A new statistical approach for the identification of outlier genes in cancer microarray data

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Abstract: The aim of microarrays technology is to discover genes, which are differentially expressed as outliers between two or more groups of patients are an important task in the genomics community. The regular pattern of genes may often breakdown due to the presence of outliers and it is essential to detect those genes whose behavior looks abnormal in experimental and biological conditions. Several statistical techniques - t-statistic, cancer outlier profile analysis (COPA), outlier sums (OS), outlier robust t-statistic (ORT), maximum ordered subset t-statistics (MOST) and least sum of ordered subset square t-statistics (LSOSS) were developed to address the problem of detecting outlier genes in microarray data but these methods are affected by some problems especially if there is an unusual observation in such dataset then the standard assumptions of distribution parameter may be violated and these techniques might not be suitable to detect outliers genes as well. For these consequences, I have developed a new statistical technique that is “Propose t-statistic (PT)”. The performance of the newly proposed method PT statistic compare with the other existing methods applied to the monte carlo simulation data, package data, and real cancer datasets. The result shows that the outlier genes are identified by using the proposed method PT as well and will give the best and identical results than other methods. The performance of the proposed approach significantly improves than the traditional methods and it can extensively contribute to the medical as well as the genomic community.

Keywords: microarray data; monte carlo simulation; package data; outlier; t-test

1. Introduction

The last couple of decades are the revolution of omic technologies like genomics, transcriptomics, proteomics, and metabolomics, which played a significant contribution and drastic changes in all biomedical sciences and therapeutic management especially disease diagnosis, prognostication and development of novel therapies (Mazumdar and Raha, 2008). The fundamental key of analyzing microarray data is to identify genes in the context of experimental and phenotypical conditions. In particular, the objective of the study is to determine the genes that are outliers between the two classes of normal tissue versus tumor tissue (Griffiths et al., 1996; Hossen et al., 2018; Rashid et al., 2017).

Several statistical algorithms for this type of analysis have been proposed earlier. Initially, the fold changes simple method was used and it performance not so good because of its statistical variability was not taken into account stated by (Chen et al., 1997). After that, some other classical statistical methods have been proposed and these methods may not be reliable and may contain high noise data based on a single array microarray experiments. Since then the most widely used method student t-test which aims to check the statistical significance of every gene where the gene is not differentially expressed in different samples as under the null hypothesis (Phipson et al., 1963).

In microarray gene expression experiments, many methods were developed for aiming to detect cancer-related active genes. Recently, Tomlins (Tomlins et al., 2005) argued that classical and widely used analytical method
two-sample t-statistic, which sometimes fails to detect cancer outlier genes that are over-expressed compared to normal genes. For this consequence, COPA (cancer outlier profile analysis) method were developed for finding the cancer genes tissue with dissimilar expression shapes within cancer tissue samples. Thereafter many statistical methods were proposed - Tibshirani and Hastie (2007) introduced the outlier sums (OS) method, Wu (2007) proposed the outlier robust t-statistic (ORT), Lian (2008) introduced the maximum ordered subset t-statistics (MOST) and a simple statistical test named least sum of ordered subset square t-statistic (LSOSS) is proposed by Wang and Rekaya (2010) for detecting cancer outlier differential gene expression. However, most of the aforementioned statistical techniques for detection of outlier genes are not robust against outliers (Barnett and Lewis, 1994; Hawkins, 1980; Kannan and Monoj, 2015; Wang et al., 2011), though it is a common problem in microarray data. Therefore, the main task is to detect outlier genes in the difficulties areas of biological conditions. To address this problem a new technique is proposed that is “Propose t-statistic (PT)” for detecting the outlier genes where it behavior shows highly different expression patterns in microarray data and compares this method with other existing methods. The helpfulness of the proposed method is then examined by Monte Carlo simulation data, package data and real gene expression cancer datasets.

