Construction of a metastasis-associated gene subtracted cDNA library of human colorectal carcinoma by suppression subtraction hybridization

Li Liang, Yan-Qing Ding, Xin Li, Guang-Zhi Yang, Jun Xiao, Li-Chun Lu, Jin-Hua Zhang

AIM: To construct a differentially-expressed gene subtracted cDNA library from two colorectal carcinoma (CRC) cell lines with different metastatic phenotypes by suppression subtractive hybridization.

METHODS: Two cell lines of human CRC from the same patient were used. SW620 cell line showing highly metastatic potential was regarded as tester in the forward subtractive hybridization, while SW480 cell line with lowly metastatic potential was treated as tester in the reverse hybridization. Suppression subtractive hybridization (SSH) was employed to obtain cDNA fragments of differentially expressed genes for the metastasis of CRC. These fragments were ligated with T vectors, screened through the blue-white screening system to establish cDNA library.

RESULTS: After the blue-white screening, 235 white clones were picked out from the positive-going hybridization and 232 from the reverse. PCR results showed that 200-700 bp inserts were seen in 98% and 91% clones from the forward and reverse hybridizations, respectively.

CONCLUSIONS: A subtractive cDNA library of differentially expressed genes specific for metastasis of CRC can be constructed with SSH and T/A cloning techniques.

INTRODUCTION

CRC is one of the most common malignant tumors in the world and its metastasis is the major cause of mortality in patients with colorectal carcinoma. More than hundreds of genes have been reported to be involved in the regulations of metastasis in colorectal carcinoma. However, they are still not sufficient for fully explaining the complexity and diversity of metastasis. Besides, current investigations of these genes that mostly focused on the expression analysis of one or several genes make it difficult to understand the genes’ interactions and find the new genes. Cloning and identification of metastasis-associated genes have been hypothesized to be beneficial to the elucidation of the molecular mechanisms underlying the metastasis and finding gene targets for metastatic forecast, therapy and prognosis of CRC patients. In our study, SSH and bacterial culture PCR screening were used to construct a differentially-expressed gene subtracted cDNA library specific for metastasis of human CRC with the hope to screen new metastasis associated genes.

MATERIALS AND METHODS

Cell lines

Highly metastatic cell line SW620 and lowly metastatic cell line SW480 from human colorectal carcinoma were purchased from ATCC with the number of CCL-227 and CCL-228 respectively. Both of the paired cell lines were from one colonic adenocarcinoma patient and had the same genetic backgrounds. Cell lines were routinely cultured with DMEM supplemented with 10% bovine serum under the atmosphere containing 5% CO₂ at 37°C.

PCR primers and adaptors
cDNA synthesis primer (5'-TTTTGATCAAGCTT_N-N-3') was used for the first-strand cDNA synthesis. Adaptor1 (5' - CTAATACGACTCACTATAGGGCGAGGCAGGGCGCCGAGGTACCCTCGGCG-3') and adaptor2R (5' CTAATACGAC TCACTATAGGCCAGCAGTGGTCCCGCGCCGGAGGTACCCTCGC CG-3') have the same sequence of the 5' region and the different palindromic structures in the 3' end. Only the differentially expressed fragments digested by RsaI restriction enzyme could be ligated with two adaptors. Primary PCR was performed to amplify these fragments with primer1 (5'-CTAATACGACTCACTATAGGGCGAGGCAGGGCGCCGAGGTACCCTCGGCG-3') and adaptor2R (5' CTAATACGAC TCACTATAGGCCAGCAGTGGTCCCGCGCCGGAGGTACCCTCGC CG-3') that corresponds to the outer common sequence of the adaptors. After that, nested primer1 (5' - CGTGGCCGCCGAGGTACCCTCGGCGCCGACGAGGGCCGAGGTACCCTCGGCG-3') and nested primer2R (5' - AGCGTGGTGCCGAGGTACCCTCGGCGCCGACGAGGGCCGAGGTACCCTCGGCG-3') was designed from the inner sequences of the adaptors respectively were used for the further enrichment of these fragments in secondary PCR. The underlines in the sequences indicated the sites of RsaI restriction enzyme. The above primers and adaptors were provided by PCR-Select™cDNA Subtraction Kit (Clontech Laboratories Inc, USA).

mRNA isolation

Total mRNA was isolated from the two cell lines respectively using QuickPrep micro mRNA purification kit (Pharmacia, USA) following the recommendations of the manufacturer.

