Mirror orientation selection (MOS): a method for eliminating false positive clones from libraries generated by suppression subtractive hybridization

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ABSTRACT

Suppression subtractive hybridization (SSH) is one of the most powerful and popular methods for isolating differentially expressed transcripts. However, SSH-generated libraries typically contain some background clones representing non-differentially expressed transcripts. To overcome this problem we developed a simple procedure that substantially decreases the number of background clones. This method is based on the following difference between target and background cDNAs: each kind of background molecule has only one orientation with respect to the two different flanking adapter sequences used in SSH, while truly differentially expressed target cDNA fragments are represented by both sequence orientations. The described method selects the molecules that arose due to hybridization of such mirror-orientated molecules. The efficiency of this method was demonstrated in both model and real experimental subtractions.

INTRODUCTION

A powerful approach for studying the genetic nature of many biological processes is to characterize genes that vary in expression level during this process. Suppression subtractive hybridization (SSH) (1,2) is a highly efficient and widely used (3–5) PCR-based method for identifying differentially expressed genes. A key feature of the SSH method is simultaneous subtraction and normalization that makes it possible to equalize abundance of target cDNAs in the subtracted population. As a result, rare differentially expressed transcripts can be enriched by ~1000-fold. Suppression PCR effect but are not confirmed by further detailed analysis. To overcome this problem we developed a simple procedure that substantially decreases the number of background clones in the libraries generated by SSH.

MATERIALS AND METHODS

Telencephalons of E15.5 and E13.5 mouse embryos were surgically extracted. After removing pial membrane, cortices were separated from the rest of the brain tissue (basal ganglia, hippocampus and olfactory bulb). Total cortex RNA was purified as described (6). Human skeletal muscle polyA+ RNA was obtained from Clontech (CA) and ϕX174 DNA was obtained from Promega (WI). Double-strand cDNA synthesis was performed using the template switch technique (Smart™ PCR cDNA Synthesis Kit; Clontech).

1000-fold diluted. The sample was subsequently amplified again up to 10–20 ng/µl in 10–12 cycle PCR using the same primer and conditions as described for primary PCR. This additional PCR step greatly decreased the portion of background molecules that could be amplified during or following secondary PCR. Such background cDNAs are flanked by one of the nested primers (NP1 or NP2R, see below) on both termini and originated from tester–tester homohybrids. Amplification of such symmetrically flanked molecules is inhibited in primary PCR due to the suppression PCR effect but is permitted in secondary PCR. This type of background is very dangerous for mirror

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clones were arrayed in 96-well microtiter dishes with 150 µl PCR-Script Cloning Kit (Stratagene, CA). Randomly selected µl was mixed with 268 µl of cDNA. To remove NP1 adapters, 5 µl of the cDNA sample was mixed with 2 µl of 10× XmaI restriction buffer, 12 µl H2O and 1 µl XmaI (10 U/µl). The reaction was allowed to proceed for 1 h at 37°C. The enzyme was then inactivated by adding 2 µl of 200 mM EDTA and incubated at 70°C for 10 min. One microliter of XmaI-digested cDNA (5–7 ng) was mixed with 1 µl of 4x hybridization buffer (2 M NaCl, 200 mM HEPES pH 8.3, 0.8 mM EDTA) and 2 µl of H2O [or 2 µl of driver skeletal muscle cDNA (300 ng/µl in some model experiments] and incubated in a thermal cycler at 98°C for 1.5 min and then at 68°C for 3–12 h. It should be noted that theoretically, duration of hybridization could strongly affect the MOS outcomes. Too short a hybridization could result in enrichment of highly abundant cDNA species and loss of rare species due to second-order kinetics of re-annealing. Complexity of typical SSH samples is no more than 10^4 independent cDNA species. Such low-complexity samples can be almost completely re-annealed during a relatively short length of time. Our practice showed that 3 h hybridization is enough and following hybridization prolongation has little effect on the MOS efficiency. After hybridization the sample was mixed with 200 µl of dilution buffer (50 mM NaCl, 20 mM HEPES pH 8.3, 0.2 mM EDTA) and heated in a thermal cycler at 70°C for 7 min. One micro-liter of diluted cDNA was taken for subsequent PCR in a total volume of 20 µl. The PCR mixture contained 1× Advantage KlenTaq Polymerase Mix with the provided buffer (Clontech), 200 µM dNTPs and 0.6 µM adapter-specific primer NP2Rs (5'-GTCGGCCGCGCATGG; this primer, which is shorter than NP2R, was designed to reduce the strong suppression PCR effect that occurs for short DNA fragments). The PCR mixture was incubated in a thermal cycler at 72°C for 2 min to extend the 3'-ends of DNA duplexes and was then immediately switched to the amplification program (25 cycles; Hybrid OmniGene thermocycler, tube control mode) 95°C, 7 s; 62°C, 20 s; 72°C, 2 min. Generaly, for each particualr sample the number of PCR cycles needed should be determined experimentally (usually this PCR consists of 18–23 cycles). Note that after XmaI digestion, a small portion of the NP1 adapter sequence remains intact. After hybridization, the target duplexes formed by annealing of DNA strands with opposite adapter orientation bear two unpaired 3'-terminal bases originating from the NP1 adapter. These bases do not impede the 3'-end extension due to the proofreading polymerase that is included in the Advantage KlenTaq Polymerase Mix.

