Ectopic expression of Id1 or Id3 inhibits transcription of the GATA-4 gene in P19CL6 cells under differentiation condition

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SUMMARY  Inhibitor of DNA binding (Id) is a dominant negative form of the E-box binding basic-helix-loop-helix (bHLH) transcription factor since it is devoid of the basic region required for DNA binding and forms an inactive hetero dimer with bHLH proteins. The E-box sequence located in the promoter region of the GATA-binding protein 4 (GATA-4) gene is essential for transcriptional activation in P19CL6 cells. These cells differentiate into cardiomyocytes and start to express GATA-4, which further triggers cardiac-specific gene expression. In this study, expression plasmids for Ids tagged with human influenza hemagglutinin (HA)-FLAG were constructed and introduced into P19CL6 cells. The stable clones expressing the recombinant Id proteins (Id1 or Id3) were isolated. The GATA-4 gene expression in these clones under differentiation condition in the presence of 1% dimethyl sulfoxide (DMSO) was repressed, with concomitant abolishment of the transcription of α-myosin heavy chain (α-MHC), which is a component of cardiac myofibrils. Thus, the increased expression of Id protein could affect GATA-4 gene expression and negatively regulate the differentiation of P19CL6 cells.

Keywords  Inhibitor of DNA binding factor, GATA transcription factor, cardiomyocyte differentiation, transcription

1. Introduction

It is well known that many gene products participate spatio-temporally in a complex manner in the developmental process (1). In mouse heart formation, transcription factor GATA-binding protein 4 (GATA-4) together with NK2 homeobox 5 (Nkx2.5) and T-box factor 5 (Tbx5), and chromatin remodeling protein SMARCD3 (BAF60c) are key upstream regulators of cardiomyocyte differentiation in the cardiogenic splanchnic mesoderm (2). Expression of GATA-4 in the extraembryonic endoderm in the embryo is required for folding morphogenesis to form a primitive heart tube and foregut (3). Consistent with the expression of GATA-4 in cardiomyocytes at all stages of cardiac development in mouse (4), it is reported that mutations of the GATA-4 gene lead to variable phenotypes of human congenital heart diseases (5).

To study cardiomyocyte differentiation at the cellular level, P19 mouse embryonic carcinoma cells and their derivative P19CL6 cells have been used because they easily start differentiating into beating cardiomyocytes upon the addition of dimethyl sulfoxide (DMSO) (6). It was further demonstrated that expression of the GATA-4 gene is required for the differentiation of cardiomyocytes (7). As for transcriptional regulation, the GATA-4 gene is transcribed from two alternative 5’ untranslated exons (E1a and E1b) (8,9), and the level of the E1a transcript of mouse heart is 20-times higher than that of E1b (8). Analyses of the promoter for E1a demonstrated that the conserved E-box sequence (CACGTG) is essential for transcription from the E1a exon (10,11), suggesting that the basic-helix-loop-helix (bHLH) transcription factor(s) may play a role in transcription of the GATA-4 gene (9,12). It is also reported that more euchromatic state of the further upstream region of the cluster of GC-boxes and E-box activates the GATA-4 promoter (13).

The inhibitor of DNA binding (Id) proteins (Id1-Id4) which do not have the amino-terminal basic region of bHLH transcription factor function as a dominant negative form of bHLH proteins, and their functions seem to be redundant or compensatory although Ids are required for heart development (14). Ids are also suggested to play a role in keeping precursor cells immature and expanding the cell population size during development (15). Id1 and Id3, although the latter is less active, are essential for specification of mesendosomal progenitors into cardiac progenitors in the first heart field (16). Biochemical studies further demonstrated that Id1, Id2 and Id3 bind GATA-4 and Nkx2.5 proteins and...
inhibit their synergistic transcriptional activation due to inhibition of DNA binding and/or mutual interaction (17). Stable expression of Id3 ectopically in P19 cells inhibited the differentiation concomitant decrease of GATA-4 expression (17). In this study we constructed expression plasmids for tagged-Id1, Id2 and Id3, and tried to express these tagged proteins stably in P19CL6 cells to determine whether their differentiation into cardiomyocytes is inhibited or not.

