Use of cDNA Tiling Arrays for Identifying Protein Interactions Selected by In Vitro Display Technologies

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Abstract

In vitro display technologies such as mRNA display are powerful screening tools for protein interaction analysis, but the final cloning and sequencing processes represent a bottleneck, resulting in many false negatives. Here we describe an application of tiling array technology to identify specifically binding proteins selected with the in vitro virus (IVV) mRNA display technology. We constructed transcription-factor tiling (TFT) arrays containing ~1,600 open reading frame sequences of known and predicted mouse transcription-regulatory factors (334,372 oligonucleotides, 50-mer in length) to analyze cDNA fragments from mRNA-display screening for Jun-associated proteins. The use of the TFT arrays greatly increased the coverage of known Jun-interactors to 28% (from 14% with the cloning and sequencing approach), without reducing the accuracy (~75%). This method could detect even targets with extremely low expression levels (less than a single mRNA copy per cell in whole brain tissue). This highly sensitive and reliable method should be useful for high-throughput protein interaction analysis on a genome-wide scale.

Methods

IVV screening

Preparation of bait template and IVV template libraries, and the procedure of IVV screening were described in detail in our previous report [21]. Details are also given in Methods S1 online.
Design and construction of the TFT array

Oligonucleotide arrays were constructed photolithographically by an oligo DNA microarray construction service (NimbleGen). The sequences of 1,562 mouse transcription regulatory factors listed by Gunji et al. [23], as well as 37 Jun-associated protein candidates found in our previous studies [14,21,22], were collected from the RefSeq (http://www.ncbi.nlm.nih.gov/RefSeq/) and Genbank (http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db = Nucleotide) databases. Both strands of the total of 1,599 mRNA sequences were doubly represented by a total of 334,372 oligo DNA probes 50-mer in length, with no gap between the probes (Figure 1).

Sample labeling, hybridization and signal detection

Biotin-labeling of the samples was performed by means of in vitro transcription from an SP6 promoter at the 5'-end of each cDNA fragment in the libraries, as described [24], with some modifications. In this process, biotin-labeled sense-strand RNA fragments were produced. Thus, only the antisense-strand probe set was further analyzed in this study. The labeled samples from the bait (+) and bait (−) screenings were hybridized separately on the tiling arrays. The hybridized tiling arrays were stained with Cy3-Streptavidin (Amersham) and detection was done with a scanner. Details are given in Methods S1 online.

Data analysis

Collected data from the tiling array were normalized with the median correction algorithm. Ratios of signal values between the two samples from bait (+) and bait (−) screenings were calculated (Data S1 online). The ratio data were expressed as log2X (X is the actual measurement). After signal measurement, specific signal peaks were identified by the “Windowed Threshold Detection” algorithm in SignalMap software (NimbleGen). This algorithm looks for at least four data points that are above a threshold value within a window. These points were grouped together and presented as a peak. We used the following parameters in the algorithm: Peak Window Size, 300 bp; Percent of Peak Threshold, 20% of maximum data in each mRNA sequence. The value of each peak was the maximum value of the data points in that peak. Only reproducible peaks in the duplicated data were collected as candidates for Jun interactors (Table S1 online). A probe set for NM_183316 was not analyzed, because the sequence of NM_183316 overlapped with that of NM_025925 on the array.

Real-time PCR analysis

Real-time PCR was performed with SYBR Premix Ex Taq (Takara) and protein-specific primer sets (Table S2 online) on the LightCycler (Roche) as previously described [22].
Sensitivity and longer probes produce non-specific signals [25]. 2) There should be no gaps between the probes. A contiguous linear series of data is required to recognize a signal peak in the algorithm for tiling array analysis (in this case, at least 4 data points are needed in a search window), so the probes must be densely arranged. 3) mRNA sequences were employed for the tiling array. Only coding regions are required for the purpose of protein-interaction analysis, so other genomic sequences, e.g., introns, control regions and non-coding RNAs, were not employed. In this study, we constructed TFT arrays containing ~1,600 ORF sequences of known and predicted mouse transcription-regulatory factors (334,372 oligonucleotides) to analyze cDNA fragments from IVV screening for Jun (a transcription factor)-interactors [21].

