of midbrain DA neurons. Nurr1-based therapies may restore DA function in Parkinson’s disease (PD) by replacing damaged cells with differentiated cells derived from stem cells. Providing tissue-specific microenvironments such as brain extract can effectively induce dopaminergic gene expression in stem cells. The present study aimed to investigate the combined effects of Nurr1 gene overexpression and a neonatal rat brain extract (NRBE) induction on dopaminergic differentiation of P19 stem cells. In order to neural differentiation induction, stably Nurr1-transfected cells were treated with 100 µg/ml of NRBE. The differentiation potential of the cells was then evaluated during a period of 1–3 weeks via various methods. The initial evaluation of the cells by direct observation under a light microscope and cresyl violet specific staining, confirmed neuron-like morphology in the differentiated cells. In addition, different molecular and cellular techniques, including real-time PCR, immunofluorescence, and flow cytometry, demonstrated that the treated cells expressed pan-neuronal and dopaminergic markers. In all experimental groups, neuronal phenotype with dopaminergic neuron-like cells characteristics mainly appeared in the second week of the differentiation protocol. Overall, the results of the present study revealed for the first time the synergistic effects of Nurr1 gene overexpression and possible soluble factors that existed in NRBE on the differentiation of P19 stem cells into dopaminergic neuron-like cells.

Keywords: stem cell differentiation, Nurr1 gene, tissue-specific microenvironment, soluble factors, dopaminergic neurons, Parkinson’s disease

INTRODUCTION

In vitro production of midbrain dopamine neurons from stem cells is a promising approach for managing and treating Parkinson’s disease (PD). The study of dopaminergic (DA) neuron development is one of the areas of interest in order to generate these neurons from stem cell sources. Although many pieces of research focused on in vivo and in vitro studies of these neurons for different purposes, still further investigations are needed to realize the factors that control
DA cell differentiation and patterning. The development of DA cells is a very complex process, and a combination of extrinsic (sonic hedgehog, fibroblast growth factor 8, transforming growth factor b and Wnts) and intrinsic (En1/2, Lmx1a/b, Foxa1/2, Pitx3, and Nurr1) factors involves in their differentiation (Yuan et al., 2015). Previous studies have demonstrated that orphan nuclear receptor Nurr1 (nuclear receptor-related 1 protein, also known as NR4A2; nuclear receptor 4A2), as a transcriptional factor, plays an essential role in the differentiation and maintenance of midbrain DA neurons. Nurr1 activates tyrosine hydroxylase (TH) gene expression via direct binding to its promoter (Kim, 2011), and Nurr1 gene defect is associated with Parkinson’s disease (Dong et al., 2016). DA neurogenesis failed in Nurr1 mutant mice and resulted in their death immediately after birth (Hermanson et al., 2003). Although Nurr1 heterozygous mice are healthy, dopamine levels reduced significantly in their ventral midbrain and striatum (Zetterström et al., 1997). Exogenous Nurr1 inserted in neural stem cells (NSCs) induced TH expression in a concentration-dependent manner (Kim et al., 2007). On the other hand, factors such as parathyroid hormone and corticotropin-releasing factor can induce Nurr1 expression in bone cells and a pituitary cell line, respectively (Murphy and Conneely, 1997; Tetradsis et al., 2001). Nurr1-based therapies through replacing damaged cells by in vitro differentiated cells might restore DA functions in PD (Bartus et al., 2013). It has been previously reported that injection of Nurr1-overexpressing cells in combination with glial-derived neurotrophic factor (GDNF) or neurturin into the midbrain might protect DA neurons against toxicity of alpha-synuclein (Decressac et al., 2012; Bartus et al., 2013).

In order to produce highly specialized cells, it will be beneficial to create an environment similar to natural in vivo conditions. Numerous reports have shown that tissue-specific microenvironment can induce stem cell differentiation into appropriate cell types with the desired characteristics (Rivera et al., 2006; Bentz et al., 2010; Han et al., 2014; Narayanan et al., 2014; Momendoust et al., 2019). Recently, several studies have indicated that the use of a neonatal rat brain extract (NRBE) alone or in combination with a three-dimensional culture system (Azizi et al., 2018) or a neuroprotective drug, deprenyl (Momendoust et al., 2019), could effectively induce DA phenotype with the ability to express TH gene in stem cells.

