Quantitative Analysis of Ras Oncogene Isoforms Using Decision Tree and Support Vector Machine

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Abstract—Ras isoform plays a significant functional role in cell growth, cell cycle progression, cytoskeletal changes, apoptosis, and senescence. Mutated Ras gene causes cell proliferation. These genes are found in various kinds of cancer, thus it has been considered that mutated Ras is oncogenic. Ras oncogenes have three isoforms, which are K-Ras, N-Ras and H-Ras. For three decades, there has been active research on Ras gene, but quantitative analysis on each isoform is unknown. For the research, our goal is to compare and contrast each Ras isoform for deeper understanding. We seek for any possibilities of relations between the results and functional role of each isoform by using two types of data-mining technologies; Decision tree and Support Vector Machine. The results show that all three Ras isoforms are fairly alike, especially in the case of K-Ras and N-Ras. Decision tree shows that position 2, 3, 9, 10 act as significant parts in each Ras Isoform. We want to point out that this is the first time to conduct quantitative classification of Ras Isoforms.

Index Terms—ras isoform, K-Ras, N-Ras, H-Ras, cancer, mutated gene, oncogene, bioinformatics, data mining, decision tree, Support Vector Machine (SVM)

I. INTRODUCTION

Ras proteins are members in a class of proteins called small GTPase, which act as a role of cellular signal transduction. When Ras proteins are activated by incoming signals, they are involved in cell growth, differentiation and survival [1]. They convert a molecule called GTP into another molecule called GDP. Ras proteins act as a switch. They must be turned on by attaching to GTP in order to transmit signal. Then, they are turned off when they convert the GTP to GDP. Ras genes code these Ras proteins. There are three Ras gene isoforms: K-Ras, N-Ras and H-Ras [2]. When Ras genes are mutated, they become oncogenic. Ras oncogene sends permanent signalling to Ras proteins leading to cell proliferation, eventually resulting in cancer. Although the incidence varies, mutated Ras genes are found in various tumor types. To name some mutated cases, there are adenocarcinomas of the pancreas (90%), the colon (50%), the lung (30%), thyroid tumors (50%), and myeloid leukemia (30%) [3]. Mutated Ras genes are found mostly in codon 12, 13, 59 and 61.

Initially, it was believed that Ras proteins were functionally overlapping and interchangeable, thus a majority of the earlier studies on Ras were done using H-Ras. However, series of experimental data began to suggest otherwise. Ras isoform were raised by observations such as the prevalent presence of specific Ras oncogenes in particular forms of human tumors; the high conservation across mammalian species of the distinct amino acid sequences of the different Ras isoforms in their C-terminal HVRs; the different patterns of expression, intracellular processing, and subcellular location displayed by the fully processed, mature protein products of the different Ras gene isoforms [4].

Even though each Ras isoform displays high degree of sequence homology and share common sets of upstream effectors and downstream activators, series of experiments and observations suggest that each has functional specificity [4].

A. K-Ras (Kristen Ras Sarcoma Viral Oncogene Homolog)

K-Ras mutations are found in colorectal cancer in 40 to 45% in codon 12 and 13. Among the three isoforms, K-Ras shows the highest frequency of mutations in human tumors. K-Ras mutations occur at very high frequency in pancreatic, colon, or lung cancer, whereas N-Ras and H-Ras mutations are extremely uncommon in those tumors;

B. N-Ras (Neuroblastoma Ras Viral(v-Ras) Oncogene Homolog)

High rate of N-Ras mutations is found in hematopoietic tumors and in malignant melanomas, whereas the rate of K-Ras or H-Ras mutations in the latter tumors is insignificant. N-Ras mutations occur in a high percentage of acute leukemias, whereas H-Ras and K-Ras mutations are much less common there (Fernandez-Medarde and Santos, current journal issue).

C. H-Ras (Harvey Rat Sarcoma Viral Oncogene Homolog)

H-Ras mutations have been reported in melanoma, bladder, thyroid, and mammary carcinoma; it has been reported that the H-Ras oncogene exhibits consistently greater transforming ability than oncogenic N-Ras or K-
Ras when tested in a range of fibroblast-based transformation assays [5], [6].