We also improved the method for labeling of cDNA samples. Usually, double-stranded DNA samples for a tiling array analysis are labeled by using random primers [26]. However, cDNA fragments selected from a randomly fragmented cDNA library [21] seem to be too short for efficient labeling by random priming. Indeed, in a test analysis with a TFT array using the random priming labeling method, we failed to detect all of the previously detected candidates (data not shown). Therefore we employed another labeling procedure [24], in which sense-strand labeled RNAs were produced by one-step in vitro transcription using a SP6 promoter attached to cDNA fragments from IVV screening.

**Results**

**Design of a transcription-factor tiling (TFT) array and sample labeling**

In affinity selection of protein interactions from randomly fragmented cDNA libraries, relatively short cDNA fragments encoding specific binding regions are often obtained. In order to detect these fragments, we adopted a tiling array strategy for the design of custom oligo DNA microarrays as follows: 1) Oligonucleotide probes of 50-mer in length were used. This is the preferred length for microarray probes, because shorter probes result in low sensitivity and longer probes produce non-specific signals [25]. 2) There should be no gaps between the probes. A contiguous linear series of data is required to recognize a signal peak in the algorithm for tiling array analysis (in this case, at least 4 data points are needed in a search window), so the probes must be densely arranged. 3) mRNA sequences were employed for the tiling array. Only coding regions are required for the purpose of protein-interaction analysis, so other genomic sequences, e.g., introns, control regions and non-coding RNAs, were not employed. In this study, we constructed TFT arrays containing ~1,600 ORF sequences of known and predicted mouse transcription-regulatory factors (334,372 oligonucleotides) to analyze cDNA fragments from IVV screening for Jun (a transcription factor)-interactors [21].

Identification of selected candidates using the TFT array

From the 5th-round DNA library of the IVV screening in the presence and absence of a bait Jun protein, called bait (+) and bait (−) screening, respectively [21], we obtained labeled RNAs and hybridized them onto the tiling array. First, the ratios of the signal intensities from the experiments in the presence and absence of bait were calculated. The ratio data are presented in Data S1 online as a GFF formatted file. Next, we searched for signal peaks in the data, as described in Methods. Only reproducible signal peaks were collected (Table S1), where the total number of peaks was 647 on 545 mRNA sequences (some of the mRNA sequences included multiple peaks). To distinguish between true positives and false positives, specific enrichment of the selected candidates was validated by real-time PCR. Among the top 10 percent of the peaks (64 regions), specific enrichment of 35 peaks was confirmed in the screening (white graph in Figure 2A, the signal intensity and peak data of the 35 candidates are presented in Figure S1). The data indicate that the appropriate threshold for distinguishing between true positives and noise in the microarray signal is a signal ratio of 3–4.

The 35 candidates identified in the present study include all of the 20 Jun-interactors identified in our previous studies using conventional cloning and sequencing (Table 1) [21,22]. Furthermore, the 35 candidates include eight well-known Jun-associated proteins, i.e., c-Maf, Fos, Jun, Atf7, Atf4, Jdp2, Atf3 and Fosl2 (Table 1), which is double the number in the previous study, in which four known Jun-interactors were obtained (white graphs of Figure 2B) [21,22]. In other words, 15 proteins including four known Jun-interactors were newly detected using the TFT arrays.

Figure 2. Data from the tiling array. (A) The top 10% candidates were confirmed by real-time PCR. White and gray indicate numbers of enriched and non-enriched candidates, respectively. (B) Numbers of known (white) and newly selected (gray) proteins from conventional sequencing and the TFT arrays.

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Verification of the newly found protein-protein interactions

To confirm the physical association of the 11 newly discovered candidates with Jun, we first performed in vitro pull-down assay. Seven of the 11 tested candidates exhibited specific interaction with the bZIP domain of Jun (Figure S2 online). The affinity of the remaining four proteins, i.e., Cutl1, Myh11, Tax1bp1 and Cebpz, for Jun may be weaker, because their enrichment ratios (excluding that of Cutl1) in the IVV screening were lower than those of others (Table 1). Thus, we next employed the surface plasmon resonance (SPR) method, a highly sensitive analysis tool for protein interactions. In this case, all of the candidates except Tax1bp1 interacted with Jun in a concentration-dependent fashion (Figure S3 online). Although most of the above tested interactions seem to be very weak, we considered that the interactions are true positives, because all of the candidates except Cebpz contain leucine-heptad repeats in the selected regions, and such repeats are an important motif for heterodimerization with Jun. Further experiments in vivo will be required to examine the physiological roles of these interactions.

