Recap

So far this class has looked at methods for computing spectra and assigning uncertainties, and we’ve identified the the lowest resolved frequency and the Nyquist frequency (the highest frequency for which we have any information.) We’ve almost managed to produce useful spectra, but we still need to be able to assign error bars and figure out how many degrees of freedom we have.

Spectral Uncertainties

We left off last time talking about how to attach error bars to spectra. We defined our spectral estimate:

$$\hat{E}(f) = \langle |a(f)|^2 \rangle = \frac{1}{n} \sum_{i=1}^{n} |a(f)|^2,$$

where $a$ is used for the complex Fourier coefficients, and $f$ is frequency. With $n$ segments to average, this gives us $\nu = 2n$ degrees of freedom.

Then we defined the ratio of the estimated spectrum to the true spectrum, which we multiplied by $\nu$:

$$y = \nu \frac{\hat{E}_\nu(f)}{E(f)}$$

So even though we don’t know $E$, we know that the ratio of $\nu \hat{E}$ to $E (\nu \hat{E}/E)$ is a $\chi^2$ quantity. The expectation value of this ratio is $\nu$, and the standard deviation is $\langle y^2 \rangle - \langle y \rangle^2 = 2\nu$ (see Bendat and Piersol, section 4.2.2). From that we can infer that with two degrees of freedom, the standard deviation $\langle \hat{E}^2 \rangle - \langle \hat{E} \rangle^2 = 2/\nu E_0^2 = E_0^2$, which means that the standard deviation is equal to the original value. Clearly we need more samples.

Now in reality, we don’t care about the details of $\chi^2$ but rather the probability that our estimate of the spectrum $\hat{E}$ is close to or far from the unknown true spectrum $E$, so we’re going to look at the probability that $\nu \hat{E}(f)/E(f)$ falls within a fixed range. So we want to use the $\chi^2$ distribution as a probability distribution and ask about the probability that $\hat{E}(f)/E(f)$ should be within \pm some range of 1. Formally, the probability that the estimated spectrum should be close in value to the true spectrum is:

$$P \left( \frac{\chi^2_{\nu,1-\alpha/2}}{\nu < \frac{\hat{E}(f)}{E(f)} < \frac{\chi^2_{\nu,\alpha/2}}{\nu} } \right) = 1 - \alpha$$

so if we want to find a 95% significance level, we set $\alpha$ to 0.05. We’ll want to know the value of $\chi^2$ that corresponds to a given point on the cdf, so for this we use the inverse $\chi^2$ function (“chi2inv” in Matlab). So we can invert this relationship to provide the probability that the true value is within a particular range of the observed value:

$$P \left( \frac{\chi^2_{\nu,1-\alpha/2}}{\nu} < \frac{\hat{E}(f)}{E(f)} < \frac{\chi^2_{\nu,\alpha/2}}{\nu} \right) = 1 - \alpha$$

$$P \left( \frac{\nu}{\chi^2_{\nu,1-\alpha/2}} > \frac{E(f)}{E(f)} > \frac{\nu}{\chi^2_{\nu,\alpha/2}} \right) = 1 - \alpha$$
Whether you use $\chi^2$ or its reciprocal, the ratio between the high and low error bars should be the same, and the ratio will be what matters.

This error formulation differs from the usual error bars that we’re used to seeing where we say for example that the true temperature should be the measured temperature plus or minus an uncertainty: $T = \hat{T} \pm \delta T$. We can develop a similar expression for the true spectrum: $E(f)$ is in the range between $\nu \hat{E}(f)/\chi^2_{\nu,1/2}$ and $\nu \hat{E}(f)/\chi^2_{\nu,1/2}$, where $\nu$ is twice the number of segments. This expression isn’t very easy to interpret, since it varies as a function of frequency, and the estimated value $\hat{E}(f)$ is not at the mid-point of the range.

Instead we’ll keep in mind that we’ve computed the uncertainty for the ratio $\hat{E}(f)/E(f)$, and the probabilities for this ratio does not depend on frequency. On a log plot, error bars defined by the range between $\nu/\chi^2_{\nu,1/2}$ and $\nu/\chi^2_{\nu,1/2}$ are the same size at all frequencies, so we can easily compare spectral peaks at different frequencies.