Given the importance of Nurr1 as a key gene for DA neuron development, the present study was designed to investigate the effects of Nurr1 overexpression on dopaminergic differentiation of P19 embryonal carcinoma (EC) stem cells. In addition, stable Nurr1-transfected cells were exposed to NRBE to evaluate the synergistic effects of combined gene overexpression technology with a natural tissue-specific medium on the more efficient production of DA cells.

**MATERIALS AND METHODS**

**Culture and Stable Transfection of P19 Cells**

A teratocarcinoma stem cell line, P19 (Cell Bank, Pasteur Institute of Iran, Tehran, Iran), was grown in alpha-MEM (alpha minimum essential medium, Gibco-BRL, Carlsbad, CA, 11900073) containing 10% FBS (fetal bovine serum, Gibco, 10270-106), 50 µg/ml penicillin (Sigma, P3032) and 50 µg/ml streptomycin (Sigma, S1277). A CaPO4 (calcium phosphate) co-precipitation method was used for transfection of the cells as previously described (Chen and Okayama, 1987). Briefly, the exponentially growing cells were trypsinized and seeded (1.5 x 10^6) onto a 60-mm culture dish 24 h before transfection. A solution of CaPO4-DNA (10-20 µg total plasmid DNA; pCMV3-C-NR4A2-GFPSpark, Sino Biological, HGI4938-ACG) was added dropwise onto the cells at 60–70% confluency. The cells were then incubated at 37°C in 5% CO2, and the growth medium was replaced by fresh alpha-MEM from 7–9 h. For stable transfection, a medium containing 200 µg/ml hygromycin B (Sigma, H0654) was applied, and the transfected cells were selected. Eight days after transfection, live cells were examined and photographed using a fluorescence microscope to detect GFP expression. P19-GFP+ cells were also generated by pCMV3-GFPSpark plasmid (empty) and considered as transfection control.

**Preparation of Neonatal Rat Brain Extract**

Wistar rat strain was obtained from Isfahan University of Medical Sciences (Isfahan, Iran) and kept under standard housing conditions with a regular dark/light cycle. The animal procedures in the present study were performed according to the rules and regulations set by the Bioethics Committee of the University of Isfahan (Code: IR.U.I.REC.1397.125), based on the National Specific Ethical Guidelines for Biomedical Research issued by the Ministry of Health and Medicinal Education (MOHME) of Iran in 2005. The rats were mated overnight, and the day neonates were born was considered as P0. One-week-old neonates sacrificed, their whole brains removed from the skull, and brain extracts prepared as described previously (Momendoust et al., 2019). Briefly, the whole brain was homogenized in a protease inhibitor (PMSF, Roche, 10 837 091 001) solution and then centrifuged at 3,000 rpm (10 min) and 12,000 rpm (20 min). All collected NRBE was pooled and total protein concentrations were assayed using the Bradford method. Finally, the extract was stored at −70°C until use.

**Neural Differentiation Induction Protocol**

To neural differentiation of P19 cells, the cells were cultured in suspension for 3 days to induce embryoid bodies (EBs) production. The resulted EBs were transferred to gelatin-coated tissue culture dishes and allowed 1 day to attach. Four experimental groups were designed as follows: (1) E (empty): EBs transfected with pCMV3-C-GFPSpark vector (alpha-MEM + 5% FBS + empty plasmid); (2) EEx: EBs transfected with pCMV3-C-GFPSpark vector and treated by NRBE (alpha-MEM + 5% FBS + empty plasmid + 100 µg/ml NRBE); (3) N: EBs transfected with pCMV3-C-NR4A2-GFPSpark vector (alpha-MEM + 5% FBS + N4A2 plasmid); (4) NEx: EBs transfected with pCMV3-C-NR4A2-GFPSpark vector and treated by NRBE (alpha-MEM + 5% FBS + NR4A2 plasmid + 100 µg/ml NRBE). The differentiation potential of the cells was evaluated during a period of 1–3 weeks.
Specific Staining of the Differentiated Cells

The initial identification of the differentiated cells with neuronal phenotype was evaluated by: (1) direct observation of cell morphology under a light microscope; and (2) cresyl violet specific staining (Fraichard et al., 1995). To confirm the existence of Nissl bodies in the cytoplasm, the cells were fixed in 70% ethanol for 10 min at room temperature, dehydrated in 95% ethanol/5% acetic acid for 20 min at −20°C, and then exposed to cresyl violet solution (0.25% cresyl violet, 0.8% glacial acetic acid, 0.6 mM sodium acetate). Cells were washed in PBS (phosphate buffer saline) and then mounted for further examination.

