

Large-scale power loss in ground-based CMB mapmaking

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ABSTRACT

CMB mapmaking relies on a data model to solve for the sky map, and this process is vulnerable to bias if the data model cannot capture the full behavior of the signal. I demonstrate that this bias is not just limited to small-scale effects in high-contrast regions of the sky, but can manifest as $\mathcal{O}(1)$ power loss on large scales in the map under conditions and assumptions realistic for ground-based CMB telescopes. This bias is invisible to simulation-based tests that do not explicitly model them, making it easy to miss. I investigate the common case of sub-pixel errors in more detail and demonstrate that this special case of model error can be eliminated using bilinear pointing matrices. Finally, I provide simple methods for testing for the presence of large-scale model error bias in the general case.

1. INTRODUCTION

CMB telescopes observe the sky by scanning their detectors across it while continuously reading off a series of samples from the detectors. Typically the signal-to-noise ratio of each sample is small, but by combining a large number of samples with knowledge of which direction the telescope was pointing at any time, it's possible to reconstruct an image of the sky. There are several ways of doing this, with the most common being maximum-likelihood, filter+bin and destriping. These all start by modelling the telescope data as (Tegmark 1997)

$$d = Pm + n \quad (1)$$

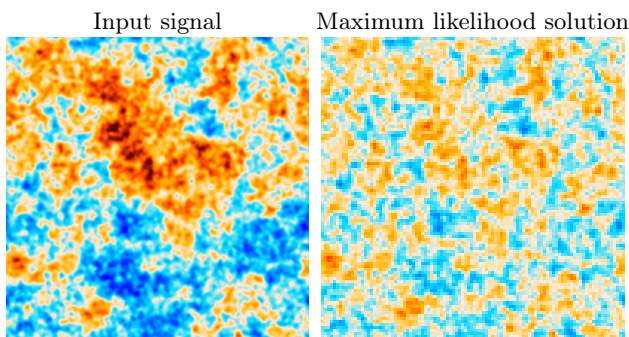


Figure 1. Preview of the model error bias we will discuss in the following sections. Despite the standard expectations that maximum-likelihood mapmaking is optimal and unbiased, the maximum-likelihood solution (**right**) for a simple toy example is strongly power-deficient on large scales compared to the input signal (**left**). As we shall see this bias is not unique to maximum-likelihood methods, and can be triggered by several subtle types of model error.

where d is the set of samples read off from the detectors (often called the time-ordered data), m is the set of pixels of the sky image we want to reconstruct, n is the noise in each sample (usually with significant correlations), and P is a pointing matrix that encodes how each sample responds to the pixels in the image.

Given this data model, it's possible to either directly solve for an unbiased map (as in maximum likelihood mapmaking or destriping), or to measure and correct for the bias in a biased estimator (as in filter+bin mapmaking). However, this fails when the model does not actually describe the data, and this turns out to be the norm rather than the exception. Previously this bias has been thought of as mainly a small-scale effect relevant only in regions of the sky with very high contrast, leading to artifacts like thin stripes or bowling around bright sources (Piazzo 2017; Naess 2019), or as a tiny correction on intermediate scales (Poutanen et al. (2006) saw a bias of 0.6% peaking at $\ell = 800$ for simulations of the Planck space telescope). The goal of this paper is to demonstrate the unintuitive and surprising result that that these effects can lead to $\mathcal{O}(1)$ bias on large scales everywhere in the map. The full scope of these errors appears to be largely unappreciated, and I fear that many CMB analyses so far suffer from uncorrected bias at low multipoles in total intensity due to these effects. As we shall see, it's mainly ground-based telescopes that are vulnerable to this bias, which usually manifests as a power deficit at large scales, as illustrated in figure 1.

2. SUBPIXEL ERRORS

Subpixel errors may be both the most common and most important class of model errors in CMB mapmak-

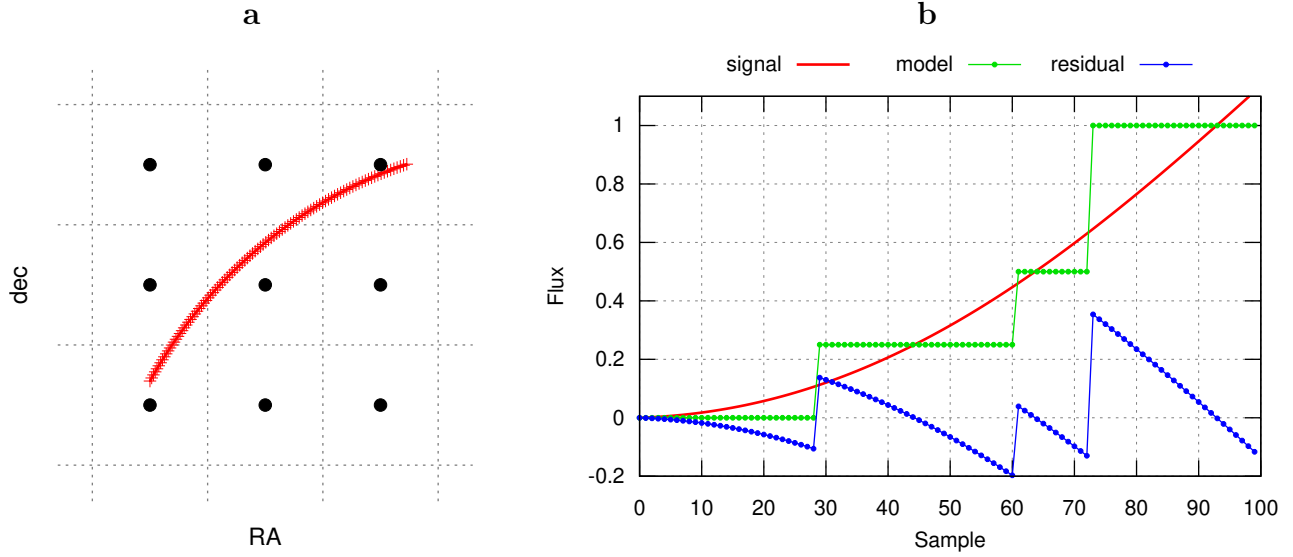


Figure 2. **a:** Example path (red) of a detector across a few pixels. The area closest to each pixel center (black dots) is shown with dotted lines. In the nearest neighbor model, the value associated with each sample is simply that of the closest pixel, regardless of where inside that pixel it is. **b:** Example detector signal (red) for the same path. The closest matching model (green) leaves a jagged residual (blue) that has power on all length scales despite the signal itself being very smooth. For comparison, if our model were a constant zero, then the residual would just be the signal itself (red), and hence smooth. **If smooth residuals are much cheaper in the likelihood than jagged ones, then a zero model will be preferred to one that hugs the signal as tightly as possible like the green curve.**

ing. For efficiency reasons P is always¹ chosen to use simple nearest-neighbor interpolation, where the value of a sample is simply given by the value of the pixel nearest to it. This means that P can be implemented by simply reading off one pixel value per sample, and its transpose P^T consists of simply summing the value of the samples that hit each pixel. However, this comes at the cost of there being a discontinuous jump in values as one scans from one pixel to the next, as illustrated in figure 2. Hence, the closest the model can get to a smooth curve is a staircase-like function that hugs it, leaving a residual full of discontinuous jumps (the blue curve in the figure).