2. Materials and Methods

2.1. Cell culture

P19CL6 cells (RIKEN Cell Bank, Tsukuba, Ibaraki, Japan) were cultured at 37°C in α-Eagle’s minimal essential medium (MEM) (Sigma, St. Louis, MO, USA) containing 10% (v/v) fetal bovine serum (FBS) (GIBCO BRL, Gaithersburg, MD, USA) and antibiotics [2.5 μg/ml fungizone (GIBCO BRL), 100 units/mL benzylpenicillin (Wako, Osaka, Japan), and 100 μg/mL streptomycin sulfate (Wako)]. Cells (1 × 10⁵ cells in a 10 cm dish) were allowed to differentiate in the presence of 1% (v/v) DMSO (Wako) (10). The medium was changed 4 days after inoculation and then every 2 days.

2.2. Analysis of RNA

Total cellular RNA was extracted with Isogen (Nippon Gene, Toyama, Japan) from a 10 cm dish according to the manufacturer's protocol, and an aliquot (5 μg) was reverse transcribed with Moloney Murine Leukemia Virus (M-MLV) Reverse Transcriptase (TaKaRa, Kusatsu, Shiga, Japan) and the oligo(dT)₁₅ primer in a volume of 20 μL. After RNase H (TaKaRa) treatment, cDNA (0.3 μL) was subjected to semi-quantitative PCR analysis (18) with GoTaq® DNA polymerase (Promega, Madison, WI, USA). The unsaturated conditions for polymerase chain reaction (PCR) comprised preheating (94°C, 3 min), followed by 17-27 cycles of denaturation (94°C, 0.5 min), annealing (55-61°C, 0.5 min), and extension (72°C, 0.5 min), and then post-incubation (72°C, 5 min): typically, GATA-4 (27 cycles, 55°C), α-myosin heavy chain (α-MHC) (24 cycles, 60°C), Id1 (21 cycles, 61°C), Id2 (21 cycles, 60°C), Id3 (21 cycles, 60°C), and β-actin (17 cycles, 60°C). The primer pairs used for PCR are shown in Table 1A. The PCR products were size-separated by 2% (w/v) agarose gel-electrophoresis, and DNA bands were visualized with ethidium bromide. Images were recorded with a FAS-III UV-imaging system (Toyobo, Osaka, Japan).

2.3. Cloning of cDNAs for mouse Id1, Id2 and Id3

P19CL6 cells were cultured for 12 days in the presence of 1% (v/v) DMSO. cDNA was synthesized from total cellular RNA as prepared in 2.2., and was subjected to PCR using a primer pair [YY001/YY002 to amplify cDNA for the coding region of Id1 cDNA, YY003/YY002 for Id2, or YY005/YY006 for Id3 (Table 1B)] and GoTaq® DNA polymerase. The PCR conditions were [preheating (94°C, 3 min), followed by 30 cycles of denaturation (94°C, 0.5 min), annealing (60°C for Id1, 51°C for Id2 and 63°C for Id3, 0.5 min), and extension (72°C, 0.5 min), and then post-incubation]

In (B), an Ndel site (underlined) was introduced at the initiation codon in the sense primer (the primer S in Figure 1). The primer A in Figure 1 showed the antisense primer. Capital letters in (B) indicate the nucleotide residues identical to those in the coding region, while lower case letters indicate unrelated residues introduced.

Table 1. Sequences of oligonucleotides used in this study

In (B), an Ndel site (underlined) was introduced at the initiation codon in the sense primer (the primer S in Figure 1). The primer A in Figure 1 showed the antisense primer. Capital letters in (B) indicate the nucleotide residues identical to those in the coding region, while lower case letters indicate unrelated residues introduced.

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with a sequence Φluc.

2.5. Transfection of expression plasmids for Ids tagged with HA and FLAG

We first expressed the tagged Id proteins (HA/FLAG-Ids) transiently by means of the diethylaminoethyl (DEAE)-dextran method (22) to verify the expression of tagged Ids derived from the plasmid constructs; Cos-1 cells (ATCC, Manassas, VA, USA) were grown in Dulbecco's modified Eagle medium (DMEM) (GIBCO BRL) supplemented with 7% (v/v) FBS. Cells (5 × 10^5 cells) in 5 mL medium were plated onto a 6 cm dish. Into each dish the plasmid construct was introduced at a concentration of 2.5 μg/dish. Cells were treated with chloroquine followed by glycerol, and then further cultured for 2 days. The whole cell extract for immunoprecipitation was prepared from duplicate samples. Protein concentrations were determined with a BioRad Protein Assay Kit (Hercules, CA, USA) using bovine serum albumin (Fraction V) (Sigma) as a standard (23).