The PCR product was cloned into pCR-Script Amp using the PCR-Script Cloning Kit (Stratagene, CA). Randomly selected clones were arrayed in 96-well microtiter dishes with 150 µl Luria–Bertani broth with ampicillin and grown overnight on the shaker. One microliter of the bacterial cultures was used for PCR in 96-well PCR plates using the NP2Rs primer. PCR products were spotted on nylon membranes. Hybridization was performed with [32P]dATP-labeled subtracted cDNA from both forward and reverse subtractions as described (11).

RESULTS AND DISCUSSION

The SSH technique is based on the suppression PCR effect that is mediated by long inverted terminal repeats attached to the ends of DNA fragments (12). By incorporating this suppression effect in a PCR amplification scheme, the SSH method normalizes sequence abundance within the amplified cDNA population and prevents amplification of undesirable DNA fragments. The SSH scheme includes the following main steps: (i) subdivision of tester cDNA into two samples and ligation of these samples with two different suppression adapters; (ii) hybridization of tester with excess driver; and (iii) amplification of the tester cDNA molecules that are flanked only with different suppression adapters (this fraction contains the enriched and normalized target cDNA) (1,2,7,8).

We propose that there are the following two main sources of background amplification in SSH. (i) Long oligonucleotides from non-ligated suppression adapters can non-specifically anneal during subtractive hybridization to cDNA molecules having similar sequences. After DNA elongation such molecules can serve as a template for the SSH primary as well as for the secondary (nested) PCR. Also, some background can be generated due to non-specific annealing of PCR primers. (ii) Some redundant cDNA molecules can by chance evade elimination by hybridization with driver and be amplified in subsequent PCRs. For any given redundant cDNA species the latter explanation is extremely unlikely. We estimate that in most cases just a single molecule of each redundant cDNA species is present among several thousand other cDNA molecules that are used for PCR after the subtractive hybridization. However, a huge excess of redundant sequences relative to target cDNAs can cumulatively result in a high number of such background molecules.

Clones representing type (i) background can be easily revealed by differential screening because they do not produce a differential signal. On the contrary, type (ii) background clones show differential signals during screening with probes prepared from two reciprocal (forward and reverse) subtracted samples. Only northern blot and RT–PCR analysis can demonstrate the equal abundance of such sequences in the initial mRNA samples. As a consequence, the elimination of this type of background is the most difficult and time-consuming step in subtracted library analysis. As an alternative to screening with subtracted probes, it is possible to use the tester and driver cDNA as probes for differential screening, but in this case many clones representing rare transcripts give no signals.