Real-Time PCR

To quantify the relative mRNA expression level of the interested genes, total RNA was extracted from all the samples, including untreated and treated groups. Complementary DNA synthesized (Parstous, Easy™ cDNA Synthesis Kit, A101161), and real-time PCR assay was performed by StepOnePlus™ Real Time PCR System using specific primers (Table 1) and SYBR Premix Ex Taq (Takara, RR081Q). To analyze PCR reaction efficiency, standard curves were used for each gene, and a melt curve analysis was performed at the end of each reaction. The gene expression profile was quantified by the convenient 2^ΔΔCt method. Triplicate samples of cells were collected at each time point, and real-time PCR was performed on the corresponding synthesized cDNA. The Ct values provided by the Q-PCR instrument were easily imported into a spreadsheet program such as Microsoft Excel. The data were analyzed using amount of target Ct values normalized to individual sample.

| Gene         | Accession number | Primers                                      | Product size |
|--------------|------------------|----------------------------------------------|--------------|
| Sox2         | NM_011443        | F:TGAACCGAGGAGCATGGACAGCCTA                      | 221          |
| Nanog        | XM_011241471     | R:CGATGGTCTGCTTCCATGGCA                       | 240          |
| Oct3/4       | NM_013633        | F:CCTCACCACTCTTAC                           | 156          |
| Nestin       | NM_016701        | R:ACGGTGGATTTTTTCTACGTCTACGTG                 | 143          |
| Synaptophysin| NM_009935        | F:TGGCAGACACAGGATGCCTGTG                      | 217          |
| CREB         | NM_009952.2      | R:AGAAGACAGGAGGAAGAGAG                       | 250          |
| Tyrosine     | NM_009377        | F:TGGCAAGCTCTACCAAGATCAAC                     | 103          |
| Nurr1        | XM_011239033.2   | R:CAGGGTACGGATGGAGGCA                       | 159          |
| Onecut       | NM_008262        | F:GGCAACCTGAGGCTTAC                           | 159          |
| GSK3β        | BC034862         | R:TGTGCTGGAGGTATGCTG                         | 181          |
| Nurr1        | XM_011239033.2   | F:CAGGTCCGAGTCTTACAC                         | 159          |
| Onecut       | NM_008262        | R:AGGGGATCATGGTCAAC                         | 159          |
| GSK3β        | BC034862         | F:GAAATCTGCTGAC                             | 181          |
| BDNF         | BC011231.1       | R:CACGGCAATTCTTCTGTG                          | 224          |
| NGF          | BC011231.1       | F:TTTCAACAGGACTCGCAG                         | 224          |
| β-actin      | NM_007393        | R:TCTCAACAGGAGGATTGAC                       | 124          |
| GAPDH        | NM_007393        | F:CAGGGCTGACATCTTCTGTG                       | 94           |
|             |                  | R:TGACCCAGGGGCCCAATAC                       |              |

Table 1: Real-time PCR primer sequences.

Immunofluorescence

Evaluation of the specific protein profiles of the differentiated cells carried out by immunofluorescence. The cells of all the samples were fixed in 4% paraformaldehyde and then permeabilized by 0.3% Triton X-100 and incubated with blocking buffer (10% normal goat serum; NGS, Sigma, G9023). The cells subsequently incubated with primary antibodies, including rabbit polyclonal nestin (Sigma, N5413), mouse anti-synaptophysin monoclonal (Abcam, ab8049), mouse anti-β-III tubulin monoclonal (Abcam, ab7751), rabbit anti-tyrosine hydroxylase–neuronal marker (Abcam, ab6211), rabbit polyclonal antibody to GFP (Abcam, Cambridge, USA, ab290), and mouse monoclonal anti-Nurr1 antibody (Abcam, ab41917), each at 1:1,000 dilutions. FITC-conjugated anti-mouse IgG (Sigma, F9137) and Cy5.29-conjugated anti-rabbit IgG (Abcam, ab6564) each at 1:1,000 dilutions applied as secondary antibodies. DAPI (4’,6-Diamidino-2-phenylindole, Sigma, D9542) utilized to counterstain the nuclei.

