A SEARCH FOR COSMIC MICROWAVE BACKGROUND ANISOTROPIES ON ARCMINUTE SCALES WITH BOLOCAM


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Received 2008 May 20; accepted 2008 September 14; published 2008 December 22

ABSTRACT

We have surveyed two science fields totaling 1 deg2 with Bolocam at 2.1 mm to search for secondary Cosmic Microwave Background (CMB) anisotropies caused by the Sunyaev-Zel’dovich effect (SZE). The fields are in the Lynx and Subaru/XMM SDS1 fields. Our survey is sensitive to angular scales with an effective angular multipole of \( \ell_{\text{eff}} = 5700 \) with FWHM \( \theta = 2800 \) and has an angular resolution of 60 arcsec FWHM. Our data provide no evidence for anisotropy. We are able to constrain the level of total astronomical anisotropy, modeled as a flat-band power in \( C_\ell \), with most frequent 68%, 90%, and 95% CL upper limits of 590, 760, and 830 \( \mu K^2_{\text{CMB}} \). We statistically subtract the known contribution from primary CMB anisotropy, including cosmic variance, to obtain constraints on the SZE anisotropy contribution. Now including flux calibration uncertainty, our most frequent 68%, 90%, and 95% CL upper limits on a flat-band power in \( C_\ell \) are 690, 960, and 1000 \( \mu K^2_{\text{CMB}} \).

1. INTRODUCTION

1.1. Background

The Sunyaev-Zel’dovich effect (SZE)6 is the inverse Compton scattering of Cosmic Microwave Background (CMB) photons with a distribution of hot electrons, causing a net increase in the energy of the photons (Sunyaev & Zel’dovich 1972). Since the background CMB is redshifted along with the SZE-induced distortion, the relative amplitude of the distortion, \( \Delta T_{\text{CMB}} / T_{\text{CMB}} \), is independent of the redshift. The distortion caused by the SZE is proportional to the Comptonization parameter \( y \), which is a measure of the integral of the electron thermal energy density along the line of sight and is given by

\[
y = \frac{\sigma_T}{m_e c^2} \int dl \, n_e k_B T_e,
\]

where \( \sigma_T \) is the Thomson cross-section, \( m_e \) is the electron mass, \( c \) is the speed of light, \( k_B \) is Boltzmann’s constant, \( T_e \) is the temperature of the electrons, and \( n_e \) is the number density of electrons. Since the scattering process conserves the photon number, the thermal spectrum of the CMB is distorted by the SZE; there is a negative temperature shift at low frequency and a positive temperature shift at high frequency. The cross-over point where there is no distortion of the CMB occurs at approximately 218 GHz. The temperature shift caused by the SZE, \( \Delta T_{\text{CMB}} \), is

\[
\frac{\Delta T_{\text{CMB}}}{T_{\text{CMB}}} = f(x) y,
\]

where

\[
f(x) = \frac{x e^x + 1}{e^x - 1} - 4,
\]

and \( x = h\nu / k_B T_{\text{CMB}} \). \( h \) is Planck’s constant, \( \nu \) is the frequency, and \( T_{\text{CMB}} = 2.73 \) K is the temperature of the CMB. For reference, excellent reviews of the SZE and its relevance to cosmology are given by Birkinshaw (1999) and Carlstrom et al. (2002).

1.2. Untargeted SZE Surveys

Recently, the South Pole Telescope discovered the first previously unknown galaxy clusters using the SZE (Staniszewski et al. 2008). However, even when no clusters are detected, unresolved objects in SZE surveys will produce anisotropies in the CMB that are expected to dominate the CMB power spectrum at small angular scales corresponding to angular multipoles above \( \ell \approx 2500 \). The overall normalization of these SZE-induced CMB anisotropies is extremely sensitive to \( \sigma_8 \) and can be used to constrain the value of this cosmological parameter (Komatsu & Seljak 2002). Several experiments have conducted SZE surveys that have produced tentative detections of the SZE-induced anisotropies in the CMB. At 30 GHz, CBI has measured an excess CMB power between \( \ell = 2000 \) and \( \ell = 3500 \) at a significance of 3.1\sigma (Mason et al. 2003). Also at 30 GHz, BIMA/OVRO has measured a CMB anisotropy of 220\(^{+140}_{-120}\) \( \mu K^2_{\text{CMB}} \) at an

5 Jansky Fellow, National Radio Astronomy Observatory.
6 Throughout this paper SZE refers to the thermal SZE.
angular multipole of $\ell = 5237$ (Dawson et al. 2006). ACBAR, at 150 GHz and at $\ell < 3000$, has measured an excess power of $34 \pm 20 \mu K_{\text{CMB}}^2$ (Reichardt et al. 2008). A joint analysis of the CBI and ACBAR excesses shows that they are six times more likely to be caused by the SZE than primordial fluctuations (Reichardt et al. 2008).

Additionally, these tentative anisotropies have been used to constrain cosmological parameters: the CBI data are consistent with $s_8 \sim 1$ (Bond et al. 2005), and the BIMA/OVRO data measure $s_8 = 1.03^{+0.20}_{-0.20}$ (Dawson et al. 2006). Reichardt et al. (2008) combine various data sets to place constraints on $s_8$ via an excess contribution to anisotropy at high $\ell$, $\ell > 1500$. ACBAR and WMAP3, plus CBI and BIMA/OVRO data at high $\ell$ and lower frequency, combine to indicate $s_8^{\text{SZ}} = 0.95^{+0.03}_{-0.04}$ when the amplitude of the SZE contribution is not slaved to its contribution at low $\ell$. Since this result is inconsistent with other constraints on $s_8$, including those from the lower $\ell$ portions of the CMB spectrum, Reichardt et al. (2008) also consider a case in which the high-$\ell$ SZE contribution is slaved to the contribution at lower $\ell$ via $s_8^{\text{SZ}} = s_8$. In this case, the excess power is produced by point sources. Fitted to the CMBall data set, which excludes CBI high-$\ell$ and BIMA/OVRO data, this model results in $s_8$ values consistent with other measurements, $\pm 0.80 \pm 0.15 \pm 0.03 - 0.04$ depending on the assumptions and data sets included. No attempt is made to explain the CBI and BIMA/OVRO excesses. Overall, the current results suggest two possibilities: there are point source contributions to all the high-$\ell$ data (ACBAR, CBI, BIMA/OVRO) that have not been properly included; or the SZE contribution calculated from theory is underestimated.

The survey presented here is the first such survey at 150 GHz and at $\ell \sim 6000$. As we shall explain, contributions from primary CMB anisotropies, SZE, radio, and submillimeter point sources are all expected to be comparable, each at a level of $C_\ell \simeq 50 \mu K_{\text{CMB}}^2$.

2. OBSERVATIONS

2.1. Instrument Description

Bolocam is a large format, 144 detector, millimeter-wave camera designed to be operated at the Caltech Submillimeter Observatory (CSO). For these observations the array was comprised of 115 optical and six dark detectors. Each detector is housed within its own integrating cavity, formed by a front short plate and a back short plate (Glenn et al. 2002). Smooth-walled conical feed horns separated by 0.7 (f/#)\,, a cold (4 K) high-density polyethylene (HDPE) lens, and a room-temperature ellipsoidal mirror are used to couple the detectors to the CSO optics. Each feed horn terminates into a cylindrical waveguide, which defines the low-frequency cutoff of the system; the final filter in a series of six cold metal-mesh filters determines the high-frequency cutoff. The resulting passband is centered at 143 GHz, and has an effective width of 21 GHz. A cold (4 K) Lyot stop is used to define the illumination of the 10.4 m primary mirror, and the resulting far-field beams have FWHMs of 60 arcsec. Bolocam can also observe at 270 GHz, and has been used in this mode for several types of observations, including surveys for submillimeter galaxies and protostellar cores (Laurent et al. 2005; Enoch et al. 2006; Young et al. 2006).

The detector array has a hexagonal geometry, and utilizes silicon nitride micromesh (spider-Web) bolometers (Mauskopf et al. 1997) which are cooled to 260 mK using a three-stage $^4$He/$^3$He/$^3$He sorption refrigerator (Bhatia et al. 2000, 2002). Junction field effect transistors (JFETs) located near the array and operated at 140 K are used to buffer the high-impedance bolometer signals from sources of current noise. In order to avoid the $1/f$ noise from the JFETs, the bolometers are biased at 130 Hz and read out using room-temperature lock-in amplifiers. More details of the Bolocam instrument can be found in S. R. Golwala et al. (2008, in preparation), Glenn et al. (1998, 2003), and Haig et al. (2004).

2.2. Observing Strategy

The data described in this paper were collected during a forty night observing run in late 2003. During the first half of each night we observed a 0.5 deg$^2$ region centered at 02h18m00s, $-5^\circ00'00''$ (J2000), which coincides with the Subaru/XMM Deep Survey (SXDS or SDS1); and during the second half of each night we observed a 0.5 deg$^2$ region centered on the Lynx field at 08h49m12s, +44d50m24s (J2000). These fields were selected because they have extremely low dust emission and a large amount of optical/X-ray data that could be used to follow up any SZE cluster candidates found in the maps.

Two nights at the start of the run were used to analyze different scan strategies for mapping the science fields. The maps were made by repeatedly raster scanning across the field, stepping perpendicular to the scan, and then rastering across the field in the opposite direction until the entire field has been covered. Our studies showed that the time-stream noise is independent of the angle of the raster scan and the turnaround time between scans, so we chose to scan parallel to right ascension or declination and turnaround as quickly as the telescope would allow ($\sim 10$ s). Additionally, we found that our sensitivity to astronomical signals is maximized when we raster scan at a speed of 240 arcsec s$^{-1}$. At this speed it takes approximately 12.5 s to complete one scan across the field, which means we were on-source approximately 56% of the time during an observation. Although scanning at this relatively quick speed reduces our time on-source because a larger fraction of time is spent on turnarounds between scans, it also puts a larger amount of our signal band above the 1/f atmospheric noise. Given the scan speed and turnaround time mentioned above, along with our step size of 162 arcsec ($\simeq 1/3$ of the field of view) a complete map of the field was made in approximately 8 minutes.

3. DATA REDUCTION

3.1. Initial Processing

After merging the bolometer time streams recorded by the data acquisition system with the pointing information recorded by the telescope, we parse the data into files that contain a single observation. Each single observation contains a set of scans that completely map the astronomical field or object, and they are typically around 10 minutes in length. Parsing the data by observation is useful because individual observations are statistically independent, have a small enough number of data samples to be easily manageable from an analysis standpoint, and provide a convenient division of the data for the sake of bookkeeping.

Once the initial merging and parsing of the data is complete, we begin the process of refining the data. The first step in this process is to remove the effects of the lock-in amplifier electronic filters and to down sample the data from 50 Hz

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7 Faster speeds were not attempted due to fears that the CSO would not function properly and/or would be damaged.
to 10 Hz. We down sample the data because essentially no astronomical signal is lost, while a large amount of 60 Hz pickup noise is removed. See Figure 1, for an illustration of the noise spectrum and the shape of the beam in frequency space.

3.2. Noise Removal

There are several forms of correlated noise present in the raw bolometer data which contaminate the astronomical signal and therefore must be modeled and removed. First, the emission from the atmosphere changes as a function of telescope elevation angle due to the changing path length through the atmosphere. The path length through the atmosphere relative to the zenith path length is called the air mass, A, and is described by

\[ A = 1 / \sin(\epsilon), \]

where \( \epsilon \) is the elevation angle. For a typical observation the range of elevation angles is approximately 1°, which corresponds to a change in air mass between 0.005 and 0.060 for elevation angles between 75° and 30°. For reference, a change of 0.060 in air mass corresponds to a change of approximately 0.5 K of optical loading from the atmosphere, or a change in surface brightness of a little less than 1 K_CMB. To remove this elevation-dependent signal, we calculate a linear fit of bolometer signal versus air mass. We build up a fit using each 12.5 s long scan within the observation, after removing the mean signal level and air mass for the scan. This process yields one set of linear-fit coefficients for each bolometer for the entire observation, which is used to create a template that is removed from the bolometer time streams.

Next, we create a template from the bias voltage monitors to account for the small amount of noise from the bias electronics. Note that the bias applied to the bolometers is monitored through amplifier electronics identical in design to those used to monitor the bolometer signals. This template is then correlated and removed from each of the bolometer time streams. A template is also created from the dark bolometer signals and removed from the bolometer time streams. Note that both the bias template and dark bolometer template have an rms of \( \lesssim 1 \) mK_CMB.