Discontinuous residuals are not necessarily problematic. The trouble arises when this is coupled with a likelihood² where some modes have much less weight than others. Ground-based CMB telescopes suffer from atmospheric emission that acts as a large source of noise on long

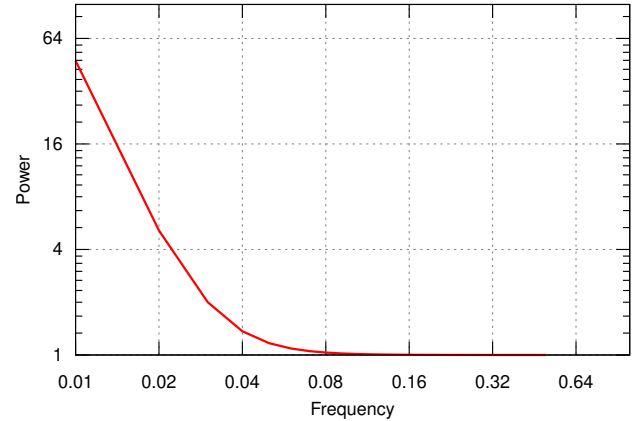


Figure 3. The noise model/inverse weights/inverse filter used in the subpixel bias demonstration in figures 4 and 5. It is a simple Fourier-diagonal $1/f$ + white noise spectrum typical for ground-based CMB observations. The frequency axis is in dimensionless units in this toy example, but for real telescopes the transition from white noise is typically a few Hz, corresponding to multipoles of hundreds on the sky. The power axis is dimensionless here, but for a real-world case could have units like $\mu\text{K}^2/\text{Hz}$.

¹ I am not aware of any published CMB analysis that has done something else. This includes destripers like MADAM (Keihänen et al. 2010) and SRoll (Planck Collaboration 2020), filter+bin map-makers like those used in SPT (Schaffer et al. 2011; Dutcher et al. 2021) and BICEP (BICEP2 Collaboration 2014) or the maximum likelihood map-makers used in ACT (Aiola et al. 2020) and QUIET (Ruud et al. 2015).

² The equivalent for filter+bin is a filter that impacts some modes more than others (which is the whole point of a filter), and for destriping it's the baseline length and any amplitude priors.

timescales. This leads to time-ordered data noise power spectra similar to the one sketched in figure 3, with a white noise floor at short timescales (high frequency) transitioning to a steep rise of several orders of magnitude as one moves to longer timescales (low frequency). In this

case long-wavelength modes have orders of magnitude lower weight in the likelihood than short-wavelength modes. Put another way, they are orders of magnitude *cheaper to sacrifice* when the model can't fully fit the data.

2.1. 1D toy example

To illustrate the interaction between a nearest neighbor pointing matrix's subpixel errors and a likelihood where large scales have low weight, let us consider a simple 1D case where 100 samples scan uniformly across 10 pixels:

```
npix = 10
nsamp = 100
pix = np.arange(nsamp).astype(float)*npix/
      nsamp
```

A standard nearest-neighbor pointing matrix for this looks like:

```
P = np.zeros((nsamp,npix))
for i, p in enumerate(pix):
    P[i,int(np.round(pix[i]))%npix] = 1
```

We assume a typical Fourier-diagonal “1/f” noise model $N(f) = 1 + (f/f_{\text{knee}})^\alpha$ with $f_{\text{knee}} = 0.03$ and $\alpha = -3.5$, and build the inverse noise matrix/filter/baseline-prior F by projecting the inverse noise spectrum N^{-1} into pixel space:³

```
freq = np.fft.rfftfreq(nsamp)
inv_ps = 1/(1+(np.maximum(freq,freq[1]/2)/0.03)
            ** -3.5)
F = np.zeros((nsamp,nsamp))
I = np.eye(nsamp)
for i in range(nsamp):
    F[:,i] = np.fft.irfft(inv_ps*np.fft.rfft(I[i
    ]), n=nsamp)
```

The signal itself consists of just a long-wavelength sine wave:⁴

```
signal = np.sin(2*np.pi*pix/npix)
```

With this in place, we can now define our map estimators.

2.1.1. Binning

The *binned map* is simply the mean value of the samples in each pixel, with no weighting:

```
map_binned = np.linalg.solve((P.T.dot(P)), P.T.
                             dot(signal))
```

This is the unweighted least-squares solution for the map.

³ In practice the “1/f” power law does not continue to infinity at zero frequency, but stabilizes at some high value at low frequency. We implement this by replacing $f = 0$ with a small, non-zero number for the purposes of computing the noise spectrum in the following.

⁴ Since all the mapmaking methods we will discuss are linear, signal and noise can be analysed independently. Since the focus in this example is bias, it's therefore enough to consider a signal-only data set, and we thus don't add any noise.

2.1.2. Maximum-likelihood

The maximum-likelihood solution of equation 1 for the sky image m is

$$\hat{m} = (P^T N^{-1} P)^{-1} P^T N^{-1} d \quad (2)$$

where N is the covariance matrix of the noise n . This is the generalized least-squares solution for the map.⁵ In our toy example, $N^{-1} = F$, so the *maximum-likelihood map* is

```
map_ml = np.linalg.solve((P.T.dot(F).dot(P)), P.
                          T.dot(F.dot(signal)))
```

2.1.3. Filter+bin

As the name suggests, filter+bin consists of filtering the time-ordered data, and then making a binned map. We'll use F as our filter, so the *filter+bin map* is

```
map_fb = np.linalg.solve(P.T.dot(P), P.T.dot(F)
                          .dot(signal))
```

The filter+bin map is biased by design, so to interpret or debias it one needs to characterize this bias. There are two common approaches to doing this: Observation matrix and simulations.