Some statistics books include look-up tables for $\chi^2$, but we can compute it directly in Matlab. For $N$ data segments and $\nu = 2 \times N$ degrees of freedom, the error limits are:

```matlab
err_high = nu/chi2inv(.05/2,nu);
err_low = nu/chi2inv(1-.05/2,nu);
```

We can plot these values as:

```matlab
semilogy([f f],[err_low err_high]*A);
```

where we set the frequency $f$ and the amplitude $A$, so that the error bar ends up positioned in a convenient spot on the plot.

Now to have $M$ data segments (and $\nu = 2M$ degrees of freedom), we have to split our long data record into shorter segments. We can do this by taking $N$ data points at a time:

```matlab
Ntotal=length(data);
N=segment_length; % define this value
for n=1:floor(Ntotal/N)
    d=data((n-1)*N+1:n*N); %select data for the nth segment
    fd(:,n)=fft(d); % compute fft
end
sd=sum(abs(fd(1:N/2+1,:)).^2,2)/N; % sum over all spectra
    % (sum over 2nd index)
sd(2:end)=sd(2:end)*2;

nu=2*floor(Ntotal/N);
err_high = nu/chi2inv(.05/2,nu);
err_low = nu/chi2inv(1-.05/2,nu);
semilogy(0:N/2,sd,[N/4 N/4],[err_low err_high]*sd(N/4))
```
Getting the units right

We’ve talked about units for spectra, but let’s lay everything out in one place. Our fundamental principle is that we want Parseval’s theorem to work. But this gets a tiny bit messy when we average multiple frequencies. Still the basic rule of Parseval’s theorem is not that the sum of the squares equals the sum of the squared Fourier coefficients, but rather that the integrated variance equals the integral under the spectrum.

Some instructions say to divide the spectral estimate by \( N^2 \) and then divide by \( df = 1/T \), in effect multiplying by \( T/N^2 \). But my sample code so far has only had a division by \( N \). And some of you complained in problem set 4 about the division by \( N \) rather than \( \sqrt{N} \). Why?

In the discrete Fourier transform, we have:

\[
\sum_{n=0}^{N-1} x_n^2 = \frac{1}{N} \sum_{k=0}^{N-1} |X_k|^2,
\]

or after spectral normalizations, maybe we write

\[
\sum_{n=0}^{N-1} x_n^2 = \frac{1}{N} \left[ |X_0|^2 + \sum_{k=1}^{(N/2)-1} 2|X_k|^2 + |X_{N/2}|^2 \right] \approx \frac{1}{N} \sum_{k=0}^{(N-1)/2} 2|X_k|^2.
\]

For this discussion, to keep the equations compact, we’ll use the approximation in the final line, neglecting the fact that the mean and the \( k = N/2 \) value should not be doubled. Many times we work with this form, since it gives us meaningful spectral slopes.

If I take the limiting case in which \( x_n \) has one frequency, then I can easily determine how to normalize \( X_0 \). Suppose \( x_n \) is a constant, so \( x_n = \bar{x} \). Then

\[
\sum_{n=0}^{N-1} x_n^2 = N \bar{x}^2 = \frac{1}{N} \sum_{k=0}^{N-1} |X_k|^2 = \frac{1}{N} |X_0|^2,
\]

since only frequency 0 has a non-zero Fourier coefficient. This implies that:

\[
\bar{x} = \frac{|X_0|}{N}.
\]

Similarly, if \( x = a \cos(\omega t) \), then

\[
\sum_{n=0}^{N-1} x_n^2 = \sum_{n=0}^{N-1} a^2 \cos^2(\omega t) = \frac{N a^2}{2} = \frac{1}{N} \sum_{k=0}^{N-1} |X_k|^2 = \frac{2}{N} |X_k|^2,
\]

which tells us how to determine the amplitude \( a \):

\[
a = \frac{2|X_0|}{N}.
\]

In continuous form, we had a form more like this:

\[
\int_{-\infty}^{\infty} x^2(t) dt = \int_{-\infty}^{\infty} |X(f)|^2 df
\]
where $f$ is frequency in cycles per unit time (or we might use $\sigma = 2\pi f$ sometimes, where $\sigma$ is frequency in radians per unit time.) If we want this integral form to work for our real data, then we have to be a bit careful with our normalizations. We’re going to want the area under the curve in our spectrum to be equal to the total variance integrated over time. So if total integrated variance is

$$\text{variance} = \sum_{n=0}^{N-1} x_n^2 \Delta t$$  \hspace{1cm} (16)

where $\Delta t = T/N$. Then the integrated spectrum should be

$$\text{variance} = \frac{\alpha}{N} \sum_{k=0}^{N/2-1} 2|X_k|^2 \Delta f,$$  \hspace{1cm} (17)