Finally, and most importantly, we remove a template describing the fluctuations in emission from the atmosphere (i.e., the atmospheric noise). Since atmospheric noise is the dominant signal in our data, and the beams from the individual detectors overlap to a high degree while passing through the atmosphere, a template for the atmospheric signal is created by averaging the signals from all the bolometers. Three different algorithms are used to construct this template, one for which the atmospheric signal is assumed to be constant over the array, one for which the atmospheric signal is allowed to vary linearly with bolometer location on the array, and one for which the atmospheric signal is allowed to vary quadratically with bolometer location on the array.

For the most basic case of an average template, the algorithm proceeds as follows. Initially, a template is constructed according to

\[ T_n = \frac{\sum_{i=1}^{N_b} c_i d_{in}}{\sum_{i=1}^{N_b} c_i}, \tag{1} \]

where \( n \) is the sample number, \( N_b \) is the number of bolometers, \( c_i \) is the relative responsivity of the bolometer \( i \), \( d_{in} \) is the signal recorded by bolometer \( i \) at time sample number \( n \), and \( T_n \) is the template. The template generally has an rms between 10 and 100 mK_CMB, depending on the observing conditions. A separate template is computed for each 12.5 s long scan. After the template is computed, it is correlated with the signal from each bolometer to determine the correlation coefficient, with

\[ \hat{c}_i = \frac{\sum_{j=1}^{N_s} T_n d_{jn}}{\sum_{j=1}^{N_s} T_n^2}. \tag{2} \]

\( \hat{c}_i \) is the correlation coefficient of bolometer \( i \) and \( N_s \) is the number of samples in the 12.5 s long scan. Next, the \( c_i \) in Equation (1) are set equal to the values of \( \hat{c}_i \) found from Equation (2), and a new template is computed. The process is repeated until the values of \( \hat{c}_i \) stabilize. We generally iterate until the average fractional change in the \( c_i \)'s is less than \( 1 \times 10^{-8} \), which usually takes five to ten iterations. If the \( c_i \)'s fail to converge after 100 iterations, then the scan is discarded from the data. For the more advanced planar and quadratic algorithms, the process proceeds in the same way except linear and quadratic variations with bolometer position are allowed when the template is constructed. These algorithms, along with adaptive PCA and time-lagged average template subtraction, are described and compared in more detail in Sayers et al. (2008).

Each of the three different atmospheric noise-removal algorithms, average, planar, and quadratic template removal, was applied to each observation. Therefore, three different atmospheric-noise-cleaned time streams are generated for each observation. A figure of merit (FOM) is calculated for each of the three files for each observation, based on the noise level of the data and the expected astronomical signal shape. Details of the calculation of this FOM are given in Section 7. For each observation, the file with the best FOM value will be the one

8 To quantify the degree of overlap, note that the far-field distance for Bolocam at the CSO is approximately 30 km; at a height of 20 km, well above most of the water vapor in the atmosphere, the ray bundles from nearby detectors have only separated by one half-width.

9 The best-fit correlation coefficients change from one scan to the next, typically by a couple percent.
used to create the final map of the data. Weather is the main criterion that determines which algorithm will be selected as optimal for a given observation; more aggressive algorithms (planar or quadratic) are selected in poor weather conditions and more benign algorithms (average or planar) are selected in good weather conditions. However, there is some dependence on the profile of the source, and observations of compact objects tend to be optimally processed using more aggressive algorithms than observations of extended objects.

4. CALIBRATION

4.1. Pointing Reconstruction

Pointing reconstruction consists of determining the location of each detector’s beam on the sky at each instant in time. We compute this location in two steps: (1) we calculate the location of each bolometer relative to the center of the array and (2) then determine the absolute coordinates of the center of the array.

To determine the relative locations of the bolometers, we observed Uranus or Neptune for approximately 15 minutes every other night. These planets are bright enough to appear at high signal-to-noise (S/N) in a map made from a single bolometer, so they can be used to determine the position of each detector relative to the array center. Since Bolocam was held at a fixed angle in the alt/az coordinate system for the entire observing run, each bolometer views the optics in the same way for the entire run and the coordinates on the sky in alt/az units remain fixed. Therefore, we combined the data from all the planet observations to determine the average position of each beam on the sky (see Figure 2). The uncertainties on these average positions were ~ 1 arcsec, which is negligible when compared to the 60 arcsec FWHM of a Bolocam beam.

We found no evidence for a systematic difference in the beam positions derived from any single observation to the average beam position found from all the observations. This indicates that the optical system was very stable over the entire observing run, including a wide range of telescope elevation angles.

To determine the absolute location of the center of the array, we observed a bright quasar with a known position near the science field for approximately 10 minutes once every 2 hr. Three different quasars were used for the SDS1 field (0106+013, 0113-118, and 0336-019), and two different quasars were used for the Lynx field (0804+499 and 0923+392). Each source was observed for 5 minutes while scanning parallel to right ascension, then for 5 minutes scanning parallel to declination. (analogous to how the science fields were observed). We found no systematic offset based on scan direction; the maps made while scanning parallel to right ascension produce the same source location as the maps made while scanning parallel to declination. The difference in the centroid location for these consecutive observations was then used to determine the measurement uncertainty for the centroided location of each source. As expected, the uncertainty in the centroided location of the five sources is inversely proportional to the flux of the source. Additionally, we found no evidence that the measurement uncertainty degrades or improves as a function of time during the night for our typical observing times between 20:00 and 07:00 local time.

The pointing data were broken up into three distinct subsets corresponding to the azimuthal position of the telescope: SDS1 was observed between azimuth angles of 90 and 270 (in the south), while Lynx was observed between azimuth angles of −90 and 90 (in the north approaching from the east), and also between azimuth angles of 270 and 360 (in the north approaching from the west). Most of the Lynx data were taken between an azimuth angle of −90 to 90, so the third subset of data is considerably smaller than the first two (about 1/5 of the size). Note that the slewing limits of the telescope are roughly equal to azimuth angles of −90 to 360. There is a correlation between the elevation angle of the telescope and the pointing offset for each of these subsets. We attempted to model this correlation with several low-order polynomials, but we found that a quadratic fit of pointing offset versus elevation was sufficient since higher order fits did not significantly reduce the scatter of the data. We found no correlation between the telescope azimuth angle and the residual offset, other than the slight difference between the pointing models determined for the three subsets. Therefore, a simple quadratic fit of pointing offsets versus telescope elevation angle served as our only pointing model. Plots of this final model can be found in Figure 3.

For each of the three subsets, we calculated the uncertainty in the pointing model by analyzing the residual offset of each centroid location from the model. Some residual scatter is expected due to the measurement uncertainty of each centroid; however the scatter we find is slightly larger. The difference between the actual scatter and the predicted scatter is consistent for all three subsets, and translates to an uncertainty in the pointing model of 4.9 arcsec. This uncertainty is small compared to our beam size and thus made a negligible difference in the beam shape used in the final science analysis.

4.2. Flux Calibration

Our flux calibration technique, summarized below, has been used previously with Bolocam to calibrate 1.1 mm data (Laurent et al. 2005). Since the amount of astronomical signal attenuation by the atmosphere is a function of opacity and air mass, the standard flux calibration technique for millimeter-wave instruments requires frequent observations of calibration sources that are close to the science field. However, we were able to use a more
advanced technique with Bolocam because we continuously monitor the operating resistance of the bolometers using the carrier amplitude measured by the bolometer voltage at the bias frequency. When the atmospheric transmission decreases, the optical loading from the atmosphere increases, which lowers the bolometer resistance. Additionally, the bolometer responsivity is a monotonically decreasing function of the bolometer resistance. Therefore, by fitting the flux calibration as a function of the bolometer-operating resistance, we can simultaneously account for changes in the atmospheric transmission and bolometer responsivity.

Six different flux calibrations were needed for our data set. The base temperature of the sub-Kelvin refrigerator was changed on 2003 November 4, and the bias voltage applied to the bolometers was changed on 2003 November 5, 8 (twice), and 10. Each of the bias changes caused a change in the responsivity of the bolometers, so a different flux calibration is needed after each change. Since the first five data sets are relatively short in duration, the observing conditions were relatively constant within each set. Therefore, a constant flux calibration, rather than a flux calibration that varies as a function of bolometer-operating resistance, was adequate to describe the data for these five sets. However, a fit of the flux calibration as a function of bolometer-operating resistance was required for the final data set.

The relative calibration of the detectors was determined from the science field observations. Since these observations covered regions of the sky with negligible amounts of astronomical flux, the fluctuations in thermal emission from the atmosphere are the dominant source of the signal recorded by each bolometer. Additionally, this signal should be the only one that is correlated among all the bolometers since the beams from all bolometers overlap to a high degree when passing through the atmosphere. Therefore, this signal should be the same in each bolometer, weighted by the responsivity of that bolometer. So, by determining how correlated the data from each bolometer is with this common signal, it is possible to determine the relative calibration of each bolometer. The uncertainties in the relative calibrations determined using this method are less than 1%.

The absolute flux calibration was determined from observations of Uranus, Neptune, 0923+392, and NGC 2071IR. Since we did not have enough observations of Uranus and Neptune to adequately determine the shape of the calibration versus bolometer-operating resistance, we used 0923+392 and NGC 2071IR as secondary calibrators. These two sources are known to have minimal variations in the emitted flux as a function of time, so they are well suited for determining the functional form of the flux calibration versus bolometer-operating resistance relationship (Peng et al. 2000; Sandell 1994). Note that we did not use any of the published fluxes for 0923+392 or NGC 2071IR, rather the fluxes were left as free parameters and they were used to determine the shape of the calibration curve versus bolometer-operating resistance. We used the peak signal and median bolometer-operating resistance from each observation to determine the fit parameters in the function

\[ V_j(R_{bolo}) = F_j(\alpha_1 + \alpha_2 R_{bolo}), \]

where \( V_j \) is the peak bolometer signal (in nV) recorded for the \( j \)th source, \( R_{bolo} \) is the bolometer-operating resistance, \( F_j \) is equal to the flux of the \( j \)th source (known for Uranus and Neptune, left as a free parameter for NGC 2071IR and 0923+392), and \( \alpha_1 \) and \( \alpha_2 \) free parameters. The planet fluxes were determined from the temperature spectra given in Griffin & Orton (1993) or Orton et al. (1986), along with the planet solid angles calculated from the planet flux calculator at the James Clerk Maxwell Telescope Web site.\(^{10} \) For reference, the absolute calibration ranges from approximately 180 nV Jy\(^{-1}\) up to 280 nV Jy\(^{-1}\) over the range of bolometer-operating resistances recorded during our observing run (see Figure 4).

Our flux calibration uncertainty was determined as follows. First, the temperature profiles of Uranus and Neptune were derived by Griffin and Orton using Mars as an absolute calibrator (Griffin & Orton 1993). To determine the surface brightness of Mars at millimeter wavelengths, Griffin & Orton used the model developed by Wright based on observations made at far-infrared wavelengths (Wright 1976), along with the logarithmic interpolation to longer wavelengths described by Griffin et al. (1986). The estimated uncertainty on this interpolated model is

\(^{10}\) http://www.jach.hawaii.edu/jacbin/planetflux.pl
Flux calibration for one of the six calibration data sets, overlaid with a linear fit of calibration vs. bolometer-operating resistance measured by the bolometer voltage at the bias frequency. Note that the bolometer voltage at the bias frequency is a monotonic function of the atmospheric opacity and the bolometer responsivity. (A color version of this figure is available in the online journal.)

Second, the uncertainties on the temperature profiles of Uranus and Neptune are estimated to be less than 1.5% relative to Mars (Griffin & Orton 1993). Additionally, the observations of Uranus and Neptune were taken with a precipitable water vapor of 1.5 ± 0.5 mm, which results in a calibration uncertainty of ∼1.4%. Finally, the error inferred by the scatter of our measurements results in calibration uncertainties between 0.6% and 3.0% for each of the data sets. The end result is an overall flux calibration uncertainty of approximately 5.5%, limited by the uncertainty in the temperature of Mars.