Observation matrix—The observation matrix approach recognizes that the whole chain of operations observe, filter, map together make up a linear system, and can therefore be represented as a matrix, called the *observation matrix* (BICEP2 and Keck Array Collaboration 2016). Building this matrix is heavy, but doable for some surveys. Under the standard assumption that the observation step is given by equation 1, the observation matrix is given by

```
obsmat = np.linalg.inv(P.T.dot(P)).dot(P.T.dot(
    F).dot(P))
```

and using it, we can define a debiased filter+bin map

```
map_fb_deobs = np.linalg.solve(obsmat, map_fb)
```

Simulations—Alternatively, and more commonly, one can characterize the bias by simulating the observation of a set of random skies, passing them through the filter+bin process, and comparing the properties of the input and output images (e.g. Sayre et al. 2020). The standard way of doing this is by assuming that equation 1 describes the observation process, and that the bias can be described

⁵ The matrix $P^T N^{-1} P$ is often poorly conditioned, making it slow for iterative solution methods to recover the large scales. If iteration is stopped early, this could result in a lack of power at large scales. This well-known phenomenon is *not* the bias I'm exploring in this paper, and since I solve the equation exactly it does not appear here.

by a *transfer function*: a simple independent scaling of each fourier mode. Under these assumptions, we can measure and correct the bias as follows.

```
nsim = 1000
sim_ips = np.zeros(npix//2+1)
sim_ops = np.zeros(npix//2+1)
for i in range(nsim):
    sim_imap = np.random.standard_normal(npix)
    sim_omap = np.linalg.solve(P.T.dot(P), P.T.
        dot(F).dot(P).dot(sim_imap))
    sim_ips += np.abs(np.fft.rfft(sim_imap))**2
    sim_ops += np.abs(np.fft.rfft(sim_omap))**2
tf = (sim_ops/sim_ips)**0.5
map_fb_detrans = np.fft.irfft(np.fft.rfft(
    map_fb)/tf, n=npix)
```

2.1.4. Destriping

Destriping splits the noise into a correlated and uncorrelated part, and models the correlated noise as a series of slowly changing degrees of freedom to be solved for jointly with the sky image itself (Sutton et al. 2010; Tristram et al. 2011). The data is modeled as

$$d = Pm + Qa + n_w \quad (3)$$

where n_w is the white noise with diagonal covariance matrix N_w , and Q describes how each correlated noise degree of freedom a maps onto the time-ordered data, typically in the form of seconds (ground) to minutes (space) long baselines. Given this, the maximum-likelihood solutions for a and m are

$$\begin{aligned} Z &= I - P(P^T N_w^{-1} P)^{-1} P^T N_w^{-1} \\ a &= (Q^T N_w^{-1} Z Q + C_a^{-1})^{-1} Q^T N_w^{-1} Z d \\ m &= (P^T N_w^{-1} P)^{-1} P^T N_w^{-1} (d - Qa) \end{aligned} \quad (4)$$

where C_a is one's prior knowledge of the covariance of a . Destriping allows a speed/optimality tradeoff in the choice of baseline length and prior, and approaches the maximum likelihood map when the baseline length is a single sample and $C_a + N_w = N$. We implement this limit below, but explore other choices in section 2.2. In our toy example $N_w = I$, so $C_a = F^{-1} - I$.

```
iCa = np.linalg.inv(np.linalg.inv(F) - I)
Z = I - P.dot(np.linalg.solve(P.T.dot(P), P.T))
a = np.linalg.solve(Z + iCa, Z.dot(signal))
map_ds = np.linalg.solve(P.T.dot(P), P.T.dot(
    signal - a))
```

2.1.5. Results

Figure 4 compares the recovered 1D sky “images” for the different mapmaking methods for this toy example. All methods are expected to have a small loss of power at small scale (called the “pixel window”) due to averaging the signal within each pixel, but this effect is well-known, easy to model, and not our focus here. We deconvolve it using the following function before plotting.

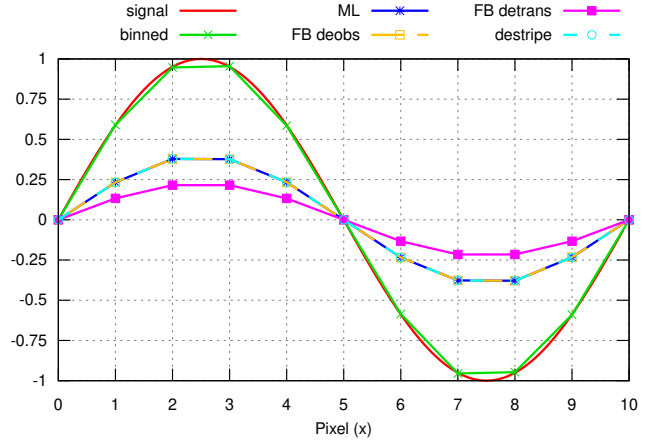


Figure 4. Demonstration of large loss of power in long-wavelength mode caused by the poor subpixel treatment in the standard nearest-neighbor pointing matrix. The vertical axis is dimensionless in this toy example, but could have units like μK or Jy/sr for a real-world case. Figure 3 shows the noise model/inverse weights/inverse filter used in the various methods. **signal**: The input signal, a smooth long-wavelength mode, sampled at 10 samples per output pixel. **binned**: Simple binned map (the unweighted average per pixel). Very suboptimal in the presence of correlated noise, but unbiased. **ML**: Maximum-likelihood map. 2/3 of the signal amplitude is lost despite the naive expectation of biaslessness for this estimator. **FB deobs**: Filter+bin map debiased using an observation matrix. Identical to ML. **FB detrans**: Filter+bin map debiased by deconvolving a transfer function measured from simulations. Even more biased than the others due to ignoring mode coupling. **destripe**: Destriper in the maximum-likelihood limit (1-sample baselines with optimal baseline prior). Identical to ML.

```
def dewin(x): return np.fft.irfft(np.fft.rfft(x)
    )/np.sinc(freq,n=len(x)).real
```

After pixel window deconvolution the binned map (green) closely matches the input signal (red). The same can not be said for the other estimators. Maximum likelihood, filter+bin with observation matrix debiasing and destriping (which are all equivalent in the limit we consider here) are strongly biased, with the signal only being recovered with 1/3 of its real amplitude.

The situation is even worse for filter+bin with simulation-based debiasing, as this suffers from an additional bias due to assuming that the Fourier modes are independent.⁶

2.1.6. Explanation

To see why inaccuracies in modelling the signal at sub-pixel scales can bias the largest scales in the map,

⁶ This additional bias disappears if the simulations have exactly the same statistical properties as the real signal we wish to recover.

