

Microarray Data Analysis for Detection and Classification of Viral Infection

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Abstract

DNA microarrays consist of collection of DNA microscopic spots that in order to form an array attached to a solid surface such as glass, plastic or silicon chip. The pieces of fixed DNA considered as a searcher. In this technology it is possible to test sample against thousands probes for specific genes. With this ability, arrays accelerate the biological investigations, gene finding, molecular detection and disease diagnosis. Microarray technology can be seen as a continued development of southern blotting. The most important stage in this technology is data analysis. To analysis such large data with high degree of confidence and reliability needs reliable bioinformatics tools. Infectious diseases still is major problem for human. One of the most important application of microarray technology is the possibility of testing for the presence of thousands micro-organism in environmental and clinical samples only in a single excrement. Thereby we take an important step in rapid and accurate detection of infectious diseases. Here, we present E-Predict algorithm and DetectiV package that is based on species identification in microarray. We demonstrate the application of E-Predict and DetectiV for viral detection in a large publicly available dataset and show that DetectiV performs better than E-Predict. DetectiV is implemented as a package for R - powerful, open source software for statistical programming - that containing visualization, normalization and significance testing functions.

Keywords: Microarray, Microarray Data Analysis, Infectious Diseases

Introduction

Old methods of studying and analyzing the gene expression such as PCR, RT-PCR, Northern blot, Southern blot and determination of the gene sequence are of the best methods for the analysis of a limited number of genes at a time [1]. But microarray technology can assess thousands of genes or proteins simultaneously. The type of microarray depends on the materials placed on the slide: If the material is DNA, it is called DNA microarray, if the material is RNA, it is called RNA microarray, if it is protein, the relevant microarray is called protein microarray and if the material has been derived from a particular tissue, it is called a tissue microarray.

The laboratory procedures of microarray technology involve production of microarray, extraction and labeling of samples using fluorescent dyes, hybridization of samples with a microarray and washing of the samples. Finally, the intensity of probes on the microarray can be measured and quantified using a scanner; and then the data analysis started [2]. DNA microarrays are among useful and emerging tools in Meta-genomics. This technology has been very successful in the detection of different infectious agents including bacteria, viruses, parasites, fungi, and also exploration of the emerging human pathogens [3-7].

Perhaps the most important application of microarray technology in infectious diseases is the rapid and simultaneous detection of different targets and differentiation of genotypes of the pathogens. Due to the pathogens have isolated genetic composition and microarrays ability in studying the sequences of all genes simultaneously, this

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technology is an ideal tool for this purpose [8, 9]. Many microarrays are designed to identify a range of microorganisms present in a sample and their application has been proved in detection of pathogenic viruses [10]. Of other applications of microarray are identification of different bioterrorist threat agents [11] and examining the quality of water and food for the detection of pathogens [12].

However, microarray data analysis in infectious diseases is not easy, because any tested sample of Nucleic Acid usually containing a mixture of DNA and RNA of different organisms, host and variety of contaminants, all this may affect the test results. Furthermore, it is possible that the hybridization process becomes complicated due to the presence of several species, even interdependent, of pathogens.

Materials and Methods

In this study, two methods of E-predict and DetectiV were used for microarray data analysis.

E-predict

One of the methods of microarray data analysis is to use the E-Predict algorithm [13]. This method is based on observed microarray hybridization patterns for the identification of species. In this algorithm, the hybridization energy profiles are first calculated for every reference viral genome as fully sequenced replication in Gen Bank (including the 1229 virus isolation).

It is expected all oligonucleotides of microarray with the genome of specified viruses identified by BLAST [14] are hybridized. Then the hybridization free energy (G) is



calculated for each alignment using the nearest neighbor method. For those oligoes that are unsuccessful in the production of the BLAST output, the hybridization energy is considered to be zero. Thus, a theoretical specific energy profile includes non-zero hybridization energies which are calculated for a subset of oligoes that produce a BLAST alignment compatible with the genome. In general, the energy profiles of all viruses form a matrix of energy. In this matrix, each row is according to a viral species and each column is based on an oligo from a microarray. Then the vector of oligo intensity is normalized and using a similarity measurement compared with each profile normalized in the energy matrix. The result of this function is a raw vector of similarity scores. Each element in this vector expresses the similarity between observed patterns and a profile predicted for a species present in the energy matrix. In the E-Predict method, the p value and the similarity score determine the answer. Using E predict, some useful results were achieved in a number of conditions. However, E-Predict is not currently a tool for imaging and there is a need for updating and costly calculations before using it for the new arrays. Also E-Predict can be used only in Unix and Linux operating systems.

