## Computational Vision U. Minn. Psy 5036 Daniel Kersten Lecture 12: Coding Efficiency, Spatial statistics

#### Initialize

```
Off[General::spell1];
SetOptions[ArrayPlot, ColorFunction → "GrayTones",
DataReversed → True, Frame → False, AspectRatio → Automatic,
Mesh → False, PixelConstrained → True, ImageSize → Small];
SetOptions[ListPlot, ImageSize → Small];
nbinfo = NotebookInformation[EvaluationNotebook[]];
```

```
dir = ("FileName" /. nbinfo /. FrontEnd`FileName[d_List, nam_, ___] :> ToFileName[d]);
```

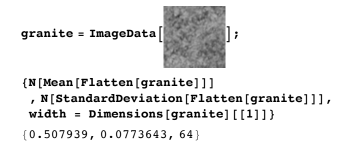
#### Histogram

```
myhistogram[image_] := Module[{histx},
    histx = BinCounts[Flatten[image], {0, 255, 1}];
    Return[N[histx / Plus@@histx]];
];
```

#### Entropy

entropy[probdist\_] := Plus @@ (If[# == 0, 0, -#Log[2, #]] & /@probdist)

#### Image data



## Outline

#### Last time

First-order intensity statistics. Explain point non-linearities in terms of histogram equalization natural image intensities

The assumption was that the cell's output range was effectively fixed to encode N levels, and that an efficient use of those levels was to not favor one over another. Limited resolution doesn't mean that the output is digital, but rather that noise limits the number of resolvable levels.

The question then was how best to allocate input contrasts to those levels, so that probability of using an output value was equal across all N levels.

The answer was to use the cumulative distribution function of the input histogram as the mapping function. If the input is bell-shaped, a sigmoidal non-linearity results.

#### Introduction to 2nd order statistics

A difference histogram is a "marginal" distribution, which means the histogram you get when you project higher-dimensional data points onto a lower-dimensional axis. If natural images were gaussian, the marginals would be too. But in general, they are not. The differences concentrate near zero ("blue sky effect"), but also get spread out in the tails (big jumps at edges). Difference histograms for natural images are said to have "excess kurtosis"--i.e. a normal distribution has kurtosis of 3, but natural images tend to have larger values.

This doesn't just show up with simple differences between nearby pixels, but more generally with any derivative-type filter in which uniform values map to zero. (i.e. because the excitatory and inhibitory contributions cancel out). For example, consider the  $\nabla^2 G$  filter:





Kurtosis [Flatten [ImageData [LaplacianGaussianFilter

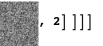


5.69827

compared with the kurtosis of gaussian noise:

igaussiannoise = RandomImage[NormalDistribution[0, .2], {64, 64}] // ImageAdjust;

Kurtosis [Flatten [ImageData LaplacianGaussianFilter]



2.97776

#### Today

#### 2nd order spatial statistics and efficient coding

We've learned about localized spatial frequency filters in early vision. We now ask: Why?

## Efficient representation of information: the range problem

When we considered the rationale for a point-wise sigmoidal non-linearity, we assumed a fixed output range. But what if we could code the input efficiently so that an even smaller range would do. This could lead to metabolic savings. First, let's see why there is a range problem.

We'll first consider the single-channel spatial filtering model and retinal coding. Lateral inhibition is pervasive in early visual coding across many species of animals, from invertebrates like the horseshoe crab to primates. We would like to know why, in other words, come up with a computational theory for lateral inhibition. We already saw an argument for lateral inhibition as a front-end for edge detection. It is also a means to reduce the dynamic range--but is there a principled way of reducing the dynamic range while avoiding discarding information? Let's look at possible explanation is in terms of efficient encoding.

The retina needs to encode a large number of levels of light intensities into a small number of effective neuronal levels. There is the huge range of physical light energy, ranging from  $10^{-6}$  to  $10^{7}$  candelas/ $m^{2}$  (a measure of luminance of a surface)--from the just visible to painfully bright. But the number of distinguishable levels can be much smaller over a large range of intensities.

A quick calculation based on Poisson statistics shows that in about a 1/5 second, *there are about 200 reliably distinguishable light levels* given a potential range of between  $10^{10}$  and  $10^{-2}$  photon-s/sec/receptor at 555 nm.