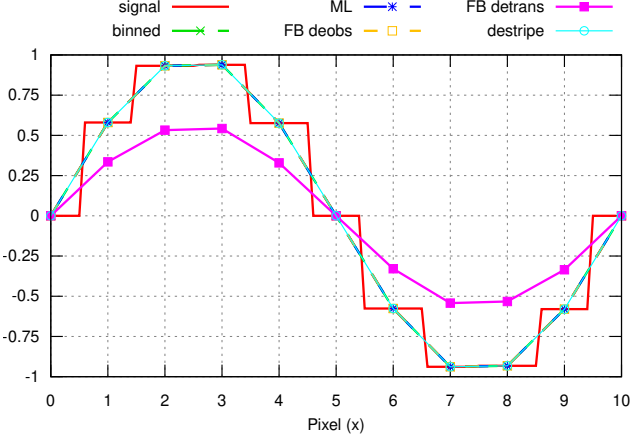


Figure 5. Like figure 4, but with the input signal having the same nearest-neighbor pixelization as the models. In this case all models except FB detrans are unbiased.

let us consider the example in figure 2, where a detector measures a smooth, large-scale signal while moving across a few pixels. With a nearest-neighbor pointing matrix it is impossible to model this smooth signal: the model for each sample is simply that of the closest pixel, regardless of where inside that pixel it is. The model therefore looks like a staircase-like function in time domain.

Given that we can't exactly match the signal, what is the best approximation? Let us consider two very different alternatives. **Model A:** The value in each pixel is the average of the samples that hits it, making the model curve trace the smooth signal as closely as it can. This is the green curve in figure 2, and has the sawtooth-like residual shown with the blue curve. It is probably the model most people would choose if asked to draw one manually. **Model B:** The value in each pixel is zero, and the residual is simply the signal itself. Model B seems like a terrible fit to the data, but under a reasonable noise model for a ground-based telescope, like the one shown in figure 3, it will actually have a higher likelihood (lower $\chi^2 = r^T N^{-1} r$ where r is the residual) than model A. The reason is that while model B has a much larger residual than model A, model B's residual is smooth and hence has most of its power at low frequency in time domain where N^{-1} is very small. Meanwhile, model A's residual extends to all frequencies due to its jagged nature, including the costly high frequencies. The actual maximum-likelihood solution will be intermediate between these two cases, sacrificing some but not all of the large-scale power in the model to make the residual smoother.

To demonstrate that the bias really is caused by subpixel errors, I repeated the simulation with a signal that follows the same nearest-neighbor behavior as the

data model, thus eliminating subpixel errors. The result is shown in figure 5. The bias has disappeared in all methods except filter+bin with transfer-function based debiasing, which has an additional source of bias due to its assumption of a Fourier-diagonal filter response.

2.2. 2D toy example

The 1D toy example is useful for understanding the origin of the bias, but its unrealistic observing pattern makes it insufficient for exploring optimality/bias trade-offs in the different methods. I therefore made a larger toy example where a single detector scans at constant speed across a square patch, sampling $N_{\text{scan}} = 400$ equi-spaced rows with N_{scan} equi-spaced samples per row, followed by a column-wise scan the same patch, leading to a total cross-linked data set $N_{\text{samp}} = 2N_{\text{scan}}^2 = 3200$ samples long. This equi-spaced grid of samples was chosen to make it easy to simulate a signal directly onto the samples without needing the complication of pixel-to-sample projection. These samples will then be mapped onto a square grid of pixels with a side length of $N_{\text{side}} = 100$ using the different mapmaking methods.

For the signal I draw realizations from a CMB-like $1/l^2$ power spectrum with a Gaussian beam with standard deviation $\sigma = 3$ pixels. Here l is the pixel-space wave-number, which I evaluate in the higher resolution sample space as

```
ly = np.fft.fftfreq(N_scan)[: ,None] * N_side /
    N_scan
lx = np.fft.fftfreq(N_scan)[None, :] * N_side /
    N_scan
l = (ly**2 + lx**2)**0.5
```

With this we can define the signal power spectrum $C_l = l^{-2}$ and beam $B_l = \exp(-l^2 \sigma^2 / 2)$, and draw signal realizations as

```
signal_map = np.fft.ifft2(np.fft.fft2(np.random
    .standard_normal((N_scan, N_scan))) * Cl**0.5 *
    B1).real
signal_tod = np.concatenate([signal_map.reshape
    (-1), signal_map.T.reshape(-1)])
```

The last step here takes into account that the simulated scanning pattern covers the field twice, first horizontally and then vertically.

For the noise I use a simple $1/f$ spectrum $N(f) = 1 + (f/f_{\text{knee}})^\alpha$, with $f = \text{np.fft.rfftfreq}(N_{\text{samp}})$ and the knee frequency f_{knee} corresponding to $1/30$ th of the side length, $f_{\text{knee}} = 0.5 \cdot 30/N_{\text{scan}} = 0.0375$ (6.7 pixel wavelength), and with an atmosphere-like exponent $\alpha = -3.5$.

The pixel coordinates of each sample are

```
pix_pat1 = (np.mgrid[:N_scan, :N_scan] * N_side /
    N_scan).reshape(2, -1)
pix_pat2 = pix_pat1[:, :-1]
pix = np.concatenate([pix_pat1, pix_pat2], 1)
```


which I use to build the nearest-neighbor pointing matrix

```
iy, ix = np.floor(pix).astype(int)%N_side
P_nn = scipy.sparse.csr_array((np.full(N_samp, 1), (np.arange(N_samp), iy*N_side+ix)), shape=(N_samp, N_pix))
```

2.2.1. Cases

We will investigate 5 classes of nearest-neighbor maps:

1. **bin**: Simple binned map, which I expect to be unbiased but very noisy
2. **ML**: Maximum-likelihood map. Ideally unbiased and optimal, but will deviate from this due to model errors.
3. **ML wX**: Maximum-likelihood maps with a “whitened” noise model, which overestimates the white noise power by a factor 10^X , $X \in \{1, 2, 3\}$, reducing the overall correlatedness of the noise model. I expect the bias to be proportional to the overall dynamic range of the noise model, so making the noise model whiter should reduce bias. The cost will be suboptimal noise weighting, leading to a noisier map, but it might be worth it.
4. **DS X**: Destriping map with a baseline of $X \in \{4, 16, 64\}$ samples and no amplitude prior. This is probably the most common type of destriping. The baseline lengths can be compared with the noise knee wavelength of 6.7 pixels ≈ 27 samples. This is the wavelength where the correlated and white noise powers are equal.
5. **DS+ X**: Like DS X, but uses the correlated part of the maximum-likelihood noise model as an amplitude prior (C_a in equation 4).
6. **FB**: Filter+bin mapmaking where a transfer function is measured using simulations based on the data model, and this is deconvolved when measuring the power spectrum. This is the standard approach for filter+bin mapmaking. In this case only the power spectrum is debiased, not the map, so I don’t show an example map for this case.

Since all the mapmaking methods are linear, the signal and noise can be mapped separately. I make 400 signal-only and noise-only data realizations, and map them using each method, computing the mean signal and noise power spectra.

