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Diagnosis of Ovarian Cancer
Based on Mass Spectra of Blood Samples*

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Abstract — The early detection of cancer is crucial for successful treatment, and medical researchers have investigated a number of early-detection techniques. Recently, they have discovered that some cancers affect the concentration of certain molecules in the blood, which allows early diagnosis by analyzing the blood mass spectrum. Researchers have developed several techniques for the analysis of the mass-spectrum curve, and used them for the detection of prostate, ovarian, breast, bladder, pancreatic, kidney, liver, and colon cancers.

We have continued this work and applied data mining to the diagnosis of ovarian cancer. We have identified the most informative points of the mass-spectrum curve, and then used decision trees, support vector machines, and neural networks to determine the differences between the curves of cancer patients and healthy people.

Keywords: Data mining, medical application, decision trees, support vector machines, neural networks.

1 Introduction

The development of tools for the early cancer diagnosis is a major open problem, and clinicians have investigated a variety of diagnosis techniques. Recently, they have discovered that cancer may affect the blood mass spectrum, and studied diagnosis methods based on the analysis of mass-spectrum data, which provide information about proteins and their fragments [3, 4, 24]. The blood mass spectrum is a curve (Figure 1), where the x-axis shows the ratio of the weight of a specific molecule to its electric charge, and the y-axis is the signal intensity for the same molecule. The mass-spectrum analysis is a fast inexpensive procedure based on a sample of a patient’s blood, and it may potentially allow cancer screening with little discomfort to a patient.

Medical researchers have developed several techniques for analyzing the mass-spectrum data, which allow the diagnosis of various cancers, including ovarian, breast, prostate, bladder, pancreatic, kidney, liver, and colon cancers. The effectiveness of these techniques varies across cancer types, methods for generating mass spectra, and algorithms for analyzing the resulting data. Clinicians use three standard measures of the effectiveness of diagnosis techniques: sensitivity, specificity, and accuracy. The sensitivity is the probability of the correct diagnosis for a patient with cancer, the specificity is the chances of the correct diagnosis for a healthy person, and the accuracy is the chances of the correct diagnosis for the overall population of healthy and sick people. The sensitivity of the mass-spectrum diagnosis techniques has varied from 64% to 99%, the specificity has been between 66% and 98%, and the overall accuracy has been between 73% and 98%.

We have continued this work and investigated techniques for the diagnosis of early-stage ovarian cancer. Specifically, we have applied decision-tree learning, support vector machines, and neural networks to identify the differences between the mass spectra of ovarian-cancer patients and those of healthy people. We have used three data sets (Table 1), available at http://clinicalproteomics.steem.com. Sets 1 and 2 include the mass spectra of 100 cancer patients and 116 healthy people, whereas Set 3 includes the data of 162 cancer patients and 91 healthy people. Each mass-spectrum curve consists of 15,155 points.

The experiments have confirmed that the mass spectra allow the diagnosis of ovarian cancer. The sensitivity of the developed technique varies from 85% to 99%, depending on the data set, its specificity is between 81% and 99%, and its accuracy is between 82% and 99%.

2 Previous work

Medical researchers have developed techniques for the detection of early cancer based on protein markers, which are certain molecules in body tissues and fluids [15], but these techniques are often inaccurate.
For example, the specificity of an antigen method for the prostate-cancer detection is only 25–30%, although its sensitivity is high [2]; as another example, the sensitivity of a similar method for breast cancer is 23%, and its specificity is 69% [10]. Recently, researchers have developed a new cancer-detection method based on the application of data mining to the mass spectra of patients’ tissue cells, blood, serum, and other body fluids [12, 14, 23].

Peaks in mass spectra. Some researchers have analyzed mass-spectrum curves using the Ciphergen System software, which helped to identify major peaks. Hlavaty et al. found that a 50.8k Dalton protein peak was present in all prostate-cancer samples, and absent in all samples of healthy people [9]. Watkins et al. used the same method to detect breast, colon, and prostate cancers [21]. They correctly identified 100% of breast cancer cases and ruled out 96% of noncancer cases. For colon cancer, they correctly identified 100% of cancer cases and ruled out 86% of noncancer cases. For prostate cancer, their results were 100% correct for both cancer and noncancer cases. Sauter et al. analyzed mass-spectrum curves of the nipple aspirate fluid over the 5–40k Dalton range, and identified five relevant peaks [19]. The most relevant peaks were 6.5k Dalton and 15.9k Dalton, and their use gave 84% sensitivity and 100% specificity.