There are a few unanswered questions regarding Ras gene studies; Current studies on Ras isoforms have not succeeded in complete classification yet. In addition, there is not much known about quantitative characterization, of the degree of functional specificity or overlapping displayed by each Ras oncogenes in the various cell lineages and tissues where they are all present and simultaneously expressed [7]. We set out our purpose of making a quantitative approach to classify and compare each Ras isoform using a few methods of data mining, which are Decision Tree and Support Vector Machine (SVM). The experiments with 9, 13, 17 Window for reliable results were conducted. Window refers to the number of amino acids sliced from the genome sequence of each Ras Isoform.

In previous research, the experiments using sequence of Rattus norvegicus neuroblastoma’s oncogene isoforms (K-Ras, N-Ras, H-Ras transcript variant 1, 2) were conducted [8]. Then, with using the result of Apriori, the frequency of each amino acids in item sets which satisfy the minimum support was compared. In this paper, the origin of genome sequence was replaced from Rattus norvegicus neuroblastoma to Homo sapiens. When the results were compared, they were noticeably different. Frequent sets in each Window are distinct which will be explained further in results. The genome sequence from National Center for Biotechnology Information (NCBI) were acquired.

II. MATERIALS AND METHODS

A. Decision Tree

Decision tree is a typical method of analysis in data mining. Decision tree is based on rules or conditions that are found in various classification conditions. This algorithm is utilized in artificial intelligence, machine learning and statistical analysis a lot. Decision tree is used in order to classify a given data, but can’t be used for prediction. It is available when the target variable is categorical, while the numerical target variable cannot be applied to decision tree algorithm. If numerical target variable is applied to decision tree, discretization process needs to be passed. The result of data analysis by decision tree is expressed in tree structure. So, the analyst can comprehend and explain the result readily.

There are steps in decision tree analysis. The first step is the formation of decision tree. In this step, classification criterion and stopping rule is defined according to data structure and the purpose of analysis. Next step is spreading branches. Remove the branches that have a high risk of classification error or inappropriate induction rule. Then, evaluate the feasibility and establish interpretation and prediction model.

B. Support Vector Machine

Support Vector Machines (SVM) are supervised learning machines used for classification and regression data between two classes. ‘Class’ means a type of data. SVM include associated learning algorithms that analyze data and recognize patterns. When a pair of data sets is given, SVM construct hyperplanes to separate the data. Among many hyperplanes, SVM tries to find the one that can classify the data sets efficiently. The most accurate hyperplane produces the largest margin between the nearest data point of any class, so called as functional margin. Larger margin leads to lower generalization error of the classifier.

When given a set of data, SVM separates the data into training and testing sets. To acquire the higher accuracy, the training of the models needs to be repeated several times while the components of each data set(training data set and test data set are changed, which is called cross-validation [9]. Initially, linear SVM was used for separation, but it was soon realized that few data could be separated with linear method [10], [11]. Therefore, non-linear method (kernel functions) was applied to SVM to solve the problem. Nonlinear kernels allow SVM to produce the maximum-margin hyperplane in a transformed feature space (n-dimensional vector space of numerical feature that represents the object). The transformation may be nonlinear and the transformed space may be high dimensional. Accordingly, although the classifier is a hyperplane in the high-dimensional feature space, it may be nonlinear in the original input space.

Here are some commonly used nonlinear kernels:

- Normal kernel: \( k(x, y) = x^T y + c \)  

- Sigmoid kernel (also regarded as hyperbolic tangent kernel):
  \( k(x, y) = \tanh(\alpha x^T y + c) \)

- Radical basis function kernel (RBF kernel, also regarded as Gaussian kernel):
  \( k(x, y) = \exp\left(-\frac{|x-y|^2}{2\sigma^2}\right) \)

- Polynomial kernel: \( k(x, y) = (\alpha x^T y + c)^d \)

III. EXPERIMENT

A. Assumptions

According to our previous study on comparison of Ras isoforms of Rattus Norvegicus Neuroblastoma, we assumed that H-Ras and N-Ras have more sequences in common than with K-Ras.