We further isolated P19CL6 clones that express tagged Ids stably by the calcium-phosphate method (21); briefly, cells (5 × 10^5 cells) were plated onto a 6 cm dish. On the next day, the DNA solution was poured onto the culture medium and left for 4 hr. The medium was removed and washed with phosphate-buffered saline [10 mM phosphate buffer (pH7.2), 137 mM NaCl, 3 mM KCl] (PBS). Cells were further cultured for fresh medium for 24 hr, and then neomycin (G418) (100 μg/mL) (Wako) was added. On the following day, cells were plated onto 2 dishes (5 × 10^5 cells) in 5 mL medium were plated onto a 6 cm dish. Into each dish the plasmid construct was introduced at a concentration of 2.5 μg/dish. Cells were treated with chloroquine followed by glycerol, and then further cultured for 2 days. The whole cell extract for immunoprecipitation was prepared from duplicate samples. Protein concentrations were determined with a BioRad Protein Assay Kit (Hercules, CA, USA) using bovine serum albumin (Fraction V) (Sigma) as a standard (23).

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antibiotic. After ~12 days, single colonies were isolated. A whole cell extract was prepared from a confluent culture of undifferentiated clones grown in two Φ 6 cm dishes.

2.6. Preparation of a whole cell extract and immunoprecipitation

After washing with ice-cold PBS three times, cells were suspended in 1 mL 20 mM tris(hydroxymethyl)aminomethane (Tris)-HCl (pH 7.5), 150 mM NaCl, 2 mM ethylenediaminetetraacetic acid containing 10 μg/mL leupeptin, 10 μg pepstatin A and 1% (v/v) NP-40 [TNE (+)] and kept on ice for 30 min. All the following procedures were carried out at 4°C. The cell suspension was sheered through a 25G needle 10 times, and then centrifuged (12,000× g, 30 min). The supernatant was used as the whole cell extract.

The Protein G Sepharose beads (GE Healthcare, Chicago, IL, USA) were pre-washed twice with TNE (+) and then centrifuged (2,000× g, 5 min). The whole cell extract was incubated for 1 hr in a Mini Disk Rotor BC-710 (BIO CRAFT, Tokyo, Japan) with a 30 μL bed volume of pre-washed Protein G Sepharose beads. The supernatant was incubated in a Rotor for 1 hr with 5 μL of Monoclonal anti-HA (HA-7) (Sigma), and then further incubated in a Rotor with a 50 μL bed volume of pre-washed Protein G Sepharose beads for 1hr. The beads were precipitated (2,000× g, 5 min) and then washed five times (three times by rotation and then twice without rotation) with 500 μL 20 mM Tris-HCl (pH 7.5), 150 mM NaCl, 2 mM ethylenediaminetetraacetic acid (TNE). The recovered immuno-complex was heated at 95°C for 5 min together with 15-20 μL of 2 × sample buffer for sodium dodecyl sulfate (SDS)-polyacrylamide gel-electrophoresis (24).

2.7. SDS-polyacrylamide gel-electrophoresis and Western blotting

All the solubilized protein was subjected to SDS-polyacrylamide gel-electrophoresis [mini-gel (10 cm × 10 cm), 1 mm thickness, consisting 3% (w/v) stacking gel (60 V) and 18% (w/v) separation gel (100 V)] (24), and then electro-blotted [200 mA, 90 min.; ATTO Model-AE6675 (ATTO, Tokyo, Japan)] onto an Immobilon™-P membrane [Millipore (Billerica, MA, USA) polyvinylidene difluoride (PVDF) membrane (0.45 μm), IPVH00010] (25). The filter was washed with PBS and then blocked 1 hr at 4°C with PBS plus 0.1% (v/v) Tween 20 (PBS-T) containing 5% (w/v) skim-milk (BD, Franklin Lakes, NJ, USA). The filter was washed with PBS-T and then reacted for 2 hr at room temperature with ANTI-FLAG M2® monoclonal antibody-peroxidase conjugate (Sigma) (× 1,500 diluted). Chemiluminescence was detected with an ECL Western blotting kit (GE Healthcare) using Scientific Imaging Film (KODAK, Rochester, NY, USA).