where $\Delta f = 1/(N\Delta t) = 1/T$, and we’ll need to figure out $\alpha$ to ensure that the spectral estimator that we compute still properly adheres to Parseval’s theorem. This implies that we might imagine normalizing our spectra to have:

$$\sum_{n=0}^{N-1} x_n^2 \Delta t = \frac{\alpha}{N} \sum_{k=0}^{N/2-1} 2|X_k|^2 \Delta f = \frac{\Delta t}{N\Delta f} \sum_{k=0}^{N/2-1} 2|X_k|^2 \Delta f = \frac{T^2}{N^2} \sum_{k=0}^{N/2-1} 2|X_k|^2 \Delta f$$  \hspace{1cm} (18)

Or maybe this makes for units that aren’t easily compared, so we could normalize our spectra to represent the average energy per unit time in the time domain, and adjust the frequency domain accordingly:

$$\frac{1}{T} \sum_{n=0}^{N-1} x_n^2 \Delta t = \frac{1}{N\Delta t} \sum_{n=0}^{N-1} x_n^2 \Delta t$$  \hspace{1cm} (19)

$$= \frac{T}{N^2} \sum_{k=0}^{N/2-1} 2|X_k|^2 \Delta f$$  \hspace{1cm} (20)

$$= \frac{(\Delta t)}{N} \sum_{k=0}^{N/2-1} 2|X_k|^2 \Delta f$$  \hspace{1cm} (21)

$$= \frac{1}{N^2 \Delta f} \sum_{k=0}^{N/2-1} 2|X_k|^2 \Delta f$$  \hspace{1cm} (22)

$$= \frac{1}{N^2 f_{\text{Nyquist}}} \sum_{k=0}^{N/2-1} 2|X_k|^2 \Delta f$$  \hspace{1cm} (23)

so we could divide our spectrum by twice the Nyquist frequency to have energy in units appropriate for comparing if we wanted to have our integrals match.

This isn’t always the way we think about this, but it serves as our reminder that we should think about the units of our spectrum. What we know is that integral of our spectrum over a certain frequency range should give a measure of the signal variance:

$$\text{variance in a band} = \int_{f-\Delta f/2}^{f+\Delta f/2} |X(f)|^2 \, df$$  \hspace{1cm} (24)
So if we expand this out, this implies that the units of $|X(f)|^2$ should be equivalent to variance divided by frequency, so it’s our reminder that we’ll label the y-axis units as the squared units of $x$ divided by frequency, with a normalization to account for the units of time in our data.

**Sidebar: Some further convolution examples**

Let’s return to convolution for one more example. We’ve pointed out that convolution in the time domain is equivalent to multiplication in the frequency domain.

What if I compute the convolution of a boxcar filter with itself?

$$y(\tau) = \int_{-\infty}^{\infty} h(t)h(\tau-t)dt \quad \text{... definition of convolution} \tag{25}$$

$$= \int_{0}^{1} h(\tau-t) dt \quad \text{for } 0 < \tau - t < 1 \quad \text{... adjust integration limits to represent } h(t) \tag{26}$$

$$= \int_{0}^{1} h(\tau-t) dt \quad \text{for } -\tau < -t < 1 - \tau \quad \text{... subtract } \tau \tag{27}$$

$$= \int_{0}^{1} dt \quad \text{for } -\tau < t < \tau \quad \text{... reverse sign from } -t \text{ to } +t \tag{28}$$

$$= \begin{cases} \int_{0}^{\tau} dt = \tau, & \text{for } 0 < \tau \leq 1 \\ \int_{1}^{\tau-1} dt = 1 - (\tau - 1) = 2 - \tau & \text{for } 1 < \tau \leq 2 \end{cases} \tag{29}$$

The final step requires some examination of the limits of integration. We require that $t$ be between 0 and 1, but $\tau$ can have a broader range. We can impose limits on $t$ or on $\tau$, somewhat separately. First, we worry only about the upper limit. If $t < \tau$, and $0 < t \leq 1$, then we can consider a case when $0 < \tau \leq 1$, while imposing the lower limit of the integral. This gives us limits of integration from 0 to $\tau$ for $0 < \tau \leq 1$. If we consider only the lower limit, so that $\tau - 1 < t$ with $0 < t \leq 1$ then $\tau$ can be between 1 and 2, with integration limits from $\tau - 1$ to 1. In any case, the results produce a triangle.