Flow Cytometry

The cells were washed with PBS and gently dissociated into a single-cell suspension using 0.25% trypsin/EDTA (Sigma, T4799). Fixation buffer (4% paraformaldehyde in PBS) was added to 100 μL cell suspension, and then the cells were permeabilized by Triton X-100 and incubated with blocking buffer (10% NGS in PBS). The cells were exposed to above mentioned primary and secondary antibodies. Un-transfected cells were used to adjust the detector settings. Fluorescent intensity was determined on FACS (BD FACSCalibur), with 10,000 events captured per sample. The interested cell population, excluding debris and dead cells, was determined by forward and side scatters gating. The
acquisition and analysis of the FACS data were performed with FlowJo software.

**Statistical Analysis**

Statistical differences were evaluated between two groups by Student’s *t*-test and multiple groups by one-way ANOVA (analysis of variance) and LSD (least significant difference) test. The experiments were performed in triplicate, and the data presented as mean ± SD (standard deviation) and *p* < 0.05 considered significant.

**RESULTS**

**Morphological Studies of the Differentiated Cells**

To induce neural differentiation of P19 cells, they were stably transfected with a vector containing Nurr1 gene and GFP reporter gene under the control of cytomegalovirus (CMV) promoter. The expression of GFP in the transfected cells was confirmed by using a fluorescence microscope (Figure 1A). The embryoid bodies resulted from the suspension culture of GFP+ cells were transferred on gelatin-coated culture dishes and exposed to four various conditions as mentioned above. Initially, the morphological changes of the differentiated cells were evaluated using a light microscope over 1–3 weeks (Figure 1B). The cells with small bodies and numerous processes which formed a cellular network with each other were considered neuron-like cells. Although cell counting was not carried out here, visual examinations of the images showed that in all experimental groups, neuronal phenotype mainly appeared in the second week of the differentiation protocol. By the third week of the differentiation, the morphology of most cells changed to a non-neuronal appearance with large polyhedral cell bodies and a few short processes. There were very few cells with neuronal phenotype in the E group that showed spontaneous cell differentiation.
Clusters of neuron-like cells were interconnected by their fine processes. A few neuron-like cells were observed in the E group, while pCMV3-GFPSpark vector-transfected cells showed no immunoreactivity to Nurr1 antibody.

Furthermore, double immunofluorescence staining of the paraformaldehyde-fixed cells with antibodies against Nurr1 and GFP confirmed co-expression of Nurr1 and GFP in the cells transfected by pCMV3-NR4A2-GFPSpark vector (Figure 2A), while the cells of E group (pCMV3-GFPSpark vector-transfected cells) showed no immunoreactivity to Nurr1 antibody.

Evaluation of Protein Expression by Immunofluorescence and Flow Cytometry

According to the preliminary results from the morphological studies and the results of gene expression evaluation, further analysis of all experimental groups focused on the second week of the differentiation protocol via various methods, including immunofluorescence and flow cytometry. Immunofluorescence assessments confirmed that the cells treated under different conditions were immunoreactive to neuroepithelial marker nestin (Figure 5A). However, there was no distinct positive response in the E group in the second week of analysis. Furthermore, the expression of nestin protein was quantified by flow cytometry (Figure 5B). The most nestin expression by the differentiated cells was in EEx and NEx groups (24.8 and 17.4%, respectively). There was no evident expression of nestin in the E and N groups (10 and 10.6%, respectively).

To evaluate the expression of neuronal-specific proteins, the cells of all the groups were qualitatively and quantitatively analyzed by immunofluorescence and flow cytometry (Figure 6). The expression of pan-neuronal markers, β-III tubulin, and synaptophysin was confirmed in all groups except E (Figures 6A, C). The results from flow cytometry analysis showed that in EEx and NEx groups, the expression of β-III tubulin (57.1 and 68.9%, respectively) and synaptophysin (69.2 and 81.5%, respectively) was at the highest level. While, the expression of β-III tubulin (20.2 and 35.1%, respectively) and

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Evaluation of Gene Expression by Real-Time PCR