2.8. Chemicals

Restriction enzymes were obtained from New England Biolabs (Ipswich, MA, USA) and Toyobo. The Klenow enzyme, T4 DNA ligase (Ligation Kit Ver2.1), and Agarose-LE Classic Type were provided by TaKaRa. Tween 20 and chloroquine diphosphate were obtained from Nacalai. DEAE-dextran, NP-40, leupeptin and pepstatin A were from Sigma. Oligonucleotides were purchased from Gene Design Inc. (Ibaraki, Osaka, Japan). All other chemicals used were of the highest grade commercially available.

3. Results

3.1. Construction of expression plasmids for Ids

The cDNAs for mouse Ids (Id1, Id2 and Id3) were cloned from P19CL6 cells by means of reverse transcription PCR (RT-PCR) and then inserted into a mammalian expression plasmid pDsRed2-N1. To detect ectopic expression, we introduced a HA/FLAG-tag at the amino terminus of each Id. The neomycin-resistance marker was also replaced with that of hygromycin (Figure 1). Plasmids were introduced into Cos-1 cells, and the Id proteins transiently expressed were detected by Western blotting to verify their molecular sizes. Since we sometimes experienced difficulty in detecting the tagged Id proteins in a whole cell extract, an immunoprecipitation method was developed, as shown under Materials and Methods. The HA/FLAG-Id in the whole cell extract was first immunoprecipitated with anti-HA antibody and then detected by means of Western blotting with anti-FLAG antibody. As shown in Figure 2, the calculated molecular size of the tagged-Id (see legend to Figure 2) was not significantly different from its molecular size on the gel.

3.2. Stable expression of Ids in P19CL6 cells

The expression plasmids for tagged Ids were introduced into P19CL6 cells and neomycin-resistance colonies were isolated. However, most of the resistant clones did not express HA/FLAG-Id in spite of several trials of transfection. Typical results are shown in Figures 3A and 3B; two clones for tagged Id1 (Id1α and Id1β) and one for tagged Id3 (Id3α) were obtained in two separate transfection experiments. However, clones for tagged Id2 could not be obtained. We also did not obtain a clone from hygromycin-resistant colonies.

3.3. Transcription level of GATA-4 under differentiation condition for P19CL6 cells

As already reported, many preceding studies involving
cells, similar (pneo-HA/FLAG-Id3), N, (pneo-HA/FLAG-Id1), N, (pneo-HA/FLAG-Id2), N, (pneo-HA/FLAG-Id3), and C (without plasmid)). Cells were grown for 2 days, and then the transiently expressed HA/FLAG-Id protein in the whole cell extract was immunoprecipitated with anti-HA antibody. HA/FLAG-Id protein was detected by Western blotting with peroxidase conjugated anti-HA antibody after SDS-polyacylamide gel-electrophoresis as described under Materials and Methods. Chemiluminescence was detected after 15 min exposure. The average molecular mass of the tagged Id protein was calculated with GENETYX-MAC GENETIC INFORMATION PROCESSING SOFTWARE (GENETYX Corporation, Tokyo, Japan); HA/FLAG-Id1, 19.1 kDa; HA/FLAG-Id2, 18.5 kDa; and HA/FLAG-Id3, 16.7 kDa. The values were the sums of the amino acid residues of the native Id (148, 134 and 119 residues for Id1, Id2 and Id3, respectively) and HA/FLAG peptide (31 residues) (20). The Id1 used in this study was a smaller isoform produced by removing a coding intron (49).

P19 and P19CL6 cells revealed that the transcription of GATA-4 gene starts at around 4 days under differentiation conditions, that of the genes for the components of cardiac muscle following later (7,26-28). Figure 4 (left) shows that GATA-4 mRNA was actually detected within 4 days after DMSO addition and that of α-MHC appeared at 8 days weakly and at 12 days clearly. However, transcription of the GATA-4 gene did not occur in Id1A5 and Id3A5 cells even in the presence of DMSO, ectopically HA/FLAG-Id1 and HA/FLAG-Id3 being expressed, respectively (Figure 4, middle and right). As the transcription of cardiac muscle genes such as α-MHC, β-MHC, cardiac troponin C (cTpc), atrial myosin light-chain 1 (MLC-1A) and ventricular MLC-1 (MLC-1V) is positively regulated by GATA-4 (29), the expression of α-MHC was not induced in Id1A5 and Id3A5 cells without GATA-4 expression (Figure 4). It must be further noted that the GATA-4 and α-MHC genes were also repressed in Id1A5 cells, similar to in Figure 4 (middle) under differentiation condition (not shown), which may suggest that ectopic Id1 reproducibly inhibits GATA-4 gene transcription under our experimental condition.