Suppression subtractive hybridization

SSH was performed by using PCR-Select™cDNA subtraction kit (Clontech Laboratories Inc, USA) according to the following protocol:
recommendations of the manufacturer. The highly metastatic cell line SW620 was used as the tester while the lowly metastatic cell line SW480 was used as the driver in the forward hybridization, and vice versa in the reverse hybridization.

Double strand cDNA synthesis

A total of 2 µg (4 µL) mRNA with 1 µL oligo (dT)\textsubscript{30} primer was heated to 70 °C for 2 min and rapidly cooled on ice. The reaction mixture was made up to 10 µL by adding 1 µL cDNA synthesis primer (10 µmol/L), 2 µL 5x the first strand reaction buffer, 1 µL dNTP mix (10 mmol/L each) and 1 µL sterile H\textsubscript{2}O. Reverse transcription was started by adding 1 µL AMV reverse transcriptase (20 U/µL). The reaction mixture was incubated at 42 °C for 1.5 h to synthesize the first strand cDNA. The second strand cDNA synthesis was performed by immediately adding 48.4 µL sterile H\textsubscript{2}O, 16.5 µL 5x the second strand buffer, 1.6 µL dNTP mix, 4 µL 20x the second strand enzyme cocktail. The reaction mixture was incubated at 16 °C for 2 h. Double strand cDNA was blunted by adding of 2 µL T4 DNA polymerase and incubated at 16 °C for 20 min. The reaction was stopped by adding 20xEDTA/glycogen mix into the reactive mixture. cDNAs were then extracted, precipitated, and resuspended in 50 µL of deionized water.

Ligation of cDNA fragments

For the ligation of cDNA fragments, 43.5 µL of ds cDNA from the tester and driver was digested respectively with 1.5 µL of RsaI (10 U/µL) at 37 °C for 1.5 h. The reaction was terminated by adding 20xEDTA/glycogen mix. The resulting fragments of cDNAs were extracted, precipitated and finally resuspended in 5.5 µL of deionized water. Adapter1 and adapter2R- ligated cDNAs were ligated separately to 2 µL of RsaI digested tester cDNA with 1:6 dilution in the presence of T4 DNA ligase at 16 °C overnight followed by heating at 70 °C for 5 min to inactivate the ligase. In order to determine whether the ligation efficiency was high or not, 1 µL of adapter1-ligated and adapter2R-ligated cDNAs of the tester was diluted into 200 µL H\textsubscript{2}O respectively and amplified in two separate 50 µL reactions. One reaction used G3PDH\textsuperscript{3'}, 5' primer, while the other used G3PDH\textsuperscript{3'} primer and PCR primer. PCR parameters were as follows: 30 cycles at 94 °C for 30 s, at 65 °C for 30 s and at 68 °C for 2.5 min. The products were examined by electrophoresis on a 20 g/L agarose/EB gel.

Two subtractive hybridizations

In the first hybridization, 1.5 µL of RsaI digested driver cDNA was mixed with 1.5 µL of diluted adapter1- or adapter2R-ligated tester cDNA. The samples were denatured at 98 °C for 1.5 min and immediately incubated in a thermal cycler at 68 °C for 8 h. In the second hybridization, two kinds of sample resulting from the first hybridization were mixed in the presence of a freshly denatured driver cDNA. The samples were incubated at 68 °C for 18 h. After 200 µL of dilution buffer was added, the samples were incubated for an additional 7 min. Analysis of the subtraction efficiency was carried out using PCR amplification of G3PDH in the diluted subtracted cDNA versus unsubtracted cDNA. PCR was performed for 33 cycles at 94 °C for 30 s, at 60 °C for 30 s and at 68 °C for 2 min. The products were monitored on a 20 g/L agarose/EB gel for an aliquot which was removed from each reaction after 18, 23, 28, 33 cycles.