In order to investigate the effects of various treatment conditions on the differentiation of P19 EC cells into neuron-like cells with emphasis on dopaminergic neuron-like cells phenotype, the gene expression profile of the cells was quantitatively analyzed by real-time PCR (Figure 4). Overall, the results showed that in the second week of the treatment, the expression levels of stem cell markers reduced significantly. At the same time, there was an enhancement in the expression of neuroepithelial, neural, and dopaminergic specific markers. At this time, the expression of Sox-2 was lowest in the N group, and the lowest level of Oct3/4 gene expression was in N, NEx, EEx, and Ex groups. Although the expression of Nanog decreased in the second week, it was at its lowest level in the NEx group in the third week (Figure 4A). Neuroepithelial (nestin) and neural-specific markers (synaptophysin, BDNF, NGF) and CREB tend to increase in the second week, and their highest expression was in the NEx group (Figure 4B). The expression patterns of dopaminergic specific genes (TH, Nurr1) and one-cut were similar to those of neuronal genes, so that they tend to elevate in the second week, and the highest level of their expression was in the NEx group (Figure 4C). The results demonstrated that with a few exceptions, nearly all the genes had a similar pattern of expression. Although stem cell markers showed a reduction pattern in their expression, the expression of Sox-2 increased significantly in the NEx group. Despite nestin, expression of synaptophysin and CREB decreased in the N group, but levels of all three genes increased in the NEx group. Interestingly, BDNF, NGF, TH, Nurr1, and one-cut genes were concert in their expression patterns, so that they had the highest expression level in the NEx group. The expression of other markers of dopaminergic neurons including Lmx1a, Lmx1b, Foxa2, and Girk2 (a marker preferentially expressed by neurons of the substantia nigra) was also evaluated (Figure 4D) in the second week of differentiation. The results confirmed the synergistic effects of Nurr1 overexpression and NRBE on Lmx1a, Lmx1b, and Girk2 level in the NEx group. The expression of Foxa2 was significantly increased in EEx and NEx groups. Since the results of Q-PCR confirmed that there was no significant difference between the two control groups, P19 and E, the subsequent analyses continued with only the E group as control.
synaptophysin (21.8 and 33.0%, respectively) was lower in E and N groups (Figures 6B,D) in comparison with the other groups.

Furthermore, in the second week of the differentiation protocol, the expression of dopaminergic specific proteins, tyrosine hydroxylase, and Nurr1 in the cells of all groups was analyzed by immunofluorescence and flow cytometry (Figure 7). The expression of these specific markers was confirmed in all groups except E (Figures 7A,C). Flow cytometry analysis showed that in EEx and NEx groups, the expression of TH (23.7 and 32.1%, respectively) and Nurr1 (76.5 and 73.9%, respectively) was at the highest level. While, the expression of TH (14.5 and 21.0%, respectively) and Nurr1 (24.6 and 27.9%, respectively) was lower in E and N groups (Figures 7B,D) compared to other groups.

Simultaneous expression of specific proteins in the differentiated cells was evaluated through double immunofluorescence (Figure 8). The results confirmed that the majority of β-III tubulin-expressing neural cells were also immunoreactive to tyrosine hydroxylase, which is a valid marker for the dopaminergic phenotype (Figure 8A). In addition, some differentiated cells expressed two dopaminergic specific proteins, Nurr1, and tyrosine hydroxylase, simultaneously (Figure 8B).

Yellow-stained cytoplasm in the representative fluorescence micrographs (arrows) illustrated co-expression of two proteins in the same cells.

