Differential display analysis of gene expression accompanied by neurite outgrowth of human neuroblastoma Cell IMR32 using non-gel molecular sieving capillary electrophoresis

N. Ishioka1,*, Y. Kurosu2,3, A. Kuhara4, T. Kogure4,5, Y. Ueno4,6, M. Saito4,6, K. Watabe7 and I. Nagaoka2

1Space Utilization Systems, Tsukuba Space Center, National Space Development Agency of Japan (NASDA), 2-1-1, Sengen, Tsukuba-shi, Ibaraki 305-8505, Japan
2The department of Biochemistry, Juntendo University, School of Medicine, 2-1-1, Hongo, Bunkyo-ku, Tokyo 113-0033, Japan
3JASCO Technical Research Laboratories Corporation, 2097-2, Ishikawa-cho, Hachioji-shi, Tokyo 192-0032, Japan
4Division of Molecular Cell Biology, Institute of DNA Medicine, The Jikei university School of Medicine, 3-25-8, Nishi-Shinbashi, Minato-ku, Tokyo 105-0003, Japan
5Department of Neuro Surgery, The Jikei university School of Medicine, 3-25-8, Nishi-Shinbashi, Minato-ku, Tokyo 105-0003, Japan
6Department of Orthopedic Surgery, The Jikei university School of Medicine, 3-25-8, Nishi-Shinbashi, Minato-ku, Tokyo 105-0003, Japan
7Department of Neuropathology, Tokyo Metropolitan Institute for Neuroscience, 2-6, Musashidai, Fuchu-shi, Tokyo 183-0042, Japan

Abstract. The induced proliferative response of the human derived neuroblastoma cell (IMR32), with the cell proliferation mediating factor, is much milder compared to other cell strains, and develops cellular clusters which are characterized by numerous extensions of dendrites. Presuming this phenomenon to be one of the induced neuronal differentiation due to genetic alterations, we preliminary studied to elucidate the changes in gene expression of the IMR32 by mRNA differential display analysis, monitoring polymerase chain reaction (PCR) products by non-gel molecular sieving capillary electrophoresis in linear polyacrylamide solution. The CE fingerprints revealed a number of peaks with differential expression patterns.


Introduction

Most cell proliferation, differentiation, and other developmental processes are characterized by changes in gene expression. Identification of such changes is very important to understand the gene regulatory mechanism. Several polymerase chain reaction (PCR)-based methods have been offered to identify such differential changes [1-10]. One of the developmental techniques of such methods is a mRNA differential display [3,4,6,9,10] which consists of two steps. The first step is the reverse transcription of total cellular RNA to single-strand cDNA using a 3'-anchored oligo-dT primer. The second step is the amplification of a subset of cDNAs using the same 3'-anchored primer and an arbitrary primer. The rapid analysis for the differences in expression between different cell types would require a highly reproducible system for peak patterns of PCR products. Currently, the slab gel electrophoresis is used, and band patterns are detected by autoradiographs or fluorescent stains in agarose and/or polyacrylamide gels. It takes relatively long time to get results and scanning of band patterns is also very time consuming. Furthermore the PCR enables the amplification of specific cDNAs that are often present in trace amounts of biological material, therefore a high-resolution tool which provides an alternative to conventional slab gel electrophoresis is required for the analysis of the PCR products.

* Correspondence and reprints
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Capillary electrophoresis (CE), with its high-resolution power, speed, ability to analyze small sample volumes and automation, is a widely utilized technique in the field of life science [11-13] and has great potential for analysis of the PCR products [14-19]. CE also provides the capability for quantitation in real-time and on-line detection, and even fractionation of peaks [12,13]. In this study, we attempted to characterize the genetic alterations and employed the molecular sieving CE in linear polyacrylamide for mRNA fingerprinting and for detecting differentially expressed genes accompanied by neurite outgrowth of human neuroblastoma cell IMR32.

In molecular sieving CE, there are two methods to realize the separation by gel filtration mode. One is to form gel, which is similar to conventional gel, in the capillary as a sieving medium. The other is to pack non-cross-linked medium in the capillary instead of conventional gel to perform gel filtration. In this method, non-branched or branched macromolecule solution such as linear polyacrylamide, cellulose derivatives, dextran are used as sieving media. The greatest advantage of use of non-gel for CE is that the sieving medium can be changed easily for every run according to analytical requirements. Considering the feature, it is expected that CE using non-gel as sieving media will be more commonly used in the future.