A similar calculation based on Poisson statistics for neural discharge indicates *only about 14-16 levels can be encoded in 1/5 of a second*. (Ganglion cell discharge is in general modeled by a Gamma distribution on inter-spike intervals, and Poisson statistics are a convenient approximation that corresponds to a first order gamma distribution; Gerstein, 1966; Robson and Troy, 1987.)

Let's make a calculation based on a first order Poisson approximation:

$$p (k \text{ spikes in } \Delta t) = \frac{e^{-\lambda \Delta t} \lambda \Delta t^{k}}{k!}$$

( $\lambda$ =average rate,  $\lambda$ (t)=f(intensity or contrast). Because of the refractory period, the maximum rate is less than 1000 Hz. In general, it is much lower for ganglion cells, and 250 would be a liberal upper bound. And 250 Hz => 50 spikes in 1/5 sec. Working down in steps of 1 standard deviation produces about 14 levels. The big challenge then is how to go from 200 levels to 14:

 $Log_2200 \rightarrow Log_214$ , with minimal loss of information?

This would require squeezing 7.6 to 3.8 bits/receptor. Of course, we don't have to handle this whole range for a given scene and using a single mechanism. A duplex receptor system (rods and cones) helps, and sigmoidal compression. We've also noted, this is not simply a matter of introducing a non-linearity--this will not work because the variability is the ultimate limit to resolution and it would still remain. What tricks that could be used to handle the range problem?

It turns out that for an arbitrary image ensemble where there are no spatial (or temporal) dependencies, one cannot construct a reversible coding scheme that could squeeze the number of bits down. But for an image ensemble with some statistical structure or redundancy, there is hope. What is meant by statistical structure or redundancy?

In a 128 x 128 x 4 bit graphics display, there are 2^(128\*128\*4) or about 10^19,728 possible pictures. Imagine a machine that started iterating through them. The vast majority would appear unnatural and look like TV "snow" or visual noise. Only a near infinitesmal small fraction would correspond to natural images...i.e. are likely to occur. So what is this fraction? Some years ago, I estimated an upper

bound on this fraction using theoretical results from Claude Shannon's famous guessing game for the predictability of written English text (Kersten, 1987;D'Antona et al., 2013). The result was that number of possible meaningful images <  $10^{6905}$ . If you could sit for multiple eons of time and view all the  $10^{19,728}$  on your 128 x 128 x 4 bit computer display, about one out of every  $10^{12,823}$  pictures and your brain would "click" and you would say "aha, that one looks natural." Why is this? One fundamental reason is that there are correlations between neighboring pixel intensities. Correlations are one simple and basic measure of redundancy in images.



We need tools for measuring correlations, and redundancy in images.

## 2nd order statistics

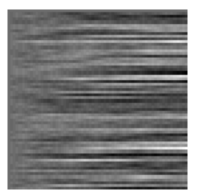
#### Example of the idea: a non-isotropic "I-D random-walk" image ensemble

We can build our intuitions be considering a space of 1-D images that, like natural images, is constrained to have similar nearby pixels. We start with a gray-level of 128, and then flip a coin to decide whether to increase or decrease the intensity of the next pixel. So nearest-neighbor pixels are close, but not identical in intensity.

#### I-D Brownian images

```
step := 2 (Random[Integer, 1] - 1 / 2);
next[x_] := Mod[x, size] + 1;
size = 64;
brown = N[Table[128, {i, 1, size }, {i, 1, size }]];
For[j = 1, j < size, j++,
For[i = 1, i < size, i++,
If[Random[] > 0.5, brown[[next[i], j]] = brown[[i, j]] + step,
brown[[next[i], j]] = brown[[i, j]] - step];
If[brown[[i, j]] > 255, 255];
If[brown[[i, j]] < 1, 0];
];
];
```

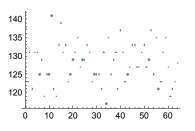
Visual each 1-D image using Image[]. Let's stack the images horizontally, one on top of the other:



ImageRotate[Image[brown, ImageSize o Small] // ImageAdjust]

Once we get far enough away from the starting value of 128, along a vertical line, the intensity samples become fairly independent of each other. The gray-levels from pixel to pixel are not correlated:

ListPlot[brown[[32]], ImageSize → Small]

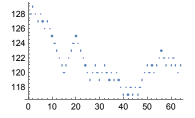


{Entropy[2, brown[[32]]] // N, entropy[myhistogram[brown[[32]]]]}

 $\{3.44411, 3.44411\}$ 

In contrast, horizontal lines show a degree of regularity:

 $\texttt{ListPlot[Transpose[brown][[32]], ImageSize} \rightarrow \texttt{Small}]$ 



{Entropy[2, Transpose[brown][[32]]] // N, entropy[myhistogram[Transpose[brown][[32]]]] // N}

{**3.38946**, **3.38946**}

```
histobrown = myhistogram[brown];
ListPlot[histobrown, PlotStyle \rightarrow PointSize[0.015], PlotRange \rightarrow {0, 0.1}]
entropy[histobrown] // N
0.00
0.04
0.02
0 50 100 150 200 250
```

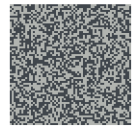
```
4.46008
```

#### Efficient encryption code for I-D brownian images

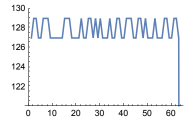
Let's encode the brownian images as the difference between neighboring pixel values:

```
codebrown = Table[0, {size}, {size}];
For[j = 1, j < size, j++,
For[i = 1, i < size, i++,
    codebrown[[i, j]] = brown[[next[i], j]] - brown[[i, j]] + 128;
];
];
```

```
ArrayPlot[codebrown, Mesh \rightarrow False, PlotRange \rightarrow {126, 130}]
```



 $\texttt{ListPlot[codebrown[[32]], PlotRange} \rightarrow \{120, 130\}, \texttt{Joined} \rightarrow \texttt{True, ImageSize} \rightarrow \texttt{Small}\}$ 



```
histocodebrown = myhistogram[Flatten[codebrown]];
ListPlot[histocodebrown, PlotStyle \rightarrow PointSize[0.015], PlotRange \rightarrow {0, 1}]
entropy[histocodebrown]
1.0
0.8
0.6
0.4
0.2
0 50 100 150 200 250
1.16837
```

#### Second order statistics in natural images

Let's now look at ways of quantifying second-order dependencies. First we look at 1D data, by analyzing dependencies across single lines of an image.

#### Autocorrelation function

ListCorrelate[ker, list] computes  $\sum_r K_r a_{s+r}$ . Autocorrelation corresponds to  $K_r \rightarrow a_r$ :  $\sum_r a_r a_{s+r}$ . Let's analyze the correlation between pixel gray levels for each line, and then average them:

#### Example with granite image

Image[granite]



```
autogranite = Table[0, {width}];
For [i = 1, i < width + 1, i++,</pre>
  autogranite += ListCorrelate[granite[[i]], granite[[i]], width / 2]];
ListPlot[autogranite/Max[autogranite],
 Joined \rightarrow True, PlotRange -> {.95, 1}, ImageSize \rightarrow Small]
1.00
0.99
0.98
0.97
0.96
      10
          20
             30
                 40
                     50
                         60
  0
```

Note how this measure of correlation drops as pairs of lines get shifted away from each other.

#### Covariance matrices, and the outer product

Recall that the covariance is:  $Cov[X, Y] = E[[X - \mu_X] [Y - \mu_Y]]$ , where X,Y are scalar random variables. The correlation gives a dimensionless measure of covariation, relative to the standard devia-

tions:  $\rho [X, Y] = \frac{Cov[X,Y]}{\sigma_X \sigma_Y}$ .

Now let X={ $x_1 \dots$ } and Y = { $y_j \dots$ } be vectors, and the lower case represent the scalar random variable elements. The average of the products  $x_i y_j$  or  $(x_i - \mu_{x_i}) (y_j - \mu_{y_j})$  give measures of how well  $x_i$  and  $y_j$  predict each other. The latter collection of average products is called the covariance matrix:

 $\operatorname{Cov}[X, Y] = E\left[[X - \mu_{X}] [Y - \mu_{Y}]^{T}\right]$ 

where  $XY^{T}$  is the notation for outer product of X and Y. Mathematica for the outer product is: Outer[-Times, **X**,**Y**]. The outer product takes two vectors and produces the matrix whose entries are all possible pair-wise products of the elements of the two vectors. Contrast the outer with the inner (or dot) product which returns a scalar given two input vectors.

