Automated analysis of protein subcellular location in time series images

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ABSTRACT

Motivation: Image analysis, machine learning and statistical modeling have become well established for the automatic recognition and comparison of the subcellular locations of proteins in microscope images. By using a comprehensive set of features describing static images, major subcellular patterns can be distinguished with near perfect accuracy. We now extend this work to time series images, which contain both spatial and temporal information. The goal is to use temporal features to improve recognition of protein patterns that are not fully distinguishable by their static features alone.

Results: We have adopted and designed five sets of features for capturing temporal behavior in 2D time series images, based on object tracking, temporal texture, normal flow, Fourier transforms and autoregression. Classification accuracy on an image collection for 12 fluorescently tagged proteins was increased when temporal features were used in addition to static features. Temporal texture, normal flow and Fourier transform features were most effective at increasing classification accuracy. We therefore extended these three feature sets to 3D time series images, but observed no significant improvement over results for 2D images. The methods for 2D and 3D temporal pattern analysis do not require segmentation of images into single cell regions, and are suitable for automated high-throughput microscopy applications.

Availability: Images, source code and results will be available upon publication at http://murphylab.web.cmu.edu/software

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1 INTRODUCTION

Subcellular distribution is an important characteristic of a protein because location is intimately related to its function. The most common method used to find a protein’s subcellular location is to fluorescently tag the protein, take images by microscopy and then visually analyze the images. Previous work has shown that automatic image analysis can outperform visual examination, providing higher sensitivity to subtle differences (Murphy et al., 2003). Proteins from major subcellular locations can be differentiated from each other with over 90% accuracy (Huang and Murphy, 2004). However, previous work has been done using static images and the challenge now is to extend these approaches to time series images that better reflect protein behavior in living cells. One objective is to distinguish proteins that have similar static but different temporal patterns. This is of obvious biological importance since many proteins change their location over time in order to carry out their functions. For example, helicases localize to the nucleus during the G1 phase of the cell cycle to repair endogenous DNA damage, and exit the nucleus during S phase (Gu, 2004).

For automated analysis of protein subcellular locations, machine learning tools for classification and clustering are well established (Glory and Murphy, 2007). A key component is feature design: good features are numerical representations of the patterns that capture the differences between classes or clusters. The goal for temporal pattern analysis is therefore to design good features from time series images that distinguish different protein movements.

Automated analysis of biological time series images has received significant attention in recent years, but most movies are of low resolution at the organ or cell level (Liebling et al., 2006; Souvenir et al., 2008), and only a few deal with high resolution microscopy images that record protein movement within a single cell (Damse and Waterman-Storer, 2006; Sigal et al., 2006; Zhou et al., 2009). We have carefully studied applicable theories and algorithms from the computer vision field and adopted/designed five sets of temporal features for application to subcellular pattern analysis for microscopy images.

The first, and perhaps most straightforward, approach to characterizing temporal behavior is to track individual objects in a cell and calculate features to describe these tracks. For example, speed, the most common numerical description of movement, requires measuring the total distance the object travels along its way. However, some proteins do not form rigid objects and it is therefore unclear what the targets for tracking should be. The best example is cytoplasmic proteins: even when they show changing fluorescence distributions, there is typically no easily characterized movement of objects. Such ‘complex and nonrigid’ motion that has statistical regularity was defined as ‘temporal texture’ (Nelson and Polana, 1992). In that work, the normal flow on each pixel was used
to represent movement along the image gradient and compared to uniform direction flow. The concept of ‘temporal texture’ has also been applied to adapt Haralick texture features for static images to movies (Bouthemy and Fablet, 1998) or to develop a series of features based on temporal slices (Ngo et al., 2002). We therefore chose co-occurrence based temporal texture features (Hua et al., 2006) and normal flow features as our second and third temporal feature sets. To design other temporal features without defining an entity for tracking, we have also explored image intensity changes in the frequency domain (Fourier transform features) and analyzed static feature changes over time (autoregression features).