Experimental

IMR32 cell culture and induced neuronal differentiation

Human neuroblastoma IMR32 cells (Dai-Nippon Pharmaceutical Co., Osaka, Japan) were cultured in RPMI1640 medium supplemented with 10% fetal bovine serum (FBS), 100 U/mL of penicillin G and 100 µg/mL of streptomycin in an atmosphere of 5% CO2-95% air at 37 °C. After the cells reached confluence in RPMI1640 with 10% FBS, cells were separated in half. Each cell fraction collected by low-speed centrifugation and washed twice with PBS to remove remaining medium components. One of the half was suspended again in RPMI1640 with 10% FBS and the other was suspended in RPMI1640 with 10% FBS for the cell proliferation mediating factor (Daigo’s GF21; Wako, Osaka, Japan) instead of FBS. The GF21 was commercially provided by adding insulin, transferrin, sodium selenite, and ethanolamine for cell growth [19] to adult bovine serum provided by adding insulin, transferrin, sodium selenite, and ethanolamine for cell growth [19] to adult bovine serum (75 µm i.d., 50 cm total length, 30 cm effective length) was mixed with 50 pmol of 3'-anchor primer and stored at –20 °C until use. One of the arbitrary primers (0.5 µM at final concentration) was added into the reaction mixture (20 µL) for PCR which contained 2 µL of the cDNA solution described above, 50 µM dNTPs and 1 unit of DNA polymerase Gene Taq in the kit. The second strand was synthesized at 95 °C for 3 min, 40 °C for 5 min and 72 °C for 5 min, and the amplification was repeated by 24 cycles of a thermal cycling program which was composed of 95 °C for 15 sec, 40 °C for 2 min and 72 °C for 1 min. An additional final extension step was done at 72 °C for 5 min. The PCR products were cleaned-up and desalted using a Microcon 100 (Amicon, Beverly, MA, USA), according to the reported method [18]. The resulting samples were subjected to CE analysis.

Non-gel molecular sieving capillary electrophoresis in linear polyacrylamide solution

Capillary electrophoresis (CE) was performed on the JASCO model CE-800 system (JASCO, Tokyo, Japan). A linear polyacrylamide solution-filled, uncoated fused-silica capillary (75 µm i.d., 50 cm total length, 30 cm effective length) was prepared as follows. The capillary was filled with 12% acrylamide in a buffer of 100 mM Tris/250 mM boric acid/7M urea, pH 8.1, containing 0.1% potassium persulfate (PPS) and 0.1% N,N,N’,N’-tetramethylethylenediamine (TEMED) as polymerization initiator and catalyst, respectively. Both ends of the capillary tube were dipped into the non-gel solution until completely polymerized (c.a. 3h). After polymerization, one end (injection side) of capillary tube was replaced into a buffer (100 mM Tris/250 mM boric acid, pH 8.1) reservoir attached to the negative side of the power supply and the other end (detection side) was placed as it is, in the non-gel solution attached to the positive side, providing new solution through electroosmotic flow. The sample was introduced into the capillary by electrokinetic injection (–5 kV for 40 s) from the cathode side, and electrophoresis was performed at –15 kV constant voltage (–300 V/cm) for 70 min at room temperature and was monitored simultaneously at 260 nm at the anode side. The DNA Hae III digests (500 µg/mL, BRL Labs., Gaithersburg, MD, USA) were used for standard DNA ladder which was dissolved in 10 mM Tris-HCl (pH 9.7), 1.0 mM EDTA, 20 mM NaCl.

Total RNA isolation

Total RNA was isolated from cultured cells by the RNAzol B RNA isolation kit (TEL-TEST, Inc., Friendwood, TX, USA) according to manufacturer’s instruction.