Decision trees. Adam et al. applied decision-tree learning to the blood mass spectra of prostate-cancer patients [1]. They used the Ciphergen System software for peak detection, and decision trees for classification based on the intensity of nine highest peaks, which gave 96% accuracy, 83% sensitivity, and 97% specificity. They also experimented with biostatistical algorithms, genetic clustering, and support vector machines, which gave accuracy between 83% and 90%. Qu et al. applied a boosted decision tree method [17] using the same data and features as Adam. They developed two new classifiers, called AdaBoost and Boosted Decision Stump Feature Selection. For AdaBoost, the sensitivity was 98.5% with the 95% confidence interval of 96.5–99.7%, and the specificity was 97.9% with the 95% confidence interval of 95.5–99.4%. For Boosted Decision Stump Feature Selection, the sensitivity was 91.1% with the 95% confidence interval of 86.9–94.6%, and the specificity was 94.3% with the 95% confidence interval of 90.7–97.1%.

Neural networks. Ball et al. applied back-propagation neural networks to determine astroglial tumor grade (1 or 2), which gave 100% accuracy [5]. Poon et al. used neural networks to distinguish hepatocellular carcinoma from chronic liver disease, which gave 92% sensitivity and 90% specificity [16].

Clustering. Petricoin et al. combined a genetic algorithm with self-organizing cluster analysis for identifying ovarian cancer [11]. The sensitivity of their technique was 100% with the 95% confidence interval of 93–100%, and the specificity was 95% with the 95% confidence interval of 87–99%. They also applied their technique to diagnose prostate cancer [13], which gave 95% sensitivity with the 95% confidence interval of 82–99%, and 78% specificity with the 95% confidence interval of 72–83%. Poon et al. applied two-way hierarchical clustering to distinguish hepatocellular carcinoma from chronic liver disease [16]; however, they did not report its sensitivity, specificity, or accuracy.

Other methods. Valerio et al. applied the statistical $\chi^2$ test to the mass spectra of thirteen pancreatic cancer patients, nine chronic pancreatitis patients, and ten healthy people, and found unique protein peaks for each of the three groups [20]; however, they did not report the sensitivity, specificity, or accuracy of their method. Cazares et al. analyzed mass spectra of prostate cancer [6]; they used the Ciphergen System software for peak detection, and logistic regression for classification, which gave 93% sensitivity and 94% specificity. Wu et al. compared several methods for classification of ovarian cancer, including linear discriminant analysis, quadratic discriminant analysis, nearest neighbors, bagging classification trees, boosting classification trees, support vector machines, and random forests [22]; they concluded that the random-forest classification was the most effective.
3 New results

We describe a technique for selecting relevant points of the mass-spectrum curve, and give results of detecting ovarian cancer based on the values of these points.

Feature selection. We view each point of a mass-spectrum curve as a feature, and the corresponding signal intensity as its value. To select relevant features, we calculate the mean intensity values for each point in the mass spectra of the cancer and non-cancer groups, $\mu_1$ and $\mu_2$, and the corresponding standard deviations, $\sigma_1$ and $\sigma_2$. The mean difference of these intensities is $|\mu_1 - \mu_2|$, and the standard deviation of this difference is $\sqrt{\sigma_1^2 + \sigma_2^2}$. For each point, we determine the ratio of the mean difference to its standard deviation, $|\mu_1 - \mu_2|/\sqrt{\sigma_1^2 + \sigma_2^2}$, and select a given number of points with the greatest ratios.

We impose a lower bound on the distance between selected points, which prevents the choice of points with correlated values. After selecting the point with the greatest ratio, we discard all points within the distance bound from it and choose the second greatest-ratio point among the remaining points. Then, we discard the points within the distance bound from the second selected point, choose the third greatest-ratio point among the remaining points, and so on.

Experiments. We have experimented with the use of decision trees, support vector machines, and neural networks for identifying cancer patients based on the selected points. We have used the C4.5 package (www.cse.unsw.edu.au/~quinlan) for learning decision trees [18], the svmfu package (five-percent-nation.mit.edu/SvmFu) for constructing support vector machines with linear kernel functions [7], and the Cascor 1.2 package (www.cs.cmu.edu/afs/cs/project/connect/code/supported) for generating neural networks using the cascade-correlation algorithm [8].

We have implemented an experimental setup that allows control over the number of features and minimal distance between selected features. We have varied the number of features from 1 to 64, and the minimal distance from 1 to 1024. For each combination of settings, we have used eighteen-fold cross-validation to evaluate the three learning techniques. In Figures 2–8, we show the dependency of the accuracy on the control variables.

In Table 2, we give the minimal and maximal sensitivities, specificity, and accuracy for decision trees, support vector machines, and neural networks.

We have determined the number of features and minimal distance between features that lead to the highest accuracy (Table 3). The optimal number of features varies from four to thirty-two, depending on the learning technique and data set. We have also constructed the learning curves for the optimal choice of parameters (Figures 9–11); these curves show the dependency of the accuracy on the training-set size. The results show that all three techniques reach the maximal accuracy after processing about one hundred learning examples.