DetectiV

DetectiV is a package for the R statistical software which includes functions for imaging, normalization and performing significance tests of pathogen detection microarray data. R is a free statistical software and available in the Windows, Unix, Linux and Mac operating systems [15]. Since the DetectiV is created in the R software, it is easily integrated with many packages available for microarray analysis [16].

The original data set is a matrix of data which its rows represent probes and its columns represent the measurements of the microarray. These data is easily created from the structures of data obtained by limma [17]. Limma includes functions for reading output formats of many scanners. Affy has functions for reading data of Affy Matrix [18].

We obtain the mean of duplicate probes and at then perform the normalization procedure using dividing corresponding values for each probe on the corresponding array mean value. Then we do the logarithm at the base 2. Therefore, data becomes normal and their mean is equal to zero (Of course, it is assumed that most probes are not hybridized with something). Then the data are grouped

based on viral species using t-test. Those data are used in later stages that their mean logarithm in the base 2 is greater than or equal to 1. These numbers are ordered based on P value.

In this section, the data obtained from the GEO are implemented using the access number of GSE2228 [19]. The array platform for these data is the GEO with the access number of GPL1834 and includes more than 11,000 oligoes representing 1000 viral and bacterial species.

Results and Discussion

Expected and proper results are pre- specified. The results obtained from the data are considered correct if the following two cases occur:

- 1) The result of analysis is similar to the pre- specified pathogen.
- 2) If the target pathogen is not present in the array, the obtained result is very similar and related to the virus.

In the existing 55 arrays of the total 56 arrays, DetectiV will lead to the correct result. In comparison, E-Predict method gives correct results in 53 arrays of 56 arrays. The results of data analysis of 3 arrays with wrong answers in E-Predict algorithm and correct answers in DetectiV are given in the tables 1 and 2. GSM40809 is hybridized with RNA composing of 26 human rhinoviruses. Here, any oligoes designed specifically for this serotype is not present in the array. However, we expect that rhinovirus sp. human and human rhinovirus B to be at the top table as a serotype of human rhinovirus B. The expected infectious agent in GSM40821 array is the virus of Hepatitis B. Finally, GSM40847 array is hybridized with RNA containing human rhinovirus 87 Virus. In this array, we are also faced with a kind of serotype with which no specific oligo has been designed. Therefore, we can expect that the group human rhinovirus sp. to gain the highest value among all results. We achieve correct result using DetectiV in all three arrays.

DetectiV is one of the proper and simple imaging techniques and is a statistical method for data analysis of pathogen detection microarrays. For example, two of the diagrams in Figures 1 and 2 are drawn using normalized data. The first diagram has been drawn based on the family and the second diagram has been drawn based on the species of the pathogens we distinguish between statistical

Table 1. Wrong results of E-Predict

GSM40809			GSM40821			GSM40847		
Virus	P-value	Similarity	Virus	P-value	Similarity	Virus	P-value	Similarity
Human enterovirus D	4.3E-05	0.258894	Orangutan hepadnavirus	2.29E-03	0.148865	Human enterovirus B	1.4E-05	0.386095
Human enterovirus B	4.5E-05	0.267815	Hepatitis B virus	2.38E-03	0.147182	Human enterovirus A	1.6E-05	0.378912
Human enterovirus C	5.2E-05	0.254504	Woodchuck hepatitis B virus	2.72E-03	0.10964	Human echovirus 1	2.2E-05	0.414618
Enterovirus Yanbian 96-83csf	9.4E-05	0.276873	Woolly monkey hepatitis B virus	2.84E-03	0.128919	Enterovirus Yanbian 96-83csf	2.2E-05	0.412299
Human echovirus 1	1.34E-04	0.253816	Arctic ground squirrel hepatitis B virus	3.22E-03	0.103357	Human enterovirus D	2.6E-05	0.296065

Table 2. Result of DetectiV

GSM40809			GSM40821			GSM40847		
Virus	P-value	Mean	Virus	P-value	Mean	Virus	P-value	Mean
Human rhinovirus B	1.0E-05	2.568949	Hepatitis B virus	9.4E-07	1.258597	Human rhinovirus sp.	3.9E-08	2.928193
Human rhinovirus sp.	2.8E-05	1.935673	Orangutan hepadnavirus	7.7E-04	1.231151	Human enterovirus 70	3.5E-06	4.995768
Human rhinovirus A	1.4E-04	1.196806	Woolly monkey hepatitis B virus	8.3E-04	1.169343	Human rhinovirus A	6.3E-06	2.257194
Human echovirus 1	4.6E-03	4.878200	70BK3-040	NA	3.469946	Human rhinovirus B	1.1E-04	1.937722
Porcine enterovirus B	5.4E-03	2.241665	aconitase 2, mitochondrial	NA	1.237789	Porcine enterovirus B	1.7E-04	3.530519

and biological concepts. In DetectiV method, the minimum p value in combination with the largest mean value, and in the E-Predict method, the minimum p value together with the maximum similarity level, determines the answers. In many cases, the right answer is concluded using automatic rules. However, inevitably there are some cases that require human intervention. The proper result of using

DetectiV on the GSE8746 dataset and achieving the correct answer in each 12 arrays is a great incentive for us. In particular, the ability of array and DetectiV to distinguish between not only the viral species but also subtypes of FMDV is very interesting and deserves attention and demonstrates its power. The results of DetectiV application on the SARS data set is also encouraging and promising. In this case, the oligoes specifically designed for SARS virus are absent on the array.

