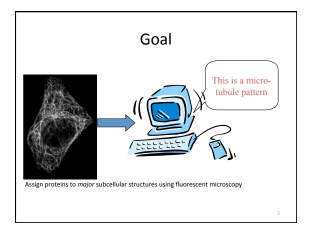
Machine Learning Approaches to Biological Research: Bioimage Informatics and Beyond

Lecture 2: Concepts of automated image analysis

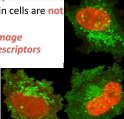
Robert F. Murphy
External Senior Fellow, Freiburg Institute for Advanced Studies
Ray and Stephanie Lane Professor of Computational Biology, Carnegie
Mellon University

September 29-October 1, 2009



The Challenge

- Problem is hard because different cells have different shapes, sizes, orientations
- Organelles/structures within cells are not found in fixed locations
- Therefore, describe each image numerically and use the descriptors



Feature-Based, Supervised Learning Approach

- Create sets of images showing the location of many different proteins (each set defines one class of pattern)
- 2. Reduce each image to a set of numerical values ("features") that are insensitive to position and rotation of the cell
- 3. Use statistical classification methods to "learn" how to distinguish each class using the features

Acquisition considerations

- For automated acquisition
 - Optimize autofocus parameters
 - Maintain constant camera gain, exposure time, number of slices
 - Select interphase cells or ensure sampling of cell cycle

Acquisition considerations

- Collect sufficient images per condition
 - For classifier training or set comparison, more than number of features
 - For classification or clustering, base on confidence level desired
- Collect reference images if possible (DNA, membrane)

Annotation considerations

- Maintain adequate records of all experimental settings
- Organize images by cell type/probe/condition

Preprocessing

- Correction for/Removal of camera defects
- Background correction
- Autofluorescence correction
- Illumination correction
- Deconvolution

Preprocessing (continued)

- Registration
 - Not critical if only using DNA or membrane references
- Intensity scaling (constant scale or contrast stretched for each cell)
- Single cell segmentation
 - Manual, semi-automated, automated
- Region finding
 - Nucleus
 - Cytoplasmic annulus
 - Cell boundary

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Segmentation of Images into Single Cell Regions

Approaches

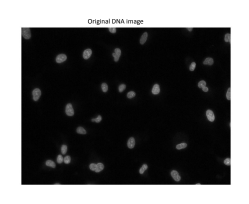
- Voronoi
- Watershed
- · Seeded Watershed
- Level Set Methods
- Graphical Models

Given a set of seeds, draw vertices and edges such that each seed is enclosed in a single polygon where each edge is equidistant from the seeds on either side.

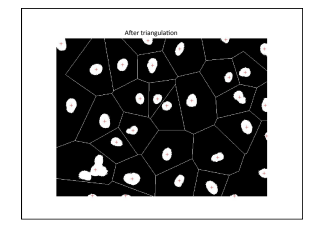
4

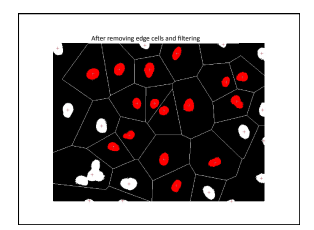
Voronoi Segmentation Process

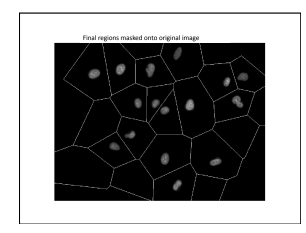
- Threshold DNA image (downsample?)
- Find the objects in the image
- Find the centers of the objects
- Use as seeds to generate Voronoi diagram
- Create a mask for each region in the Voronoi diagram
- Remove regions whose object that does not have intensity/size/shape of nucleus











Watershed Segmentation

- Intensity of an image of elevation in a landscape
 - Flood from minima
 - Prevent merging of "catchment basins"
 - Watershed borders built at contacts between basins



http://www.ctic.purdue.edu/KYW/glossary/whatisaws.html

Watershed Segmentation

- If starting image has intensity centered on the cells (e.g., DNA) that you want to segment, invert image so that bright objects are the sources
- If starting image has intensity centered on the boundary between the cells (e.g., plasma membrane protein), don't invert so that boundary runs along high intensity

Seeded Watershed Segmentation

- Drawback is that the number of regions may not correspond to the number of cells
- Seeded watershed allows water to rise only from predefined sources (seeds)
- If DNA image available, can use same approach to generate these seeds as for Voronoi segmentation
- Can use seeds from DNA image but use total protein image for watershed segmentation