Try this: Outer[Times, {x, y, z}, {a, b, c}] // MatrixForm

Given M vector samples indexed by s,  $\{X^s, Y^s\}$ , we can estimate the covariance matrix as:

$$\frac{1}{M}\sum_{s=1}^{M} [\boldsymbol{X}^{s} - \boldsymbol{\mu}_{\boldsymbol{X}}] [\boldsymbol{Y}^{s} - \boldsymbol{\mu}_{\boldsymbol{Y}}]^{T}.$$

When X=Y, an covariance matrix is called an autocovariance matrix, and similarly for autocorrelation. A covariance matrix is a symmetric matrix, and thus has orthogonal eigenvectors with real eigenvalues--a property that will become useful later.

Try this: Outer[Times, {x, y, z}, {x, y, z}] // MatrixForm

#### Multivariate gaussian (See ProbabilityOverview.nb, Lecture 3)

If the distribution is assumed to be multivariate gaussian, then the vector mean and covariance matrix fully determines the distribution. The multivariate gaussian is a generalization of the gaussian distribution to higher dimensions, in which the standard deviation is replaced by the covariance matrix. The multivariate gaussian plays a central role in statistics, and provides a crude approximation as a generative model for natural images. The probability density for vector x of dimension p is given by:

$$\mathbf{p}(\mathbf{x}) = \frac{1}{\sqrt{(2\pi)^{p} |\Sigma|}} e^{-\frac{1}{2}(\mathbf{x}-\mu)^{T} \Sigma^{-1}(\mathbf{x}-\mu)}, \text{ where } |\Sigma| = \mathsf{Det}[\Sigma].$$

where  $\mu$  is the vector mean, and  $\Sigma$  is the covariance matrix. *Mathematica* has an add-in package that extends the normal routines to the multivariate case:

A two-dimensional example.

```
\Sigma = \{\{1, .6\}, \{.6, 1\}\};
\mu = \{1, 1\};
ndist = MultinormalDistribution [\mu, \Sigma];

ContourPlot[PDF[ndist, {x, y}], {x, -1, 3}, {y, -1, 3}, ImageSize \rightarrow Tiny, ColorFunction \rightarrow "DarkRainbow", ContourStyle \rightarrow None]

3^{2}
1^{-1}
-1^{-1}
0
1^{-2}
3^{-1}
```

Going to higher dimensions, an exponential drop-off in correlation can be modeled as a covariance matrix with diagonal elements equal to 1, and an exponential drop-off away from the diagonal. So the first row would be:

#### $row1[\rho_{-}] := Table[\rho^{i}, \{i, 0, 15\}];$

Later we show how the covariance matrix can be used to find a new basis set for images such that when we project images onto the basis elements, the projections are no longer correlated. One way to do this is through the classical statistical technique called Principal Components Analysis or PCA.

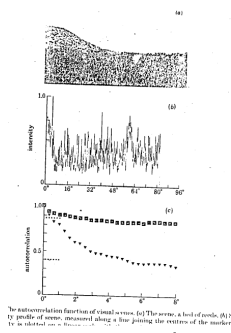
But first, let's look at some early and recent research that has sought to explain receptive field structure in terms of redundancy reduction.

## Efficient coding by the retina

#### Predictive coding & retina

Srinivasan et al. (1982) were the first to make quantitative predictions of how the retina makes use of inherent spatial and temporal correlations between light intensities found in natural images to reduce the output range required to send information about images. They showed that the autocorrelation of a natural image could be fit with an exponential curve.