2. Materials and Methods
2.1. Existing Outlier Detection Methods
2.1.1. t-statistic
Let $x_{ij}$ be the values of expression for the genes $i=1,2,3,..., m$ and samples $j=1,2,3,..., n$ and consider that the samples are divided into two groups and group 1 denoted by normal or reference group, where group 2 is referred as disease group. The two sample t-statistic for gene $i$ is defined as:

$$ t_i = \frac{\bar{y}_i - \bar{x}_i}{s_i} $$

(1)

Where $\bar{y}_i$ is the mean expression value in cancer tissues and $\bar{x}_i$ is the mean expression value in normal tissues for gene $i$ and $s_i$ is the standard deviation.

$$ s_i^2 = \frac{\sum_{1 \leq j \leq n}(x_{ij} - \bar{x}_i)^2 - \sum_{1 \leq j \leq m}(y_{ij} - \bar{y}_i)^2}{n + m - 2} $$

(2)

When most of the cancer tissues are activated the t-statistic is then powerful.

2.1.2. COPA (Cancer Outliers Profile Analysis statistic)
Tomlins et al. (2005) proposed cancer profile outlier analysis (COPA) for detecting cancer outlier genes. They show that usual t-statistic is less powerful than COPA statistic in these cases. It defines the COPA statistic as:

$$ copa_i = \frac{q_r\{(y_{ij}: 1 \leq j \leq m)\} - med_i}{mad_i} $$

(3)

Where $q_r(.)$ the rth, and $med_i$ is the median expression value for each samples $med_i = median\{(x_{ij}: 1 \leq j \leq n)\} \{y_{ij}: 1 \leq j \leq m\}$ and $mad_i$ is the median absolute deviation that is $mad_i = 1.4826 \times median\{(x_{ij} - med_i): 1 \leq j \leq m\}$.

2.1.3. Outliers Sums (OS) Statistic
Tibshirani and Hastie (2007) introduced a method called outlier sums for detecting the outlier’s genes. The limitation of COPA statistic is fixed rth sample percentile was considered by users. To overcome this problem OS statistic used and defined as

$$ OS_i = \frac{\sum_{y_{ij} \in R_i}(y_{ij} - med_i)}{mad_i} $$

(4)

Where,

$$ R_i = \{y_{ij}: y_{ij} > q_{75}\{(x_{ij}: 1 \leq j \leq n)\} \{y_{ij}: 1 \leq j \leq m\}) + IQR\{(x_{ij}: 1 \leq j \leq n)\} \{y_{ij}: 1 \leq j \leq m\}\}$$

and IQR(•) is the inter-quartile range $IQR\{(x_{ij}: 1 \leq j \leq n), \{y_{ij}: 1 \leq j \leq m\}) = q_{75}\{(x_{ij}: 1 \leq j \leq n), \{y_{ij}: 1 \leq j \leq m\}) - q_{25}\{(x_{ij}: 1 \leq j \leq n), \{y_{ij}: 1 \leq j \leq m\})$.

2.1.4. Outliers Robust t (ORT) statistic
ORT is usually similar to OS, and it uses the median value of normal sample instead of median of total data. Also estimates the absolute error using the median of several groups instead of the square error as in COPA for performing more robust and consistent estimation. Wu (2007) introduced this method as given-
Where, \( R_i = \{y_{ij} : y_{ij} > q_{75}(\{x_{ij} : 1 \leq j \leq n\}) + IQR(\{x_{ij} : 1 \leq j \leq n\})\} \) \( mad'_i = 1.4826 \times \text{median}(\{(x_{ij} - \text{med}_{ix}) : 1 \leq j \leq m\}, \{(y_{ij} - \text{med}_{iy}) : 1 \leq j \leq m\}) \)

### 2.1.5. Maximum Ordered Subset t (MOST) statistic
To overcome the used of arbitrary outliers in OS and ORT statistic statistic Lian (2008) proposed MOST statistic because all possible values for outlier thresholds are considered. This statistic is calculated as

\[
\text{MOST}_i = \max_{1 \leq k \leq m} \left( \frac{\sum_{1 \leq j \leq m} (y_{ij} - \text{med}_{ix})^2}{\text{mad}'_i} - \mu_k \right) / \delta_k
\]

Where \( k \) is unknown value, the data are normalized by \( \mu_k \) and \( \delta_k \) standard normal distribution.