We evaluated the ability of these feature types to improve our ability to discriminate protein patterns in a collection of 4D images (three spatial dimensions over time) for 12 cell lines expressing fluorescently tagged proteins.

2 METHODS

2.1 Image acquisition

NIH 3T3 cell lines previously generated as part of a proteome-scale tagging project were used in this study (Chen et al., 2003; Garcia Osuna et al., 2007; Jarvik et al., 2002). In each cell line, a particular protein is tagged by CD tagging (Jarvik et al., 1996) which introduces a green fluorescent protein (GFP) tag into the gene coding for the protein using self-inactivating retroviral vectors.

Cells were plated on glass-bottomed culture dishes in Dulbecco’s modified Eagle’s media. After 48h, the media were changed to Opti-MEM to avoid the interference by phenol red and provide pH stability. Three-dimensional movies of GFP tagged proteins in 3T3 cells were taken by a spinning disk confocal microscope. The imaging system consists of a LaserPhysics Reliant 100 × 488 Argon laser, a Yokogawa CSU10 Confocal Scanner Unit and an Olympus IX50 microscope with a 60 ° 100 s 488 Argon laser, a Yokogawa CSU10 Confocal Scanner Unit and an Olympus IX50 microscope with a 60 × 1.4NA objective. Images (1280 × 1024) were collected with a Roper Scientific/Photometrics CoolSnap HQ Cooled CCD camera, with a final resolution of 0.11 microns per pixel in the sample plane. A stack of 15 images was collected using a 3s exposure time and a spacing of 0.5 μ between slices. Stacks were taken every 45s, with the total period of time for each movie varying depending on the extent of fluorescence photobleaching. Before starting acquisition for each field, the focus was manually adjusted using the GFP fluorescence channel to ensure that cells were centered in the stack.

Time series images were taken for 12 cell lines, each having a different protein labeled with GFP. The proteins (genes) were as follows: cytochrome b-5 reductase (dih1) and annexin A5 (anxa5) in cytoplasm, serum deprivation response protein (sdrp) in vesicles and cytoplasm, adipose differentiation-related protein (adip) in vesicles, ATP-ATP translocase 23 (timm23), ATP synthase (atp5a1) and mitochondrial stress-70 protein (hspa9a) in mitochondria, catalase (cat) in mitochondria and vesicles, glucose transporter 1 (glut1) in plasma membrane and t-complex testis expressed 1 (hspa9a), alpha-actinin-4 (actn4), caldesmon 1 (cald1) in cytoskeleton. Figure 1 illustrates their 2D static patterns and Figure 2 shows a sample movie for ATP synthase. The full movies are available as described in the Abstract.

2.2 Image preprocessing

Background fluorescence was removed by subtracting the most common pixel value in the image. This was based on the assumption that an image contains more pixels outside the cell than inside it and that background is roughly uniform, both of which hold true for our images. Then a threshold was chosen by an automated method (Ridler and Calvard, 1978), and pixels below the threshold were set to zero. For simplicity, thresholded images were used to calculate all features, although thresholding is not required for calculating texture features. No segmentation was performed because no

Fig. 1. Sample fluorescence microscope images of 12 GFP-tagged proteins. Each image is the most typical center slice at the first time point among all 3D time series images of the protein, selected by TypIC (Markey et al., 1999) which finds the image closest to the mean in feature space. Images are preprocessed with background removal and thresholding. For display purposes, only a rectangular region containing a single cell is shown.