The Ids were transcribed even in an undifferentiated state, as shown at Day 0 in Figure 4, as the Ids were usually detected in the control samples without DMSO (17,28,30). Regarding the behavior of the expression pattern in our study, the mRNA level of Id2 was clearly increased in a later stage of culture in the presence of DMSO independently of the forced expression of Id1 and Id3 (Days 8 and 12 in Figure 4). The mRNA levels of Id1 and Id3 seemed to increase or to be maintained during culture, although those of the transcripts for Ids in Id1A5 cells decreased immediately after differentiation.

4. Discussion

Transcription of the GATA-4 gene in heart could be activated by ubiquitous E-box binding proteins (9) and heart-specific upstream enhancer (G2), which is a direct target of Forkhead box (Fox) and GATA transcription factors (31). Ids are known to directly inhibit ubiquitous E-box binding bHLH transcription factors such as Transcription factor 3 (Tcf3 known as E2A) (16) and UCF2 (32), and indirectly Foxa2 (16). Furthermore, Id proteins bind to GATA-4 and inhibit its DNA binding to the GATA motif (17). These observations suggest that
the increased expression of Id1 and Id3 likely inhibits GATA-4 transcription and the GATA-4 function, which results in inhibition of expression of cardiac muscle genes and differentiation of P19CL6 cells.

A negative feedback regulatory loop between Ids and cardiac transcription factors such as GATA-4 and Nkx2.5 is proposed since transiently expressed these transcription factors induce Id2 expression in rat and mouse cardiomyocyte precursor cell lines (28). Such induction of Id2 was detected in our experiment (Figure 4, left), although increases of Id2 transcripts were detected in the presence of DMSO without GATA-4 expression (Figure 4, middle and right), possibly due to participation of DMSO-inducible factor(s) hitherto unidentified (26).

P19-derived cardiomyocytes resemble embryonic cardiomyocytes (6). The E-box binding Hey1 and Hey2 bHLH transcription factors (different nomenclature summarized in ref. 33), which are expressed developmentally in atrial and ventricular compartments (33), are repressors of GATA-4 (34). Although Id1 could interact with Hey1 (35), it is unlikely that Ids cancel repression of the GATA-4 gene through binding to Hey1 and Hey2 upon DMSO addition since P19CL6 cells do not express Hey1 and Hey2 in the absence of DMSO (Figure S1, http://www.ddtjournal.com/action/getSupplementalData.php?ID=78). Actually, the transcript of Hey1 increased starting from 4 days of differentiation (27).

Although transcription factors GATA-4, myocyte-specific enhancer factor 2C (Mef2c), Nkx2.5 and Tbx5, which mutually interact (6), could stimulate cardiomyocyte differentiation of P19 and P19CL6 cells (7,36,37), signaling pathways further govern the expression and function of these transcription factors. Bone morphogenetic protein (BMP) stimulates differentiation of P19CL6 cells through the Smad and TAK1 (a member of the mitogen-activated protein kinase kinase kinase) pathways together with DMSO-inducible factors independent of BMP signaling (26). Furthermore, the hedgehog, Wnt and Notch signals may participate in cardiac development (33,38,39). BMP-mediated inhibition of Wnt signal stimulated the commitment of cardiac precursor cells to become cardiomyocytes (39). Although the BMP signal also induces Ids (35), Ids can stimulate cardiomyocyte proliferation (30,40,41). The relative contents of Ids and relevant transcription factors would govern the growth and differentiation of cardiac cells since single cell analysis of fate conversion of fibroblasts into cardiomyocytes suggested a strong anti-correlation of Id genes with those of GATA-4, Mef2c and Tbx5 (42).

Ectopic p204 could partially substitute for DMSO in inducing the differentiation of P19 cells and concomitantly stimulate nuclear export of Ids resulting in their degradation by proteasomes (17,43). Thus, not only transcription but also cellular localization of Id proteins should be carefully examined. Several modes of regulation by bHLH proteins and Ids have been also postulated (44), and their context-dependent regulatory circuits including miRNAs were proposed (35,45). Culture conditions might also influence the regulatory circuits since Id expression and cardiac differentiation could be affected by serum (46) and the insulin dose (47), respectively. Since cardiomyocyte regeneration is clinically important for the repair of a damaged heart (48), further study of the molecular mechanisms underlying fine-tuning of such regulatory circuits in which Ids and transcription factors including GATA-4 participate may be helpful for improvement of cardiac cell therapy.

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