2.2.2. Results

Example maps are shown in figure 6, while the bias and noise are quantified in figures 7 and 8 respectively.

As expected the binned map is unbiased but extremely noisy. The maximum-likelihood map is low-noise, but measurably biased on all scales, with a power deficit of a few percent at the smallest scales which grows to almost 100% on the largest scales. The deficit appears to fall proportionally with the whitening, with ML w1 and w2 being respectively 10x and 100x as close to unbiased. Sadly this comes at the cost of 40% and 350% higher noise power respectively. These numbers may differ for real-world cases, but this still seems like a very expensive bias mitigation method. All but the longest-baseline destriped maps are also strongly biased, with the shortest baseline being considerably worse than ML for almost all scales. Much like we saw with the ML variants, the less biased destriping versions come at a high cost in noise. Finally, the filter+bin map is biased even after simulation-based debiasing due to the simulations not capturing the subpixel behavior of the real data. Both the bias and noise levels are the same as maximum-likelihood in this example.

To test whether the observed biases are truly caused by subpixel errors, I repeat the simulations with only one sample per pixel ($N_{\text{scan}} = N_{\text{side}}$). As expected this results in an unbiased power spectrum for all methods.⁷

2.2.3. Effective mitigation of subpixel errors

There will be no subpixel errors in the limit of infinitely small pixels, so it’s tempting to simply reduce the pixel size to solve the problem. This does work, but since subpixel errors are proportional to $|\nabla s \cdot \vec{\Delta}|$, where s is the true, smooth signal on the sky and $\vec{\Delta} = [\Delta_x, \Delta_y]$ is the pixel shape, the improvement is only first order in the pixel side length. A much more feasible solution is to instead improve the subpixel handling in the pointing matrix. Going from nearest neighbor to bilinear interpolation is enough to practically eliminate subpixel errors without reducing pixel size. We can implement this in the toy example by defining

```
pix_left = np.floor(pix).astype(int)
ry, rx = pix - pix_left
iy1, ix1 = pix_left % N_side
iy2, ix2 = (pix_left + 1) % N_side
weights = np.concatenate([
    (1-ry)*(1-rx), (1-ry)*rx,
    ry*(1-rx), ry*rx])
samps = np.tile(np.arange(N_samp), 4)
inds = np.concatenate([
    iy1*N_side+ix1, iy1*N_side+ix2,
    iy2*N_side+ix1, iy2*N_side+ix2])
```

⁷ Even filter+bin, which ignores mode coupling, ends up having an unbiased power spectrum because the simulations used to build the transfer function followed the same distribution as the real signal.

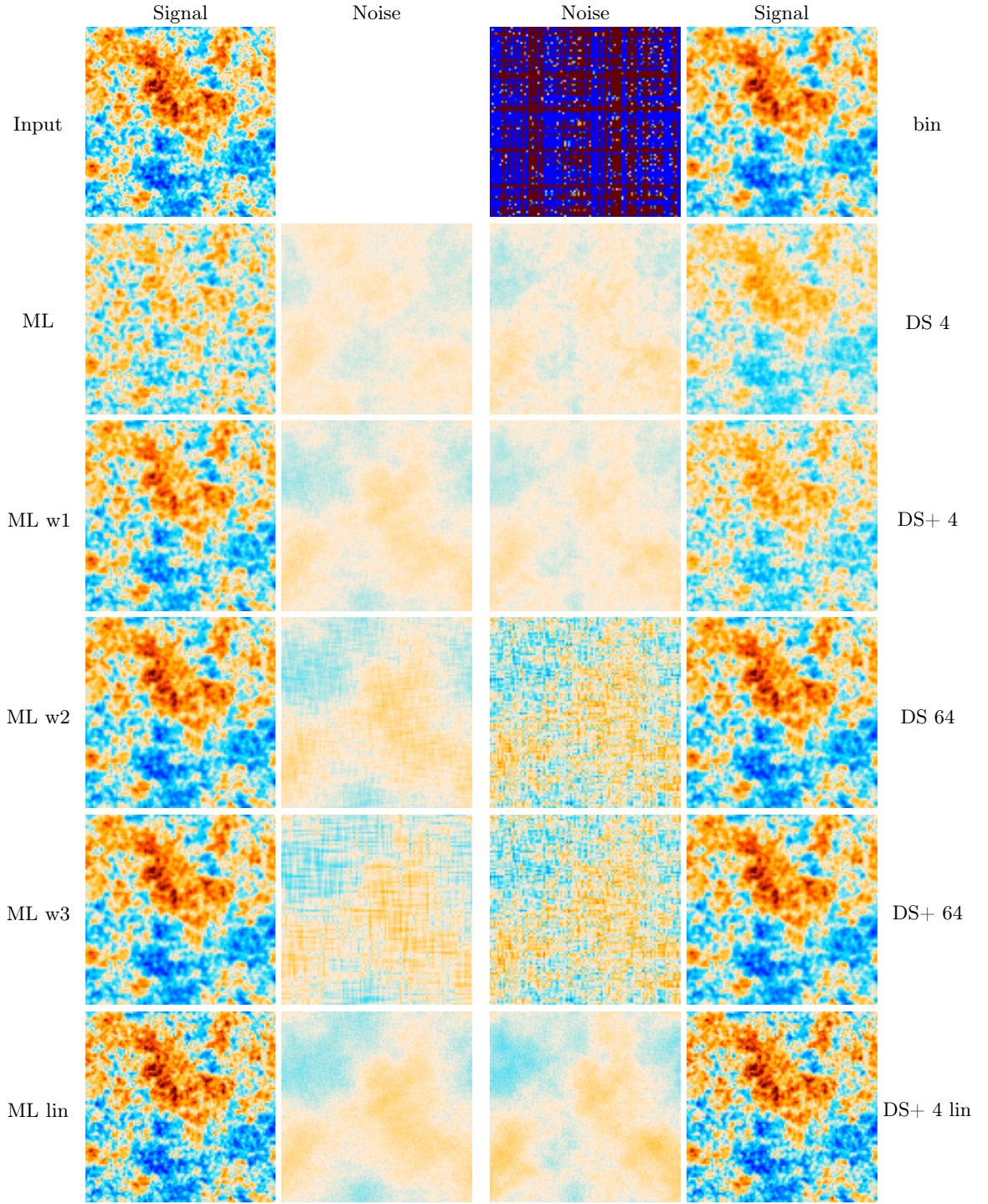


Figure 6. Example signal and noise maps for the different mapmaking methods. The top left map is the input signal, which is directly evaluated at 4x higher resolution than what is used for reconstructing the output maps. All output maps were built from the same data and assume a nearest neighbor pointing matrix, except for the last row where a bilinear pointing matrix was used. The color range is the same for all panels. The binned map (top right) is bias-free but has uselessly high noise. Maximum likelihood (ML) and destriping with short baselengths (with (DS+) and without (DS) baseline amplitude prior, which only matters for the shortest baseline) are all low-noise but biased on large scales. This bias goes away as the noise model is artificially whitened (ML wX) or the baseline length is increased (for destriping), but this comes at a cost of increased noise on all scales. Bilinear mapmaking (last row) instead avoids the bias by eliminating most of the model error. It can be difficult to judge how significant the bias and noise levels are from these images. See figures 7 and 8 for easier to interpret comparisons of the signal and noise power spectra respectively.