### 2.1.6. The Least Sum of Ordered Subset Variance t-statistic (LSOSS)
When we check the distribution pattern of gene expression data it shows two peaks in cancer tissues. To address this problem of change of points or break down of peaks Wang and Rekaya (2010) proposed LSOSS statistic. Instead of median values, this method uses mean values in gene expression data and it defined as follows.

The expression values in cancer related tissues are ordered in magnitude and then divided into two groups for each gene \( i \)

\[ S_{ik1} = \{y_{ij} : 1 \leq j \leq m\}, S_{ik2} = \{y_{ij} : k + 1 \leq j \leq m\} \]

(a) Each gene \( i \) the mean and sum of squares are calculated for these two groups

\[ \bar{y}_{S_{ik1}} = \text{mean}\{y_{ij} : 1 \leq j \leq m\}, \quad \bar{y}_{S_{ik2}} = \text{mean}\{y_{ij} : k + 1 \leq j \leq m\} \]

\[ SS_{ik1} = \sum_{1 \leq j \leq k} (y_{ij} - \bar{y}_{S_{ik1}})^2, \quad SS_{ik2} = \sum_{k+1 \leq j \leq m} (y_{ij} - \bar{y}_{S_{ik2}})^2 \]

The remaining task is to solve the value of \( k \) which divides the two groups. By minimizing the pooled sum of squares the best value of \( k \) is determined for cancer tissues ranging from 1 to \( m - 1 \) are as-

\[ \text{arg min}_{1 \leq k \leq m - 1} (SS_{ik1} + SS_{ik2}) \]

Let \( S^2_{ix} \) is the sum of squares and \( S^2_i \) be the pooled standard error for the estimated gene \( i \) for normal samples is given by

\[ \sum_{1 \leq j \leq m} (x_{ij} - \bar{x}_i)^2 \quad \text{and} \quad S^2_i = \frac{S^2_{ix} + SS_{ik1} + SS_{ik2}}{n + m + 2} \]

(b) The LSOSS statistic for stating the outlier genes is computed as given below where \( k \) could be noted as the number of outlier samples for each gene \( i \)

\[ LSSV_i = k \frac{\bar{y}_{S_{ik1}} - \bar{x}_i}{S_i} \]

### 2.2. Proposed outlier detection method
In the earlier section of method discussion, most of the outlier gene detection techniques covers some non-robust statistic such as the mean and standard deviation and also observe that these statistic may not become effective to serve the purpose of detection genes. In this case, proposed new outlier technique developed to find the appropriate outlier genes and it described below-

#### 2.2.1. Propose t-statistic (PT)
From the equation no (1) that t-statistic of two-condition for gene \( i \) is

\[ t_i = \frac{\bar{y}_i - \bar{x}_i}{S_i} \]

Where \( \bar{y}_i \) mean expression value in cancer tissues is, \( \bar{x}_i \) is the mean expression value in normal tissue for gene \( i \), since both \( \bar{y}_i \) and \( \bar{x}_i \) non-robust, therefore develop the propose t-statistic as

\[ PT_i = \frac{\text{med}_{iy} - \text{med}_{ix}}{S_i} \]

\[ i \]
Where,

\[
\text{med}_{x} = \text{median} \left( x_{i} : 1 \leq j \leq m \right); \quad \text{med}_{x} = \text{median} \left( x_{i} : 1 \leq j \leq n \right) \text{ and }
\]

\[
S_{i}^{2} = \frac{\sum_{1 \leq j \leq n} (x_{ij} - \overline{x})^{2} - \sum_{1 \leq j \leq m} (y_{ij} - \overline{y})^{2}}{n+m-2}
\]

2.3. Data Sets

In this study colon cancer package datasets is used as a stable basis for checking performance and comparison of different method for real cancer gene expression data. The data sets analyzed in this study are in Table 1 and available at (Hossen et al., 2015).