### 4.3. Beam Calibration

Since the astronomical signals in our maps are inherently smoothed based on the profile of the Bolocam beams, it is important to understand their shapes. Additionally, our flux calibration is based on observations of point sources, so our maps have units of flux density. However, since in the CMB the SZE signal we are looking for is a surface brightness or temperature, we need to know the area of our beam in solid angle to convert our maps to surface brightness units. Therefore, any error in our determination of the beam area will show up as a surface brightness or a temperature calibration error. To determine the profile of our beam, we used the observations of Uranus and Neptune. These planets are well suited for measuring our 60 arcsec FWHM beams; they have semidiameters of ≃ 1 arcsec, which means they are essentially pointlike and thus will appear in our maps with shapes given by our beam profile.

Based on simulations, we expected all of the beams to have a similar profile. However, we first calculated the beam for each bolometer separately to validate this expectation. There were not enough data from a single planet observation to make a high S/N measurement of the beam for an individual bolometer, so we averaged the data for groups of four bolometers that are close to each other on the focal plane. Nearby bolometers have beams with similar paths through the optics, so they should also have similar profiles. Each bolometer was grouped into four distinct sets, each of which contained four nearby bolometers, and the average profile from these four sets was determined. The measurement uncertainty on these profiles can be quantified by the standard deviation of the areas of the beam profiles, which was approximately 3.1% (see Figure 5). Within our measurement uncertainty, all of the individual bolometer beam profiles were consistent, so a single beam profile can be used to describe every bolometer. To measure this single beam profile, we averaged the data from all of the planet observations for all of the bolometers. The peak-normalized area of this profile is 3970 arcsec², which is the area of a Gaussian beam with an FWHM of 59.2 arcsec. However, the beam profile is not exactly Gaussian, and the measured profile was used for all of our analyses. Since we cannot rule out systematic variations in the beam area from one bolometer to the next at the level of our single bolometer measurement uncertainty, we have conservatively estimated the uncertainty in this beam area measurement to be 3.1%.

### 5. MAP MAKING

#### 5.1. Least-Squares Map-Making Theory

The astronomical signals we seek can be thought of as two-dimensional objects, which can be represented by a map with finite pixelization. For simplicity, this two-dimensional map can be thought of as a vector, \( \vec{m} \). This map is stored in the bolometer
time streams, \( \tilde{d} \), according to
\[
\tilde{d} = \mathbf{p}\tilde{m} + \tilde{n},
\]
where \( \mathbf{p} \) is a matrix containing the pointing information and \( \tilde{n} \) is noise. Note that we represent matrices with a bold symbol and vectors with an arrow. Since \( \tilde{m} \) is what we are fundamentally interested in obtaining, we need to find a solution to Equation (3) that yields the optimum unbiased estimate of \( \tilde{m} \) given \( \tilde{d} \). There are several methods that can be used to estimate \( \tilde{m} \), including the commonly used least squares method described below (Tegmark 1997; Wright 1996).

Solving the least squares problem for Equation (3) requires minimizing
\[
\chi^2 = (\tilde{d} - \mathbf{p}\tilde{m})^T \mathbf{w}(\tilde{d} - \mathbf{p}\tilde{m}),
\]
where \( \mathbf{w} \) is the inverse of the time-stream noise covariance matrix, \( \langle \tilde{m} \tilde{m}^T \rangle^{-1} \). The estimator for \( \tilde{m} \) derived from Equation (4) is
\[
\tilde{m}' = \mathbf{c}\mathbf{p}^T \mathbf{w}\tilde{d},
\]
where \( \mathbf{c} = (\mathbf{p}^T \mathbf{w})^{-1} \) is the map space noise covariance matrix. If the time-stream noise, \( \tilde{n} \), has a white spectrum, then the various terms in Equation (5) are easy to understand because \( \mathbf{w} \) and \( \mathbf{c} \) are both diagonal. The inverse of the time-stream noise variance is \( \mathbf{w} \), and applies the appropriate weight to each sample in the time stream. The data time stream is then binned into a map by \( \mathbf{p}^T \), and \( \mathbf{c} \) corrects for the fact that \( \mathbf{p}^T \) sums all of the data in a single map bin instead of averaging it. The general idea is the same for nonwhite time-stream noise, but \( \mathbf{w} \) will mix time samples and \( \mathbf{c} \) will mix map pixels.

If the time-stream noise is stationary, then the time-stream noise covariance matrix can be diagonalized by applying the Fourier transform operator, \( \mathbf{F} \). In this case, any element of the inverse time-stream noise covariance matrix can be described by
\[
\mathbf{w}(t_1, t_2) = \mathbf{n}(t_1)\mathbf{n}(t_2)^T \rightarrow \mathbf{W}(\Delta t),
\]
where \( t_1 \) and \( t_2 \) are any two time samples separated by \( \Delta t \). The corresponding elements of the Fourier transform of the inverse covariance matrix, \( \mathbf{W} = \mathbf{F}\mathbf{w}\mathbf{F}^{-1} \), can be written as
\[
\mathbf{W}(f_1, f_2) = \mathbf{W}(f_1)\delta_{f_1,f_2},
\]
where \( \delta_{f_1,f_2} \) represents a Kronecker delta and \( f \) is the frequency in Hz.\(^{13}\) The diagonal elements of \( \mathbf{W} \) are equal to \( 1/(\text{PSD} \times \Delta f) \), where PSD is the noise power spectral density and \( \Delta f \) is the frequency resolution of the time stream. The Kronecker delta ensures that all of the off-diagonal elements are equal to zero. Returning to Equation (5), the estimate for \( \tilde{m} \) can be rewritten as
\[
\tilde{m}' = ((\mathbf{p}^T \mathbf{F}^{-1})(\mathbf{F}\mathbf{w}\mathbf{F}^{-1})(\mathbf{F}\mathbf{p}^T))^{-1}((\mathbf{p}^T \mathbf{F}^{-1})(\mathbf{F}\mathbf{w}\mathbf{F}^{-1})(\mathbf{F}\tilde{d})),
\]
using the fact that \( \mathbf{F}^{-1}\mathbf{F} = 1 \). Finally, taking the Fourier transform of the various terms in Equation (6) yields
\[
\tilde{m}' = (\mathbf{p}^T \mathbf{w} \mathbf{p})^{-1}\mathbf{p}^T \mathbf{W}\tilde{D}
\]
as an alternate expression to estimate the value of \( \tilde{m} \), where \( \mathbf{p} = \mathbf{F}\mathbf{p}, \tilde{D} = \mathbf{F}\tilde{d}, \) and \( \mathbf{W} = \mathbf{F}\mathbf{w}\mathbf{F}^{-1} \). Note that \( \mathbf{c} = (\mathbf{p}^T \mathbf{w})^{-1} = (\mathbf{p}^T \mathbf{W})^{-1} \) does not in general simplify as a result of Fourier transforming.

5.2. The Bolocam Algorithm: Theory

The science field maps produced by Bolocam each contain \( n_p \approx 20000 \) pixels, and an extremely large matrix must be inverted to calculate \( \mathbf{c} \) since \( \mathbf{p}^T \mathbf{w} \mathbf{p} \) has dimensions of \( n_p \times n_p \). Direct inversion of such a matrix is possible, but is not practical on a typical high-end desktop computer. The map could be determined on a desktop computer via a conjugate gradient solver, but determining the covariance matrix would require a significant amount of simulation power. Therefore, we developed an algorithm to approximate \( \tilde{m} \) by exploiting the simplicity of our scan pattern, which involved raster scanning parallel to either the right ascension or declination axis. This approximation allows us to make maps in a relatively short amount of time using a standard desktop computer, which is extremely convenient.

To illustrate this simplification, consider the map made from a single bolometer for a single scan within an observation. This scan will produce a one-dimensional map at a single declination value (for a right ascension scan) or a single right ascension value (for a declination scan). Each data point in the time stream is separated by 24 arcsec in map space since our data are sampled at 10 Hz and the telescope scans at 240 arcsec s\(^{-1} \). Therefore, our data are approximately Nyquist sampled for Bolocam’s \( \approx 60 \) arcsec FWHM beams. The maps are binned with 20 arcsec pixels (1/3 of the beam FWHM, and is slightly finer than Nyquist sampled), so \( \mathbf{p}^T \) will map either 1 or zero time-stream samples to each map pixel. Note that \( n_s \), the number of time-stream samples, will be slightly less than \( n_p \), the number of map-space pixels. Since \( \mathbf{p}^T \) has dimensions of \( n_s \times n_p \), the sum of each row in \( \mathbf{p}^T \) is either 1 or zero and the sum of each column is 1. Consequently, we will make the approximation that \( \mathbf{p}^T = 1 \).

From Equation (5), this means that
\[
\mathbf{c} = \mathbf{w}^{-1},
\]
and therefore \( \tilde{m} = \mathbf{W}^{-1} \) for a single scan of time-stream data. If we Fourier transform Equation (7), then we find that
\[
\mathbf{C} = \mathbf{W}^{-1}.
\]
Since \( \mathbf{W} \) is diagonal, the inversion is trivial, and the result is that the Fourier transform of the map-space noise covariance matrix is diagonal with elements equal to the time-stream PSD\( \times \Delta f \).

The next step is to consider a map made from a single bolometer for a full observation, which contains 20 scans. We move the telescope in the orthogonal direction to the scan between scans by more than the size of a single map pixel, so we can still approximate \( \mathbf{p}^T \approx 1 \). There are almost no correlations between scans because the atmospheric-noise subtraction coefficients are calculated scan by scan along with subtraction of the mean signal level. The covariance of maps made for a single observation from alternate scans is negligible, supporting this assumption that individual scans are uncorrelated. Therefore, the time-stream data and map-space data for different scans are essentially independent.\(^{14}\) Consequently, the noise in map space will be stationary, which means that the noise covariance matrix can be diagonalized.

\(^{13}\) Note that physical space values are denoted with a lower case letter, and the corresponding frequency space values are denoted with an upper-case letter.

\(^{14}\) To verify that the data from different scans are independent, we created maps for each observation from all the odd-numbered (right-going) scans and from all the even-numbered (left-going) scans. The cross-PSDs of the right-going maps with the left-going maps were consistent with noise, indicating that the data from separate scans are independent.
by Fourier transforming it. The Fourier transform of the full-map noise covariance matrix, $C$, can be visualized by noting that each diagonal element corresponds to a single Fourier-space map pixel (or equivalently, a single Fourier-space time-stream sample). So, this visualization of $C$ will be equal to the single scan time-stream $\text{PSD} \Delta f$ for rows of map-space pixels that are parallel to the scan direction, and will have a white spectrum for columns of map-space pixels that are perpendicular to the scan direction. Alternatively, since there is a one-to-one correspondence between time-stream samples and map-space pixels, this visualization of the diagonal elements of $C$ is equal to the full map-space $\text{PSD} \Delta f \Omega$, where $\Delta f \Omega$ is the angular frequency resolution of the map.

At this point, we need to add together all of the individual observations to make a single map. Since we have shown that the map-space data are equivalent to the time-stream data for a single observation, the easiest way to co-add data from separate observations is to use the single observation maps. Since the noise in separate observations is uncorrelated, the maps can be co-added according to

$$\tilde{m} = \left( \sum_i c_i^{-1} \right)^{-1} \sum_j c_j^{-1} \tilde{m}_j,$$

where the subscripts $i$ and $j$ refer to the observation number. The easiest way to evaluate Equation (8) is to Fourier transform it so that the noise covariance matrices are all diagonal. The result is

$$\tilde{M} = \left( \sum_i c_i^{-1} \right)^{-1} \sum_j c_j^{-1} \tilde{M}_j,$$

where $\tilde{M}$ is the Fourier transform of the map and $C$ is the Fourier transform of the noise covariance matrix, with diagonal elements equal to the $\text{PSD} \Delta f \Omega$ of the map. Since all of the $C$s are diagonal, we can simplify Equation (9) to

$$M = \left( \sum_i \frac{1}{P_i} \right)^{-1} \sum_j \frac{M_j}{P_j},$$

where $M$ is the two-dimensional Fourier transform of the map and $P$ is the two-dimensional PSD of the noise in the map. At this point, we have dropped the vector and matrix notation since $M_i$ and $P_i$ have the same dimensions. Note that $\Delta f \Omega$ is the same for every map, so the constant factor of $\Delta f \Omega$ from the first sum in Equation (10) cancels the factor of $1/\Delta f \Omega$ from the second sum in Equation (10).