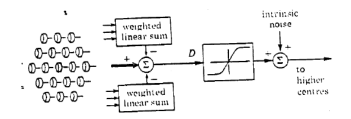
#### Autocorrelation measurements & model



$$E(L_i L_j) = \sigma^2 exp\left(-\frac{d_{ij}}{D}\right) + M^2 + N^2 \delta_{ij}$$

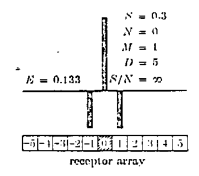
#### Linear neural network

They assumed a linear model:

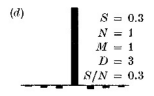


#### The result

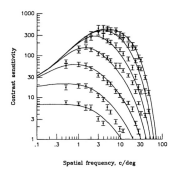
Given the autocorrelation function, and the linear model,  $R_j = \sum_i w_{ji} L_i = L_j - \sum_{i \neq j} H_{ji} L_i$ , they were able to show that the receptive field weights that minimized  $E(R_j^2)$  predicted a "center-surround" receptive field:



They also showed that one would expect the inhibitory side lobes to get smaller at low light levels. Compare with the CSF functions for various light levels below.



The results of Srinivasan et al. were a "proof of concept". Ten years later, Atick & Redlich (1992) showed how the exact shape of the human CSF as a function of mean light level could be accounted for in terms of efficient coding given the statistics of natural images (see below).



The left figure shows contrast thresholds for various light levels (from van Nes, & Bouman, M. A. (1967). Spatio modulation transfer in the human eye. J Opt Soc Am, 57(3), 401-406). The right figure is a replot of the left figure from: Atick, J. J., & Redlich, A. N. (1992). What does the retina know about natural scenes? Neural Computation, 4(2), 196-210. The solid lines show fits by Atick & Redlich based on an efficient coding model.

There has been considerable work since the above early studies. For example, see the review by Simoncelli and Olshausen, and a more recent review by: Fairhall, A., Shea-Brown, E., & Barreiro, A. (2012). Information theoretic approaches to understanding circuit function. Current Opinion in Neurobiology, 22(4), 653–659. http://doi.org/10.1016/j.conb.2012.06.005

### Principal components analysis

In the next two sections we will look at ways of encoding natural images that take advantage of statistical regularities. First describe a classical technique in statistics called principal components analysis (PCA). Then in the following section, describe how the notion of sparse coding provides an elegant explanation for how the visual cortex exploits redundancy in natural image patterns.

#### Introduction to PCA

Principal components analysis (PCA) is a statistical technique that is applied to an ensemble of ndimensional measurements (vectors or in our case images). To do PCA, all one needs is the autocovariance matrix and a good PCA algorithm. Good because images are big enough (p=mxn), and the covariance is much bigger (p^2).

PCA finds a matrix that transforms the input vectors into output vectors, such that output elements are *no longer correlated* with each other. There is more than one matrix that will do this however, and PCA find the matrix which is a rigid *rotation* of the original coordinate axes, so it preserves orthogonality. (The Fourier transform is also a rotation.) Further, the new coordinates can be ordered in terms of how much variance is captured when the data is projected on to each coordinate. The new coordinates turn out to be eigenvectors of the covariance matrix. The directions or eigenvectors with the biggest variances are called the principal components. So the dominant principal component has the most variance, and so forth. For data that are highly redundant, PCA can be used to eliminate dimensions that account for little of the total variance.

PCA is important in computational models of visual processing (See Wandell, pages 254-258). For example, PCA has been used to account for and model:

opponent color processing visual cortical cell development efficient representation of human faces face recognition given variability over illumination internal model of objects for visual control of grasping

There is a literature on theoretical neural networks and PCA. An introduction to some of the ideas is given in the optional section below. High-level languages and standard computer statistical packages provide the tools for doing PCA on large data sets. Below we try to provide intuition and background into the computation of principal components.

#### Statistical model of a two-variable input ensemble

Consider a two variable system whose inputs are correlated--e.g. an ensemble of 2 pixel images. The random variable, **rv**, is a 2D vector. To be concrete, we'll give the scatter plot for this vector an average

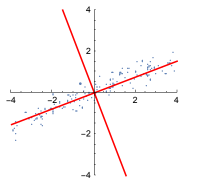
slope of Tan[theta=Pi/8] = 0.41. The variances along the axes are  $4^2$  and  $.25^2$  (16 and .0625). **gprincipalaxes** is a graph of the principal axes which we will use for later comparison with simulations.

```
ndist = NormalDistribution[0,1];theta = Pi/8;
bigvar = 4.0; smallvar = 0.25;
alpha = N[Cos[theta]]; beta = N[Sin[theta]];
rv :=
{bigvar x1 alpha + smallvar y1 beta,
bigvar x1 beta - smallvar y1 alpha} /.{x1-> Random[ndist],y1-> Random[ndist]};
gprincipalaxes = Plot[{x beta, x (-1/beta)}, {x,-4,4},
PlotRange->{{-4,4},{-4,4}},
PlotStyle->{RGBColor[1,0,0]},
AspectRatio->1];
```