Two suppression PCRs (nested PCR)

A 1 µL of diluted subtraction mixture was amplified with PCR primer1 in the primary PCR. The reaction mixture was incubated at 75 °C for 5 min to extend the adaptors and followed in turn at 94 °C for 25 s, 30 cycles at 94 °C for 10 s, at 64 °C for 30 s and at 71 °C for 1.5 min. The primary PCR mixture was diluted 10-fold and 1 µL from that was used in secondary PCR with nested PCR primer1 and primer2. The conditions of the reaction were 15 cycles at 94 °C for 10 s, at 68 °C for 30 s and at 72 °C for 1.5 min. Eight µL products from each PCR reaction of the secondary PCR was analyzed on a 20 g/L agarose/EB gel.

Screening for subtraction library

The second PCR products were purified and their concentrations were measured by a spectrophotometer. The TA/A cloning method was performed by using pGEM-T vector system I (Promega, USA) according to the recommendations of the manufacturer. The PCR product was cloned into the vector with a molar ratio of 6:1 (insert: vector). A 2 µL purified PCR products (25 ng/µL) was used in a 10 µL ligation-reaction system including 5 µL 2× Rapid ligation buffer, 1 µL pGEM-T Easy Vector (50 ng/µL), 1 µL T4 DNA ligase (3 Weiss units/µL) and 1 µL deionized water. The mixture was incubated at 4 °C overnight. The positive control was made by using the control insert DNA from the kit instead of the PCR products. The background control was made without inserting any fragments. The recombinant plasmids were transformed respectively into the competent JM109 E-coil cell with the CaCl\textsubscript{2} method. A 100 µL of transformants was grown on 8 cm×8 cm agar plates containing 100 µg/mL ampicillin, 100 µg/mL IPTG and 100 µg/mL X-gal at 37 °C for 20 h when the blue/white staining could be clearly distinguished. White clones were counted, inoculated into 3 mL of LB liquid medium containing ampicillin and shaken overnight at 37 °C. The resulting bacterial culture was directly used as PCR templates to amplify the inserts in 50 µL reaction mixture containing 10xPCR buffer 5.0 µL, Mg\textsuperscript{2+} (25 mmol/L) 3 µL, nested primer 1 (20 µmol/L) 1 µL, nested primer 2R (20 µmol/L) 1 µL, dNTP mix (10 mmol/L) 1 µL, Taq enzyme mix (5 U/µL) 0.5 µL, and bacterial culture 0.5 µL. PCR consisted of an initial denaturation step at 94 °C for 5 min, followed by 25 cycles at 94 °C for 30 s, at 68 °C for 30 s and at 72 °C for 2 min each. The PCR products were analyzed on an 10 g/L agarose/EB gel.

RESULTS

Quality identification of extracted mRNA

The extracted mRNA that was electrophoresed on 10 g/L agarose gel exhibited as a clear smear with over 0.5 kb length (Figure 1). Their absorbances (A\textsubscript{260}/A\textsubscript{280}>1.8) in spectrophotometry suggested that extracted mRNAs were in a highly purified quality.

![Figure 1](image)

Figure 1 Quality identification of mRNA extracted from two cell lines, Lane 1: SW480 cell line, Lane 2: SW620 cell line.

RsaI digestion analysis of synthesized cDNA

The results of 10 g/L agarose electrophoresis showed that cDNA before digested with RsaI, displayed a zonal smear with a length from 0.5 kb to 10 kb. After the digestion, the cDNA length became shorter with fragments from 0.1 kb to 2 kb, suggesting that cDNAs were completely digested (Figure 2).

Analysis of ligation efficiency

The high ligation efficiency was the most important factor for
success of SSH. Figure 3 shows that the intensity of band 2/band1, band4/band3 was over 80%, indicating that subtracted cDNA library had the high ligation efficiency, 80% of the tester’s ds cDNA fragments were ligated with adaptors1 or 2R.

Figure 2  RsaI digestion analysis of synthesized cDNA, Lane 1: Synthesized cDNA digested with RsaI, Lane 2: Synthesized cDNA, Lane M: Marker DGL2000.

Figure 3  Ligation efficiency analysis of ds cDNA, Lane 1: PCR products using tester1-1 (adaptor1-ligated-cDNA fragment) as the template and the G3PDH3', 5' primer, PCR primer1; Lane 2: PCR products using tester1-1 as the template and the G3PDH3' primer, PCR primer1; Lane 3: PCR products using tester1-2 (adaptor2R-ligated-cDNA fragments) as the template and the G3PDH3' primer, PCR primer1; Lane 4: PCR products using tester1-2 as the template and the G3PDH3' primer, PCR primer1; Lane M: Marker DGL2000.