3. Results and Discussion

Gene expression data can be detected outlier on both genes and samples and try to identify the outlier genes in microarray data based on the samples. The performance of the newly proposed method PT statistic compares with the other existing methods applied to the monte-carlo simulation data, package data and real gene expression cancer datasets.

3.1. Identification of outlier genes from monte carlo simulated data

Before applying to real data sets, I have to check the contribution of the newly proposed technique in simulation studies and compare it with the other existing methods. The simulation study was considered for different stages. The simulated data were generated from the standard normal distribution and generated 80 genes. Out of 80 genes, 50 genes were generated and consider the equal number of normal and cancer tissue samples with uniform condition for both groups. Remaining 30 genes were generated with two different conditions and assume that outliers do exist in these genes. The simulation process is taking for 7 times by changing the number of normal and cancer tissue sample sizes. In the first set of simulation n = 50 and m = 50 are generated as the number of samples from normal and cancer groups respectively. In other simulations, chose (n = 10, m = 90), (n=35, m=65), (n=65, m=35), (n=75, m=25), (n=85, m=15) and (n=90, m=10) and the simulation results of the number of detected outlier genes are given in Table 2.

Table 2 shows that the result of the t-test performs well but in some situations, it fails to identify the original outliers. The methods of COPA, OS, ORT and MOST performance are not very satisfactory level rather than other methods. It also sees that the performance of LSSOS method gives the worst because most of the time it fails to detect not a single outlier properly. Besides, the proposed method PT performs best and gives identical results for all considered situations in simulation studies.

3.2. Identification of outlier genes from package data

The details of colon cancer package data sets consist of 2,000 genes where 40 tumor cancer tissues and 22 normal tissues exist (Merk, 1999). The ranges of tumor samples are 5.89 to 20903.18 whereas the ranges of normal samples are 5.82 to 14173.05. As I know that the genes with highly expressions pattern are identified as responsible for tumor tissue. For finding responsible tumor genes, I applied the classical outlier detection methods along with the newly proposed method. An indexed plot was considered to get an idea about tumor sample gene expression which is presented in Figure 1.

In Figure 1, the tumor sample genes are identified through cut-off points and the genes above the cut-off point 3 considered outlier genes. I get 27 common outlier genes that are detected all of the methods including the proposed method. Additionally, find the accuracy (the ratio of common genes and number of detected outlier genes) for all of the existing methods and proposed methods that are presented in Table 3.

Table 3 states the number of outliers detected by several methods along with accuracy level and show that the classical t-test can identify 39 genes as outliers. The COPA, OS, ORT, MOST and LSSOS statistic identify 47, 38, 52, 57 and 68 genes respectively. Whereas the newly proposed statistic PT can identify 32 genes with highest accuracy level than other methods.

3.3. Identification of Outlier Genes from Real Data

The newly proposed method and other outlier detection methods were applied to the six sets of real cancer datasets. The number of outliers is detected through the index plot at a certain cut-off points 3 for all methods. From the output genes (above cuff of points) try to calculate the common genes to check the accuracy of all methods. The common genes for the singh, golub v1, gordon, laiho, pomeroy-v1 and west data are 3, 6, 150, 150, 14 and 52 respectively. The number of the outlier and their corresponding accuracy are given in Table 4.
Table 4 shows that the LSOSS method cannot identify a single outlier in four datasets also the traditional t-test cannot identify a single outlier gene in two datasets whereas the COPA, OS, ORT, MOST, LSOSS and proposed statistic PT can identify outlier genes successfully. I also investigated the performance of proposed method compared with the existing methods through the accuracy measurement level and the results give that the accuracy of traditional t-test is 63.34%, and the accuracy of COPA, OS, ORT, MOST and LSOSS’s are 26.90%, 26.67%, 30.21%, 26.24% and 22.38% respectively, whereas the accuracy level of PT is 70.17%. The LSOSS method gives the worst results followed by the OS, COPA, MOST, ORT and t-test methods. The Proposed method PT which can identify the outlier genes successfully and achieve the highest accuracy level in all datasets except Laiho data. So I may conclude that the proposed method gives the best results among all other methods.