0.25

**x1** and **y1** are correlated. Let's view a scatterplot of samples from these two correlated Gaussian random variables.

```
npoints = 200;
rvsamples = Table[rv,{n,1,npoints}];
g1 = ListPlot[rvsamples,PlotRange->{{-4,4},{-4,4}},
AspectRatio->1];
Show[g1,gprincipalaxes,ImageSize→Small]
```



#### Standard Principal Components Analysis (PCA)

Let E[•] stand for the expected or average of a random variable, •. The covariance matrix of a of vector random variable, **x**, is:

E[  $[\mathbf{x}-E[\mathbf{x}]][\mathbf{x}-E[\mathbf{x}]]^T$ ]. Let's compute the autocovariance matrix for **rv**. The calculations are simpler because the average value of **rv** is zero. As we would expect, the matrix is symmetric:

```
autolist = Table[
    Outer[Times,rvsamples[[i]],rvsamples[[i]]],
        {i,Length[rvsamples]}];
MatrixForm[auto=
        Sum[autolist[[i]],
        {i,Length[autolist]}]/Length[autolist]]
Clear[autolist];
( 13.8811 5.77958
    5.77958 2.48708 )
```

The variances of the two inputs (the diagonal elements) are due to the projections onto the horizontal and vertical axis of the generating random variable.

Now we will calculate the eigenvectors of the autocovariance matrix

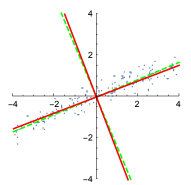
#### MatrixForm[eigauto = Eigenvectors[auto]]

 $\left(\begin{array}{ccc} -0.922497 & -0.386004 \\ 0.386004 & -0.922497 \end{array}\right)$ 

The rows of a symmetric matrix are orthogonal. Verify that.

Let's graph the principal axes corresponding to the eigenvectors of the autocovariance matrix together with the scatterplot we plotted earlier.

```
Show[g1, gPCA, gprincipalaxes, ImageSize \rightarrow Small]
```



The eigenvalues give the ratio of the variances of the projections of the random variables **rv[[1]]**, and **rv[[2]]** along the principal axes:

eigvalues = Eigenvalues[auto]

```
\{16.2995, 0.0687114\}
```

The projections along the principal axes are now **decorrelated**. We can see this by calculating the autocovariance matrix of the projected values:

These values are close to the true population variances of the generative model.

Note that the off-diagonal elements (the terms that measure the covariation of the transformed random variables) are zero. Further, because the variance of one of the projections is near zero, one can in fact dispense with this component and achieve a good approximate coding of the data with just one coordinate.

#### PCA and natural images

#### Break a large image into a series of subimages.

The idea is that each subimage will be used as a statistical sample. We compute the outer product of each, and then average all 16 to get an estimate of the autocovariance matrix.

```
subfacelist [i + 16 * (j - 1)] = N[Flatten[subface[[i, j]]]], {i, 1, 16}, {j, 1, 16}];
```

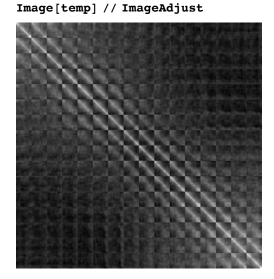
Subtract off the mean.

```
subfacelist2 = Table[subfacelist[[i]] - Mean[subfacelist[[i]]], {i, 1, 256}];
```

#### Calculate the autocovariance matrix

```
{Dimensions[subfacelist2],
Dimensions[Outer[Times, subfacelist2[[1]], subfacelist2[[1]]]]}
{{256, 256}, {256, 256}}
```

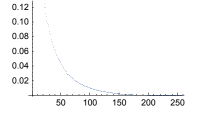
```
temp = Table[0.0, {256}, {256}];
For[i = 1, i < Dimensions[subfacelist][[1]], i++,
   temp = N[Outer[Times, subfacelist2[[i]], subfacelist2[[i]]]] + temp;
];</pre>
```