B. Decision Tree

With decision Tree, we did the experiment for ten times for the accuracy, and omitted the rules below the frequency of 0.75 judging that they have no value. Rule refers to group of amino acid which was distracted in Ras.
We experimented for 9 Windows, 13 Windows and 17 Windows in order to find more reliable rules. According to Fig.1, the value had no balance since the length of K-Ras was too long when we used the initial length of sequence. Therefore all of the result converged to K-Ras and we duplicated the N-Ras for 3 times and the H-Ras for 10 times, of which the amount is same with K-Ras.

![Figure 1. Length of ras isoforms.](image)

Table II shows the result of decision tree with 13 Window. Due to the regularity of H-Ras in 13 Window and since many amino acids appeared in pos10, we could infer that H-Ras plays a significant role in Ras gene distribution.

**Table II. The Result of 13 Window Decision Tree**

<table>
<thead>
<tr>
<th>RAS</th>
<th>Amino acid</th>
<th>frequency</th>
<th>N/A</th>
<th>R/C</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-RAS</td>
<td>pos2 = W</td>
<td>0.909</td>
<td>9</td>
<td>0.016</td>
</tr>
<tr>
<td></td>
<td>pos4 = R</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H-RAS</td>
<td>pos2 = Q</td>
<td>0.966</td>
<td>9</td>
<td>0.021</td>
</tr>
<tr>
<td></td>
<td>pos14 = R</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K-RAS</td>
<td>pos2 = K</td>
<td>0.933</td>
<td>8</td>
<td>0.01</td>
</tr>
<tr>
<td></td>
<td>pos17 = L</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table III shows the result of decision tree with 17 Window. Because 17 Window has the longest length of sequence, we can guarantee its accuracy among 3 Windows. Identically, H-ras showed the most regularity and the amino acid located in pos2 seems to be important in Ras gene distribution.

**Table III. The Result of 17 Window Decision Tree**

<table>
<thead>
<tr>
<th>RAS</th>
<th>Amino acid</th>
<th>frequency</th>
<th>N/A</th>
<th>R/C</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-RAS</td>
<td>pos7 = D</td>
<td>0.857</td>
<td>7</td>
<td>0.006</td>
</tr>
<tr>
<td></td>
<td>pos10 = N</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H-RAS</td>
<td>pos2 = Q</td>
<td>0.967</td>
<td>9</td>
<td>0.016</td>
</tr>
<tr>
<td></td>
<td>pos10 = A</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K-RAS</td>
<td>pos6 = C</td>
<td>0.889</td>
<td>7</td>
<td>0.007</td>
</tr>
<tr>
<td></td>
<td>pos10 = _</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Overall, the experiment displayed the result clearly. Although the length of K-Ras was the longest of all, 1797 Rules in N-Ras, 783 Rules in H-Ras, and 1364 Rules in K-Ras were came out. And we thought that this was the result of duplicating the length of Ras genes to make them as the same length of K-Ras. However, because we extended them maintaining their ratios using certain function, it doesn’t seem to affect the result largely.

### C. Support Vector Machine

1) SVM-multiclass

During the experiment with SVM, multi-class support vector machine (SVM multiclass) was used, which uses the multi-class formation [12]. Among all the results, it is clear that the numbers of support vectors were the lowest with RBF kernel, compared to those with normal and polynomial kernel. In addition, average loss was recorded lowest in all Windows with RBF kernel. Average loss indicates the discordance with the kernel. Thus, high percentage of average loss means poor separation of the data sets. By applying the characteristics of each kernel to this result values, it can be inferred that the overall structure of the Ras isoforms have nonlinearity.