Finally, to make a map using all of our data, we need to consider every bolometer, not just a single detector. To properly weight the data from each bolometer prior to co-adding, we calculate the expected variance, $(\sigma_p)^2$, in measuring the peak flux of a pointlike source from a single scan through the center of the source for bolometer $i$. This variance is calculated using the scan-averaged time-stream PSD for each bolometer, $\text{PSD}(f)$, and the Fourier transform of the expected signal shape of a pointlike astronomical signal, $S(f)$, according to

$$(\sigma_p)^2 = \left( \int df \frac{S(f)^2}{\text{PSD}(f)} \right)^{-1},$$

where $f$ is temporal frequency. Note that $S(f)$ is the beam profile, not a delta function. Then, the data from each bolometer are weighted by a factor proportional to $1/(\sigma_p^2)$ prior to co-adding it with data from other bolometers. This is the optimal way to co-add the data for pointlike signals; it is nearly optimal for signals of any shape if the PSDs have similar profiles for every bolometer, which is largely true for our bolometer signals since they are dominated by atmospheric noise.

However, due to atmospheric noise, along with our noise-removal algorithms, there are correlations between the bolometers, but most of these correlations are instantaneous in time and constant over the observation. Additionally, the relative positions of the bolometers do not change during the observation, so the map-space separation of the correlations does not change. Therefore, the correlations are stationary in time with separations that are fixed in the map space, so the correlations are an additional time-independent covariance between map pixels that are sampled at the same time by different bolometers. This additional covariance is approximately stationary over the entire map, except where it breaks down near the edges because part of the focal plane is outside the map region. Since this additional covariance between map pixels is approximately stationary in space, its contribution to $W^{-1}$ will be diagonal. Since $W^{-1}$ is still diagonal, co-addition of the maps of individual observations can proceed according to Equation (10). Therefore, Equation (10) can be used as the algorithm to produce our final science field maps.

Note that we were forced to make several simplifying assumptions in order to develop Equation (10). We have assumed that the pointing matrix, $p_\Omega^i$, is equal to 1. We have also assumed that the noise in our time streams is stationary for each 8 min long observation. Additionally, we have assumed that the PSD of the correlations between bolometers is white, and that all of the correlations are time instantaneous. Finally, we have assumed that the map coverage (i.e., the number of time-stream samples that are binned in each map-space pixel) is uniform, so that the Fourier transform of the map is a valid description of the time-stream data. Deviations from these assumptions will alter the map estimate we compute from the optimal least-squares map estimate. But, these deviations only affect how each time-stream sample is weighted before it is mapped. This means our final map will have more noise than an optimal map, but it will not be biased in any way. In other words, since the map-making operation is linear, the resulting map will be unbiased no matter what weightings are used to co-add the data, as long as the weights are properly normalized. We have confirmed this lack of bias via simulation, as we discuss below.

5.3. The Bolcam Algorithm: Implementation

To start, we must first produce a map from the time-stream data for each 8 minute observation. As mentioned in Section 5.2, this is done by calculating the variance in measuring the peak flux of a pointlike source under the assumption that the profile of the time-stream PSD is similar for every bolometer. To determine this variance, we calculate the PSD for each bolometer for each scan. These spectra are then averaged over all 20 scans for each bolometer, thereby making the assumption that the noise properties do not change over the course of the observation. Then, we determine the expected shape of a pointlike source using our measured beam profile and scan

15 We have been able to find a small amount of correlated atmospheric signal that is not time-instantaneous. However, the time lag of these correlations is generally much less than one time sample, which means they will also be less than one map pixel.
are multiplied by the statistical properties of the noise will be unchanged when half of the data then the distribution of signals it will produce is symmetric about 0. Therefore, bolometers within a single scan are multiplied by −1. To zero at the edges of the map (see Figure 6). To obtain sufficient uniform coverage, we restrict our map to have sides of 42 arcmin; the fractional rms variations in coverage within this region for a single 8 minute observation are only about 8%–9%. Since the coverage variations are minimal, we will assume that this square central region has uniform coverage, and therefore uniform noise properties. This assumption of uniform coverage allows us to directly compute the Fourier transform and noise properties of the map. We emphasize that, even if the assumption of uniform coverage fails and our algorithm is nonoptimal, it is never biased because Equation (10) is linear in the map-space maps.

We now have a uniform coverage map for each observation, which can easily be Fourier transformed to produce the $M_j$s needed in Equation (10). But, we still need to determine the two-dimensional PSD of each single observation map. Due to residual correlations between bolometers, we do not understand the noise properties of our data well enough to determine the map PSD from simulation, so we instead estimate the PSD by generating a large number of jackknifed maps from our real data. In each jackknifed map, a different subset of the time streams from half of the scans within each observation was multiplied by −1. Note that the data from all of the bolometers within a single scan are multiplied by −1, so the residual atmospheric noise that is correlated between bolometers is preserved. This multiplication leaves the noise properties of the map unchanged, while allowing us to produce a large number of noise realizations for each map. Note that the residual atmospheric noise correlations are time instantaneous, so they remain in the jackknifed realizations. We then generate 100 realizations for each observation, and we set the true PSD for each observation equal to the average of the map-space PSD computed for each realization. See Equation (13). Examples of the PSDs we calculated are given in Figure 7. This method of determining the map-space PSDs assumes that the time-stream data for each scan is uncorrelated with the data from all other scans, which we argued in Section 5.2.

To determine the validity of the map-space PSDs we estimated from the jackknifed map realizations, we examined the distribution of PSD values for each realization. If the noise properties of the data are Gaussian, as we have assumed, then the PSD measured at any given Fourier map-space pixel will be drawn from

$$f(X_{i,\tilde{v}}) = (1/P_\tilde{v})e^{-(X_{i,\tilde{v}}/P_\tilde{v})},$$

where $X_{i,\tilde{v}}$ is the measured PSD for realization $i$ at pixel $\tilde{v}$, $P_\tilde{v}$ is the true PSD for pixel $\tilde{v}$, and $f(X_{i,\tilde{v}})$ is the probability density function of $X_{i,\tilde{v}}$. Note that $\tilde{v}$ has units of spatial frequency (i.e., rad$^{-1}$), and describes a pixel in the spatial Fourier transform of the map. See the Appendix for a derivation of $f(X_{i,\tilde{v}})$. The true PSD is estimated from

$$\hat{P}_\tilde{v} = \frac{1}{N_r} \sum_{i=1}^{N_r} X_{i,\tilde{v}},$$

where $N_r = 100$ is the number of realizations. To compare our measured PSDs to the probability density function (PDF) given in Equation (12), we created the dimensionless value

$$Y_{i,\tilde{v}} = \frac{X_{i,\tilde{v}}}{\hat{P}_\tilde{v}},$$

with the associated PDF

$$f(Y_{i,\tilde{v}}) = e^{-Y_{i,\tilde{v}}}. \tag{15}$$

Then, we compared our measured values of $Y_{i,\tilde{v}}$ to the PDF in Equation (15). In general, we found that our measured $Y_{i,\tilde{v}}$ follow a distribution extremely close to $f'(Y_{i,\tilde{v}})$, except that the number of $Y_{i,\tilde{v}}$ with values near zero is slightly less than expected. Therefore, the map-space PSDs estimated from the jackknife realizations should be a good estimate of the true map-space PSDs.

Let us consider the possible effects of imperfect signal removal in the single observation jackknife maps. Because we only use the single observation noise estimates as weights for co-adding, the result of residual signal in the jackknife maps will be co-addition weights that are nonoptimal. This nonoptimality may degrade the noise of the final maps, but will not cause them to be biased. We may estimate the size of the signal leakage to determine how large the deviation from optimality could be. Our final flat-band power anisotropy upper limit is approximately $1000 \mu K^2_{CMB}$. With our effective $\Delta \ln(\epsilon)$ of 0.63 (derived in Section 6), this upper limit corresponds to an excess variance in our final maps of $\simeq 500 \mu K^2_{CMB}$. These final maps have a variance of $10,000 \mu K^2_{CMB}$ (see Section 8 where we find that the map rms per beam is approximately $90 \mu K_{CMB}$). Given that $\simeq 500$ observations contribute to each map, the single-observation map variance is $\simeq 5 \times 10^6 \mu K^2_{CMB}$, or approximately $10,000$ times larger than our upper limit on the astronomical signal contribution. Even if we did not remove the astronomical signal using the jackknifing procedure, it would affect the single-observation PSDs, and therefore the weights, at only the 0.01% level. Using jackknife-generated PSDs reduces the effect of signal contamination further. Therefore, the effect of signal leakage into the single observation jackknife maps is negligible.

6. TRANSFER FUNCTIONS

The transfer function describes the fraction of the astronomical signal that remains after processing as a function of map-space Fourier mode. In order to determine the transfer function of our data-processing algorithms, we first generate a simulated map of the expected astronomical signal. This map is then reverse mapped into a time stream using the pointing information...
in a real observation. Next, this simulated time stream is added to the real bolometer time streams from the observation, and then processed and mapped in the standard way. A map made from data that did not have a simulated signal added to it is then subtracted from this map, producing a map with the simulated signal after processing. A map made from data that did not have a simulated signal added to it is then subtracted from this map, producing a map with the simulated signal after processing. Finally, this map is divided by the PSD of the original simulated signal map to determine how much of the signal remains. Note that we are computing the transfer function for a PSD because we are interested in measuring an excess noise and not a specific signal shape, which means we do not need the phase of the transfer function.

This transfer function was computed for 20 randomly selected observations, ten taken while scanning parallel to right ascension and ten taken while scanning parallel to declination. Realizations of the expected flat-band power anisotropy signal were used as the simulated signal. These realizations were generated in Fourier map-space assuming Gaussian fluctuations and a flat-band power in \( C_\ell \approx C_\ell (\ell + 1)/2\pi \) of 50 \( \mu K^2 \). Note that although a flat-band power of 50 \( \mu K^2 \) was used for the simulated signal maps, we found that the transfer function is independent of the amplitude of the flat-band power anisotropy signal. For each observation, we averaged the transfer function obtained from 100 different signal realizations to determine the average transfer function. We then compared the average transfer function for each of the ten observations taken with a similar scan pattern. The result is that the transfer functions were the same within our measurement uncertainty for all of the observations. Therefore, we averaged the transfer function from all ten observations to produce a high S/N measurement for each atmospheric-noise-removal method: average, planar, and quadratic (see Figure 8).

Since all of the data processing is performed on the time streams, the attenuation caused by the processing has a preferred orientation based on the scan strategy. The result is that the transfer function is not azimuthally symmetric because of the large amount of attenuation at low frequencies parallel to the scan direction due to atmospheric noise removal. Additionally, there is massive attenuation on scales larger than the Bolocam focal plane (\( \approx 500 \text{ rad}^{-1} \)) because of the atmospheric noise-removal algorithms. This occurs because these algorithms are designed
to remove all time-instantaneous signals at each data sample, which is equivalent to subtracting any signals that vary slowly compared to the size of the focal plane.

In addition to the signal attenuation caused by the data processing, the Bolocam system also attenuates some of the astronomical signal. By scanning across the sky, we are effectively convolving any signal with the profile of a Bolocam beam; since the beams have a nonzero width, this convolution will act like a low-pass filter on all of the astronomical signals. This filter will be approximately symmetric because the Bolocam beam profiles have a high degree of rotational symmetry. Additionally, since the beams are nearly Gaussian, the filter will be approximately Gaussian with an HWHM in variance of about 1000 rad\(^{-1}\) (which is equivalent to an HWHM\(_{\nu}\) \(\simeq 6000\) in angular multipole space) (see Figure 8).

In order to quantify the amount of signal attenuation by each atmospheric noise-removal algorithm, it is useful to determine the effective bandwidth (BW\(_{\text{eff}}\)) of the transfer function. The BW\(_{\text{eff}}\) describes the range of angular multipoles to which we are sensitive, as quantified by the transfer function, and can be used to convert an angular power, \(C_\ell\), to a map-space variance in \(\mu K^2_{\text{CMB}}\). In general, the BW\(_{\text{eff}}\) is calculated by integrating the transfer function over all angular multipoles. However, since the expected SZE power spectrum is approximately flat in \(C_\ell\), which results in a spectrum in \(T_\nu\) that falls like \(1/\ell(\ell + 1)\), it is more useful to weight the transfer function by a factor of \(1/\ell(\ell + 1)\). This weighting will produce an effective logarithmic, rather than linear, bandwidth approximately equal to

\[
\text{BW}_{\text{eff}} \propto \ln(\ell_{\text{max}}) - \ln(\ell_{\text{min}}) = \Delta \ln(\ell).
\]

Although the Bolocam transfer functions are not azimuthally symmetric, it is still useful to determine the effective \(\Delta \ln(\ell)\) for each of the atmospheric noise-removal algorithms, with \(\Delta \ln(\ell) = 0.98, 0.58, \text{ and } 0.37\) for average, planar, and quadratic subtraction. Note that our final map, which consists of observations processed with different atmospheric noise-removal algorithms as described in Section 7, has a bandwidth of \(\Delta \ln(\ell) \simeq 0.63\).