Seeded Watershed Segmentation

Original image

Seeds and boundary

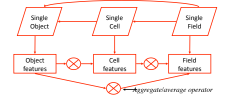
Applied directly to protein image (no DNA image) Note non-linear boundaries

Feature Extraction for Subcellular Pattern Analysis

Subcellular Location Features (SLF)

- Combinations of features of different types that describe different aspects of patterns in fluorescence microscope images have been created
- Motivated in part by descriptions used by biologists (e.g., punctate, perinuclear)
- To ensure that the specific features used for a given experiment can be identified, they are referred to as <u>Subcellular Location Features</u> (SLF) and defined in sets (e.g., SLF1)

Feature levels and granularity



Granularity: 2D, 3D, 2Dt, 3Dt

Thresholding

- First type of feature is morphological
- Morphological features require some method for defining objects
- Most common approach is global thresholding
- Methods exist for automatically choosing a global threshold (e.g., Riddler-Calvard method)

Ridler-Calvard Method

- Find threshold that is equidistant from the average intensity of pixels below and above it
- Ridler, T.W. and Calvard, S. (1978) Picture thresholding using an iterative selection method. *IEEE Transactions on Systems, Man, and Cybernetics* 8:630-632.

Ridler-Calvard Method Blue line shows histogram of intensities, green lines show average to left and right of red line, red line shows midpoint between them or the RC threshold

Ridler-Calvard Method original original thresholded

Otsu Method

- Find threshold to minimize the variances of the pixels below and above it
- Otsu, N., (1979) A Threshold Selection Method from Gray-Level Histograms, IEEE Transactions on Systems, Man, and Cybernetics, 9:62-66.

Adaptive Thresholding

- · Various approaches available
- Basic principle is use automated methods over small regions and then interpolate to form a smooth surface

Suitability of Automated Thresholding for Classification

- For the task of subcellular pattern analysis, automated thresholding methods perform quite well in most cases, especially for patterns with well-separated objects
- They do not work well for images with very low signal-noise ratio
- Can tolerate poor behavior on a fraction of images for a given pattern while still achieving good classification accuracies

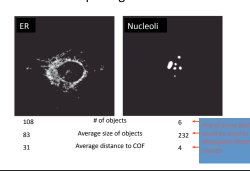
Object finding

 After choice of threshold, define objects as sets of touching pixels that are above threshold

2D Features Morphological Features

SLF No.	Description
SLF1.1	The number of fluorescent objects in the image
SLF1.2	The Euler number of the image
SLF1.3	The average number of above-threshold pixels per object
SLF1.4	The variance of the number of above-threshold pixels per object
SLF1.5	The ratio of the size of the largest object to the smallest
SLF1.6	The average object distance to the cellular center of fluorescence(COF)
SLF1.7	The variance of object distances from the COF
SLF1.8	The ratio of the largest to the smallest object to COF distance

2D Features Morphological Features



Suitability of Morphological Features for Classification

- Images for some subcellular patterns, such as those for cytoskeletal proteins, are not well-segmented by automated thresholding
- When combined with non-morphological features, classifiers can learn to "ignore" morphological features for those classes

2D Features DNA Features

DNA features (objects relative to DNA reference)

SLF No.	Description
SLF2.17	The average object distance from the COF of the DNA image
SLF2.18	The variance of object distances from the DNA COF
SLF2.19	The ratio of the largest to the smallest object to DNA COF distance
SLF2.20	The distance between the protein COF and the DNA COF
SLF2.21	The ratio of the area occupied by protein to that occupied by DNA
SLF2.22	The fraction of the protein fluorescence that co-localizes with DNA

2D Features Skeleton Features

Skeleton features

SLF No.	Description
SLF7.80	The average length of the morphological skeleton of objects
SLF7.81	The ratio of object skeleton length to the area of the convex hull of the skeleton, averaged over all objects
SLF7.82	The fraction of object pixels contained within the skeleton
SLF7.83	The fraction of object fluorescence contained within the skeleton
SLF7.84	The ratio of the number of branch points in the skeleton to the length of skeleton