2) SVM

Since SVM multiclass collects all the data given and produces the result, it is possible that all the data is mixed, making the process become error-prone. Therefore, in...
order to identify the structural relationship in a further
level, 3 different datasets comparing two isoforms were
made; they are (K-Ras, N-Ras), (N-Ras, H-Ras), (H-Ras,
K-Ras). SVM was employed to reveal the structural
relationship between two isoforms. The experiment
methods are the same as SVM multiclass. The results are
in Fig. 2.

![Figure 2. Results of SVM by 9 Window, 13 Window, 17 Window.](image)

Fig. 2 shows the relationship between two Ras
isoforms. Their genome sequences were analyzed by 9
Window, 13 Window and 17 Window. The y-axis in the
graphs (Fig. 2) indicates classification accuracy between
two isoforms using three kernel functions; normal, RBF
and polynomial. High classification accuracy rate means
that the couple is able to be divided easily, meaning that
the two isoforms are not similar.

Among all the functions, the accuracy rates in all
couples are the highest when RBF function was used.
This validates the unique characteristics of RBF function,
presenting that the sequence of each Ras isoform is quite
similar to the shape of Gaussian function, therefore “non-
linear”.

In all the results, it is clearly shown that the
classification accuracy of K-Ras and N-Ras is lower than
that of the others. The result shows that K-Ras and N-Ras
have similar amino acids in common than H-Ras.

IV. CONCLUSION

The result of decision tree shows that amino acids in
position 2, 3, 9, 10 are significant in distinguishing each
Ras Isoform. Moreover, we discovered 1797 rules in N-
Ras which had the shortest sequence in experiment
although frequency, N/A and R/C pointed H-Ras has the
most regular rules. We can infer that this is the result of
amplifying length of N-Ras and H-Ras which we should
make up for next paper.

Experiments done by SVM multiclass show that all the
Ras isoforms have non-linear dispersion since RBF. In
addition, the results of SVM demonstrate K-Ras and N-
Ras resemble each other more than H-Ras.

In this paper, we concentrated on comparing three Ras
Isoforms. Since we conducted three experiments
individually, it was hard to find demonstrate relationship
among the tools. Therefore, we would like to develop a
unifying tool of three experiments to acquire better
accuracy, which will be the groundwork for analyzing
genome sequence of various biological data. Finding their
functional roles will devote to the target therapy of
cancers developed by each Ras isoform.

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Sin Hye Hwang was born in Seoul, Kore, in 1997. She is a student in science major of Hankuk Academy of Foreign Studies. She is interested in medical science and applying bioinformatics to analyzing genome sequence of genes. Although it was hard to approach medical science in an innovative way, she didn’t give up, kept working and succeeded in reaching the conclusion. Her previous work is “Original Comparison of Rattus Norvegicus Neuroblastoma’s Mutated Ras Isoforms Using Apriori Algorithm” (ICEBS 2014).

Su Jin Jang was born in Republic of Korea. She was very curious about all the things, especially about the nature. So she attended HAFS and started to write a paper that she studied and research. Genetics was choice of her major and she tried to discover the secrets of DNA and other genes. In 2015, she continued working on the project and finished the paper above. She, with 3 fellow workers, is currently working on a paper, concerning Ras isoforms.

Dong Won Lee was born in Republic of Korea. He is good at badminton, playing flute, and solving math problems. He entered HAFS in 2013 and started to write a paper that he studied and researched. He kept writing about genes with 3 fellow students and in 2015, he is currently working on a paper, concerning Ras isoforms.

Taeseon Yoon was born in Seoul, Kore, in 1972. He was Ph.D. Candidate degree in Computer education from the Korea University, Seoul, Kore, in 2003. From 1998 to 2003, he was with EJB analyst and SCJP. From 2003 to 2004, he joined the Department of Computer Education, University of Korea, as a Lecturer and Ansan University, as a Adjunct professor. Since December 2004, he has been with the Hankuk Academy of Foreign Studies, where he was a Computer Science and Statistics Teacher. He was the recipient of the Best Teacher Award of the Science Conference, Gyeonggi-do, Korea, 2013.