**DISCUSSION**

In the present study, we have shown that combined molecular cloning and tissue-specific induction strategies could efficiently enhance the neural differentiation of stem cells with dopaminergic neuron-like phenotypes. In order to differentiation induction, P19 EC stem cells were stably transfected by a plasmid containing a dopaminergic key transcription factor, Nurr1, and then exposed to a medium supplied by a developing rat brain extract NRBE. The results demonstrated that although Nurr1 overexpression or NREB individually induced dopaminergic neuron-like differentiation, a combination strategy had more significant effects. Results from morphological, cellular, and molecular studies confirmed DA-like cell characterizations in the differentiated cells.
FIGURE 4 | Evaluation of gene expression profile in the experimental groups by quantitative Q-PCR. (A) The expression of stem cell markers tend to decrease in treatment groups. Sox-2 expression was lowest in the N group, and the lowest level of Oct3/4 gene expression was in N, NEx, EEx, and Ex groups during the second week of differentiation. Nanog gene expression was at its lowest level in the NEx group in the third week. (B) The level of neuroepithelial (nestin) and neural-specific genes increased in the second week with their highest expression in the NEx group. (C) TH, Nurr1 and one-cut followed nearly the expression pattern of neuronal genes, and their expression increased in the NEx group compared to the other groups. Despite Oct3/4 and Nanog, the expression of Sox-2 significantly enhanced in the NEx group. (D) During the second week of differentiation the expression of other specific markers of dopaminergic neurons including Lmx1a and Lmx1b was significantly increased in the NEx group. Foxa2 gene expression was at its highest level in the EEx group. Furthermore, Girk2; a marker preferentially expressed by neurons of the substantia nigra, showed significant enhancement in the NEx group. The experiments were carried out at least in triplicate. Bars represent standard error and different lowercase letters indicate significant differences by LSD test at \( P < 0.05 \), \( n = 3 \). For the variables with same letters, the difference is not statistically significant, and for the variables with different letters, the difference is statistically significant. P19, EBs with no treatment (α-MEM + 5% FBS); Ex, EBs treated by brain extract (α-MEM + 5% FBS + 100 µg/ml NRBE); E (empty), EBs transfected with pCMV3-GFPspark vector; EEx (empty/extract), EBs transfected with pCMV3-GFPspark vector and treated by extract; N (Nurr), EBs transfected with pCMV3-NR4A2-GFPspark vector; NEx (Nurr/extract), EBs transfected with pCMV3-NR4A2-GFPspark vector and treated by extract; W1, W2, W3, week 1, 2, 3, respectively.
Degeneration of DA neurons which are critical for controlling key functions of the brain, is associated with neurological diseases such as PD. Using drugs such as monoamine oxidase-B inhibitors, L-DOPA, and dopamine receptor agonists to treat this disease is not always possible for a long time due to their side effects. An alternative method for PD treatment is the replacement of degenerated cells by DA cells produced from stem cells (Politis and Lindvall, 2012). So far, various protocols have been reported to generate such cells for different aims including, studies on mechanisms of DA neuron development, PD pathogenesis, pharmaceutical approaches, and cell therapy.

Applying extrinsic growth factors, i.e., small molecules and soluble reagents in combination with molecular cloning strategies during in vitro differentiation of stem cells, provides an important tool to generate specific cells for PD therapy. Both extrinsic and intrinsic factors impact the determination of cell fate during development. Soluble extrinsic factors secreted by developing tissues in concert with intrinsic transcription factors provide a unique microenvironment for cell differentiation (Kim, 2011). For instance, differentiation of DA neurons requires the concerted action of extrinsic (Shh, FGF8, and Wnt1) and intrinsic (Foxa1/2, En1/2, Lmx1a/b, Pitx3, and Nurr1) signals (Joksimovic et al., 2009; Chakrabarty et al., 2012). To induce a certain differentiation fate in stem cells, essential intrinsic factors can be introduced into the cells through cloning strategies, and or soluble extrinsic factors can be added to the culture media as supplements. Kawasaki et al. (2000) showed that SDIA (stromal cell-derived inducing activity) was able to mediate stem cell differentiation into tyrosine hydroxylase-positive neurons without the use of either retinoic acid (RA) or EBs (Kawasaki et al., 2000). Furthermore, it reported that exogenous expression of Pitx3 could raise the gene and protein level of two neurotrophic factors, BDNF and GDNF (glial-derived neurotrophic factor) (Peng et al., 2007).
study, we showed that neonatal whole brain extract could induce DA phenotype with high expression of TH gene in P19 cells. In addition, LC-MS/MS analysis confirmed the presence of factors related to neuronal differentiation in NRBE content (Momendoust et al., 2019). The existence of cytokines, growth factors, and some other peptides in the developing and adult rat brain tissue has been proved previously (Pettmann et al., 1986; Bartlett et al., 1991; Lindholm et al., 1992; Lee et al., 1993; Mogi et al., 1994). This study aimed to address whether or not the overexpression of Nurr1 with a combination of tissue-specific microenvironments could efficiently induce DA neuronal differentiation of P19 cells. Therefore, Nurr1 as an intrinsic factor, and NREB as a source of soluble extrinsic factors applied to generate the desired neuronal phenotype in vitro.