7. OPTIMAL ATMOSPHERIC NOISE SUBTRACTION

Each of the science field observations were processed with average, planar, and quadratic sky subtraction, creating three separate files for each observation. Quadratic subtraction removes the most atmospheric noise, while average subtraction retains the most astronomical signal, so there is an optimal sky subtraction algorithm for each observation based on the type of astronomical signal we are looking for. To determine which algorithm is optimal, we computed a FOM for each subtraction method. Since the anisotropy signal appears as a variance in the map, the variance on the amplitude of the anisotropy signal will be proportional to the square of the map PSD divided by the transfer function of the experiment. This can be seen in Equations (A7) and (A8). Therefore, the FOM is defined as the inverse of this variance on the anisotropy signal summed over all angular scales according to

\[
\text{FOM} = \sum_\nu \frac{(S^2_\nu)^2 T^2_\nu (B^2_\nu)^2}{P^2_\nu},
\]

where \(\nu\) is a two-dimensional spatial frequency with units of rad\(^{-1}\), \(S^2_\nu\) is the expected anisotropy power spectrum, \(T_\nu\) is the transfer function of the data processing in squared units, and \(B_\nu\) is the profile of the Bolocam beam. Since the expected anisotropy signal has a flat-band power in \(C_\ell\),

\[
S^2_\nu \propto \frac{1}{\ell(\ell + 1)}
\]

for \(\ell = 2\pi |\nu|\).\(^{17}\) Assuming this spectrum for \(S^2_\nu\), a top-hat window between \(\ell = \ell_{\text{min}}\) and \(\ell = \ell_{\text{max}}\) will produce a bandwidth approximately equal to

\[
\text{BW}_{\text{eff}} \propto \ln(\ell_{\text{max}}) - \ln(\ell_{\text{min}}) = \Delta \ln(\ell).
\]

\(^{17}\) We have used the small-scale flat sky approximation, \(\ell = 2\pi |\nu|\).
the profile of the Bolocam beam, and $P_i$ is the PSD of the noise in the map in squared units. Note that we have included the \( \approx 5 \) arcsec uncertainty in our pointing model in $B_i$, and this pointing uncertainty effectively broadens the beam. To be precise,

$$B_i = \mathbf{B} e^{-|\nu|^2/2\sigma_p^2},$$

where $\mathbf{B}$ is the measured beam profile, and $\sigma_p = 1/2\pi \sigma_p$ for a pointing uncertainty of $\sigma_p$. For the anisotropy spectrum, we assumed a flat-band power in $C_\ell$, so

$$S^2 = \frac{1}{\ell(\ell+1)}$$

for $\ell = 2\pi |\nu|$. The FOM is inversely proportional to the variance on an estimate of the anisotropy amplitude (in $\mu K^2_{\text{CMB}}$), so it characterizes the $S/N$ of the map.

In the end, average subtraction was the optimal method for just over 50% of the observations, planar subtraction was the optimal method for just over 40% of the observations, and quadratic subtraction was the optimal method for just under 10% of the observations. We can calculate how much the observations optimally cleaned by each method contribute to our final $S/N$ from

$$S/N = \sqrt{\frac{\sum_i FOM_i}{\sum_i FOM_i^2}},$$

where $T$ denotes the set of observations optimally cleaned by a given method and $FOM_i$ is the FOM from Equation (16). The $S/N$ contributed by the average/planar/quadraic observations is 70%/29%/1%. These ratios are different from the number of observations optimally cleaned by each method because the amount of atmospheric noise in the data generally determines which subtraction algorithm is optimal, and the observations optimally cleaned with average subtraction were made in the best observing conditions. Note that quadratic subtraction is the optimal method only when the weather conditions are extremely poor. This is because the anisotropy power spectrum falls quickly at high frequency, and the quadratic subtraction algorithm attenuates a large amount of signal at low frequency. For pointlike sources, whose spectra are flatter, quadratic subtraction is the optimal processing method slightly more often.

8. FINAL MAP PROPERTIES

Once the FOM is determined for each subtraction method for each observation, we can then produce a map of all of the data using the optimally processed map for each observation. To produce this final map, we need to make a slight modification to Equation (10) to account for the transfer function of the data processing and the Bolocam beam. We need to account for these effects because the transfer function depends on the scan direction and optimal sky subtraction algorithm for each observation. Therefore, the amount of astronomical signal in the map is in general different for each observation. To account for the amount of signal attenuation in each observation, the map PSD needs to be divided by the transfer function and the Fourier transform of the map needs to be divided by the square root of the transfer function. After making these modifications to Equation (10), we have

$$M' = \sqrt{\mathcal{R}} M = \frac{1}{\sqrt{\mathcal{R}}} \sum_i \left( \frac{M_i}{\sqrt{T_i B_i^2}} \right) \left( \frac{B_i^2}{F_i} \right),$$  \hspace{1cm} (18)

for

$$\sqrt{\mathcal{R}} = \frac{\sum_i \sqrt{T_i B_i^2}}{\sum_j \left( \frac{B_j^2}{F_j} \right)}.$$

Although $M'$ will be biased (i.e., it is not the Fourier transform of the true map of the sky), this bias is accounted for by the final transfer function we calculate in Section 8.2.\textsuperscript{19} Note that $M'$ can be Fourier transformed back to map space to produce a map $m'$, although $m'$ will be biased. The maps, $m'$, for each science field are given in Figure 9.

8.1. Noise PSDs

Analogous to the case of a single observation, we used jackknifed realizations of our data to estimate the noise PSD of $m'$. In this case, each realization is generated by multiplying a randomly selected set of half the observations in $m$ by $-1$. The map-space PSD from 1000 realizations were averaged to determine the best estimate of the noise PSD for each science field, with the results shown in Figure 10. In this section, we establish that the noise PSD estimated in this way is statistically well behaved (Gaussian) and unbiased. These characteristics are critical to the remainder of our analysis.

We analyzed the distribution of individual realization PSDs to determine if the following probability distribution describing the noise is Gaussian. As in the single observation case, we computed a dimensionless PSD value according to Equation (14), and compared the distribution of these values to the PDF given in Equation (12). In general, the agreement is good, indicating that the underlying noise distribution is well approximated by a Gaussian (see Figure 11). The Gaussianity of the noise PSDs of the jackknife maps is important because it justifies the form

\textsuperscript{18} We have not included any phase information in the factor $\sqrt{T_i B_i^2}$ because both the signal and the noise PSD contain only noise; the phase is irrelevant.

\textsuperscript{19} In Equations (18) and (19), $T_i B_i^2/F_i$ acts as a weighting factor for each observation. Therefore, $M'$ represents the weighted mean of the Fourier transform of each single observation map divided by the square root of the transfer function for that map, $(M_i/\sqrt{T_i B_i^2})$. Similarly, $\sqrt{\mathcal{R}}$ represents the weighted mean of the square root of the transfer function for each observation, $\sqrt{T_i B_i^2}$. So, $M' = (\sqrt{T B^2})^{-1}(M/\sqrt{T B^2})$, which reduces to the weighted mean of all the single observation map Fourier transforms, $M' \geq M$, in the limit that all of the single observation transfer functions, $T_i B_i^2$, are the same.
of the likelihood function we use, Equation (A5) (presented in Section 10.1). That form assumes the Fourier coefficients of the final map are Gaussian-distributed random variables with variance given by the noise PSD estimated from the jackknife maps.

Next, we show that the noise PSD estimated from the jackknives is unbiased under two assumptions: (1) the covariance of any pair of distinct observations vanishes on average; and (2) negligible signal leaks into the jackknife maps. The first assumption is equivalent to the statement that there is no scan-synchronous or fixed-pattern noise in the maps. We have checked this assumption empirically and find that the average fractional covariance of distinct observations is \( \sim 2 \times 10^{-5} \), which is consistent with noise. We will discuss below how our nondetection of signal in the final map further justifies this assumption. With these assumptions, we can prove lack of bias of the noise estimate in a straightforward fashion by a simulation that obeys the assumptions. We generate a set of \( N_{\text{obs}} = 515 \) single observation maps using the single observation noise PSDs; these obviously have no signal and are uncorrelated with one another. Then, we construct \( N_{\text{jack}} = 1000 \) simulated jackknife final maps and calculate the noise PSD of these maps, which also obviously have no signal. Next, we average these noise PSD estimates over all of the jackknife final maps, and divide by the input noise spectrum to determine how accurately we have recovered that input spectrum. The average (over all Fourier space pixels) of this normalized PSD is \( 0.9997 \pm 0.0006 \), showing that we indeed recover the input noise spectrum within the measurement uncertainty of our simulation. This exercise thus shows that the simulated final map noise PSD estimated by jackknife maps is an unbiased estimate of the simulated final map noise PSD. Note that we do not claim that the noise PSD generated in this fashion is the noise PSD of our true final map; the simulation is intended only to show that the jackknife noise PSD estimate method is unbiased.

Let us now justify the assumption that negligible signal leaks into the jackknife final maps. In generating jackknife final maps, negative signs were applied to exactly one half of the observations. The subset of observations multiplied by \( -1 \) varies from one jackknife realization to the next, but the total number of observations multiplied by \( -1 \) is always the same. Therefore, residual signal can arise in the jackknife final maps in only two
ways: (1) if the relative calibrations of the different observations are imperfectly known or (2) if the weights of the different observations are unequal.

In the first case, consider a single observation fractional relative calibration error of $\psi$, but assume all the component observations would otherwise be weighted equally (i.e., no variation in noise between observations). This fluctuating relative calibration error does not cause a bias; in an ensemble of experiments, the final map has an expected signal value equal to the signal value of the true final map and the jackknife final map has an expected signal value of exactly zero. But, the calibration fluctuations will cause an imperfect co-addition or cancellation of the signal in any given final map or jackknife final map realization, which will produce a fractional spread in the signal level of $\psi/\sqrt{N_{\text{obs}}}$ relative to the true signal. Given that $\psi \lesssim 3\%^{20}$ and $N_{\text{obs}} \approx 500$ observations per map, this error in both the final map and jackknife final maps is very small compared to the signal. Since we are not attempting a high S/N measurement, the vanishingly small size of the error relative to the signal is thus not a concern in the final map. The error affects the jackknives in a more subtle way because it effectively adds noise to the jackknife final maps, which means our noise estimate is slightly higher than the true noise level of our final map. However, this bias is negligible; the magnitude of the error is of order the signal times $\psi/\sqrt{N_{\text{obs}}}$. We know $\psi \lesssim 3\%$, $N_{\text{obs}} \approx 500$, and the signal is less than 2% of the noise in the final maps (in rms units, see Section 5.3), so this bias is less than 0.003% of the final map noise level in rms units, or $< \cdot 0.006\%$ in variance units. The small size of the effect is not surprising: it is proportional to the signal size, and we have no detection of signal. Moreover, even if the effect were not negligible, it would result in an overestimate of the noise PSD and thus would result in an overly conservative upper limit. This kind of effect would only be problematic if signal were visible at high significance.

In the second case, the argument is very similar, but now what matters is the fractional variation in observation weight. This fractional variation is large, $\lesssim 2$, due to the significant differences in atmospheric noise between observations. Here, the large number of observations and the fact that the spread is proportional to the signal size render the effect negligible. The rms spread of the residual signal in the jackknives will be $\lesssim 2/\sqrt{N_{\text{obs}}} = 0.09$ times the signal size. Again, because of the small size of the error relative to the signal and the lack of detected signal, the error due to this effect is insignificant.

If the noise estimation approach has underestimated the noise (for example, by failing to account for nonstationarity or correlations, or in any other manner) then there will be more noise in the final map than expected from the jackknives. However, because we find in Section 10.5 that our 90% CL interval on the amplitude of astronomical anisotropy, $A$, includes $A = 0$, we do not see any significant excess of noise in the true map above what is expected from the noise PSD estimate. This explicitly rules out scan-synchronous or fixed-pattern noise that would be averaged away in jackknife maps but would remain in the co-added map at the level of interest for this analysis. Had there been an excess above the expectation from the noise estimate, we would have had to show more explicitly that the noise estimate was correct in order to claim a detection.