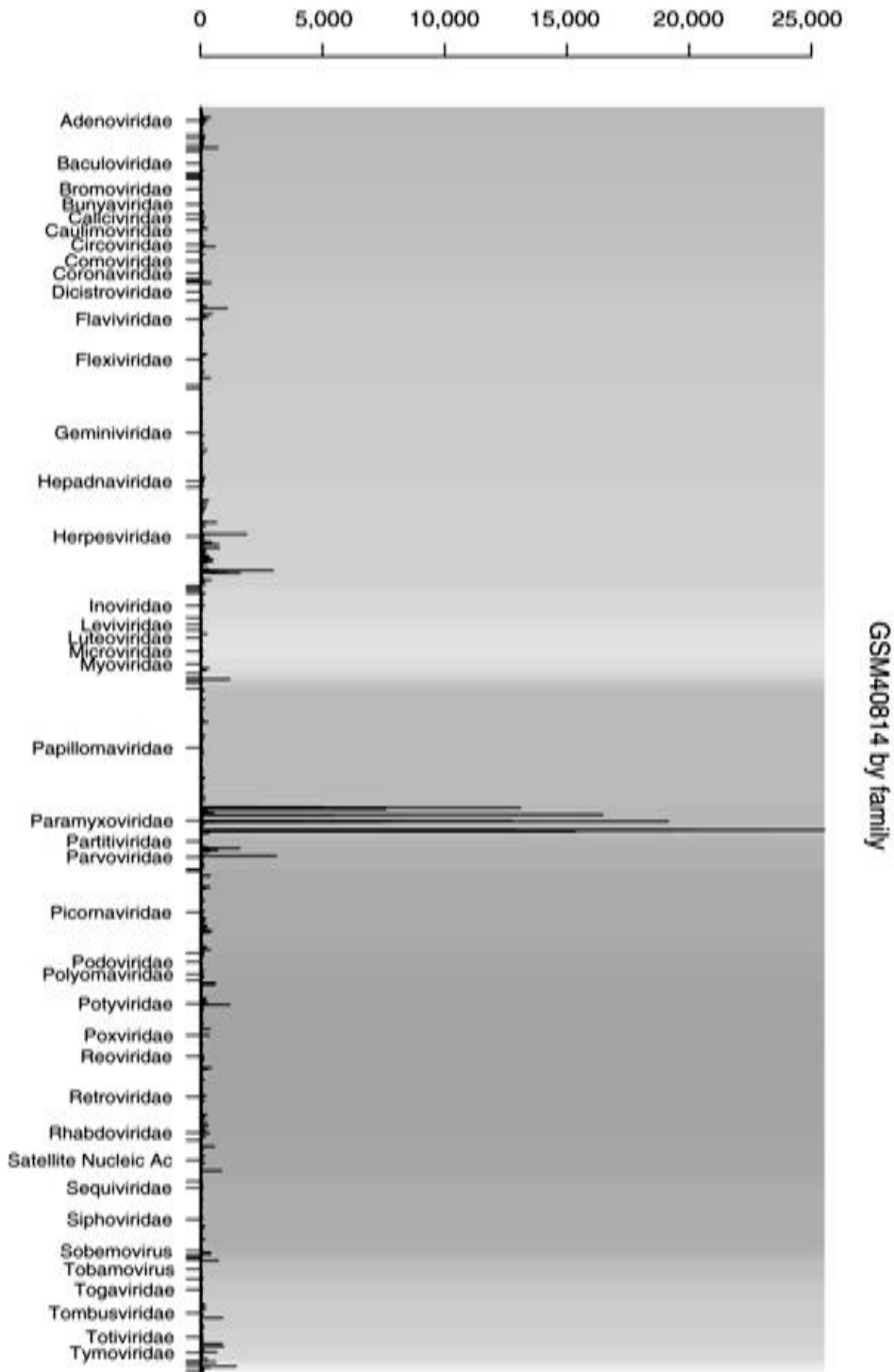


Figure 1. GSM40814 Families; This figure shows an example of barplot prepared by DetectiV, indicating microarray data for the detection of viruses. Oligoes are clustered based on viral family and their mean is obtained according to the replications. Each oligois shownby abar and each viral family has a background color. Asspecifiedin the figure, the Paramyxoviridae family is introduced as the answer.