Table 1. Data Description.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Chip</th>
<th>Tissue</th>
<th>Sample No.</th>
<th>Dist. Classes</th>
<th>Gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Golub-V1</td>
<td>Affy</td>
<td>Bone marrow</td>
<td>72</td>
<td>n=47, m=25</td>
<td>1877</td>
</tr>
<tr>
<td>Gordon</td>
<td>Affy</td>
<td>Lung</td>
<td>181</td>
<td>n=31, m=150</td>
<td>1626</td>
</tr>
<tr>
<td>Pomeroy-V1</td>
<td>Affy</td>
<td>Brain</td>
<td>34</td>
<td>n=25, m=9</td>
<td>857</td>
</tr>
<tr>
<td>Laiho</td>
<td>Affy</td>
<td>Colon</td>
<td>37</td>
<td>n=8, m=29</td>
<td>2202</td>
</tr>
<tr>
<td>Singh</td>
<td>Affy</td>
<td>Prostate</td>
<td>102</td>
<td>n=50, m=52</td>
<td>339</td>
</tr>
<tr>
<td>West</td>
<td>Affy</td>
<td>Breast</td>
<td>49</td>
<td>n=25, m=24</td>
<td>1198</td>
</tr>
</tbody>
</table>

Table 2. Outliers in different methods in Simulation Study.

<table>
<thead>
<tr>
<th>Methods</th>
<th>n=35 m=65 gene=40</th>
<th>n=65 m=35 gene=40</th>
<th>n=75 m=25 gene=40</th>
<th>n=50 m=50 gene=40</th>
<th>n=10 m=90 gene=40</th>
<th>n=90 m=10 gene=40</th>
<th>n=85 m=15 gene=40</th>
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</thead>
<tbody>
<tr>
<td>T</td>
<td>18</td>
<td>19</td>
<td>20</td>
<td>18</td>
<td>17</td>
<td>20</td>
<td>20</td>
</tr>
<tr>
<td>COPA</td>
<td>14</td>
<td>11</td>
<td>14</td>
<td>16</td>
<td>16</td>
<td>8</td>
<td>3</td>
</tr>
<tr>
<td>OS</td>
<td>13</td>
<td>12</td>
<td>11</td>
<td>14</td>
<td>0</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>ORT</td>
<td>15</td>
<td>12</td>
<td>11</td>
<td>17</td>
<td>17</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>MOST</td>
<td>16</td>
<td>13</td>
<td>10</td>
<td>15</td>
<td>16</td>
<td>9</td>
<td>4</td>
</tr>
<tr>
<td>LSOSS</td>
<td>25</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>26</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>PT</td>
<td>20</td>
<td>19</td>
<td>20</td>
<td>20</td>
<td>20</td>
<td>19</td>
<td></td>
</tr>
</tbody>
</table>

Table 3. Number of Detected Outliers for Package Data.

<table>
<thead>
<tr>
<th>Method</th>
<th>Number of Outliers (n = 22, m = 40, g=2000)</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>t-test</td>
<td>39</td>
<td>69.23%</td>
</tr>
<tr>
<td>COPA</td>
<td>47</td>
<td>57.44%</td>
</tr>
<tr>
<td>OS</td>
<td>38</td>
<td>71.05%</td>
</tr>
<tr>
<td>ORT</td>
<td>52</td>
<td>51.92%</td>
</tr>
<tr>
<td>MOST</td>
<td>57</td>
<td>47.36%</td>
</tr>
<tr>
<td>LSOSS</td>
<td>68</td>
<td>39.70%</td>
</tr>
<tr>
<td>PT</td>
<td>32</td>
<td>84.37%</td>
</tr>
</tbody>
</table>

Table 4. Summery for the accuracy value of real cancer datasets.

