Isolation of Reporter Cells That Respond to Vitamin A and/or D Using a piggyBac Transposon Promoter-Trapping Vector System

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Abstract: Previously, we established a highly sensitive promoter-trapping vector system using the piggyBac transposon for the efficient isolation of reporter cells. Herein, we examine whether this screening system can be applied to obtain vitamin-responsive cells. As a result, one and two reporter cells that responded to bexarotene (vitamin A) and calcitriol (vitamin D), respectively, were isolated from $4.7 \times 10^6$ seeded HeLaS3 cells. 5′ RACE analyses identified the well-known CYP24A1 gene as a calcitriol-responsive gene, as well as two new bexarotene- or calcitriol-responsive genes, BDKRB2 and TSKU, respectively. TSKU, interestingly, also responded to bexarotene. Endogenous levels of the TSKU and BDKRB2 transcripts displayed only slight changes and were not detected in the comprehensive analyses performed to date. Dose–response analyses of BDKRB2 and TSKU reporter cells in parallel revealed a differential profile in response to each vitamin A agonist, suggesting a bioanalyzer. The present study demonstrates that producing multiple reporter cells by a type of random screening can efficiently identify novel genes with unusual characteristics and be used for the profiling of the properties of vitamin compounds. Similar approaches to the method shown here may be useful for identifying new markers and for the analysis or diagnosis of nutrients, toxins, metabolites, etc.

Keywords: trap vector; reporter cell; gene mining; transposon; vitamin A; vitamin D

1. Introduction

We previously developed a trap vector system for the efficient isolation of cells that can express reporter genes in response to a certain condition of interest [1] based on transposon-mediated genomic manipulation [2]. In general transposon-mediated vector systems, the transposase gene located between the terminal repeat sequences in natural transposons is replaced with DNA modules, which then can be integrated into the host genome via a cut-and-paste mechanism by a transposase that is expressed via its co-introduction as a helper vector. In our promoter trap system, the DNA modules were designed to express GAL4FF (an extremely trimmed minimal DNA-binding domain of the yeast GAL4 transcription factor with two repeats of the minimal transcription activation module from VP16 [3]), only after they are inserted into the genome, where transcription occurs. Because GAL4FF can lead to the expression of the reporter genes EGFP and firefly luciferase (Fluc), which lie downstream of the UAS at another site in the vector, even a small amount of promoter activity is amplified and can be detected with high sensitivity [1,4]. Among the transposon systems that are applicable to mammalian cells, we employed the piggyBac transposon system because its transposase mutant, called hyPBase, has extremely efficient transposition activity [5]. This piggyBac transposon system replaces the DHFR gene in the genome during transposition. The 5′−TTAA−3′ sequences are innumerable in the genome,
and it is known that the selectivity of the insertion position is random, even though some bias toward transcriptionally active regions has been recognized [6,7]. Therefore, it is possible to deeply explore the trapping of a gene promoter on the genome via a simple vector-introduction operation.

Our trap vector system efficiently produces cells expressing EGFP and Fluc as reporters in response to arbitrary conditions. We reported the production of reporter cells that respond to the expressions of the c-Myc gene [1], ER stresses [1], or glucocorticoids [4]. It is not always necessary to identify the responding gene in the reporter cell; however, if so, an unexpected gene is occasionally found as a novel marker. For example, OSBPL9 was identified as a novel ER-stress-responsive gene, but this gene was overlooked in public data because it ranks fairly low in score [1]. In this way, the almost random screening is highly attractive in identifying marker genes without being bound by past data and models, and without narrowing down multiple candidates. Because of the high signal-to-noise ratio afforded by the amplification mechanism of the GAL4-UAS system, slight changes in responsive gene expressions are extremely highlighted. Therefore, it is possible to select genes as important markers, despite the fact that they show only slight changes in the endogenous levels, which could not be noticed in comprehensive analyses, such as a microarray analysis.

In this study, to examine whether this vector system is also applicable in the field of nutrition, we tried to obtain reporter cells that respond to vitamin A or D, and identified novel responsive genes with low levels of endogenous transcript changes. Among them were genes that responded specifically to each vitamin; interestingly, we also identified a gene that responded to both vitamins. The procedure and results reported here may be an important prototype for other nutrition factors or conditions in the future.