**Discussion**

To evaluate the quality of the interaction data, coverage and accuracy were calculated as follows. Jun interacts with many bZIP superfamily proteins and structurally unrelated transcription

| Gene symbol | Cloning & Sequencing Accession no. | Signal ratio | Locus on mRNA sequence (base) | Abundance ratio in the initial library (%) | Abundance ratio in the 5th round library (%) | Enrichment |
|-------------|-----------------------------------|--------------|-------------------------------|-------------------------------------------|---------------------------------------------|------------|
| Nrb2        | + NM_025307.2                     | 8.255        | 651...950                     | 5.7E-4                                    | 4.8E-1                                      | 840        |
| 4732436F15Rik| + XM_143418.3                     | 8.134        | 2051...2300                   | 1.1E-4                                    | 1.5E+0                                      | 14,000     |
| c-Maf*      | - S74567.1                       | 7.796        | 1805...2004                   | 6.4E-5                                    | 5.2E-1                                      | 8,000      |
| SNAP19      | + NM_025925.1                     | 7.734        | 1...250                       | 1.6E-3                                    | 2.2E+1                                      | 14,000     |
| Fos*        | + NM_010234.2                     | 7.720        | 501...800                     | 7.4E-5                                    | 1.5E+0                                      | 20,000     |
| Mapre3      | + NM_133350.1                     | 7.714        | 601...950                     | 9.2E-4                                    | 1.4E+0                                      | 1,600      |
| Csg6        | + NM_007790.2                     | 7.577        | 2151...2700                   | 4.1E-4                                    | 1.0E+0                                      | 2,500      |
| Kif5A (region C) | + NM_008447.2    | 7.491        | 2251...2900                   | 3.3E-3                                    | 4.3E+0                                      | 1,300      |
| 9130229H14Rik| + XM_135706.4                  | 7.487        | 51...400                      | 1.6E-3                                    | 1.2E+0                                      | 730        |
| Jun*        | + NM_010591.1                     | 7.411        | 1701...1950                   | 1.4E-4                                    | 4.3E-1                                      | 3,100      |
| Mapk8ip3    | + NM_013931.1                     | 7.282        | 1351...1800                   | 7.3E-4                                    | 5.3E-1                                      | 720        |
| Creb3       | - XM_131375.2                    | 7.172        | 601...800                     | 3.3E-4                                    | 2.6E-3                                      | 8          |
| Kif5B (region C) | - NM_008448.2        | 7.141        | 2801...3200                   | 1.3E-3                                    | 1.2E-1                                      | 95         |
| Nef3        | + NM_008691.1                     | 7.129        | 951...1300                    | 8.2E-3                                    | 1.5E+0                                      | 190        |
| Kif5C (region C) | + NM_008449.2       | 7.106        | 2701...3200                   | 2.2E-3                                    | 6.4E+0                                      | 2,900      |
| Eef1d       | + NM_029663.1                    | 7.083        | 1301...1750                   | 5.5E-3                                    | 2.3E+0                                      | 420        |
| Atf7*       | - NM_146065.1                    | 6.999        | 1051...1300                   | 2.7E-5                                    | 1.4E-1                                      | 5,000      |
| Atf4*       | + NM_009716.1                    | 6.991        | 1001...1300                   | 5.4E-4                                    | 1.9E+0                                      | 3,500      |
| Cutl1       | - NM_009986.2                    | 6.850        | 301...500                     | 1.4E-3                                    | 3.2E-1                                      | 230        |
| Jdp1*       | + NM_030887.2                    | 6.768        | 451...700                     | 7.1E-4                                    | 2.1E-1                                      | 3,000      |
| Odf1        | - NM_177429.2                    | 6.692        | 1752...2001                   | 1.4E-4                                    | 2.6E-1                                      | 1,800      |
| GFAP        | + NM_010277.1                    | 6.551        | 901...1100                    | 1.1E-3                                    | 8.8E-2                                      | 77         |
| Kif5C (region N) | + NM_008449.2        | 6.098        | 1201...1450                   | 1.2E-2                                    | 5.2E+0                                      | 450        |
| Psmc5       | - NM_008950.1                    | 5.961        | 51...300                      | 3.5E-3                                    | 6.4E-2                                      | 19         |
| Kif5B (region N) | + NM_008448.2        | 5.937        | 1301...1550                   | 6.4E-4                                    | 1.5E-1                                      | 230        |
| Atf3*       | - NM_007498.2                    | 5.574        | 401...950                     | 1.1E-7                                    | 2.0E-2                                      | 180,000    |
| B130050I23Rik| + NM_153536.2                  | 5.213        | 1151...1450                   | 1.9E-4                                    | 1.5E-2                                      | 80         |
| Cebpq       | - XM_133383.2                    | 5.122        | 401...750                     | 3.5E-5                                    | 2.7E-3                                      | 78         |
| 1200008A14Rik| + NM_028915.1                  | 4.623        | 1501...1750                   | 3.6E-4                                    | 2.1E-1                                      | 600        |
| Myh11       | - NM_013607.1                    | 4.343        | 3251...3650                   | 1.4E-4                                    | 2.9E-4                                      | 2          |
| Tax1bp1     | - NM_025816.1                    | 4.176        | 501...750                     | 2.7E-3                                    | 1.0E-2                                      | 4          |
| Myt1        | - NM_008665.2                    | 4.084        | 3251...3550                   | 1.1E-5                                    | 1.3E-4                                      | 12         |
| Fosl2*      | - NM_008037.3                    | 3.935        | 401...750                     | 2.1E-5                                    | 1.4E-2                                      | 670        |
| Tef         | - NM_017376.2                    | 3.467        | 851...1000                    | 1.3E-4                                    | 1.1E-3                                      | 8          |
| Cebpz       | - NM_009882.1                    | 3.012        | 1451...1750                   | 1.2E-4                                    | 3.7E-4                                      | 3          |