Fig. 2. Movie of ATP synthase. The first 8 center slices across 315 s are shown. Images are preprocessed by background removal, thresholding and photobleaching correction. For display purposes, only a rectangular region containing a single cell is shown.

nuclear channel image was available to provide a reference for automatic segmentation algorithms. A major problem with time series images is that photobleaching can occur over time. To minimize the influence of photobleaching, pixel intensities of images at each time point were stretched so that the 95% quantile (and above) was set equal to 256. We chose to stretch the 95% quantile instead of the brightest pixel, because this is less sensitive to noise from artifacts such as fluorescent debris. Figure 3 illustrates the extent of bleaching and correction. The original intensity (dotted line) dropped to 50% after 10 time points. After photobleaching correction, the total intensity (solid line) at the 10th time point is about 80% of the original. Although the correction cannot fully restore the total intensity, it largely reduces the influence of photobleaching. The other advantage of stretching the intensity to a fixed maximum for all proteins is that effect of protein abundance or brightness of images is largely removed.
We collected 3D time series images, in order to have as much information as possible on protein behavior. However, we started our exploration with 2D temporal patterns, by using the center slices over time. We used only the first eight time points in each movie, limited by the shortest movie in our image collection. The total period analyzed is therefore 6 min.

2.4 Object tracking features

Calculating features based on object properties, the change is small. The similarity between an object at time point \( t \) and the object at time point \( t_2 \) is calculated by the distance of their features:

\[
\sqrt{\frac{(S_{t_2} - S_{t_1})^2}{N_{t_2}}} + \frac{(Y_{t_2} - Y_{t_1})^2}{N_{t_2}} + \frac{(X_{t_2} - X_{t_1})^2}{N_{t_2}} + \frac{(I_{t_2} - I_{t_1})^2}{I_{t_2}}
\]

The distance is a normalized Euclidean distance so that features with large values do not dominate the result. Distances were calculated between all objects in each frame and all objects in the next frame, but object pairs that exceeded a distance threshold of 4 were discarded.

The next step is to compare the distance between all object pairs across neighboring images, and find the matching with the minimal sum of distances. Assume there are \( N_1 \) objects in the image at \( t_1 \) and \( N_2 \) objects in the image at \( t_2 \). \( N_1 \) and \( N_2 \) are not necessarily equal because objects can appear or disappear due to movement. The matching is a bipartite matching or an assignment problem; \( N_1 \) people available for \( N_2 \) jobs with different costs, where each person can do only one job and each job only needs one person. The best assignment is the one that minimizes the total cost. The naive way to compare different matchings is to calculate total distance for all possible matches, the computation of which is the factorial of the minimum of \( N_1 \) and \( N_2 \). We used the Hungarian Algorithm, a classic algorithm that reduces the computation to complexity \( O(N_1 N_2^2) \) (Kuhn, 1955).

After matching we know the position of each object in every time point and we can build a trajectory of each object. From these, we calculated:

**Speed**: the distance traveled over adjacent time points, divided by the time interval.

**Velocity**: the distance between an object’s position when it first appears and its position when it disappears (or at the end of the movie), divided by the total time. Velocity differs from speed in that it only reflects the absolute distance that the object has traveled. If the object travels back and forth around the same area, the speed is high but velocity is low.

**Acceleration**: the change in speed over adjacent time points, divided by the time interval.

**Angles between successive movements**: the angle \((0-180^\circ)\) between the two directions the object travels in adjacent images.

We used the mean and variance of these four features across the time series as the final features of the image (eight in total).

2.4.2 Temporal texture features

Temporal texture features were calculated as described previously (Hu et al., 2006). These were inspired by the Haralick texture features, which capture the intensity correlation of neighboring pixels in space (Haralick, 1979), and designed to capture the value correlation of neighboring pixels in time. We build the temporal gray-level co-occurrence matrix, whose element in row \( i \) and column \( j \) is the frequency that a pixel with value \( i \) changes to value \( j \) in the same position at the next time point. Co-occurrence matrices capture information of protein movement, for example, for proteins that show no movement between the two time points, the temporal co-occurrence matrix contains only zeros except on the diagonal.

We calculated 13 statistics described by Haralick (Haralick, 1979) based on the co-occurrence matrix for each pair of images separated by a certain time interval. We used the mean and variance across the time series as the final features of the image. By varying the time interval, we collected 130 features in total: means and variances of the 13 statistics for images 45, 90, 135, 180 and 225 apart.