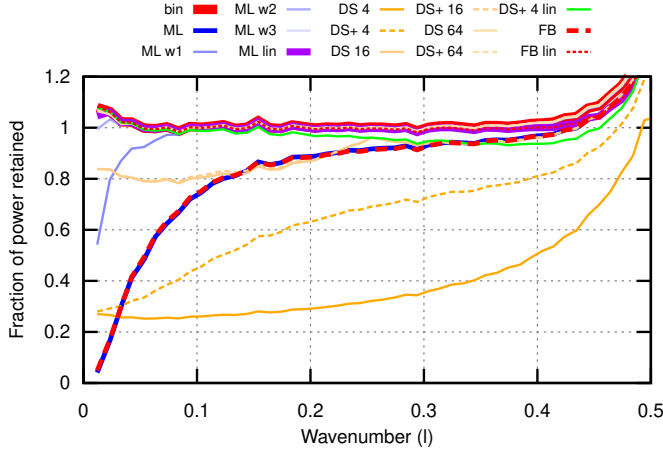


Figure 7. Comparison of the subpixel bias of different mapmaking methods described in section 2.2.1 and discussed in section 2.2.2. Standard maximum-likelihood (thick blue) is strongly biased, as are all but the (very noisy) longest destriping baseline. This bias disappears (ML) or is greatly reduced (DS) when switching to bilinear interpolation in the pointing matrix. See figure 8 for the corresponding noise spectra. The wavenumber l is dimensionless and with a Nyquist frequency of 0.5. I interpret the upturn at $l > 0.4$ as aliasing, which is expected and not relevant for the biases we consider here.

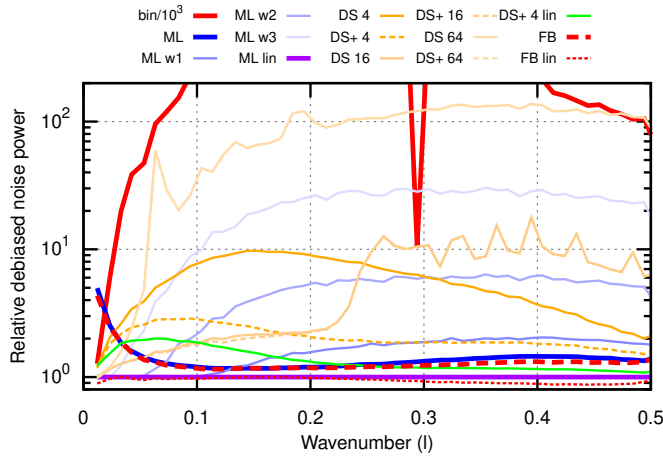


Figure 8. Comparison of the optimality for the methods defined in section 2.2.1, measured as the debiased noise power spectrum (using the bias measurements from figure 7) for each method relative to that of bilinear maximum-likelihood mapmaking. Note the logarithmic vertical axis, and that binned case was divided by 10^3 to bring it partially inside the plot bounds. The wavenumber l is dimensionless and with a Nyquist frequency of 0.5. It's clear that simple bias mitigation methods like artificially whitening the noise model or increasing the baseline length are too costly to be practical.

```
P_lin = scipy.sparse.csr_array((weights,(samps,
inds)),shape=(N_samp,N_pix))
```

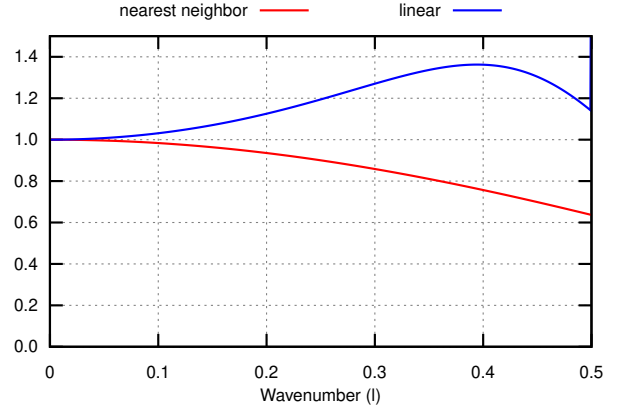


Figure 9. Comparison of the 1D pixel window for nearest neighbor mapmaking (red, $\text{sinc}(l)$) and linear mapmaking (blue). The 2D pixel window is the outer product of the 1D one along each axis. The pixel windows model the response of the Fourier coefficients to binned (unweighted) mapmaking. Square it to get the effect on the power spectrum.

Bilinear mapmaking results in a different pixel window than the standard `pixwin_nn = sinc(ly)[: ,None]*sinc(lx)[None, :]` of nearest neighbor mapmaking. I measure this empirically using simulations of simple binned mapmaking, and find it to have the separable form `pixwin_lin = linwin1d(ly)[: ,None]*linwin1d(lx)[None, :]`, with `linwin1d` being plotted in figure 9.

Since each sample in bilinear mapmaking gets contributions from the four closest pixels, the pointing matrix is about four times slower than the standard nearest neighbor version. However, as shown in figures 7 and 8 this cost is worth it, with bilinear maximum-likelihood mapmaking being bias-free and even lower noise than standard ML. The bias is also greatly reduced for destriping, but not eliminated.