*Previously reported interactors with Jun.
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Table 1. Thirty-five selected proteins
factors in a binary fashion. Chinenov and Kerppola comprehensively collected reported Jun-interactors in their review, and the number of mammalian Jun-interactors was 51 at that time [27]. Of the 51 interactors, some lack the potential to bind with the bait Jun protein in our experiment for various reasons. For example, the SMAD interacting region of Jun [29] was deleted from our bait protein construct, and NFAT family proteins require a DNA fragment including the AP-1 sequence [TGA(C/G)TCA] and NFAT recognition element (GGAAA) for stable interaction with Jun [29]. Also, the expression of some of the interactors, e.g., JunD, Fra1, Batf, MafA, Nrl and NF-IL6, was not confirmed in the cDNA library used in this study (data not shown). In all, 29 Jun-interactors were expressed in the cDNA library and were expected to bind with the bait Jun protein used here. Of these 29 proteins, four (14%) and eight (28%) were detected by the conventional sequencing and by the newly introduced TFF array method, respectively. This is a remarkable increase and confirms the value of our new methodology as a screening tool for protein interactions. While the coverage was increased considerably, the accuracy did not decrease. Specifically, the number of false positives did not increase: the rates of confirmation of proteins by in vitro pull-down assays in the previous and present studies were 75% and 74%, respectively. Further, we confirmed by SPR that most of the unbound candidates in the pull-down assay actually interacted with Jun. These results indicate that generation of false positives in this novel method is low, and that the method is practical. Undetected remaining interactors were considered to be false negatives. Mismatching of the selection conditions, e.g., salts, detergents, and pH, or the bait construct, e.g., length, region, and tags, might inhibit these interactions.

For quantitative analysis, the abundance ratios of 35 specifically selected candidates in the initial and screened cDNA libraries were determined by real-time PCR, and the enrichment rates (abundance ratio in the 5th round library per that in the initial library) were also calculated (Table 1). The abundance of the 15 newly found candidates (excluding c-Maf, Cbl1 and Odf11) was less than the theoretical threshold determined from the results of our previous study (an analysis of 451 clones) [21,22]. In order to detect the least abundant candidate (Myt1; 1.3×10^{-4}% of the screened cDNA library) by cloning and sequencing, it would have been necessary to analyze at least 1.0×10^{6} clones. These results indicate that our new method is more sensitive, higher-throughput and more cost-effective than the previous method.