2.4.3 Normal flow features

In a series of images taken over time, the intensity of each pixel in each image \( I(x, y, t) \) is a function of position \( x, y \) and time \( t \). To understand the movement, we need to infer from the observed \( I(x, y, t) \), the direction and velocity of each pixel. We use \( u \) to represent the vector of movement in the \( x \) direction and \( v \) to represent the vector of movement in the \( y \) direction.

\[
u = \frac{dx}{dt} \quad \text{and} \quad v = \frac{dy}{dt}
\]

Optical flow is a motion field with vectors \((u, v)\) at each pixel. From the definition, we know \((u, v)\) is pointing to the direction that the pixel moves...
and the length of the vector is the speed. Let us consider a pixel \( I(x, y, t = 1) \) that moves to a new position at \( t \). We here assume that the brightness of this pixel does not change:

\[
I(x, y, t = 1) = I(x + u(x, y, t) \Delta t, y + v(x, y, t) \Delta t, t)
\]

We further assume that the movement is not large. So the right-hand side formula can be approximated with first order:

\[
I(x, y, t = 1) \approx I(x, y, t) + u(x, y, t) \frac{dI}{dx} + v(x, y, t) \frac{dI}{dy}
\]

Rearranging the equation we have:

\[
u(x, y, t) \frac{dI}{dx} + v(x, y, t) \frac{dI}{dy} + I(x, y, t) - I(x, y, t - 1) = 0
\]

This is the fundamental equation of motion. \( u \) and \( v \) are the components of the gradient, which together with the difference of the intensity, \( I \), can be directly calculated from the two images. However we are left with one equation with two unknowns (\( u \) and \( v \)). This is the aperture problem: motion along the edge or perpendicular to the gradient can never be recovered. Meanwhile the motion \((u, v)\) along the gradient is

\[
u = u_I = \frac{I_y}{I_x^2 + I_y^2}
\]

and the speed of a pixel along the gradient is

\[
v = v_I = \frac{I_x}{I_x^2 + I_y^2}
\]

The flow field containing flows at the gradient direction is called normal flow. Using it, the following 34 features were calculated:

1. 13 Haralick texture features of \( U_\phi \)
2. Mean of \( U_\phi \)
3. SD of \( U_\phi \)
4. Mean/SD of \( U_\phi \)
5. Difference of direction from uniform distribution
6. Mean positive divergence
7. Mean negative divergence
8. Mean positive curl angular velocity
9. Mean negative curl angular velocity

For the texture features of direction, eight directions were used. We grouped them into eight directions, each representing one of the 0°, 45°, 90°, 135°, 180°, 225°, 270°, 315° directions. The speed of a voxel is defined similarly to 2D. Normal flow on \( x \) and \( y \) directions. The speed of a voxel along the gradient is

\[
u = \frac{I_y}{I_x^2 + I_y^2}
\]

and the length of the vector is the speed. Let us consider a pixel \( I(x, y, t = 1) \) that moves to a new position at \( t \). We here assume that the brightness of this pixel does not change:

\[
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6. Mean positive divergence
7. Mean negative divergence
8. Mean positive curl angular velocity
9. Mean negative curl angular velocity

The following is a list of the 3D temporal features that were calculated:

- \( U_\phi \) - Haralick texture features of the voxel intensity difference at each time point.
- Mean of \( U_\phi \)
- SD of \( U_\phi \)
- Mean/SD of \( U_\phi \)
- Difference of direction from uniform distribution
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For the texture features of direction, eight directions were used. We averaged the 0°, 45°, 90°, 135°, 180°, 225°, 270°, 315° directions. The speed of a voxel along the gradient is

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v(x, y, t) \frac{dI}{dx} + v(x, y, t) \frac{dI}{dy} + I(x, y, t) - I(x, y, t - 1) = 0
\]

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We further assume that the movement is not large. So the right-hand side formula can be approximated with first order:

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\]
series signal. Total of 40 features including 8 statistics of 5 coefficients, same as 2D, were calculated.