mRNA differential display

The differential display was performed by the mRNA fingerprinting kit (Wako, Osaka, Japan) based on the reported method [9]. Total RNA (2.5 µg) was mixed with 50 pmol of 3'-anchor primer and incubated at 70 °C for 10 min, and then the reaction mixture (10 µL) was chilled on ice for several minutes. To this solution, 10 µL of 20 mM Tris-HCl, pH8.4, containing 1 mM dNTPs, 5 mM MgCl2, 20 mM DTT and 200 units of reverse transcriptase in the kit was added. The mixture was incubated at 25 °C for 10 min and at 42 °C for 50 min in thermocycler (ASTEC PROGRAM TEMP CONTROL PC-800; ASTEC, Fukuoka, Japan). At the end of this incubation, the temperature was raised to 70 °C for 15 min and then reduced to 4 °C. The reaction mixture (cDNA solution) was diluted with 80 µL of TE buffer (10 mM Tris-HCl/1 mM EDTA, pH7.5) containing 9 µl of the same 3'-anchor primer and stored at –20 °C until use. One of the arbitrary primers (0.5 µM at final concentration) was added into the reaction mixture (20 µL) for PCR which contained 2 µL of the cDNA solution described above, 50 µM dNTPs and 1 unit of DNA polymerase Gene Taq in the kit. The second strand was synthesized at 95 °C for 3 min, 40 °C for 5 min and 72 °C for 5 min, and the amplification was repeated by 24 cycles of a thermal cycling program which was composed of 95 °C for 15 sec, 40 °C for 2 min and 72 °C for 1 min. An additional final extension step was done at 72 °C for 5 min. The PCR products were cleaned-up and desalted using a Microcon 100 (Amicon, Beverly, MA, USA), according to the reported method [18]. The resulting samples were subjected to CE analysis.
Results and discussion

Effects of GF21 on IMR32 cells

Human neuroblastoma IMR32 cells were cultured 7 days in FBS- and GF21-media, and cell numbers were determined after 7 days. On day 7 cell numbers in GF21-medium was about one and third of cell numbers in FBS-medium. Proliferation of IMR32 cells was modest in GF21, although GF21 was commercially sold as a cell proliferation mediating factor. From day 4 onwards, by contrast, the effect of GF21 on cell differentiation in term of neurite outgrowth became obvious, and the culture displayed quite different morphologies after 7 days (Fig. 1B). In GF21-medium, IMR32 cell bodies formed large clusters and grew a dense network of neuritic processes. From the cluster with cell aggregates, long and thick fiber bundles extended and contacted other cell clusters. Under FBS-medium, changes in cellular morphologies were not apparent on day 7 (Fig. 1A). These results presumes that the morphological change on IMR32 cultured in GF21-medium was one of induced neuronal differentiation.

Analysis of mRNA differential display by molecular sieving CE

The CE electrophoregram of the φx174 DNA Hae III digests for standard DNA ladder was shown in figure 2. The CE conditions were designed to enhance the separation efficiency for fragments in the 72 – 872 bp range, because, according to the instruction of the kit, DNA polymerase gene Taq in the kit was suitable for amplification of the cDNA ranging from 100 to 800 bp in this study. The R.S.D. values of migration and peak areas for 194 bp fragment, were 0.60% (n = 5) and 1.30 (n = 5), respectively. Thus, this separation system produced highly reproducible results, and was a useful tool for DNA analysis [20].

A series of primer combinations were utilized to generate gene expression profiles for normal cells and morphologically changed cells. The three sets of primers are shown in figure 3 as examples. Examination of these CE patterns revealed many developmental changes in gene expression and some of these were indicated. Total 51 amplicon peaks in figure were indicated, and the 41.2% of the total amplicons was diminished and 19.6% was enhanced in morphologically changed cells. The expression of the others (39.2%) was essentially constant at each cell. The enhanced amplicons were most prominent in the morphologically changed cells and were absent in the normal cells. Especially, a new peak, which was an about 260 bp amplicon (an asterisk in Fig. 3), possibly was a candidate for differentially expressed gene. However, for morphological
changes, it might also be an important factor that many dominant gene expressions are suppressed in cells cultured in GF21-medium. In any case, these may represent maternal mRNAs, the identity and function of which is almost unknown in the differential cells.

**Conclusion**

We have attempted to use molecular sieving CE as an alternative to conventional slab gel electrophoresis for the mRNA differential display. The results shown in figure 3 demonstrate the potential of CE for mRNA differential display analysis. Furthermore, in molecular sieving CE, micro-preparation of characteristic amplicons is possible for further study. Considering the features, it is expected that CE analysis for the mRNA differential display will be more commonly used in the future.

However, at present, there are still several problems to be solved: (1) Better resolution will be required in the region from low to high molecular weight; (2) the sensitivity of CE must be improved; (3) the difference of mRNA fingerprints between by slab gel electrophoresis and by molecular sieving CE must be investigated in detail.

At present, we are trying to find solutions to the problem outline above.

**References**