2. Results

2.1. Cloning of Calcitriol- or Bexarotene-Responsive Cells Using a Transposon-Mediated Promoter-Trapping Vector System

HeLaS3 cells were co-transfected with promoter/enhancer trapping vectors [4] and the transposase (hyPBase, [5]) helper vector. To obtain cells that respond only when reagents are added, cells in which EGFP was constitutively expressed were removed by three rounds of fluorescence-activated cell sorting. Bexarotene or calcitriol was then added, and the cells that expressed EGFP at high levels were single-cell sorted. Finally, three independent clones with a low background level and showing a strong shift of the EGFP reporter signal, as revealed by flow cytometry analysis, were selected (Figure 1).

![Figure 1](image-url)  
*Figure 1. Confirmation of reporter cells. The isolated clones were treated without (−) or with the indicated vitamin compound for 18 h, and the expression of EGFP, one of the reporter proteins, was analyzed by flow cytometry. EGFP-positive cells are shown in green.*
2.2. Identification of Responsive Genes by 5′ Rapid Amplification of cDNA Ends (5′ RACE)

To identify the genes that responded to each vitamin, the transcript of the upstream exon sequence that was fused to GAL4FF was analyzed by 5′ RACE (Supplementary Materials, Figures S1 and S2, and Data S1–S5), which led to the identification of responsive genes (Table 1). Among them, cytochrome P450 family 24 subfamily A member 1 (CYP24A1), which was identified in clone L5#11, is a mitochondrial monoxygenase that initiates the degradation of 1,25-dihydroxyvitamin D3, i.e., calcitriol (the physiologically active form of vitamin D3), by hydroxylation of the side chain, regulates the level of vitamin D3, and is one of the best-known vitamin D-responsive marker genes [8,9]. Conversely, the bradykinin receptor B2 (BDKRB2) (clone L20#4) and TSKU (TSK or TSUKUSHI) (clone L3#5) were identified as novel responsive genes to each vitamin. BDKRB2 encodes a receptor for the 9-aa peptide bradykinin, one of the physiologically active substances that exerts a blood pressure-lowering effect, which is released upon activation by pathophysiological conditions, such as trauma and inflammation [10,11]. The B2 receptor is associated with G proteins that stimulate a phosphatidylinositol–calcium second messenger system [12]. TSKU encodes a small leucine-rich proteoglycan and has been predicted or partially demonstrated to be involved in animal organ development, cholesterol homeostasis, bone formation [13], and glycolipid metabolism [14]; however, little is known about its exact functions.

Table 1. List of bexarotene- or calcitriol-responsive reporter cell clones and responsive genes, as revealed by 5′ RACE.

| Clone# | Reagent Used for Screening | Responsive Gene |
|--------|---------------------------|-----------------|
| L20#4  | Bexarotene (vitamin A)     | BDKRB2          |
| L3#5   | Calcitriol (vitamin D)     | TSKU            |
| L5#11  | Calcitriol (vitamin D)     | CYP24A1         |

2.3. Low Induction Levels of the Newly Identified Responsive Genes Revealed by Real-Time PCR Analysis

To analyze the endogenous expressions of the newly identified genes, BDKRB2 and TSKU, parent HelaS3 cells, in which the trap vector was not introduced, were stimulated using bexarotene or calcitriol, and their mRNA was analyzed by real-time PCR (Figure 2). Consistent with the responsiveness of reporter cells, their genes responded to each vitamin used for screening. However, the fold change in both genes was very low, whereas CYP24A1 is known to have fold changes that are thousands of times stronger. Interestingly, TSKU was found to also be induced by bexarotene, as well as calcitriol (Figure 2, left).

Figure 2. Real-time PCR analysis of the expressions of endogenous TSKU and BDKRB2 upon stimulation by 10 µM bexarotene or 1 µM calcitriol. HPRT1 was used as the internal control. The graphs show the mean of two experiments per gene with three measurements each. Error bars indicate SD.