Analysis of secondary PCR products
Nested primers from the inner sequences of adaptors were used to amplify the primary PCR products, and the smear was increased obviously after 15 cycles, in which some obscure bands with the lengths between 0.2 and 1.5 kb were observed (Figure 4).

Figure 4  Electrophoresis of secondary PCR products, Lane C: Positive control cDNA supplied with the kit, Lane 1: Unsubtracted cDNA, Lane 2: Subtracted cDNA.

Subtractive efficiency
Figure 5 shows that obvious bands were seen after 23 cycles in unsubtracted cDNA and after 33 cycles in diluted subtracted cDNA. The amount of G3PDH was significantly decreased after subtraction, indicating that the subtracted cDNA library had the high subtraction efficiency.

Figure 5  Reduction of G3PDH by PCR subtraction, Lanes 1-4: Unsubtracted secondary PCR products, Lanes 5-8: Subtracted secondary PCR products, Lanes 1, 5: 18 cycles, Lane 2, 6: 23 cycles, Lanes 3, 7: 28 cycles, Lane 4, 8: 33 cycles, Lane M: Marker DGL2000.

Screening for subtraction cDNA library
On the agar plates with 100 µL of transformants, 235 white clones were obtained in the forward hybridization (group A) while 232 white clones were obtained in the reverse hybridization (group R) (Figure 6). White clones accounted for more than 80% of the total clones. Ninety-eight percent and 91% of the white clones were demonstrated by PCR having the inserts with a length of 200-700 bp in groups A and R, respectively (Figure 7), indicating that a subtracted cDNA library specific for metastasis in CRC was successfully constructed.

Figure 6  Blue/white screening for target clones, Two hundred and thirty-five white clones (86%) and 37 blue clones were seen on the agar plates in group A, and 232 white clones (91%) and 21 blue clones were seen in group R.

Figure 7  Different lengths of cDNA fragments from white clones amplified by PCR, Lanes 1-8: Randomly-selected white clones, Lane M: Marker DGL2000.
DISCUSSION

Metastasis is the major cause of mortality in CRC patients. The molecular basis for metastasis has not been fully elucidated yet. Available studies have shown that many specific genes are involved in the process of metastasis, which may accelerate or suppress metastasis by a mechanism of cooperative or inhibitive interactions between genes[11,12]. Till now hundreds of genes have been reported to be involved in the regulation of metastasis in colorectal carcinoma. However, the exact methods for investigating the molecular mechanism of metastasis are mostly limited to the analysis of expressions of only one or several genes, which makes it difficult to understand the genes’ interactions and their relations to the metastasis.

One of the most important ways to understand the carcinogenesis and development of tumors is to clone and identify the differentially expressed genes. In recent years several cloning techniques for the differentially expressed genes have been established, including differential display PCR (DD-PCR), representational difference analysis (RDA), SSH and cDNA microarray. Until now, these techniques have been widely applied in the study of pathology[13-15], immune[16], embryo development[17], as well as the specific expression of tissue protein[18]. Although it is rapid, easy to perform and sensitive, DD-PCR has also some limitations, such as the higher false positive rate and the truncated length of obtained specific cDNA that is usually located in non-translated regions of 3’ poly(A). Likewise, RDA needs less amount of mRNAs samples and has lower false positive rate. But it is troublesome to perform more complicated steps of hybridization or PCR. Microarray-based methods have the potentials to revolutionize the study of differential gene expression under parallel conditions with higher automation. While a number of drawbacks of this approach have been found such as the higher expense and less sensitive than that of PCR-based techniques.