Many studies demonstrated that orphan nuclear receptor Nurr1 plays a critical role in development, differentiation, and functional maintenance of midbrain DA neurons. It has been reported that Nurr1 gene overexpression in conjunction with factors derived from type 1 astrocytes of midbrain induced differentiation of a mouse neural stem cell line into DA neurons (Wagner et al., 1999). Furthermore, overexpression of paired like homeodomain 3 (Pitx3) enhanced the expression of BDNF and GDNF (Peng et al., 2007) and also induced TH expression (Messmer et al., 2007). A combination of Nurr1 and Pitx3 overexpression could differentiate mouse induced pluripotent stem cell (iPSC) into functional DA neurons (Salemi et al., 2016). The results of the present study revealed that the expression of neuronal and dopaminergic genes is upregulated in Nurr1-overexpressing cells (N group) compared to controls (P19 and E groups). However, this upregulation was significantly higher when P19 cells were exposed to NRBE (Ex group). The expression of those genes was even more remarkable in the NEx group (Nurr1-overexpressing cells treated by the extract) compared to all control and experimental groups. Indeed, using combined strategy had synergistic effects on DA differentiation of stem cells. Interestingly, the expression pattern of these genes was highly coordinated. The results confirmed the efficient expression of all DA neuron markers in the NEx group in the second week of the differentiation protocol. Previously, it has been reported that brain extracts have several activities concerning central (CNS) and peripheral (PNS) nervous system neurons. Combinations of NGF and rat brain extract have synergistic effects on the

FIGURE 6 | Qualitative and quantitative assessment of neuronal-specific proteins, β-III tubulin, and synaptophysin in the second week of the differentiation protocol. (A) The cells of all groups except E were immunoreactive to β-III tubulin. (B) Flow cytometry results showed a higher level of β-III tubulin protein in NEx, EEx, and N groups compared to the E group. (C) The cells of all groups except E were immunoreactive to synaptophysin. (D) Flow cytometry results showed a higher level of synaptophysin protein in NEx, EEx, and N groups compared to the E group. DAPI was applied to counterstain the nuclei. BT, β-III tubulin; SYN, synaptophysin; E (empty), EBs transfected with pCMV3-GFPSpark vector; EEx (empty/extract), EBs transfected with pCMV3-GFPSpark vector and treated by extract; N (Nurr), EBs transfected with pCMV3-NR4A2-GFPSpark vector; NEx (Nurr/extract), EBs transfected with pCMV3-NR4A2-GFPSpark vector and treated by extract.
survival of chick embryo DRG (dorsal root ganglion) cells (Sensenbrenner et al., 1978). Initial studies have shown that rat brain extract promoted neurite outgrowth from explants of fetal rat retina in culture (Turner et al., 1982). Brain extract contains many soluble factors, including glial growth factor (GGF), N-Acetylaspartate (NAA), N-acetyltransferase (NAT), C-type natriuretic peptide (CNP), and protein kinase C (PKC). GGF stimulates the proliferation of purified rat Schwann cells and astrocytes in tissue culture (Brain extract). NAA was initially identified in brain extracts in 1956 (Tallan et al., 1956). In addition, NAT and its corresponding gene, Nat8l, are expressed in the brain (Truckenmiller et al., 1985; Ariyannur et al., 2010; Wiame et al., 2010). PKC, first identified in bovine brain extracts (BBE), can regulate cell proliferation, differentiation, and motility. BBE stimulates the growth and expression of cell-specific genes in specialized cell types, including endothelial cells, smooth muscle cells, keratinocytes, melanocytes, and hybridomas, primary neuronal cells. It also promotes the differentiation of pluripotent stem cells (Hoshijima et al., 2004).