It is known that vitamin A- and D-responsive genes are regulated by nuclear receptor hetero- or homocomplexes, i.e., RAR/RXR and VDR/RXR, respectively, via their recognition of DNA elements, which are termed RARE and VDRE, respectively. We then examined whether the promoter regions of BDKRB2 and TSKU carry these response elements. According to previous reports [15,16], we searched 5′[GA][GT]TCA.[0,8][GA][GT]TCA3′ as RARE, where [AB] indicates “A” or “B” nucleotide, “.” indicates any nucleotide, and “.[0,8]”
indicates 0–8 of any nucleotide. Similarly, by referring to Ramagopalan et al., we searched 5′[AG][AG][GT][TC][CG]A..[GAC][AG][TG][TGC][CT][AGT]3′, which is a top-scoring VDRE for calcitriol [17]. It was revealed that BDKRB2 has one RARE composed of the two direct repeats of the core hexanucleotide motif (PuG(G/T)TCA), spaced by four nucleotides (DR4) (Figure 3, BDKRB2_RARE). Conversely, TSKU had not only one RARE with no spacer (DR0) (Figure 3, TSKU_RARE1), but also three hexanucleotide repeats, each spaced by four nucleotides (Figure 3, TSKU_RARE2+3). To the best of our knowledge, genes with this pattern, i.e., two fused RAREs (or three PuG(G/T)TCA core repeats), are rare. Moreover, in this case, the rare 4-nt spacer observed between the cores was repeated. In addition to RARE, TSKU also had three positions that matched the consensus sequence of VDRE (Figure 3, TSKU_VDRE1~3). Regardless of these numbers and rarities, the existence of the above response sequences was consistent with the reactivity to bexarotene and calcitriol (Figure 2).

Figure 3. Presumed vitamin-responsive genomic sequences of BDKRB2 and TSKU. The known consensus sequences of RARE or VDRE were searched in a promoter area of about 20 kb, and the position of the hit sequence was mapped using the UCSC genome browser.

2.4. Luciferase Assay of Reporter Cells for Responsiveness to Various Vitamin A and D Analogs

To investigate the responsiveness of reporter cells to analogs of vitamins A or D, including drugs approved by the Food and Drug Administration (FDA), we performed a dose–response analysis of these cells using a luciferase assay (Figure 4).
Figure 4. Dose–response analysis of reporter cells (Table 1) against various vitamin analogs using a luciferase assay. The plots are the mean from three independent experiments with three measurements each. Error bars indicate SD. The fold-induction value for 0 M was defined as 1.

The responsiveness of the L20#4 clone, in which BDKRB2 was identified, was specific to drugs in the vitamin A group, whereas that of the L5#11 clone (CYP24A1) was specific to vitamin D compounds. Conversely, the responsiveness of the L3#5 clone (TSKU) was highly interesting, as it responded to both vitamins A and D, as expected. Moreover, the L3#5 clone (TSKU) showed a great number of variations in the shape of the dose-dependency curve obtained using vitamin A compounds compared with L20#4 (BDKRB2) (Figure 4, left half panels, bold lines). By comparing the fold changes, the order of the vitamin A compounds that reacted strongly differed between clones L3#5 (TSKU) and L20#4 (BDKRB2) (Figure 5, upper panels). It was also observed that the various vitamin A reagents that exhibited a fold change of approximately 100 times at 1 µM in the L20#4 clone hardly reacted in the L3#5 clone under the same conditions. These results clarify that the drug can be profiled in greater detail than before by using two or more vitamin A-responsive cell lines.

Because TSKU showed both vitamin A and D responses, its responsiveness when both vitamins coexisted was finally quantitatively evaluated. It is known that both A and D vitamins bind to RXR; thus, the responsiveness to one vitamin is competitively inhibited by the other vitamin in the case of genes having only a single RARE or VDRE [18]. Accordingly, in both the luciferase assay for reporter clones (Figure 6a) and the real-time PCR for endogenous mRNA expression (Figure 6b), BDKRB2 and CYP24A1 exhibited an inhibitory effect on the response expression when a nonresponsive vitamin coexisted at a high concentration (Figure 6, right panels). In contrast, it was observed that both vitamins additively regulated the responsive expression of TSKU (Figure 6, left panels).
Figure 5. Comparison of the responsiveness of reporter cells to vitamin reagents. The data presented in Figure 4 was redrawn for comparison of responsiveness between the reagents in each reporter cell.