3. DON'T THINK YOU'RE SAFE JUST BECAUSE YOU AVOID SUBPIXEL ERRORS

By no means all model errors have the strong impact on large scales that subpixel errors do. For example, gain errors typically have a scale-independent response while pointing errors broaden the beam, damping small scales but not large ones. It is, in fact, quite difficult to come up with other mechanisms by which model errors can cause large-scale power bias. One example I found requires a multi-detector system with large gain error differences between the detectors, combined with a noise model where the noise is dominated by a long-wavelength common mode (strong correlation between detectors at long wavelengths). This is implemented in the program `common_mode_test.py` (see section 5) with the output shown in figure 10. In this example the large

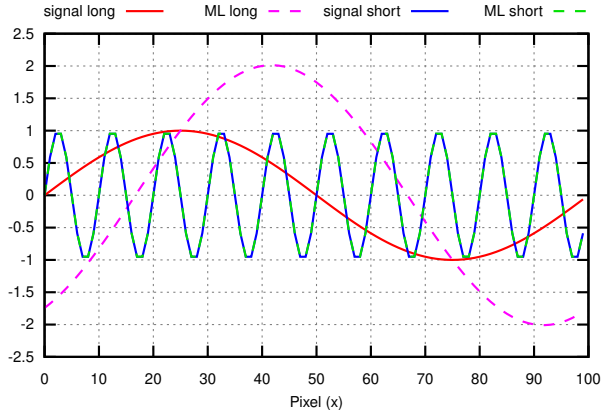


Figure 10. Demonstration of large-scale bias in a multi-detector system due to an interaction between strong large-scale detector correlations in the noise model and large relative gain errors between the detectors. **signal long:** An input long-wavelength signal, with the same pixelization as the output to avoid subpixel bias. **ML long:** Corresponding maximum-likelihood map, which exhibits both an amplitude and phase error. **signal short** and **ML short:** The same, but for a short-wavelength mode. Here the bias is negligible, despite the model’s gain errors being scale-independent.

scales are biased *high* and also subject to a phase shift, while the small scales have negligible bias.

While the common-mode/bias-interplay works as a demonstration, it is a much less robust error mechanism than subpixel errors, and requires quite extreme parameters to produce appreciable bias. In my tests it also tends to disappear as more data is added and crosslinking increases, and I have not been able to provoke large-scale bias using this method for full-scale realistic datasets (unlike subpixel errors!).

It would be convenient if the difficulty in finding model error mechanisms that bias large scales meant that subpixel errors are the only ones one needs to worry about, but sadly that does not seem to be the case. In my practical experience with ground-based CMB mapmaking there have been cases of large-scale power loss where subpixel errors were responsible for less than half of the total bias. In these cases the difficulty in finding these mechanisms becomes a curse rather than a blessing, since it makes it very difficult to track down the source of the bias.

What makes model error bias especially insidious is that it is completely invisible to any simulation that does not explicitly include that particular type of model error. For example, a simulation where the CMB is read off from a simulated map using the same pointing matrix as is used in the mapmaking itself would be blind to subpixel errors. Given the many possible ways the real data might deviate from one’s model of it, it is hard to be

sure that one has included all the relevant types of model error in the simulations. It is therefore very easy to trick oneself into believing that one has an unbiased analysis pipeline while there are in fact $\mathcal{O}(1)$ biases remaining.

4. USEFUL TESTS

The unintuitive large-scale effects of model errors rely on the interaction between model errors and non-local weighting/filtering. A noise model (or filter) with a large dynamic range, such as one capturing the huge ratio between the long-wavelength and short-wavelength noise power for ground-based CMB telescopes, is therefore much more vulnerable to large-scale power loss than one appropriate for a space-based telescopes which have almost flat noise power spectra. This suggests the following tests for large-scale model error bias:

1. Compare power spectra with those from a space-based telescope. This is reliable but comes at the cost of being able to make an independent measurement.
2. Split the data into subsets with different ratios of correlated noise to white noise and check their consistency. This could for example be a split by the level of precipitable water-vapour (PWV) in the atmosphere. High-PWV-data would be expected to have a higher dynamic range and therefore be more vulnerable.
3. Map the same data both using the standard noise model/filter and a less optimal one an artificially reduced lower dynamic range, e.g. one that underestimates the amount of correlated noise. The latter should result in less a biased but noisier map, as we saw in figures 7 and 8. If the maps are consistent, then large-scale model error bias is probably not an issue.

5. SOURCE CODE

The source code and data files behind these examples is available at https://github.com/amaurea/model_error2. The 1D and 2D subpixel simulations are available in `subpix/model_error_toy.py` and `subpix/model_error_toy.py` respectively, while the common-mode/gain-error interplay example is in `common_mode/common_mode_test.py`.

6. CONCLUSION

1. **All CMB current mapmaking methods are vulnerable to model error bias**, including maximum-likelihood mapmaking, destriping and filter+bin mapmaking.

2. **The most common model error is subpixel errors** due to the assumption that the signal is constant inside each pixel (a nearest-neighbor pointing matrix), but **other types of model error can also be important**.
3. Model error bias can manifest in unintuitive ways, with a common symptom being a large ($\mathcal{O}(1)$) **loss of long-wavelength power** in the maps.
4. The size of the bias is proportional to the dynamic range of the filter/noise model. Hence it is **important for ground-based measurements of the unpolarized CMB** due to the presence of large-scale atmospheric noise there, but is **much less important for polarization measurements or space-based telescopes**.
5. **Simulations are blind to these biases unless specifically designed to target them**. There

is a large risk of ending up thinking one has an unbiased pipeline despite there being large bias in the actual CMB maps.

6. An effective way of testing for this bias is to **also map the data using a (much) lower dynamic range noise model/filter**⁸ and checking if this leads to consistent power spectra.

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⁸ Or much longer baselines for destriping

APPENDIX

A. LINEAR PIXEL WINDOW

Let us start by considering the 1D case. Given a set of pixels $\{i\}$ with values $\{v_i\}$ we define the linear interpolation value d at some pixel coordinate $i + x$ where $x \in [0, 1)$ as

$$d = (1 - x)v_i + xv_{i+1} \quad (\text{A1})$$

This is illustrated in figure 11.

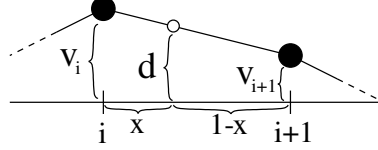


Figure 11. Definition of the quantities involved in linear interpolation. Given values $\{v_i\}$ sampled at locations $\{i\}$, we define the linear interpolation value d at some location $i + x$ where $x \in [0, 1)$ as $d = (1 - x)v_i + xv_{i+1}$.

We can generalize this to a set of samples d_a with

$$d_a = (1 - x_a)v_a^{\text{left}} + x_a v_a^{\text{right}} \quad (\text{A2})$$

where v_a^{left} and v_a^{right} are the values of the pixel to the left and right of where sample a points, and x_a is its subpixel position. This linear system can be written as the following matrix equation

$$d = Pv \quad (\text{A3})$$

P 's form will depend on the exact scanning pattern, but the final pixel window should only depend on each pixel being hit equally much at all subpixel locations. For simplicity, I will assume a simple scanning pattern that scans a single time across a row of pixels from left to right, with a constant scanning speed and infinite sample rate. This results in the pointing matrix P taking the form shown in figure 12.

Figure 12. Structure of a linear-interpolation pointing matrix P and its transpose P^T for the case of a set of samples making a single scan across a row of pixels in 1D, assuming wrapping boundary conditions. $x \in [0, 1)$ is the subpixel coordinate of each sample.