20 In Section 4.2 we calculated our flux calibration uncertainty to be approximately 5.5%. However, most of this uncertainty is due to systematics that will not change from one observation to the next. The uncertainty caused by fluctuations in the atmospheric opacity and the fit of our model are $\lesssim 2\%–3\%$.
The amount of galactic dust emission can be estimated from maps of our science fields taken from the full-sky 100 μm DIRBE/IRAS dust map (DIRBE Website; Schlegel et al. 1998). To extrapolate the 100 μm data to our band at 143 GHz ≃ 2.1 mm, we have used the “model 8” extrapolation given in Finkbeiner et al. (1999). At 100 μm, the typical surface brightness of the dust emission in our science fields is just over 1 MJy/ster, which corresponds to a surface brightness of around 5–15 nK_CMB for Bolocam. Using the maps that have been converted to a thermodynamic temperature at 143 GHz, we determined the map-space PSD of the dust emission, which corresponds to a C_ℓ less than 10^{-6} μK_CMB^2 for ℓ ≳ 1000. Since this is well below the expected SZE-induced CMB anisotropy we are looking for, it is safe to conclude that the signal from the dust emission in our maps is negligible.

Emission from radio point sources will also contribute to the astronomical signal in our maps. The power spectrum from these sources can be calculated from

$$C_\ell = \int_0^{S_{\text{cut}}} S^2 N(S) dS + w_\ell f^2, \quad (20)$$

where S is the flux of the source, N(S) is the differential number of sources at a given flux in a given solid angle, S_{cut} is an estimate of the source-detection threshold in the map (i.e., the level at which sources may be detected and removed), C_ℓ is the angular power spectrum, w_ℓ is the Legendre transform of the two-point correlation function of the sources, and

$$I = \int_0^{S_{\text{cut}}} SN(S)dS$$

is the background contributed by the sources (White & Majumdar 2004; Scott & White 1999). We will assume w_ℓ = 0, since there is a large amount of uncertainty in the clustering of these sources. Differential number counts have been determined from measurements at 1.4, 5, and 8.44 GHz (Toffolatti et al. 1998; Danese et al. 1987), with

$$N(S)_{\text{5GHz}} = 150 S^{-2.5} \text{ Jy}^{-1} \text{ster}^{-1}. \quad (21)$$

Since the spectrum of the sources is nearly flat (i.e., S_\ell \propto \nu^{\beta}$ with $\beta = 0$), this equation is valid over a wide range of frequencies. Additionally, the WMAP K, Ka, and Q bands have been used to determine the differential number counts at 22, 30, and 40 GHz (Bennett et al. 2003). N(S) is similar for all three WMAP bands, and is ≲ 70% of the value of the model in Equation (21). The differential number counts at 40 GHz are described by

$$N(S)_{\text{40GHz}} = 32 S^{-2.7} \text{ Jy}^{-1} \text{ster}^{-1}. \quad (22)$$

To extrapolate this equation to the Bolocam band center at 143 GHz, we will use the method described in White & Majumdar (2004). Since there is evidence of the power law for N(S) flattening at higher frequencies, they describe the differential number counts according to

$$N(S)_{\text{143GHz}} = (20 - 32) S^{-2.3} \text{ Jy}^{-1} \text{ster}^{-1}. \quad (23)$$

Note that the total number of sources and total integrated power in ℓ-space will not change if w_ℓ is nonzero; the clustering modeled by w_ℓ will only shift power from high-ℓ to low-ℓ.

22 There is some uncertainty in the spectrum of S_\ell for these radio sources between 40 GHz and 143 GHz. White and Majumdar quote two spectra, one with $\beta = 0$, and one with $\beta = -0.3$. This uncertainty in the spectrum of the radio point sources results in a finite range for the normalisation of the number counts after extrapolating to 143 GHz.

21 Note that the resolution of the DIRBE/IRAS dust map is 6.1 arcmin, which corresponds to HWHM in ℓ-space of ≲ 2000. Therefore, we have no direct knowledge of the power spectrum on scales smaller than ≳ 6 arcmin, which are the angular scales Bolocam is most sensitive to. However, the power spectrum of the dust falls rapidly at small angular scales, so the estimate at ℓ < 2000 should provide a reasonable upper limit.
We also need to estimate $S_{\text{cut}}$ in order to evaluate the power spectrum in Equation (20). This cutoff flux will necessarily be somewhat arbitrary, but, since $C_\ell$ is only weakly dependent on $S_{\text{cut}}$, it will not significantly alter our result. We have chosen $S_{\text{cut}} = 10 \text{ mJy}$, which is approximately four times the rms fluctuations per beam in maps made from data that have been optimally filtered for point sources.\textsuperscript{24} Inserting this value of $S_{\text{cut}}$ into Equation (20), along with Equation (22), yields $C_\ell \simeq 1.1-1.9 \text{ Jy}^2 \text{ ster}^{-1}$, or $C_\ell \simeq 7-12 \times 10^{-6} \mu K^2_{\text{CMB}}$. To compare this angular power spectrum to the expected SZE-induced CMB anisotropies, we determine the amplitude of a flat-band power, $C_\ell^\text{eff}$, that is required to cause the same temperature fluctuation as $C_\ell$ given our transfer function, $T_\ell B_\ell^2$, according to

$$C_\ell^\text{eff} = \sum_\ell \frac{\nu^2}{C_{\ell}^2} T_\ell B_\ell^2 / \sum_\ell \frac{\nu^2}{C_{\ell}} T_\ell B_\ell^2. \quad (23)$$

For the radio point sources with $C_\ell = 7-12 \times 10^{-6} \mu K^2_{\text{CMB}}$, the effective $C_\ell$ given the Bolocam transfer function is $C_{\ell}^\text{eff} \simeq 35-60 \mu K^2_{\text{CMB}}$, which is comparable to the expected signal from the SZE-induced CMB anisotropies.

Additionally, emission from dusty submillimeter galaxies will be present in our maps. The same method used to determine the power spectrum from radio point sources can also be used to estimate the power spectrum of these sources. We used the number counts distribution determined by J. E. Aguirre et al. (2008, in preparation) with

$$N(S)_{350\text{GHz}} = 1619 S^{-2.26} e^{-3035 S} \text{ Jy}^{-1} \text{ ster}^{-1}. \quad (23)$$

The spectrum of these objects can be described by $S_\nu \propto \nu^\beta$, where $2.5 \lesssim \beta \lesssim 3.5$ (Borys et al. 2003), which gives a differential number count at 143 GHz of

$$N(S)_{143\text{GHz}} = (100 - 220) S^{-2.26} e^{-(2730-1460)S} \text{ Jy}^{-1} \text{ ster}^{-1}. \quad (23)$$

Inserting the above formula into Equation (20) gives $C_\ell = 0.4-1.2 \text{ Jy}^2 \text{ ster}^{-1}$, or $C_\ell = 3-9 \times 10^{-6} \mu K^2_{\text{CMB}}$.

Equation (23) can again be used to convert this to an effective constant $C_\ell$ for our transfer function, giving $C_{\ell}^\text{eff} \simeq 15-45 \mu K^2_{\text{CMB}}$. Alternatively, we can compute a power spectrum using the differential number counts derived from SHADES data at 350 GHz (Coppin et al. 2006), which is described by

$$N(S)_{350\text{GHz}} = 2.2 \times 10^4 [S^2 + (5.9 \times 10^7)S^{5.8}]^{-1} \text{ Jy}^{-1} \text{ ster}^{-1}. \quad (23)$$

Converting this $N(S)$ to a differential number count at 143 GHz using the average spectrum of $\nu^2$ yields a similar power spectrum, with $C_\ell = 1.0 \text{ Jy}^2 \text{ ster}^{-1}$, or $C_\ell = 8 \times 10^{-6} \mu K^2_{\text{CMB}}$, which is consistent with the result from the number counts given by J. E. Aguirre et al. (2008, in preparation).

Finally, there will also be a signal in our map due to the primary CMB anisotropies, which are distinct from the SZE-induced anisotropies we are searching for. The power spectrum of the primary CMB anisotropies has been measured to high precision by WMAP at $\ell \lesssim 800$ (Nolta et al. 2008), and by ACBAR at $500 \lesssim \ell \lesssim 2500$ (Reichardt et al. 2008). This measured power spectrum is well fitted by theory, with only a small number of free parameters. Therefore, we have generated a template of the primary CMB power spectrum using the theoretical prediction generated by CMBFAST (Seljak & Zaldarriaga 1996; Zaldarriaga et al. 1998; Zaldarriaga & Seljak 2000), with the best-fit values to the free parameters from the WMAP five-year data (Dunkley et al. 2008). Since the CMBFAST routine only computes the power spectrum up to $\ell = 3000$, we fit a decaying exponential to the $C_\ell$ versus $\ell$ to extrapolate the primary CMB power spectrum to higher $\ell$. We can again use Equation (23) to convert this power spectrum to an effective constant $C_\ell$ given our transfer function, with $C_{\ell}^\text{eff} \simeq 45 \mu K^2_{\text{CMB}}$. This band power is similar to what is expected from the SZE-induced CMB anisotropies. A summary of the expected signal from the various astronomical sources is given in Figure 13.

10. SCIENCE ANALYSIS

10.1. Overview of Analyses

In addition to instrumental noise from the bolometers, electronics, etc., our maps will contain an excess noise from astro-

\textsuperscript{24} From Equation (22), we only expect 1–2 sources brighter than 10 mJy in our entire survey of 1 deg$^2$, which is why we have not attempted to subtract out any sources prior to our anisotropy analysis. Additionally, the largest excursions in our maps are $\simeq 10$ mJy, further justifying our choice to set $S_{\text{cut}} = 10$ mJy.
nominal sources, including anisotropies due to primary CMB fluctuations, fluctuations due to the SZE, and fluctuations due to unresolved astronomical point sources. It is our goal to constrain the level of these astronomically sourced noises, which we will specify as the amplitude of flat-band power anisotropy power spectrum contributions in $C_\ell$. To obtain such a constraint, we must calculate the difference between the observed and expected power spectra of our maps and obtain a best estimate of the excess noise, goodness-of-fit of the data to the model, including any possible excess noise, and confidence intervals for the amount of excess noise. This section describes how we obtain the estimate and intervals.

The first analysis we perform will simply constrain the total astronomical anisotropies in the maps, without any interpretation of the source, assuming only that the astronomical noise has a spectral shape flat in $C_\ell$.

The second analysis will statistically subtract the primary CMB anisotropy power spectrum by using the precise constraints placed on it by a variety of measurements (Reichardt et al. 2008; Nolta et al. 2008). The result will be a constraint on the nonprimary-CMB contributions to anisotropy, and will be mildly more sensitive because of the subtraction. We will do this by adding the expected “noise” from the primary CMB to our model of the instrumental noise. This expectation will fully take into account cosmic variance on the primary CMB anisotropy in a manner that we will explain below.

In the end, this analysis will yield an upper limit on the astronomical noise. Because it yields an upper limit, it is conservative to immediately interpret the constraint as a limit on SZE anisotropy: if there are point source contributions, as we expect there are, then the SZE contribution will be smaller than the upper limit we obtain by the assumption that the point source contributions are negligible. The situation would of course be different, and that assumption would not be conservative, were we claiming a detection of excess nonprimary-CMB anisotropy.

One could extend this methodology to statistical subtraction of the non-negligible submillimeter and radio point source contributions, but the large uncertainties in those contributions as well as the possibly unknown systematic uncertainties lead us to conclude that the improvement in sensitivity will be negligible and somewhat untrustworthy.