Illustration – Skeleton









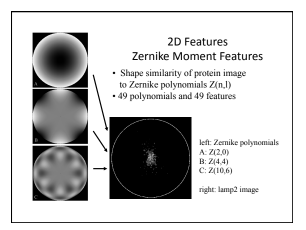




2D Features Edge Features

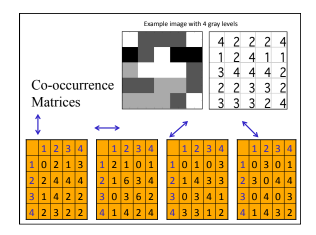
Edge features

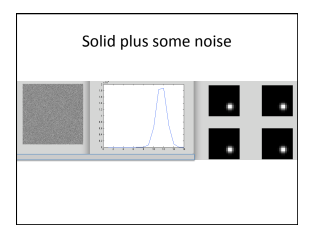
SLF No.	Description	
SLF1.9	The fraction of the non-zero pixels that are along an edge	
SLF1.10	Measure of edge gradient intensity homogeneity	
SLF1.11	Measure of edge direction homogeneity 1	
SLF1.12	Measure of edge direction homogeneity 2	
SLF1.13	SLF1.13 Measure of edge direction difference	
	·	

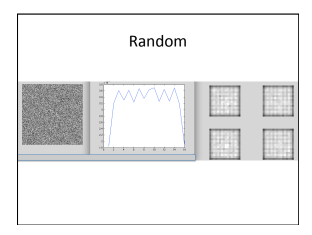


2D Features Haralick Texture Features

- Correlations of adjacent pixels in gray level images
- Start by calculating co-occurrence matrix P:
 N by N matrix, N=number of gray level.
 Element P(i,j) is the probability of a pixel with value i being adjacent to a pixel with value j
- Four directions in which a pixel can be adjacent
- Each direction considered separately and then features averaged across all directions







Checkerboard

Pixel Resolution and Gray Levels

- Texture features are influenced by the number of gray levels and pixel resolution of the image
- Optimization for each image dataset required
- Alternatively, features can be calculated for many resolutions

Wavelet Transformation - 1D X A1 D1 A2 D2 A: approximation (low frequency) D: detail (high frequency) X=A3+D3+D2+D1

2D Wavelets - intuition

• Apply some filter to detect edges (horizontal; vertical; diagonal)





After Christos Faloutsos

2D Wavelets - intuition

• Recurse







Slide courtesy of Christos Faloutsos

2D Wavelets - intuition

- Many wavelet basis functions (filters):
 - Haar
 - Daubechies (-4, -6, -20)
- http://www331.jpl.nasa.gov/public/wave.html



Slide courtesy of Christos Faloutso

Daubechies D4 decomposition





Original image

Wavelet Transformation

2D Features Wavelet Feature Calculation

- Preprocessing
 - Background subtraction and thresholding
 - Translation and rotation
- Wavelet transformation
 - The Daubechies 4 wavelet
 - 10 level decomposition
 - Use the average energy of the three high-frequency components at each level as features

3D Features Morphological

- 28 features, 14 from protein objects and 14 from their relationship to corresponding DNA images
 - Based on number of objects, object size, object distance to COF
- Corresponding DNA image required

3D set

- 14 SLF-9 features that do not require DNA images
- · 2 Edge features
 - Ratio of above threshold pixel along an edge
 - Ratio of fluorescence along an edge
- 26 3D Haralick texture features
 - Gray level co-occurence matrix for 13 directions
 - Calculate 13 Haralick statistics for each direction
 - Average each statistic over 13 directions and use mean and range as separate features: result is 26 features

Object level features (SOF)

• Subset of SLFs calculated on single objects

Index	Feature Description
SOF1.1	Number of pixels in object
SOF1.2	Distance between object Center of Fluorescence (COF) and DNA COF
SOF1.3	Fraction of object pixels overlapping with DNA
SOF1.4	A measure of eccentricity of the object
SOF1.5	Euler number of the object
SOF1.6	A measure of roundness of the object
SOF1.7	The length of the object's skeleton
SOF1.8	The ratio of skeleton length to the area of the convex hull of the skeleton
SOF1.9	The fraction of object pixels contained within the skeleton
SOF1.10	The fraction of object fluorescence contained within the skeleton
SOE1 11	The ratio of the number of branch naints in skalaten to length of skalaten

Field level features

- Subset of SLFs that do not require segmentation into single cells
 - Average object features
 - Texture features (on whole field)
 - Edge features (on whole field)

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2Dt or 3Dt Features Temporal Texture Features

- Haralick texture features describe the correlation in intensity of pixels that are next to each other in space.
 - These have been valuable for classifying static patterns.
- Temporal texture features describe the correlation in intensity of pixels in the same position in images next to each other over time.