Given the set of samples d , the least squares estimator for the pixel values v under uniform weighting is

$$\hat{v} = (P^T P)^{-1} P^T d \quad (\text{A4})$$

The denominator $P^T P$ consists of two types of non-zero cases:

$$(P^T P)_{00} = \begin{bmatrix} \cdots & 1-x & \cdots \end{bmatrix} \begin{bmatrix} \vdots \\ 1-x \\ \vdots \end{bmatrix} + \begin{bmatrix} \cdots & x & \cdots \end{bmatrix} \begin{bmatrix} \vdots \\ x \\ \vdots \end{bmatrix} = S \int_0^1 (1-x)^2 dx + S \int_0^1 x^2 dx = \frac{2}{3} S \quad (\text{A5})$$

$$(P^T P)_{01} = \begin{bmatrix} \cdots & 1-x & \cdots \end{bmatrix} \begin{bmatrix} \vdots \\ x \\ \vdots \end{bmatrix} = S \int_0^1 (1-x)x dx = \frac{1}{6} S \quad (\text{A6})$$

Here S is the number of samples per pixel, and appears due to the summing implied by P^T . $S \rightarrow \infty$ in the integral limit we used here, but it will end up cancelling. With this, we end up with the denominator taking the Toeplitz form

$$P^T P = \frac{1}{6} N \begin{bmatrix} 4 & 1 & & & 1 \\ 1 & 4 & 1 & & \\ & 1 & 4 & 1 & \\ & & & \ddots & \\ & & & & 1 & 4 & 1 \\ 1 & & & & & 1 & 4 \end{bmatrix} \quad (\text{A7})$$

Toeplitz matrices are diagonal in Fourier space, so $(P^T P)^{-1}$ becomes a simple element-wise division there. The Fourier-diagonal is given by the Fourier transform of the first row of the Toeplitz matrix. Defining $A = P^T P$, it's Fourier diagonal as \tilde{a} is given by

$$\tilde{a}(k) = \frac{1}{N} \sum_{j=0}^{N-1} e^{\frac{-2\pi i k j}{N}} A_{j0} = \frac{S}{N} \frac{1}{6} \left(4 + e^{\frac{2\pi i k}{N}} + e^{\frac{-2\pi i k}{N}} \right) = \frac{S}{N} \frac{1}{3} \left(2 + \cos \frac{2\pi k}{N} \right) \quad (\text{A8})$$

where N is the number of pixels, i is the imaginary unit, k is the zero-based index into \tilde{a} and j is the zero-based row index of A .

Turning to the numerator $b = P^T d$, we recognize that it is simply the convolution of d with the triangle function $T(x) = (1 - |x|)S$, $x \in [-1, 1]$, evaluated at each pixel center.⁹ Using the convolution theorem, we can therefore directly write down

$$\tilde{b}(k) = \tilde{T}(k) \cdot \tilde{d}(k) \quad (\text{A9})$$

We're left with computing the Fourier-space representation of the triangle function T .

$$\begin{aligned} \tilde{T}(k) &= \frac{S}{N} \int_{-1}^1 e^{\frac{-2\pi i k x}{N}} (1 - |x|) dx \\ &= \frac{S}{N} \left[\int_{-1}^1 e^{\frac{-2\pi i k x}{N}} dx - \int_0^1 e^{\frac{-2\pi i k x}{N}} x dx - \int_{-1}^0 e^{\frac{-2\pi i k x}{N}} (-x) dx \right] \\ &= \frac{S}{N} \left[2 \operatorname{sinc} \frac{2k}{N} - 2 \operatorname{sinc} \frac{2k}{N} + 2 \left(\frac{2\pi k}{N} \right)^2 \left(1 - \cos \frac{2\pi k}{N} \right) \right] \\ &= \frac{S}{N} \left(\operatorname{sinc} \frac{k}{N} \right)^2 \end{aligned} \quad (\text{A10})$$

Combining the numerator and denominator we have

$$\hat{v} = \frac{\tilde{b}(k)}{\tilde{a}(k)} \tilde{d}(k) = \frac{\left(\operatorname{sinc} \frac{k}{N} \right)^2}{\frac{1}{3} \left(2 + \cos \frac{2\pi k}{N} \right)} \tilde{d}(k) = W_{\text{lin}}(k/N) \tilde{d}(k) \quad (\text{A11})$$

⁹ This means that $T * d$ is the same function as b , just S times more densely sampled. Hence, it has the same Fourier representation up to b 's Nyquist frequency.

where the linear interpolation pixel window $W_{\text{lin}}(f)$ is as

$$W_{\text{lin}}(f) = \frac{\text{sinc}(f)^2}{\frac{1}{3}(2 + \cos(2\pi f))} \quad (\text{A12})$$

This pixel window is compared to the standard nearest neighbor pixel window $W_{\text{nn}} = \text{sinc}(f)$ in figure 13. Their behavior is quite different. While W_{nn} falls monotonically to around 0.6 at the Nyquist frequency, W_{lin} rises to about 1.4 at $f = 0.4$, and only then starts falling.

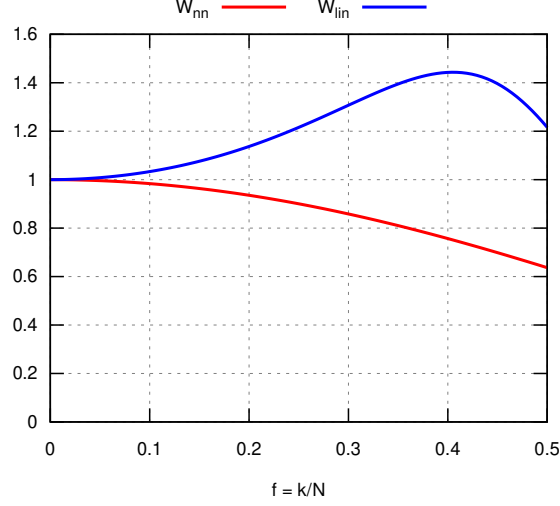


Figure 13. Comparison of the 1D pixel windows for nearest neighbor mapmaking (W_{nn}) and linear mapmaking (W_{lin}), plotted up to the Nyquist frequency, which is $\frac{1}{2}$ in these units.

The natural generalization of linear interpolation to two dimensions is bilinear interpolation, which is the composition of a horizontal and vertical one-dimensional interpolation. Because this operation is separable, so is the resulting pixel window. Hence

$$W_{\text{lin}}^{2\text{D}}(f_y, f_x) = W_{\text{lin}}(f_y)W_{\text{lin}}(f_x) \quad (\text{A13})$$