Calculate the eigenvectors and eigenvalues of the autocovariance matrix

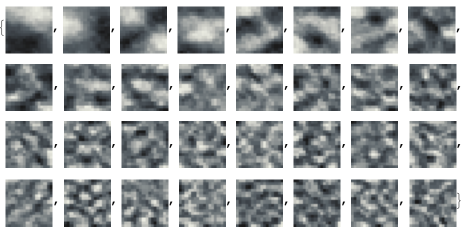
Calculate the eigenvectors and eigenvalues of the autocovariance matrix

```
eigentemp = Eigenvectors[temp];
eigenvaluestemp = Eigenvalues[temp];
ListPlot[Chop[eigenvaluestemp]]
```



#### Display the first 32 eigenvectors as "eigenpictures"

#### Table[ArrayPlot[Partition[eigentemp[[i]], 16], Mesh → False, ImageSize → Tiny, PixelConstrained → {3, 3}], {i, 1, 32}]



 How do point-wise non-linearities affect the principal components? Try replacing alpine with squashedalpine, and repeate the above analysis.

```
\begin{aligned} & \text{squash}[\texttt{x}_{,}, \mu_{,}, \gamma_{-}] := \mathbb{N}\Big[\frac{1}{1 + e^{-\gamma \ (\texttt{x}-\mu)}}\Big];\\ & \text{gain} = 0.045; \ \mu 0 = \text{Mean}[\texttt{Flatten}[\texttt{alpine}]]; \ \gamma = .5;\\ & \text{squashedalpine} = \text{squash}\Big[\big(\texttt{alpine} - \mu 0\big) + \mu 0, \ \mu 0, \ \gamma\Big]; \end{aligned}
```

## Efficient, sparse coding in VI

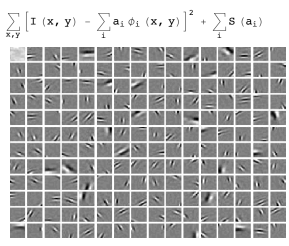
#### Olshausen & Field: Primary cortex

Fourier coefficients for natural images tend to be uncorrelated and indeed there is a close relationship between Fourier rotations and Principal Components Analysis (or Karhunen- Loeve transformations) (e.g. Appendix A in Andrews, 1983). There is also a long history to the study of the relationship between self-organizing models of visual cortex, as well as efficient coding of image information. For early work on this, see: Yuille et al., 1989; Linsker, R. (1990) and Barlow, H. B., & Foldiak, P. (1989). Linsker's computational studies show, for example, that orientation tuning, and band-pass properties of simple cells can emerge as a consequence of maximum information transfer (in terms of variance) given the constraint that the inputs are already band-pass, and the receptive field connectivity is a priori limited.

In a highly influential study in 1996, Olshausen and Field provided an elegant explanation for the spatial filtering properties of V1 simple cells. They showed that one could derive a set of basis functions that have the same characteristics as the ensemble of visual simple cells in primary visual cortex by requiring two simple constraints:

1) One should be able to express the image I(x,y) as a weighted sum of the basis functions,  $\{\phi_i\}$ 

2) The total activity across the ensemble should, on average, be small. This latter constraint is called "sparse coding". That is, a typical input image should activate a relatively small fraction of neurons in the ensemble. S() for example could be the absolute value of the activity  $a_i$ . Using a database of natural patches {I(x,y)}, they estimated the values of { $\phi_i$ } that minimized the following cost function:



We will see later that cells in the visual cortex send their visual information to an incredibly complex, and yet structured collection of extra-striate areas. Any hypothesized function of striate cortex must eventually take into account what the information is to be used for. In the next lecture, we will give a quick overview of extra-striate visual cortex, and introduce the computational problem of estimating scene properties from image data.

#### Side note: adaptation & learning

Human orientation and spatial frequency selectivity changes with adaptation. Adaptation has been interpreted as an optimal change to new conditions in the input image statistics. (e.g. see, Wainwright, M. J. (1999). Visual adaptation as optimal information transmission. Vision Research, 39, 3960--3974.)

Barlow argued that a decorrelated representation of sensory information is important for efficient learning (Barlow, 1990).