Temporal Textures based on Co-occurrence Matrix

- Temporal co-occurrence matrix P:
 N_{level} by N_{level} matrix, Element P[i, j] is the probability that a pixel with value i has value j in the next image (time point).
- Thirteen statistics calculated on P are used as features

Image at t0 Image at t1 4 2 2 2 4 4 2 2 2 4 1 2 4 1 1 1 2 4 1 1 3 4 4 4 2 3 4 4 4 2 2 2 3 3 2 2 2 3 3 2 3 3 Temporal 0 0 0 co-occurrence matrix (for 0 9 0 0 image that does 0 0 6

0

not change)

-		

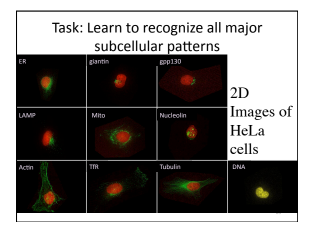
Image at t0		Ima	age	at t1		
4 2 2 2 4	2	1	4	4	3	
1 2 4 1 1	1	4	2	3	3	
3 4 4 4 2	2	3	3	2	2	
2 2 3 3 2	4	4	2	2	3	
3 3 3 2 4	2	4	2	1	4	
Temporal		1	2	3	4	
co-occurrence	1	1	0	2	0	
matrix (for	2	2	1	1	5	
image that	3	0	5	0	1	
changes)	4	0	3	3	1	

Implementation of Temporal Texture Features

 Compare image pairs with different time interval ,compute 13 temporal texture features for each pair.

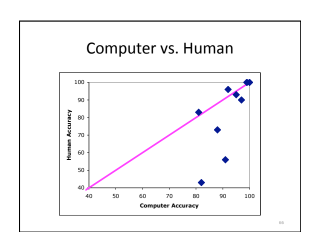


 Use the average and variance of features in each kind of time interval, yields 13*5*2=130 features



2D Classification Results True Clas Output of the Classifier DNA ER Gia Gpp Lam Mit Nuc Act TfR Tub DNA ER Gpp Lam Mit 0 Nuc Act Tub Overall accuracy = 92%

Human Classification Results										
True	True Output of the Classifier									
Clas s	DNA	ER	Gia	Gpp	Lam	Mit	Nuc	Act	TfR	Tub
DNA	100	0	0	0	0	0	0	0	0	0
ER	0	90	0	0	3	6	0	0	0	0
Gia	0	0	56	36	3	3	0	0	0	0
Gpp	0	0	54	33	0	0	0	0	3	0
Lam	0	0	6	0	73	0	0	0	20	0
Mit	0	3	0	0	0	96	0	0	0	3
Nuc	0	0	0	0	0	0	100	0	0	0
Act	0	0	0	0	0	0	0	100	0	0
TfR	0	13	0	0	3	0	0	0	83	0
Tub	0	3	0	0	0	0	0	3	0	93



3D Classification Results

True	Output of the Classifier										
Clas	DNA	ER	Gia	Gpp	Lam	Mit	Nuc	Act	TfR	Tub	
DNA	98	2	0	0	0	0	0	0	0	0	
ER	0	100	0	0	0	0	0	0	0	0	
Gia	0	0	100	0	0	0	0	0	0	0	
Gpp	0	0	0	96	4	0	0	0	0	0	
Lam	0	0	0	4	95	0	0	0	0	2	
Mit	0	0	2	0	0	96	0	2	0	0	
Nuc	0	0	0	0	0	0	100	0	0	0	
Act	0	0	0	0	0	0	0	100	0	0	
TfR	0	0	0	0	2	0	0	0	96	2	
Tub	0	2	0	0	0	0	0	0	0	98	

Overall accuracy = 98%

Supervised vs. Unsupervised Learning

- Work discussed so far demonstrates the feasibility of using classification methods to assign all proteins to known major classes
- Do we know all locations? Are assignments to major classes enough?
- Need approach to discover classes

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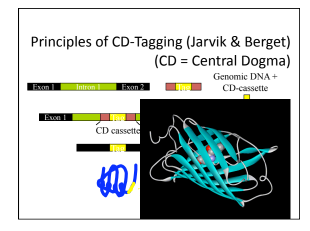
Location Proteomics

Tag many proteins
We have used CD-tagging
(developed by Jonathan Jarvik and
Peter Berget): Infect population of
cells with a retrovirus carrying DNA
sequence that will "tag" in a random gene in a





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Location Proteomics

Tag many proteins lag many proteins

- We have used CD-tagging
(developed by Jonathan Jarvik and
Peter Berget): Infect population of
cells with a retrovirus carrying DNA
sequence that will "tag" in a random gene in
Isolate separate clones, each of which produces express one tagged



Use RT-PCR to **identify tagged gene** in each clone
Collect many live cell images for each clone using spinning disk
confocal fluorescence microscopy