### 2.6 Feature selection

The large numbers of features described above could potentially overwhelm the ability of a classifier to identify meaningful decision boundaries. Therefore, we used stepwise discriminant analysis (SDA) to select the features that have the greatest power to discriminate the classes. While many feature selection methods have been described, SDA has performed well in previous studies of subcellular pattern classification (Huang et al., 2003).

### 2.7 Supervised learning (Classification)

A well established SVM package LIBSVM (Chang and Lin, 2001) was used for classification. LIBSVM uses the pair-wise algorithm to classify multiple classes and it has support for choosing optimal parameters. Ten-fold cross-validation was used to evaluate the classification accuracy: the images for each class were randomly divided into 10 groups of equal sizes. Nine of them were used to train a classifier and the remaining one was used for testing. Both feature selection and parameter tuning were done on the training set. SDA selected features were used to adjust the parameters until the classifier reached the best accuracy within the training set. The parameters were the radius of the radial basis function kernel (which was varied from 2\(^{-5}\) to 2\(^{5}\)) and the penalty for overlap between two classes (which was varied from 2\(^{-5}\) to 2\(^{5}\)). After training, the features for the test images were supplied to the classifier, and prediction was made on each image. By comparing the prediction with the class labels of the testing set, we obtain the classification accuracy. This process was repeated for each test fold, and the final overall accuracy is the average accuracy from the 10 tests. Since SVM can inherently handle a large number of correlated features without overfitting, we also repeated each classification without SDA feature selection.

### 3 RESULTS

By comparing classification accuracy using individual feature sets and combinations of them, we achieved two goals: evaluating the power of each feature set and finding the best way of differentiating our 12 proteins of interest. Three-dimensional time series images are 4D images. Feature sets we have acquired are based on 2D (2D static features of first center slice), 3D across \(z\) (3D static features), 3D across time (2D temporal features of center slices) and 4D images (3D temporal features).

#### 3.1 Classification using 2D static features

To set a reference for comparison, classification results of proteins in 3T3 cells were obtained using 21 2D static features. The overall accuracy was 63% with SDA feature selection and 66% without SDA. The confusion matrix without SDA is shown in Table 1. We can see that Cald1 has the lowest classification accuracy (25%).

#### 3.2 Classification using 2D temporal feature sets

To evaluate each temporal feature set, we trained classifiers using it with and without the static feature set. When object tracking features were combined with the 2D static features, the average classification accuracies were 61% and 68%, with or without SDA, showing a very small improvement over static pattern classification. From previous work we know when calculating static Haralick texture features, one can change the gray level or resize the image to different resolution and get different classification results (Murphy et al., 2003). Since the same effect would be expected in the temporal domain, we evaluated temporal texture features calculated using different pixel sizes and numbers of gray levels. Classification accuracies are summarized in Table 2. Accuracies fall within the range of 52–77% with SDA feature selection and 66–77% without SDA. The best accuracy of 77% was obtained for 64 gray levels and no downsampling. Apparently, accuracy is higher without SDA, and SVM inherently handled redundancy of features. To further test the power of temporal texture features, the same classification procedure was done without static features. The overall accuracies were combined with the 2D static features, the average classification accuracies were 61 and 68%, with or without SDA, showing a very small improvement over static pattern classification. The number before comma is the accuracy in percentage without SDA.