We developed a special procedure to decrease the portion of background clones in the subtracted samples. This technique is based on the rationale that after PCR, each species of background molecule has only one orientation relative to the adapter sequences. This directionality corresponds to the orientation of the progenitor molecule. On the contrary, the target cDNA fragments are involved in PCR amplification due to efficient enrichment in the SSH procedure. As a result, each specific sequence has many progenitors and is represented by both sequence orientations. We call our method MOS because this difference between target and background populations is used for specific amplification of target molecules (Fig. 1). The
procedure includes removing one adapter (adapter B in Fig. 1) by restriction endonuclease, heat-denaturation and re-annealing of the SSH sample. Some of the newly formed hybrids from target cDNAs bear adapter A at both termini. Such molecules are generated as a result of hybridization of molecules with mirror orientation of adapters A and B. Thus, they can only be derived from target cDNA fraction. Next, the 3'-ends are filled in and PCR with primer corresponding to adapter A is performed. In this PCR only molecules bearing adapter A at both termini can be amplified exponentially. Thus, the final PCR product is enriched for target sequences.

This scheme was verified in model experiments. To create artificial tester samples, we added various amounts of bacteriophage ϕX174 DNA as a target for subtraction in human skeletal muscle double-stranded cDNA. The amount of ϕX174 DNA added was 0.01% or 0.001% of the human cDNA. Both the tester and the driver were digested by HaeIII and ligated with adapters A and B. Small divergences in length of the fragments presenting in SSH-generated and control ϕX174/HaeIII samples on the one hand and in MOS-generated samples on the other hand are due to the different PCR primers used.

In our practice, we repeatedly used the MOS technique successfully, including the following example of a real MOS application. The structural heterogeneity of neural tissues and the consequent highly complex gene expression profiles presents a great challenge in isolating genes that are developmentally regulated in the mammalian brain. In order to identify genes that are involved in the establishment of cellular identity in murine cortical neurons, we compared two cDNA samples (E13 and E15) prepared from cerebral cortex on 13- and 15-day embryos. A detailed description of this comparison and the isolated sequences will be published elsewhere. The statistical analysis of this experiment, presented below, illustrates the utility of MOS in cases where subtraction yields a small portion of target clones (Table 1). Subtraction was performed by the SSH method in both directions: using E13 as a tester and E15 as a driver (E13–E15) and vice versa (E15–E13). Subtracted sample E15–E13 was cloned and 192 clones from this library were analyzed by differential screening with two subtracted cDNA probes mentioned above. Screening revealed 17 differential clones (9% of analyzed clones), and further analysis (by means of Southern blot hybridization with initial

**Figure 1.** Schematic representation of the MOS method. Rectangles represent DNA molecules (broad rectangles, double-stranded DNA; narrow rectangles, single-stranded DNA). Yellow rectangles, adapter A; green rectangles, adapter B. Pink molecules, target cDNA; blue, background cDNA. The gradient of filling shows the orientation of the cDNA molecules.
E13 and E15 amplified cDNA samples) confirmed only four of these clones (24% of putative differential clones) to have differential expression patterns. After application of the MOS technique to subtracted samples, the primary screening of 480 clones from the E15-specific library revealed 87 differentially expressed clones (18% of analyzed clones), and 71 of these (82% of putative differential clones) were confirmed by further analysis. In this case MOS increased the portion of truly differential clones 7.5-fold and decreased the portion of false positive clones in the enriched sample 4-fold. It should be noted that the complexity of confirmed differential clones in the MOS library is rather high, 62 out of 71 clones represent different cDNA species. So, one of the main advantages of the SSH method, simultaneous isolation of many differentially expressed sequences, is conserved.

Table 1. Comparison of the enriched cDNA libraries generated by the SSH and MOS techniques

<table>
<thead>
<tr>
<th></th>
<th>Analyzed clones</th>
<th>Putative differential clones</th>
<th>Confirmed differential clones</th>
<th>Different target cDNA species</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSH</td>
<td>192</td>
<td>17 (9%)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>4 (24%)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>4</td>
</tr>
<tr>
<td>MOS</td>
<td>480</td>
<td>87 (18%)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>71 (82%)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>62</td>
</tr>
</tbody>
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<sup>a</sup>Percentage of all analyzed clones.  
<sup>b</sup>Percentage of putative differential clones.

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