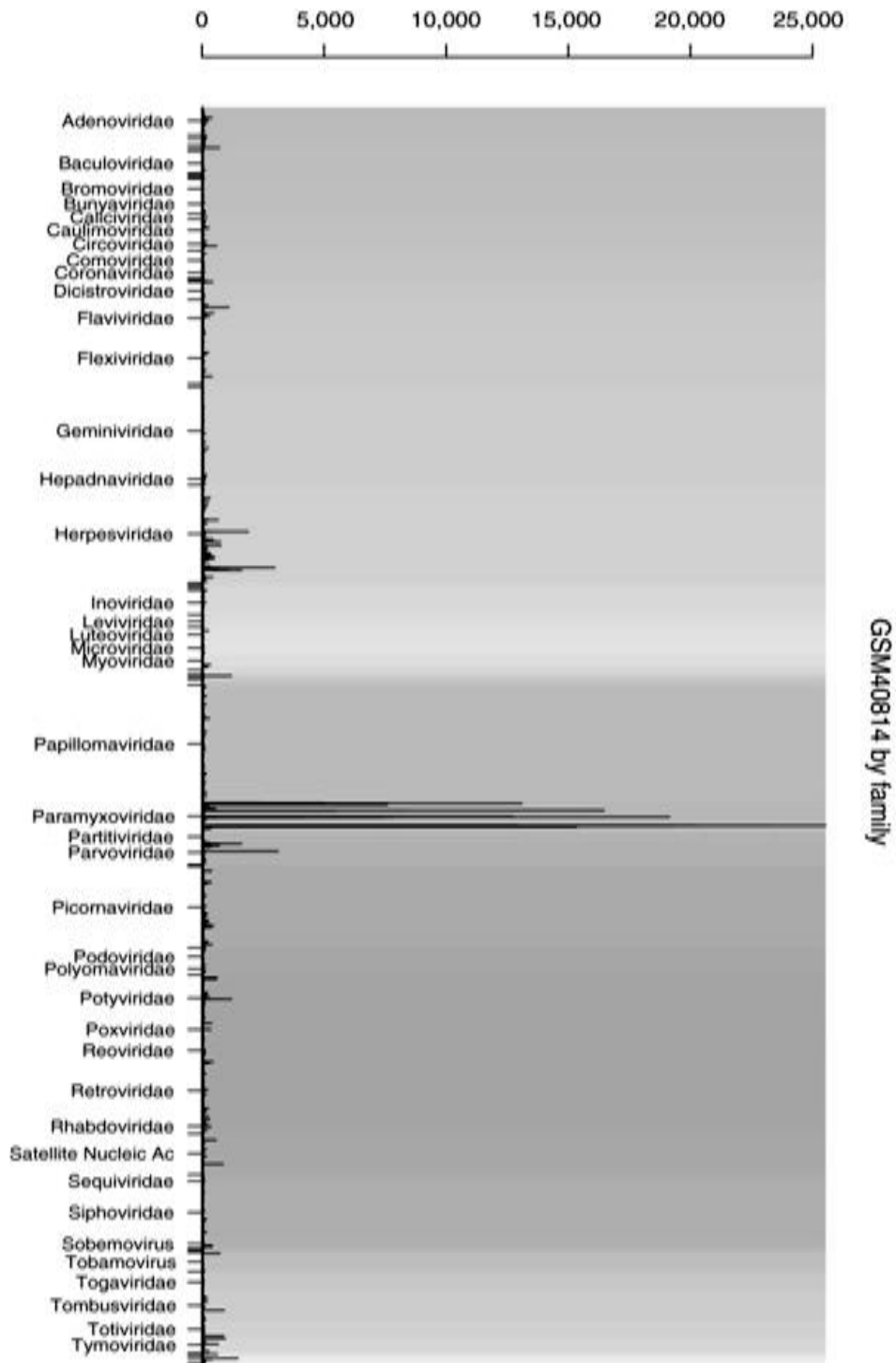


Figure 2. Paramyxoviridae species, this figure shows an example of barplot prepared by DetectiV, indicating microarray data for the detection of viruses. Those oligoes indicating the Paramyxoviridae species are shown in the figure. Oligoes are clustered based on viral family and their mean is obtained according to the replications. Each oligois shown by a bar and each viral family has a background color. As showing the figure, the Respiratory syncytial virus species is introduced as the answer.

It is worth noting that for the use of DetectiV in other data collection that uses arrays completely different from the first dataset, the only requirement is to change the user of GEO access number and the number of arrays in the data set.

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