### Table 1. Confusion matrix for classification using only 2D static features

<table>
<thead>
<tr>
<th>True classes</th>
<th>Prediction by classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dia1</td>
<td>45 35 5 0 5 0 5 0 0 0 0 0</td>
</tr>
<tr>
<td>Anxa5</td>
<td>0 61 0 0 0 0 0 0 0 0 0 0</td>
</tr>
<tr>
<td>Sdpr</td>
<td>26 4 56 0 4 0 0 0 0 0 0 8</td>
</tr>
<tr>
<td>Adfp</td>
<td>0 1 0 90 1 1 0 0 0 0 0 0</td>
</tr>
<tr>
<td>Timm23</td>
<td>2 2 2 2 72 0 0 12 2 0 0 2</td>
</tr>
<tr>
<td>Agp5a</td>
<td>10 0 5 0 0 65 10 0 0 0 0 0</td>
</tr>
<tr>
<td>Hepat4a</td>
<td>4 0 0 4 8 4 70 0 0 0 8 0</td>
</tr>
<tr>
<td>Cat</td>
<td>0 0 0 43 6 6 37 6 0 0 0 0</td>
</tr>
<tr>
<td>Glat1</td>
<td>0 0 5 11 11 11 0 5 52 0 0 0</td>
</tr>
<tr>
<td>Tctex1</td>
<td>0 6 0 0 0 0 0 0 0 0 0 0</td>
</tr>
<tr>
<td>Actn4</td>
<td>0 20 3 3 0 0 3 0 0 3 58 6</td>
</tr>
<tr>
<td>Cald1</td>
<td>0 0 0 0 6 12 12 0 0 6 37 25</td>
</tr>
</tbody>
</table>

The overall accuracy is 66%.

#### Table 2. Summary of classification accuracy of 21 static combined with 130 temporal texture features, with different gray level and resolution

<table>
<thead>
<tr>
<th>Resolution (micron/pixel)</th>
<th>Gray level</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>0.22</td>
</tr>
<tr>
<td></td>
<td>0.86</td>
</tr>
<tr>
<td></td>
<td>1.1</td>
</tr>
</tbody>
</table>

The number before comma is the accuracy in percentage without SDA, and the number after comma is the accuracy in percentage without SDA.
classification accuracy was 67% with SDA and 66% without SDA. The results show that temporal texture features themselves capture a significant amount of information.

We next evaluated the normal flow features. Three hundred and forty normal flow features were combined with the 2D static features. Since the normal flow features include texture features calculated on U₀ and the normal flow direction, a similar exploration of number of gray levels and resolution was done. Classification accuracy ranged from 66% to 75% with SDA and 59% to 73% without SDA (data not shown). The best accuracy of 75% was achieved with the original resolution (0.11 μm/pixel) and 64 gray levels, with SDA. When the same classification procedure was done without static features, the overall classification accuracy was 75% with SDA and 74% without SDA. This indicates that these features were more informative than temporal texture features.

When 40 Fourier transform features were combined with the static features, the overall classification accuracy was 69% with SDA and 67% without SDA, a slight increase over using static features alone (data not shown). When using Fourier transform features only, the overall accuracy was 60 and 57% with and without SDA (data not shown).

We next combined 189 autoregression features with the static features. The overall classification accuracy was 59% with SDA and 48% without SDA (data not shown). These are both lower than using static features alone.

From this analysis of each temporal feature set, we found that temporal texture features, normal flow features and Fourier transform features are capable of increasing classification accuracy. We combined these 3 sets of temporal features to search for the best way to differentiate the 12 proteins in 2D time series images. If combined with 2D static features, the overall accuracy is 75% with SDA and 78% without SDA. With temporal features only, very good classification accuracy of 73 and 77% (with and without SDA) can be achieved, proving again the power of the temporal features.

The confusion matrix of the best classification is shown in Table 3. Comparing the results with Table 1, the overall accuracy is 12% higher than using just 2D static features. Tctex1 was now classified perfectly. The lowest accuracy of 25% for cald was increased to 56%, and accuracy for Sdpr increased from 56% to 95%. Two other proteins, Anxa5 and Actn4, also have over 20% improvement in accuracy. While the accuracy with SDA selection was lower than without, it is interesting to note that SDA selected 7 static, 15 temporal features, 13 normal flow and 4 Fourier transform features, from the total of 531 features. This suggests that all of these feature types provide useful information.