#### Side note: PCA and ICA

PCA decorrelates the projected values between dimensions. A stronger condition would be to require that the projections be independent. This requirement is met by independent component analysis (ICA). There is a close relationship between ICA and sparseness (see: Hyvärinen, A. (2010). Statistical Models of Natural Images and Cortical Visual Representation. Topics in Cognitive Science, 2(2), 251–264. http://doi.org/10.1111/j.1756-8765.2009.01057.x)

# Using synthesis: How good is a 2nd order model of natural images?

As we will see later, building a generative model for an ensemble of natural images (e.g. for a particular texture class or a "generic" class) can provide insights into how well the model is capturing the regularities of interest. We will see more of this later when we study texture perception.

Let's construct a 2nd order generic generative statistical model of images and see what the samples look like. We will use our fourier tools.

#### **Random Fractals**

Random fractals are a crude but good statistical models for the amplitude spectra certain classes of natural images. Random fractals can be characterized by the fractal dimension D (3<D<4) and amplitude spectrum,  $1/(f_x^2 + f_y^2)^{\Lambda}(4-D)$ . The amplitude spectrum is a straight line when plotted against frequency in log-log coordinates. The condition If[] is used to include a fudge term  $(1/(2)^{\Lambda}(q))$  to prevent blow up near zero in the Block[] routine later.

Here is a function to make a low-pass filter with fractal dimension D. (D, here should be between 3 and 4). Note that we first make the filter centered in the middle, and then adjust it so that it is symmetric with respect to the four corners.

ArrayPlot[RotateLeft[fractalfilter[3.5], {hfwidth, hfwidth}], Mesh → False]



Here is the amplitude spectrum plot for a random fractal image:

```
randomspectrum = Abs[temp = Fourier[Table[Random[], {size}, {size}]]];
randomphase = Arg[temp];
```

```
ffilt = fractalfilter[3.5] randomspectrum;
ArrayPlot[RotateRight[ffilt, {hfwidth, hfwidth}], Mesh -> False, Frame -> False]
```



#### Here is a random fractal image, with D = 3.2

ArrayPlot[Chop[
InverseFourier[
fractalfilter[3.2] randomspectrum Exp[I randomphase]]]]



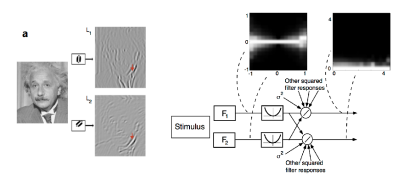
So what regularites are being captured and what ones are missing? The above gaussian fractal is lowpass, which of course we built in. This is good. But natural images tend to have edges, and somewhat sharp patches over a range of scales. Can one do better? Yes. See the sample below from the paper by: Zhu, S. C., & Mumford, D. (1997). Prior Learning and Gibbs Reaction-Diffusion. IEEE Trans. on PAMI, 19(11), 1236-1250.



## Higher order redundancies & contrast normalization

#### Contrast normalization

There are higher order redundancies in natural images, and a major challenge is to characterize them, and understand how the visual system exploits these redundancies. For example, the figure below shows that the output response of one spatial filter (receptive field (RF) responses) influences the variability in a second spatial filter. Odelia Schwarzt and Eero Simoncelli have shown how a non-linearity called "contrast normalization" serves to remove this redundancy; explaining a number of non-linearities observed in Vq cortical neurons.



See Figure 8 in Simoncelli, E. P., & Olshausen, B. A. (2001). Natural image statistics and neural representation. Annu Rev Neurosci, 24, 1193-1216.(pdf).

See also Figure 5 in Geisler, W. S. (2008). Visual perception and the statistical properties of natural scenes. Annu Rev Psychol, 59, 167-192. (pdf).

## Next time

#### Edge detection

## **Appendices**

#### Neural networks and principal components

See: Pehlevan, C., Hu, T., & Chklovskii, D. B. (2015). A Hebbian/Anti-Hebbian Neural Network for Linear Subspace Learning: A Derivation from Multidimensional Scaling of Streaming Data. Neural Computation, 27(7), 1461–1495. http://doi.org/10.1162/NECO\_a\_00745

## Neural network model using Hebb together with Oja's rule for extracting the dominant principal component

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