10.2. Deficiencies of a Bayesian Analysis

We have chosen to model astronomical anisotropies using a flat-band power in $C_\ell$, which corresponds to $C_\ell = A S^2_{\ell}$, for $\ell = 2\ell' [\ell]$ and $S^2_{\ell} = 2\pi/\ell(\ell + 1)$. With these definitions, and assuming the noise PSD, $P_\nu$, fully describes the noise properties of the data for the reasons we have explained in Section 5.3, the best-fit amplitude for an astronomical anisotropy signal is determined by maximizing Equation (A5),

$$\log(\mathcal{L}) = \sum_{i \in V} \left( -\log(P_\nu + A S^2_{\ell} B^2_{\ell'} T_{\ell'}) - \frac{X_{\nu}}{P_\nu + A S^2_{\ell} B^2_{\ell'} T_{\ell'}} \right),$$

with respect to $A$, where $X_{\nu}$ is the measured PSD of the science field map in squared units, $T_{\ell'}$ is the transfer function of our data processing in squared units, and $B_{\ell'}$ is the profile of our beam.\textsuperscript{25}

Since our maps are real, $X_{\nu} = X_{-\nu}$, $P_\nu = P_{-\nu}$, etc., so the sum only includes half the $\nu$-space pixels, denoted by the set $V$. For reference, a detailed derivation of the above equation is given in the Appendix. Note that Equation (A5) allows for $A < 0$. Although such values are not physical, fluctuations in the noise can cause the most likely value of $A$ to be less than zero when the expected value of $A$ is small compared to the nonastronomical noise.

The above expression is incorrect at some level because the $\nu$-space pixels are slightly correlated, approximately 1%–4% for nearest-neighbor pairs of pixels and less than 1% for all other pairs of pixels, while Equation (A5) treats all Fourier modes as independent. Note that the correlation function, $c_{\nu,\nu',\nu''}$, is largely

\textsuperscript{25} Note that we are calculating the anisotropy amplitude for a single bin in $\ell$-space. However, the technique can be applied to multiple bins in $\ell$-space by windowing the appropriate terms in Equation (A5) (i.e., if an $\ell$-space bin is described by the transfer function $T_{\ell'}$, then $P_\nu \rightarrow T_{\ell'} P_\nu$, $S^2_{\ell} \rightarrow T_{\ell'} S^2_{\ell}$, and $X_{\nu} \rightarrow T_{\ell'} X_{\nu}$.)
translation invariant ($c_{\nu,\nu'} \approx c(\nu - \nu')$). This error due to pixel correlations raises three questions: (1) Does maximization of the likelihood given above result in an unbiased estimator of $A$? (2) Is this an approximately minimum variance estimator? and (3) Can we derive Bayesian credibility intervals on $A$ from it? We have demonstrated using simulations that Equation (A5) remains an unbiased and approximately minimum variance estimator for $A$ in spite of these correlations, presumably because ignoring these fairly uniform correlations does not shift the peak of $\mathcal{L}$. See Table 2. However, the width of $\mathcal{L}$ is certainly dependent on these correlations: we are essentially overcounting the number of independent data points entering the likelihood and thus assuming more statistical power than we really have.

We can make an approximate, unrigorous correction for the effective number of independent modes by calculating

$$N_{\text{eff}} = N_{\text{true}} \left( \frac{1}{2N_{\text{true}}} \sum_{i,\nu} \sum_{\nu' < \nu} c_{\nu,\nu'} + \sum_{\nu} c_{\nu,\nu'} \right)^{-1},$$

where $N_{\text{true}}$ is the total number of $\nu$-space pixels in $V$, $N_{\text{eff}}$ is the effective number of $\nu$-space pixels, and $c_{\nu,\nu'}$ is the correlation between pixel $\nu$ and pixel $\nu'$. The factor of 1/2 inside the parentheses arises from the fact that we have double counted the correlations with our sums over $\nu$ and $\nu'$; the factor of $1/N_{\text{true}}$ is a normalization factor. $c_{\nu,\nu'}$ is calculated from the Fourier transform of the map, $M$, according to

$$c_{\nu,\nu'} = \left| \frac{\langle M_{\nu}^* M_{\nu'} \rangle}{\langle |M_{\nu}|^2 \rangle \langle |M_{\nu'}|^2 \rangle} \right|,$$

where the averages are taken over jackknife realizations. Equation (A5) is then multiplied by $N_{\text{eff}}/N_{\text{true}}$ to account for these correlations when calculating the Bayesian likelihood, with $N_{\text{eff}}/N_{\text{true}} \approx 0.43$ for our data. When $\ln(\mathcal{L})$ is exponentiated, this scaling factor will cause $\ln(\mathcal{L})$ to fall off less quickly than it would with $N_{\text{eff}} = N_{\text{true}}$, thereby increasing the width of $\mathcal{L}$. Plots of $\mathcal{L}$, corrected by the factor $N_{\text{eff}} = N_{\text{true}}$, are given in Figure 14.

The lack of rigor behind the above correction implies that there will be statistical problems in placing constraints using the above likelihood function. Were the above likelihood function correct, we could use it to set a $\alpha\%$ Bayesian credibility interval on the parameter $A$ by finding an interval $[A_1, A_2]$, $A_1, A_2 > 0$, such that

$$\alpha = 100 \frac{\int_{A_1}^{A_2} dA \mathcal{L}(X|A)}{\int_0^\infty dA \mathcal{L}(X|A)},$$

where we have assumed a flat prior $A > 0$ because of the nonphysical nature of $A < 0$. If the likelihood function’s width in $A$ is not to be trusted, then such credibility intervals are not valid. Not even a simulation permits one to set a credibility interval because $\mathcal{L}$ is simply not the correct likelihood, even if its distribution can be characterized by simulation.

Additionally, determining the goodness-of-fit for the best-fit value of $A$, $\hat{A}$, will require simulation. That is, if Equation (A5) was correct, then we should be able to determine an analytic expression for the distribution of $\ln(\mathcal{L}) - \ln(\mathcal{L}(\hat{A}))$ for $\hat{A}$ that would allow us to calculate the goodness-of-fit of the data to the model. But, since the above likelihood function is incorrect, we must simulate an ensemble of measurements, with appropriate correlations in the Fourier modes, to determine the distribution of $\ln(\mathcal{L})$ for the value $\hat{A}$. Therefore, the Bayesian approach offers no simplifications or reductions in computing time relative to the simulation-based frequentist technique we employ below.

### 10.3. Overview of the Frequentist, Feldman-Cousins Analysis Technique

It is possible to deal with all of the above problems approximately correctly with a frequentist technique for establishing goodness-of-fit confidence levels and frequentist confidence (as opposed to Bayesian credibility) intervals on $A$ that incorporate the prescriptions of Feldman & Cousins (1998) for dealing with a physical boundary. The technique has two main features:

1. First, we use jackknife maps with signal added based on an input value $A_{\text{sim}}$ in the physically allowed region $A_{\text{sim}} \geq 0$ to determine the distribution of $\mathcal{L}(X_{\nu,i}(A_{\text{sim}})|A_{\text{sim}})$ as defined in Equation (A5) for an ensemble of experiments with outcomes $X_{\nu,i}(A_{\text{sim}})$. With this distribution, we may determine whether $\mathcal{L}(X_{\nu,i}(A_{\text{sim}})|A_{\text{true}}, A_{\text{sim}})$ is among the $\alpha\%$ most likely outcomes for that input value $A_{\text{sim}}$, thereby determining a goodness-of-fit confidence level. In doing this, we make the reasonable approximation that, although $\mathcal{L}$ is not a rigorously correct likelihood, it maps in a one-to-one, monotonic fashion to the true likelihood function $\mathcal{L}_{\text{true}}$. Specifically, if we consider two realizations $X_{\nu,1}$ and $X_{\nu,2}$, we assume that the sign of $\ln(\mathcal{L}(X_{\nu,1}|A_{\text{sim}})) - \ln(\mathcal{L}(X_{\nu,2}|A_{\text{sim}}))$ is the same as that of $\ln(\mathcal{L}_{\text{true}}(X_{\nu,1}|A_{\text{true}})) - \ln(\mathcal{L}_{\text{true}}(X_{\nu,2}|A_{\text{true}}))$. This assumption is far looser than the assumption that rescaling $\mathcal{L}$ by $N_{\text{eff}}/N_{\text{true}}$ is correct; we only assume that the ordering of realizations in $\mathcal{L}$ and $\mathcal{L}_{\text{true}}$ is the same, even if the numerical values are not the same.

2. Second, we want to define a confidence interval of confidence level $\alpha\%$ on $A$. Since we are taking a frequentist approach, these confidence intervals are defined to include the values of $A$ for which, if $A$ is the true value of the anisotropy amplitude, then the observed outcome $X_{\nu,i}$ is within the $\alpha\%$ most likely outcomes (as defined below, a definition that is different than the usual likelihood) for that value of $A$. We use the same set of simulations with the following procedure based on the Neyman construction as modified by Feldman & Cousins (1998). We now calculate for each simulation realization $i$ for each input parameter value $A_{\text{sim}}$ the ratio

$$R(A_{\text{sim}}) = \frac{\mathcal{L}(X_{\nu,i}(A_{\text{sim}})|A_{\text{sim}})}{\mathcal{L}(X_{\nu,i}(A_{\text{true}})|\hat{A})},$$

where $A_{\text{sim}}$ is the simulation input parameter value ($A_{\text{sim}} \geq 0$) and $\hat{A}_i$ is the best-fit value of $A$ in the physically allowed region $A \geq 0$ for the given realization $i$. We order the realizations in order of decreasing $R(A_{\text{sim}})$ until $\alpha\%$ of the realizations have been included; the value of $R(A_{\text{sim}})$ defining this boundary is denoted by $R_{\alpha}(A_{\text{sim}})$. The input parameter value $A_{\text{sim}}$ is then included in the $\alpha\%$ confidence interval if the likelihood ratio for the real data, $R_{\text{data}}(A_{\text{sim}})$, is among the $\alpha\%$ largest $R_i(A_{\text{sim}})$ values. $R_{\text{data}}(A_{\text{sim}}) \geq R_{\alpha}(A_{\text{sim}})$. The interpretation is that, for values $A_{\text{sim}}$ belonging to the confidence interval of confidence level $\alpha\%$, the data are among the $\alpha\%$ most likely outcomes, where “likely” is quantified by $R_{\text{data}}(A_{\text{sim}})$ instead of $\mathcal{L}(X_{\nu,i}|A_{\text{sim}})$. The above procedure can be conveniently visualized as follows. The simulations indicate that there is a smooth relationship between $R(A_{\text{sim}})$ and $\hat{A}_i$ at a given value of $A_{\text{sim}}$. This is generically true, not specific to this analysis. Therefore, each simulation realization may be labeled by its value of $\hat{A}$ and we may write $R(A_{\text{sim}}, \hat{A})$ in place of
Figure 14. Bayesian likelihood given by Equation (A5) for each science field. The likelihoods have all been normalized to one at the peak. These plots should only be considered as rough estimates for determining confidence intervals because the cosmic variance of the CMB spectra, correlations among map pixels, and the physical boundary that the anisotropy amplitude must be greater than or equal to zero have not been fully accounted for in the likelihood function.

We comment on two important aspects of this construction of the confidence intervals. First is the ordering of the simulations by \( R(A_{\text{sim}}, \hat{A}) \) as a function of \( \hat{A} \) for a given value of \( A_{\text{sim}} \); the cutoff value \( R_u(A_{\text{sim}}) \) is a horizontal line in this plot, and so points with \( R(A_{\text{sim}}, \hat{A}) > R_u(A_{\text{sim}}) \) map to a set of intervals in \( \hat{A} \); in fact, in our case, there is a single interval for each \( A_{\text{sim}} \). These intervals, called confidence belts, can be displayed as intervals \([\hat{A}_1, \hat{A}_2]\) at the given value \( A_{\text{sim}} \) in a plot of \( A_{\text{sim}} \) versus \( \hat{A} \), as illustrated in Figure 15. Do not confuse confidence belts, which are intervals along the \( \hat{A} \) axis, with confidence intervals, which are intervals along the \( A_{\text{sim}} \) axis as defined below.

Then, to determine the confidence interval of confidence level \( \alpha \% \) on \( A \) given a data set \( X_{\nu,i} \), one finds \( A_{\text{data}} \), draws a vertical line on the plot of confidence belts at the value \( \hat{A}_{\text{data}} \) on the horizontal (\( \hat{A} \)) axis, and includes all values of \( A_{\text{sim}} \) for which the vertical line lies inside the confidence belt at that value of \( A_{\text{sim}} \). This confidence belt construction is equivalent to the above description based on \( R(A_{\text{sim}}, \hat{A}) \) because the smooth relationship between \( R(A_{\text{sim}}, \hat{A}) \) and \( \hat{A} \) ensures that, for a given \( A_{\text{sim}} \), if \( \hat{A}_{\text{data}} \) is inside the confidence belt at a given value \( A_{\text{sim}} \), then \( R_{\text{data}} \geq R_u(A_{\text{sim}}) \) for that value \( A_{\text{sim}} \).