These findings are in accordance with previous studies demonstrating that tissue-specific microenvironment could induce tissue-specific gene expression in stem cells (Bentz et al., 2010; Ebrahimie et al., 2014; Narayanan et al., 2014; Mansouri et al., 2016; Mehrfarjam et al., 2016). Our results showed that although overexpression of Nurr1 could induce neuronal and dopaminergic markers in stem cells, exposure of the cells to brain extract had synergistic effects on this process. While Nurr1 expression is critical for DA neuronal development, solely its expression is not sufficient to stimulate this phenotype. It has demonstrated that to reach full mature dopaminergic neurons from in vitro differentiation of stem cells, additional factors are required (Kim et al., 2003). Therefore, it suggested that the presence of possible factors/proteins in NRBE is involved in regulating Nurr1 gene function. Soluble factors such as BDNF, GDNF, and TGFβ (transforming growth factor β) all play an essential role in the differentiation and survival of dopaminergic neurons and also maintenance of their normal functions. Soluble extract derived from striatum could enhance in vitro development of mesencephalic DA neurons (Tomozawa and Appel, 1986). The production of DA neurons from stem cells has been stimulated by fetal midbrain astrocytes (Roy et al., 2006). GDNF, neurturin, TGF-β, and dibutyryl-cAMP as soluble
factors promoted the differentiation, maturation, and survival of TH-positive neurons derived from ES cells (Rolletschek et al., 2001). Furthermore, interleukin-1 could significantly increase the number of TH-expressing cells isolated from the midbrain of rat embryos (Ling et al., 1998).

As mentioned above, the expression of neuronal and dopaminergic markers significantly increased in the second week of differentiation. It is possible that BDNF, which is involved in the neuroprotective pathways of dopaminergic neurons, is highly associated with Nur1. Among the genes that CREB controls are BDNF, TH, and Nur1. Interaction of CREB with its transcriptional binding site of Nur1 promoter induces the upregulation of this gene (Lonze and Ginty, 2002; Lei et al., 2011; Dong et al., 2016; Jang et al., 2021). Expression analysis of Onecut1 (Oc1, also called Hnf6) transcription factor during our differentiation system showed the coordinated expression pattern of this gene with other DA-specific genes. The role of Oc transcription factors in dopaminergic development is not yet clear exactly. Matched Oc and Lmx1a expression profiles suggested the involvement of Oc in early DA development (Chakrabarty et al., 2012). The exact functions of the Oc family and their interaction with other critical factors in the development of DA are not yet fully understood and require further investigation.

It is interesting to note that concomitant with an enhancement of neuronal and dopaminergic markers, the expression of stem cell markers was down-regulated. However, the expression of Sox-2 in the NEx group increased exceptionally compared to the other groups. The core transcription factors for regulating pluripotency are Oct3/4, Sox-2, and Nanog (Chen et al., 2008). Furthermore, Sox-2 plays an essential role in the neural differentiation of stem cells and maintaining the properties of neural progenitor cells. Sox-2 can control pluripotency and neural differentiation of human pluripotent stem cells (Khoo et al., 2013; Zhang and Cui, 2014). Downregulation of Sox-2 in the differentiated cells derived from stem cells has been well-established previously (Ebrahimie et al., 2014; Mehrfarjam et al., 2016; Momendoust et al., 2019). However, it is reported that depending on the environmental input, Sox-2 could function as a pluripotency and, or neural differentiation factor (Zhang et al., 2019). Therefore, the high expression of Sox-2 here might reflect its possible role in neural differentiation rather than stem cell proliferation.
CONCLUSION

To the best of our knowledge, this work is the first attempt to evaluate the combined effects of Nurr1 gene overexpression with a natural tissue-specific medium on dopaminergic differentiation of P19 EC stem cells. To this end, NREB as a source of extrinsic factors, and Nurr1 as an intrinsic factor, was applied for differentiation induction. Overall, the results of the present study demonstrated the synergistic effects of Nurr1 gene overexpression and possible soluble factors that existed in NREB on differentiation induction of P19 stem cells into neuronal cells with a dopaminergic neuron-like phenotype. Although therapies based on molecular targets for PD sound promising, the feasibility and safety of Nurr1 gene therapy still need further improvement.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

ETHICS STATEMENT

The animal study was reviewed and approved by Bioethics Committee of the University of Isfahan, based on the National Specific Ethical Guidelines for Biomedical Research issued by the Ministry of Health and Medicinal Education (MOHME) of Iran in 2005.

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AUTHOR CONTRIBUTIONS

FE conceived and designed the experiments, analyzed the data, and wrote the paper. RB, MK, FE, and FD performed the experiments. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

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