(a) Luc assay using the reporter clones
Figure 6. Responsiveness in the case of co-existence of vitamin A (9-cisRA) and vitamin D (calcitriol). A quantitative analysis was performed using two assays when the concentration of 9-cisRA was increased in the absence or in the presence of calcitriol and vice versa. (a) Luciferase assay of the indicated reporter cell clone in which the gene indicated in parentheses responded and (b) real-time PCR analysis of the endogenous expression of the indicated mRNA. The plots are the mean from three independent experiments with three measurements each. Error bars indicate SD.

3. Discussion

In this study, we created reporter cells stimulated by vitamins to test whether the previously developed, highly sensitive trap vector system can be applied in the field of nutrition. This was a challenge because innumerable vitamin-responsive genes have been reported to date by transcriptome analysis, chromatic immunoprecipitation analysis using relevant nuclear receptors, such as VDR, RAR, RXR etc., and in silico analyses which explore response sequences [15,19–21]. Nevertheless, among the three causative genes identified in the isolated reporter cells by 5’ RACE, two (BDKRB2 and TSKU) were novel genes regulated by vitamins (Table 1). These genes both had a DR4-type RARE, which is a rare pattern in the literal sense, in their promoters (Figure 3). Thus, it would probably have been difficult to perform in silico sequence searches based on conventional wisdom (for example, the so-called 3-4-5 rule (refs in [22]) might have been an obstacle). Additionally, the changes in their endogenous mRNA levels were at most only about 2–3-fold, even with the most inducible concentration of the reagent (Figure 2, compared to Figure 4), whereas that of the well-known CYP24A1 gene was changed by several thousand-fold (Figure 6b). If we tried to narrow them down from comprehensive analyses, such as a microarray analysis, using such a low fold change, it would be absolutely impossible to classify them (BDKRB2 and TSKU) as markers. Therefore, we consider that the method of almost randomly inserting a reporter gene into the intrinsic gene region and letting it discover responsive genes is a more efficient and attractive method for identifying new marker genes in any condition of interest.

The observation that the reactivity to vitamin A and its analogs varied between L3#5 (TKSU) and L20#4 (BDKRB2) (Figure 4, left panels and Figure 5, upper panels) led to several interpretations. One was that the number of RARE, their positions, and cis elements, as well as the variation in the distance of the spacer between the cores in the RARE in each promoter, may confer different “recognition eyes” to substances. Models analogous to this have been reported in which individual genes differ in their responsiveness to
vitamin D due to differences in multiple VDREs and chromatin structure [23]. Another
was that the responding cells rigorously recognized the “difference” among them, not
only as vitamin A. It is unclear at this stage how this difference was recognized, but the
following scenarios can be imagined. First, the complex of each vitamin A with nuclear
receptors and RARE forms a slightly different structure, which may change transcriptional
activity or efficiency. Second, it is possible that only minor structural differences act on
different targets other than vitamin receptors, and feedback from them may exert effects on
promoters/enhancers or epigenetics. Nevertheless, we established that, using the two cell
lines, a profile characteristic of each compound was obtained as a pattern of dose–response
curves. This is reminiscent of the unique spectrum of each of the fluorescent proteins. By
increasing the repertoire of these reporter cell lines, it may be possible to create something
like a bioanalyzer system. In this study, we obtained reporter cells limited to vitamins;
however, this system could also be applied to nutrients, toxins, metabolites, etc.

4. Materials and Methods

4.1. Establishment of Reporter Cells Responsive to Bexarotene or Calcitriol

HeLa cells were plated into 6-well or 12-well plates at a concentration of 1.3 × 10^5/mL.
On the following day, the cells were incubated with a mixture of 1 µg of DNA of the
hyPBase, and the transposon donor vector (at a ratio of 1:3) and 4 µg of polyethylenimine
(Polysciences, Warrington, PA, USA) in 100 µL of Opti-MEM for 1 mL of the culture
volume. After cell propagation, the EGFP-positive population, which was considered to
be expressed constitutively, was removed by three rounds of cell sorting on an SH800Z
cell sorter (Sony, Tokyo, Japan). Next, the cells propagated on a 10-cm dish scale were
stimulated with 10 µM bexarotene (No. 11571, Cayman Chemical, Ann Arbor, MI, USA)
or 150 nM calcitriol (D1530, Sigma, Tokyo, Japan) overnight at 37 °C in a CO_2 incubator;
subsequently, those that appeared in the EGFP-high fraction were single-cell sorted into
96-well plates. After propagation of those cells, their reagent responsiveness was checked
under a fluorescence microscope. Whenever multiple clones were obtained from the same
parental cells, the genomes were subjected to a splinkerette PCR analysis, as described
in [1], to select independent clones as assumed from their band patterns.