SSH was first put forward by Diatchenko et al. in 1996. The basic principle was that common cDNAs in the paired materials were subtracted by subtractive hybridization and then suppressive PCR was carried out to amplify cDNA fragments especially expressed in tester. The advantage of SSH was the design of two different adaptors and the introduction of suppression PCR with the result that the differentially expressed cDNA fragments were amplified. It allows two subtractive hybridizations in the forward or reverse direction. Some mRNA expressed with low abundance could be detected, which provided clues for the further gene sequencing and identification[19-21]. With the development and improvement of SSH technique, it has become one of the most effective techniques for cloning differentially expressed genes. Since it was established, more than two hundred papers have been published concerning about carcinogenesis[10], tumor metastasis[11], signal transduction[12], apoptosis[13] and tumor therapy[14]. Combined with other molecular biological techniques such as DNA chip, it could clone tumor associated genes[15] and metastasis-associated genes in lung carcinoma, breast carcinoma and other malignancies[16-20]. Zhang et al.[21] carried out SSH in human NCI-H69 cell line and its variant, a more aggressive NCI-N417 SCLC cell line, to characterize the variable expression of genes between the two cell lines, in which 42 genes were screened by the forward and reverse hybridizations. Wang et al.[22] have identified five cDNA fragments with SSH that were expressed at much higher levels in the poorly metastatic cancer cell clone than in the highly metastatic variety. Until now studies on separating metastasis-associated genes in human CRC have not been reported.

In our study, SSH and bacterial culture PCR were performed to construct a subtracted cDNA library specific for the metastasis-related genes of human CRC, in which a pair of human colorectal carcinoma cell lines SW620 and SW480 was used. The SW480 cell line was derived from a Dukes’ stage B colon carcinoma that arose in a 50-year-old male patient with liver and mesenteric lymph node metastases. The SW620 cell line was derived from one of the lymph node metastases. The monoclonal origin of the two cell lines that was confirmed by the presence of shared maker chromosomes on cytogenetic analysis makes it suitable to study the genetic basis of their phenotypic differences. Hewitt et al.[23] have also found that the phenotypic differences of both SW620 and SW480 cell lines were retained, despite long-term culture in vitro.

Both the forward and reverse hybridizations were performed in this study. In the forward hybridization (group A), the highly metastatic cell line SW620 was treated as the tester, while the lowly metastatic cell line SW480 was treated as the tester in the reverse hybridization (group R). As a result, the total cDNAs in the tester and driver were subtracted efficiently. cDNA fragments appearing only in SW620 cells were amplified specially in group A and those only in SW480 cells were amplified in group R. After that, the second PCR products were inserted into T vectors and then screened by the blue/white screening system. Two hundred and thirty-five or 232 white clones were obtained in group A or group R respectively. The resultant subtractive cDNA library that was rich in metastasis-associated gene fragments of CRC was superior to that obtained by other methods for 4 reasons: (1) High specificity. Two subtractive hybridizations and two suppressive PCR were carried out to amplify the differentially expressed cDNA fragments, and suppress the amplification of unspecific cDNA fragments. Thus, the specificity of the subtractive library was greatly increased. (2) High sensitivity. The normalization of tester ss cDNA by SSH could make the differentially expressed genes with high or low abundance separated efficiently. (3) High efficiency. One SSH reaction could separate hundreds and thousands differentially expressed genes simultaneously. (4) More information. The differentially expressed fragments in the forward hybridization might have a role in accelerating metastasis, and those in the reverse hybridization might be metastasis suppressor genes. The forward and reverse hybridizations could provide more information of the screening results.

However, the subtracted cDNA library constructed by SSH had some limitations. One of them was the requirement of large quantities of mRNA samples. The other was that differentially expressed cDNAs obtained from SSH were cDNA fragments digested by restrictive enzyme and needed to be amplified to get the full length. These drawbacks could be overcome by other methods such as the multi-cycle linear amplification based on cDNA synthesis and templet-guided in vitro transcription, in which enough mRNA for the continual research could be amplified with less than 1-50 ng RNA samples[24,25]. Thus, it is especially suitable for tissues that are hard to get or too little to provide enough RNAs. In addition, the full length of differential fragments could be obtained by the rapid amplification of cDNA ends (RACE)[25].

After the blue/ white screening, bacterial culture PCR was used in the present study to screen the positive recombinants. Compared with the routine methods, bacterial culture PCR was not only simple but also helpful for the further identifications. Our results showed that there were 98% clones with 200-700 bp inserts in group A and 91% positive clones of recombinants in group R, suggesting that these clones may contain metastasis-related cDNA fragments. The construction of this library is therefore of significance in screening and identifying the genes associated with metastasis of CRC, which would be beneficial to the elucidation of the molecular mechanism of metastasis and provide new gene targets for metastasis forecast, gene therapy and prognosis estimation in clinical practice.
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