<table>
<thead>
<tr>
<th>Methods</th>
<th>Singh NO Ac</th>
<th>Golub V1 NO Ac</th>
<th>Gordon NO Ac</th>
<th>Laiho NO Ac</th>
<th>Pomeroy-V1 NO Ac</th>
<th>West NO Ac</th>
<th>Av. Ac</th>
</tr>
</thead>
<tbody>
<tr>
<td>t-test</td>
<td>6 50%</td>
<td>18 33.33%</td>
<td>0 0</td>
<td>0 0</td>
<td>16 87.5%</td>
<td>63 82.54%</td>
<td>63.34%</td>
</tr>
<tr>
<td>COPA</td>
<td>14 21.43%</td>
<td>456 1.316%</td>
<td>273 54.95%</td>
<td>227 66.07%</td>
<td>133 10.52%</td>
<td>303 17.16%</td>
<td>26.90%</td>
</tr>
<tr>
<td>OS</td>
<td>14 21.43%</td>
<td>530 1.113%</td>
<td>310 48.38%</td>
<td>216 69.45%</td>
<td>325 4.30%</td>
<td>338 15.38%</td>
<td>26.67%</td>
</tr>
<tr>
<td>ORT</td>
<td>27 11.11%</td>
<td>497 1.22%</td>
<td>254 59.06%</td>
<td>179 83.79%</td>
<td>324 4.32%</td>
<td>240 21.67%</td>
<td>30.21%</td>
</tr>
<tr>
<td>MOST</td>
<td>14 21.43%</td>
<td>264 2.27%</td>
<td>315 52.38%</td>
<td>264 56.81%</td>
<td>214 6.54%</td>
<td>289 17.99%</td>
<td>26.24%</td>
</tr>
<tr>
<td>LSOSS</td>
<td>14 21.43%</td>
<td>0 0</td>
<td>0 0</td>
<td>0 0</td>
<td>60 23.33%</td>
<td>0 0</td>
<td>22.38%</td>
</tr>
<tr>
<td>PT</td>
<td>5 60%</td>
<td>16 37.5%</td>
<td>165 90.91%</td>
<td>302 49.66%</td>
<td>15 93.33%</td>
<td>58 89.67%</td>
<td>70.17%</td>
</tr>
</tbody>
</table>

[N.B.: NO= Number of Outlier, Ac= Accuracy and it calculate by the ratio of common outlier genes and total detected outlier genes by the methods, Av. Ac= Average accuracy and it calculate the mean value of all datasets accuracy.]
4. Conclusions
In microarray gene expression data, discovering the highly expressed genes of the cancer tissue sample is an important growing field of medical research because it can optimize gene expression for cell growth for which it is easy to understand which tissue samples are involved or responsible for cancerous tissues. The performances of proposed statistic PT are better than other existing methods in simulation results suggest that. For package data, proposed methods PT can identify 32 genes as outlier genes among 27 common outlier genes whereas the existing method could not perform as well. For real cancer gene expression datasets, the average accuracy of the existing methods are respectively for t-statistics, COPA, OS, ORT, MOST and LSOSS are 63.34%, 26.90%, 26.67%, 30.21%, 26.24% and 22.38%. Whereas the proposed method PT will give the highest accuracy of 70.17%. Therefore I may conclude that my proposed method which is used to identify the outliers genes in the simulated data, package data and gene expression microarray real dataset gives the best and identical results than other methods. This method will afford useful information, which can help to develop the performance and could assist to detect outlier genes.

Conflict of interest
None to declare.

References