### 3.3 Classification using 3D static features

Static features based on 3D images capture information of images with one more dimension than 2D images. Since we were not able to perform automatic segmentation due to lack of nuclear channel, we selected those features that do not compare to center of cells, thus do not require segmentation. Classification accuracy using 33 field level 3D static features was 74% with SDA, and 71% without SDA. It is 6% higher than using 2D static features and 4% lower than using combined 2D static and temporal features. When 3D static and 2D static features were combined, classification accuracy was 73 and 70%, with and without SDA. As might be expected, the 3D features apparently contain all of the information in the 2D features, and adding redundant 2D static features only made classifier learning more difficult.

### 3.4 Classification using 3D temporal feature set

Similar to the way we tested 2D temporal features, we combined 3D temporal features with 3D static features to see if the classification accuracy was improved. The result is disappointing. Classification accuracies with or without SDA for 3D temporal textures were 74 and 75%, for 3D normal flow were 71 and 72% and for 3D Fourier transform were 73 and 71%. Only 3D temporal texture features, combined with 3D static features, performed slightly better than using 3D static features alone. None of the accuracies were higher than the accuracy we achieved combining 2D temporal features with 2D static features. It appears that calculating features over the full 3D images diluted the critical information in the central slices.

### 4 CONCLUSION AND DISCUSSIONS

Location proteomics as a branch of proteomics study has grown over the last 10 years. To systematically analyze large amounts of data, automatic algorithms have been developed. This article presented...
our effort to extend the analysis of static patterns with an additional dimension: the temporal domain. Time series microscopy image of 12 proteins were collected. 5 sets of 2D temporal features and 3 sets of 3D temporal features were implemented and classifications were performed to validate their usefulness. The best 2D temporal feature sets, in the order of their ability to improve classification accuracy, were normal flow, temporal texture and Fourier transform features. Combining 2D static features with 3 sets of 2D temporal feature sets gave the best accuracy of 78%, compared with 66% for static features alone. Accuracy using 3D static and/or temporal features was lower than for 2D features.

If limited acquisition time requires deciding whether to collect 3D static images or 2D time series images, our results suggest that 2D time series images have higher potential of delivering better differentiation. Although not all of the 2D or 3D temporal feature sets improved classification accuracy for our dataset, we still presented them here because they may be useful for future datasets.

While each protein in a proteome is unique, its location patterns might not be. Thus while increasing accuracy of distinguishing the 12 proteins is an indication of feature value, the ability to distinguish them all perfectly is not expected. If proteins interact or colocalize with each other, they cannot be differentiated either by static or temporal pattern. In our dataset, alpha-actinin-4 (actn4) and caldesmon 1 (cald1) both bind to actin, and over 30% of caldesmon 1 is misclassified as alpha-actinin-4. These two proteins are always observed in the same cluster using cluster analysis (data not shown). Similarly, ADP-ATP translocase 23 (timm23) and catalase (cat), which have both been described as mitochondrial proteins, are difficult to distinguish (50% of catalase is classified as ADP-ATP translocase 23). The results suggest that there are only 10 distinguishable patterns in our 12 protein set, and this agrees with prior knowledge about these proteins.

For each classification, we compared accuracy with or without SDA feature selection, because SVM is known to be highly robust with large number of correlated features. Our result shows when the number of features is large, 340 normal flow and 189 autoregression, SDA outperforms no feature selection. When the number of features is small, 21 static and 130 temporal textures, SVM does well without SDA feature selection. Many different feature selection algorithms and classifiers could be tried in order to achieve higher classification accuracy, but such an analysis is beyond the scope of this study.

Since temporal texture features and Fourier transform features can be calculated within 25 and 50 s for each time series, they are readily applicable to many high throughput applications. On the other hand, calculating normal flow features, object tracking features and autoregression features take 8, 22 and 2 min per time series, respectively.

Given the dramatic increase in automated microscopy over the past decade, we anticipate that methods for analyzing temporal changes in protein patterns such as those we have described here will be of significant utility both for basic research in systems biology and for drug screening and development purposes.

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REFERENCES