From the standpoint of the detection sensitivity, the combinatorial use of the IVV method with TFF arrays provides an extremely sensitive method for protein-interaction analysis, because even a very weakly expressed target, Atf3, could be detected in this study. In the cDNA library before IVV screening, the content of fragments of the selected region of Atf3 was 1.2×10^{-7}%: If one mRNA molecule existed per cell, the content of a fragment of the gene would be about 1.2×10^{-3} to 5.9×10^{-1}% (we employed the parameters from a reference for this calculation [30], and the details are given in Methods S1 online). Thus, the content of Atf3 mRNA in the initial library corresponds to about one molecule per 20 to 100 cells. This suggests that Atf3 is expressed at a very low level in a cell type that is a minor component of whole mouse brain tissue. It is noteworthy that targets expressed at such low levels can be detected without the need for a cell purification procedure, e.g., collection of somatic stem cells by flow cytometry. The high sensitivity of our method may allow access to targets which would be hard to analyze with other existing tools, e.g., the TAP method [18].

Among the newly detected Jun-associated protein candidates, Cebpγ, Creb3, and Tef are intriguing proteins as Jun-associated transcriptional regulators, because they contain basic regions near the leucine heptad motifs, which are necessary for binding with regulatory sequences on the genomic DNA; many known Jun-associated proteins contain such structures. Cebpγ is a member of the CAAT/enhancer-binding protein (C/EBP) family, which is one of the largest and most highly conserved groups of eukaryotic transcription factors. Cebpγ is known to interact with Cebpb, a member of the C/EBP family, but the function of the protein is not well understood [31]. Davydov et al. indicated that Cebpγ binds to the positive regulatory element-I (PRE-I) of the human interleukin-4 gene by forming a heterodimer with Fos protein. However, the interaction between Cebpγ and Jun was not clearly delineated [32]. Creb3 is also a transcriptional regulator belonging to the cyclic AMP response element-binding (CREB) activating transcription factor (ATF) protein family. The bZIP region of Creb3 is strikingly similar to that of ATFα, a known partner of Jun [33]. Tef is a member of the proline and acidic amino acid-rich basic leucine zipper (PAR bZIP) transcription factor family. The PAR bZIP proteins control circadian rhythms in tissues such as the suprachiasmatic nucleus and the liver. Mice deficient in all three PAR bZIP proteins are highly susceptible to generalized spontaneous and audiogenic epilepsies that are frequently lethal [34]. No information is available about the functional relationship between Tef and Jun. More detailed studies in vitro may reveal novel and unexpected functions of Jun in combination with these proteins.

In summary, we have applied tiling array technology, which has previously been used for ChIP-chip assays [26] and transcriptome analyses [20], to protein-interaction analysis with an in vitro display technique for the first time. Compared with previous results obtained with cloning and sequencing [21,22], use of the tiling array greatly increased the coverage of known Jun-interactors from 14% to 28% without any decrease of accuracy (~75%). The new method can also detect targets expressed at extremely low levels. We believe that this highly sensitive and reliable method has the potential to be used widely, because the tiling array method can easily be extended to genome-wide scale, even though the search space is limited in tiled sequences, and the method can also be used in combination with other display technologies, such as phage display and ribosome display.

Supporting Information

Data S1 All ratio data from the tiling arrays
Found at: doi:10.1371/journal.pone.0001646.s001 (2.36 MB ZIP)

Figure S1 Signal ratio and peak data of selected candidates
Found at: doi:10.1371/journal.pone.0001646.s002 (2.14 MB PDF)

Figure S2 In vitro pull-down assay
Found at: doi:10.1371/journal.pone.0001646.s003 (1.80 MB PDF)

Figure S3 SPR analysis
Found at: doi:10.1371/journal.pone.0001646.s004 (0.46 MB PDF)

Table S1 All detected reproducible signal peaks
Found at: doi:10.1371/journal.pone.0001646.s005 (0.09 MB XLS)

Table S2 Oligonucleotides for PCR
Found at: doi:10.1371/journal.pone.0001646.s006 (0.16 MB DOC)

Methods S1
Found at: doi:10.1371/journal.pone.0001646.s007 (0.05 MB DOC)
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Author Contributions

Conceived and designed the experiments: HY KH. Performed the experiments: KH. Analyzed the data: KH. Wrote the paper: KH ND.