4 Concluding remarks

We have considered the problem of diagnosing ovarian cancer based on the blood mass-spectrum curve, and identified the relevant points of the curve. We have then applied decision trees, support vector machines, and neural networks to determine the values of these points that indicate ovarian cancer. The effectiveness of these techniques varies across the available data sets; the accuracy of decision trees is between 82% and 99%, the accuracy of support vector machines is between 83% and 99%, and the accuracy of neural networks is between 82% and 99%.

References


Table 2: Effectiveness of ovarian-cancer diagnosis. We show the minimal (Min) and maximal (Max) accuracy, sensitivity, and specificity for decision trees, support vector machines, and neural networks.

<table>
<thead>
<tr>
<th></th>
<th>Decision trees</th>
<th>SVM</th>
<th>Neural nets</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Min</td>
<td>Max</td>
<td>Min</td>
</tr>
<tr>
<td>Data Set 1</td>
<td>Accuracy</td>
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<tr>
<td></td>
<td>Sensitivity</td>
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<tr>
<td></td>
<td>Specificity</td>
<td></td>
<td>72%</td>
</tr>
<tr>
<td>Data Set 2</td>
<td>Accuracy</td>
<td></td>
<td>81%</td>
</tr>
<tr>
<td></td>
<td>Sensitivity</td>
<td></td>
<td>81%</td>
</tr>
<tr>
<td></td>
<td>Specificity</td>
<td></td>
<td>77%</td>
</tr>
<tr>
<td>Data Set 3</td>
<td>Accuracy</td>
<td></td>
<td>96%</td>
</tr>
<tr>
<td></td>
<td>Sensitivity</td>
<td></td>
<td>96%</td>
</tr>
<tr>
<td></td>
<td>Specificity</td>
<td></td>
<td>91%</td>
</tr>
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</table>

Table 3: Control-variable values that lead to the maximal accuracy.

<table>
<thead>
<tr>
<th></th>
<th>Num. of features</th>
<th>Minimal distance</th>
<th>Accu-</th>
<th>Sensi-</th>
<th>Speci-</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>racy</td>
<td>tivity</td>
<td>city</td>
</tr>
<tr>
<td>Data Set 1</td>
<td>Decision trees</td>
<td>4</td>
<td>1</td>
<td>82%</td>
<td>86%</td>
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<tr>
<td></td>
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<td>32</td>
<td>16</td>
<td>83%</td>
<td>82%</td>
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<tr>
<td></td>
<td>Neural nets</td>
<td>32</td>
<td>256</td>
<td>82%</td>
<td>80%</td>
</tr>
<tr>
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<td>8</td>
<td>4</td>
<td>94%</td>
<td>92%</td>
</tr>
<tr>
<td></td>
<td>SVM</td>
<td>4</td>
<td>2</td>
<td>94%</td>
<td>96%</td>
</tr>
<tr>
<td></td>
<td>Neural nets</td>
<td>32</td>
<td>1</td>
<td>96%</td>
<td>93%</td>
</tr>
<tr>
<td>Data Set 3</td>
<td>Decision trees</td>
<td>8</td>
<td>64</td>
<td>99%</td>
<td>98%</td>
</tr>
<tr>
<td></td>
<td>SVM</td>
<td>16</td>
<td>8</td>
<td>99%</td>
<td>100%</td>
</tr>
<tr>
<td></td>
<td>Neural nets</td>
<td>16</td>
<td>2</td>
<td>99%</td>
<td>100%</td>
</tr>
</tbody>
</table>

Figure 2: Experiments with one feature. We show the results for Data Set 1 (left), Data Set 2 (middle), and Data Set 3 (right). For each set, we plot the accuracy of decision trees (solid lines), support vector machines (dotted lines), and neural networks (dashed lines).

Figure 3: Experiments with two features; the legend is the same as in Figure 2.
Figure 4: Experiments with four features; the legend is the same as in Figure 2.

Figure 5: Experiments with eight features; the legend is the same as in Figure 2.

Figure 6: Experiments with sixteen features; the legend is the same as in Figure 2.

Figure 7: Experiments with thirty-two features; the legend is the same as in Figure 2.
Figure 8: Experiments with sixty-four features; the legend is the same as in Figure 2.

Figure 9: Learning curves for decision trees. We show the results of experiments with Data Set 1 (left), Data Set 2 (middle), and Data Set 3 (right). For each set, we plot the accuracy (solid lines), sensitivity (dotted lines), and specificity (dashed lines).

Figure 10: Learning curves for support vector machines; the legend is the same as in Figure 9.

Figure 11: Learning curves for neural networks; the legend is the same as in Figure 9.