4.2. Identification of Responsive Genes by 5′ Rapid Amplification of cDNA Ends

5′ RACE was performed exactly as described previously [4].

4.3. Real-Time PCR Analysis

RNA was extracted from the cultured cells using the ISOGEN reagent (Nippon Gene,
Tokyo, Japan). Reverse transcription was performed using the SuperScript III First-Strand
Synthesis Kit (Thermo Fisher Scientific, Waltham, MA, USA) with random hexamers as
the primer mix. The THUNDERBIRD SYBR qPCR Mix (TOYOBO, Osaka, Japan) was used
for real-time PCR, which was executed on a StepOnePlus system (Applied Biosystems,
Waltham, MA, USA). The primers used in this study are listed in Table 2. The human
HPRT1 gene was used as the internal control.

Table 2. Primers used for the real-time PCR.

| Gene     | Primer Sequence (5′→3′)                          |
|----------|--------------------------------------------------|
| TSKU     | CTGAGCGACGTAAGCTTTAAGC                           |
|          | CCGTACCTGTCGGTGAGGAAG                           |
| BDKRB2   | GTACAGGGAGCGCATGAAG                             |
|          | GGCCAAAGGCTCCGTTAAAGA                          |
| CYP24A1  | CTCATGCTAAATACCAGCATG                           |
|          | TGGGAGACCGGATGGG                                |
| GAPDH    | GAAGGTTGAAAGTGGAGCT                            |
|          | GAAGATGTTGATGGG                                |
| HPRT1    | TGGACACTTGCGAAACAAATACG                         |
|          | GGTCCCTTTACCAAGCAAGCT                         |
4.4. Luciferase Assay

The luciferase assay was performed using a slightly modified version of the protocol originally described in the study by Siebring-van Olst [24]. Briefly, cells were spread on a white-colored 96-well cell culture plate. Eighteen hours after the addition of the stimulating reagent (for a total volume of 90 µL), the plate was set on a plate reader (Berthold Technologies, TriStar® S LB942) and the following program was executed: 90 µL of 2 × FLAR (40 mM Tricine [pH8], 2.2 mM MgCO3, 5.3 mM MgSO4·7H2O, 500 µM ATP, 2% TritonX-100, 40 mM DTT, and 500 µM D-luciferin) was dispensed to all wells of the plate, which was then shaken for 180 s, followed by photon counting for 3 s per well after a 0.2 s delay. The means of three independent experiments (in each independent experiment, three samples were measured for one condition \(n=9\)) were graphed using GraphPad Prism software. Vitamin drugs (tretinoin (ATRA) (S1653), isotretinoin (S1379), acitretin (S1368), adapalene (S1276), tamibarotene (S4260), tazarotene (S1569), alfalcacidol (S1468), and doxorcaliferol (S1569)) were purchased from Selleck. Calcitriol (1α,25-dihydroxyvitamin D3) (D1530) was purchased from Sigma. Bexarotene (No. 11571) and 9-cis retinoic acid (9-cis RA) (No. 14587) were purchased from Cayman Chemical. In the experiment reported in Figure 6, which aimed to investigate the responsiveness in the presence of both vitamin A and vitamin D, the addition of the reagents was performed simultaneously.

5. Conclusions

Our previously developed, highly sensitive promoter trap system was shown to facilitate the production of vitamin-responsive reporter cells, which led to the identification of two new vitamin-A-responsive genes, TSKU and BDKRB2. Both of these endogenous genes exhibited only slight changes in response to vitamin A and were not recognized as markers in the past. By analyzing the obtained multiple reporter cells in parallel, various agonists could be classified based on the dose–response curves. Because this screening system is highly versatile and easy to execute and has a high potential to be applied to other physiologically active substances and drugs, the results reported here seem to be of importance and may be a prototype for future research in various fields.

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