This is definitely easier to consider graphically, and it will show that boxcar filter convolved with itself becomes a triangle filter.

What if we repeat this again and again? It’s actually analogous to what we considered for the central limit theorem. Applying a filter to a filter to a filter will eventually give us a Gaussian.

**The Fourier Transform of a Boxcar**

When we deal with data records of finite length, we’re always taking finite sized segments of data, sort of like Fourier transforming a boxcar filter. Usually we think of a Fourier transform of this form:

$$X(f) = \int_{-\infty}^{\infty} x(t)e^{-i2\pi ft} dt. \tag{30}$$

If $x(t)$ is a boxcar filter, this becomes:

$$\int_{-\infty}^{\infty} h(t)e^{-i2\pi ft} dt = \int_{-1/2}^{1/2} e^{-i2\pi ft} dt = e^{-i2\pi ft}\frac{1}{i2\pi f}\left|_{-1/2}^{1/2}\right. = \frac{e^{i2\pi f/2} - e^{-i2\pi f/2}}{i2\pi f} = \frac{\sin(\pi f)}{\pi f} \tag{31}$$

This is the sinc function, which can be written as:

$$\text{sinc}(x) = \frac{\sin(x)}{x} \tag{32}$$
or (for digital signal processing)
\[
sinc(x) = \frac{\sin(\pi x)}{\pi x}.
\]
(33)

We’ll try to use the second definition, the “normalized sinc function”, since we’re trying to use a frequency \( f \) that needs to be multiplied by \( 2\pi \).

Remember that convolution in the time domain corresponds to multiplication in the frequency domain. In this case, we’re doing the opposite. We multiplied our data by a boxcar window in the time domain, and that’s equivalent to convolving with a sinc function in the frequency domain:
\[
\hat{X}(f) = \int_{-\infty}^{\infty} X(g) W(f - g) \, dg
\]
(34)

where
\[
W(f) = \frac{\sin(2\pi f T)}{2\pi f T} = \text{sinc} \left( \frac{2fT}{2} \right)
\]
(35)

for a record length of \( 2T \).

Correspondingly, the spectrum is convolved with \( W(f)^2 = \text{sinc}(2fT)^2 \). This has a central peak of width \( \Delta f = \pm 1/(2T) \). The window will have a maximum value
\[
|W(f)|^2 < (2\pi f T)^{-2}
\]
(36)

The side lobes of the window are definitely problematic. And it turns out that we’re not stuck with them. If we can widen the central peak of \( W \) (the Fourier transform of our window), we can lower the impact of the side lobes. To do this, we’ll want to forego the rectangular window in the time domain and choose something that lets us attenuate the beginnings and ends of each segment of our data. What if we chose a triangle window? That will already give us fewer side lobes.

But we can keep going to find a window that looks more like an exponential. Leading possibilities:

1. Cosine taper:
\[
w(t) = \cos^\alpha \left( \frac{\pi t}{2T} \right)
\]
(37)

with \( \alpha = [1, 4] \).

2. Hanning window or “raised cosine” window (developed by von Hann):
\[
w(t) = \cos^2 \left( \frac{\pi t}{2T} \right) = \frac{1 + \cos(\pi t/T)}{2} = 0.5 + 0.5 \cos(\pi t/T)
\]
(38)

3. Hamming window. This variant of the Hanning window was developed by Hamming.
\[
w(t) = 0.54 + 0.46 \cos(\pi t/T)
\]
(39)

The Hamming window has less energy in the first side lobe but more in the distant side lobes.

Some other options include a Blackman-Harris window or a Kasier-Bessel window, and Harris (1978, Use of windows for harmonic analysis, Proc. IEEE) provides detailed discussion of options.

So how do you use a window?
1. First you must demean your data—otherwise, the window will shift energy from the mean into other frequencies. If you’re working in segments, you should demean (and detrend) each segment before you do anything further.

2. Second, for a segment with $N$ points, multiply by a window that is $N$ points wide.

3. Since the window attentuates the impact of the edge of each segment, you can use segments that overlap (typically by 50%). This will give you (almost) twice as many segments, so instead of $\nu$ degrees some larger number.

4. Now Fourier transform, scale appropriately (e.g. by $\sqrt{8/3}$ for a Hanning window, to account for energy attenuation) and compute amplitudes.

Will the window preserve energy in your system? Not necessarily. You can normalize it appropriately, but windowing can shift the background energy level of your spectrum relative to the spectral peaks, and you’ll want to keep track of this.