We comment on two important aspects of this construction of the confidence intervals. First is the ordering of the simulation realizations by \( R \), not by \( L(X_{\nu,i}|A_{\text{sim}})A_{\text{sim}} \). Feldman & Cousins (1998) discuss both possible constructions (the latter originally proposed by Crow & Gardner 1959) and argue that the latter has a serious deficiency in that it ties the confidence level of the confidence interval to the goodness-of-fit confidence level; essentially, it is possible for the confidence interval to not provide the advertised frequentist coverage if the goodness-of-fit is poor. This typically happens when the experimental outcomes yield best-fit parameter values near or outside a physical boundary. In our application, this can occur if the simulation realization has a bit less anisotropy than expected, which would yield a best-fit \( \hat{A} \) that is negative. In such a case, the (approximate) likelihood of the data set, \( L(X_{\nu}|A_{\text{sim}}) \), will in general be small. However, the approximate likelihood of that data set may not be small compared to the approximate likelihood, \( L(X_{\nu}|0) \), of the most probable physically allowed alternative hypothesis of \( \hat{A}_i = 0 \). Feldman & Cousins show that, with this ordering principle, the confidence intervals never contain unphysical values for the observable. Additionally, there is a smooth transition from the case of an upper limit to a central confidence region, eliminating intervals that undercover due to choosing between an upper limit and a central region based on the result. There is no room here to reproduce their arguments in detail; we refer the reader to Feldman & Cousins (1998).

The second important aspect is that the construction is done entirely by simulation so that the only way in which we depend on \( L \), which we know to be deficient, is in the ordering it provides. We have assumed above that, in spite of its inaccuracy, \( L \) provides the same ordering of points as \( L_{\text{true}} \), and hence \( \hat{A} \) and \( R(A_{\text{sim}}, \hat{A}) \) for a given (simulated or real) data realization and \( R_u(A_{\text{sim}}) \) for a given \( A_{\text{sim}} \) will be the same regardless of whether we use \( L \) or \( L_{\text{true}} \).

10.4. Construction of Simulated Data Sets for the Frequentist Technique

To apply this method to our data, we first create a simulated map of the astronomical anisotropy for a given value of the astronomical anisotropy amplitude, \( A_{\text{sim}} \), using our assumed profile \( \hat{S}_2 \). This simulation is produced by drawing a value for each pixel, \( \hat{v} \), from an underlying Gaussian distribution,\(^{26}\) then multiplying it by \( A_{\text{sim}} \hat{S}_2 \). The PSD of this simulated map is multiplied by our full transfer function and added to the jackknifed realization of our data, \( X_{\nu,i} \).\(^{27}\) Note that a different simulated map is created for each jackknifed realization of the data to allow for cosmic variance. Then, we use Equation (A5) to determine the most likely value of the astronomical anisotropy amplitude, \( \hat{A}_i \), for realization \( i \). By using jackknives of our actual data, we are including all of the correlations between pixels, and by simulating the astronomical anisotropy maps we are accounting for cosmic variance in the astronomical anisotropy. For a given value of \( A_{\text{sim}} \), we repeat this process for each jackknifed realization of the data.

The data sets are then ordered based on the ratio of their likelihood to the likelihood of the most probable physically allowed outcome, \( R_i(A_{\text{sim}}) \), as defined above. The procedure outlined in

\(^{26}\) We have also determined confidence intervals using non-Gaussian distributions for the SZE-induced anisotropy signal. The results are described in Section 10.6.

\(^{27}\) The reason we add the simulated astronomical anisotropy map to the jackknifed realization map instead of the time streams is to reduce the amount of computational time required. Since the transfer functions of the maps are well measured, there is no reason to go all the way back to the time streams to add the simulated signal.
Figure 15. First plot shows the Bayesian likelihood for a range of anisotropy amplitudes for the full data set, which includes all of the observations of both science fields. The remaining three plots show the frequentist confidence belts for the full data set for confidence levels of 68%, 90%, and 95%. (A color version of this figure is available in the online journal.)

Figure 16. Band power window function, $W^B_ℓ/ℓ$, for the full data set. We have arbitrarily peak the normalized window function.

To determine the goodness-of-fit of our data to the model given by $S^2_ν$, we compare the likelihood of the actual data at the best-fit value of $A_{\text{data}}$, $L(X_ν | A_{\text{data}})$, to the likelihoods of a set of jackknifed realizations of our data with simulated spectra added according to $S^2_ν$ with amplitude $A_{\text{data}}$, $L(X_ν | A_{\text{data}})$. For the observations of the Lynx field, $L(X_ν | A_{\text{data}})$ is greater than $L(X_ν | A_{\text{data}})$ for 17% of the realizations, and for the observations of the SDS1 field $L(X_ν | A_{\text{data}})$ is greater than $L(X_ν | A_{\text{data}})$ for 43% of the realizations. Therefore, we can conclude that our model provides an adequate description of the data.

10.5. Total Anisotropy Amplitude Results

To determine the confidence intervals for the full data set, we make a joint estimate of $A$ using both the Lynx and SDS1 data sets. A plot of the Bayesian likelihood, along with confidence belts computed using the Feldman & Cousins (1998) method are given in Figure 15. Uncertainties in our pointing model have already been included in these calculations by an effective broadening of the Bolocam beam. Our upper limits on the total anisotropy amplitude are equal to 590, 760, and 830 $\mu K^2_{\text{CMB}}$ at confidence levels of 68%, 90%, and 95%. Note that the uncertainty on these limits due to the finite number of simulations we have run is $\simeq 10–15$ $\mu K^2_{\text{CMB}}$.

To determine the effective angular scale of our anisotropy amplitude measurements we have computed our band power window function, $W^B_ℓ/ℓ$, using the method given by Knox.

---

28 This band power window function is defined such that
This simulation is produced by drawing a value for each pixel, data processing, and equal to 5700. Additionally, the full width at half-maximum function for the full data set is given in Figure 16. From this (1999). A plot of the peak-normalized band power window function for all of the current CMB anisotropy measurements above \( \ell = 2000 \). The solid lines represent observations made near 150 GHz, and the dashed lines represent observations made near 30 GHz. The primary CMB anisotropies were calculated using the parameters given in Section 9; the analytic model of Komatsu & Seljak (2002), along with the best estimate of \( \sigma_8 \) from Dawson et al. (2006), were used to estimate the SZE-induced CMB anisotropies. All of the data are plotted with 1σ error bars, except for the Bolocam upper limit at \( \ell = 5700 \) and the BIMA upper limit at \( \ell = 8748 \), which are given as 90% and 95% confidence level upper limits, respectively. The ACBAR data were taken from Reichardt et al. (2008), the BIMA data were taken from Dawson et al. (2006), and the CBI data were taken from Mason et al. (2003). (A color version of this figure is available in the online journal.)

(1999). A plot of the peak-normalized band power window function for the full data set is given in Figure 16. From this band power window function we have calculated an effective angular multipole for our data set, \( \ell_{\text{eff}} \), given by

\[
\ell_{\text{eff}} = \frac{\sum \ell (W^B_\ell / \ell)}{\sum \ell W^B_\ell / \ell},
\]

and equal to 5700. Additionally, the full width at half-maximum of the window function, FWHM, is equal to 2800. A plot comparing our result to other measurements of the CMB on similar scales is shown in Figure 17.

10.6. SZE-Induced CMB Anisotropy Results and Constraints on \( \sigma_8 \)

In order to determine the amplitude of the SZE-induced CMB power spectrum, we follow the same methods described above to determine the total amplitude of the anisotropy power spectrum. However, we now have to statistically subtract the signal due to the primary CMB anisotropies by accounting for both the amplitude and fluctuations of its expected power spectrum in the likelihood; these primary CMB anisotropies are effectively an additional noise in the map. The noise contributed to the map from the Bolocam system is given by \( P_\gamma \). Since the spectrum of the primary anisotropies in the CMB is well understood, we can calculate the expected noise from the primary CMB anisotropies. To calculate this noise we first create a simulated map of the primary CMB, assuming that the underlying distribution of \( \vec{v} \)-space pixel values is Gaussian. This simulation is produced by drawing a value for each pixel, \( \vec{v} \), from an underlying Gaussian distribution, then multiplying it by the best-fit primary CMB spectrum given in Section 9. The PSD of this map is then multiplied by \( T_\gamma B^2 \) and added to a jackknifed realization of our data, \( X_{\vec{v}}^\text{[SZE]} \), to give \( X_{\vec{v}}^\text{[SZE]} \). A different simulated map is generated for each jackknifed realization of the data to account for the cosmic variance in the CMB spectrum. These modified jackknifed realizations of the data are then be used to determine the expected PSD, \( P_\gamma^\text{[SZE]} \), for the noise contributed by the Bolocam system and the primary CMB anisotropies. We note that such a simulation of the primary CMB anisotropies is more correct than simply adding the primary CMB power spectrum to the nonastronomical noise power spectrum because it correctly reproduces pixelization and Fourier-mode correlation effects.

Next, we select a model spectrum for the SZE anisotropies, \( S_{\nu}^\text{[SZE]} \). Using these new definitions, the Bayesian likelihood function in Equation (A5) can be written as

\[
\log(\mathcal{L}) = \sum \nu \left( -\log \left( P_\nu^\text{[SZE]} + A^\text{[SZE]} \left( S_{\nu}^\text{[SZE]} \right)^2 B^2 T_\gamma \right) - \frac{X_{\nu}^\text{[SZE]}}{P_\nu^\text{[SZE]} + A^\text{[SZE]} \left( S_{\nu}^\text{[SZE]} \right)^2 B^2 T_\gamma} \right),
\]

where \( A^\text{[SZE]} \) is the amplitude of the SZE-induced CMB anisotropies, \( B^2 \) is the profile of our beam, and \( T_\gamma \) is the transfer function of our data processing in squared units. As before, we create simulated SZE maps with an amplitude \( A_{\text{sim}}^\text{[SZE]} \), add these to our jackknifed realizations after multiplying by \( \left( S_{\nu}^\text{[SZE]} \right)^2 B^2 T_\gamma \), then use the ordering method developed by Feldman & Cousins (1998) to determine the width of the confidence belt at \( A_{\text{sim}}^\text{[SZE]} \). By repeating this procedure for a range of physically allowed values of \( A_{\text{sim}}^\text{[SZE]} \), we can construct a full confidence belt that can be used to determine our confidence intervals. We emphasize that, while we use the Bayesian likelihood to construct our best estimator for \( A \), a procedure that we have already demonstrated by simulation is unbiased and approximately minimum variance, we in no way rely on the Bayesian likelihood to determine confidence intervals on \( A \). The frequentist Feldman–Cousins method is used for the latter task.

Additionally, we need to account for the flux calibration uncertainty. The uncertainty in the flux calibration model derived from point sources is 5.5%, and the uncertainty in the area of our beam is 3.1%. Therefore, the uncertainty in our surface brightness calibration is 6.3%. To determine the effect of this flux calibration error on our confidence intervals, we multiplied each simulated primary and SZE-induced CMB map by \( \phi_i = 1 + y \), where \( y \) is drawn from a Gaussian distribution with a standard deviation equal to our flux uncertainty of 0.063. A different \( \phi_i \) was generated for each simulated CMB map. This means that each simulated map has a different flux calibration, distributed according to our uncertainty in the calibration. New confidence belts were then calculated using the same procedure described above. We have also determined the confidence intervals assuming that there is no uncertainty in the known flux of Uranus and Neptune (i.e., the only flux calibration uncertainties are due to our measurement errors and observational techniques). In this case, the flux calibration uncertainty is 3.5% instead of 6.3%.

These flux calibration uncertainties produce non-negligible changes to the confidence intervals we determine for the anisotropy amplitude since it is a variance (i.e., it depends quadratically on the flux calibration). Therefore, for a simulated amplitude \( A_{\text{sim}} \), a fractional flux calibration uncertainty of \( \sigma_f \)
will increase/decrease the upper/lower bounds of the 68% CL confidence belt by an amount roughly proportional to $A_{\mathrm{int}}(1 + \sigma_f)^2$. The resulting fractional change to the confidence interval limits will in general be nontrivial, but should be approximately equal to $(1 + \sigma_f)^2$ for 68% CL limits. So, for a 3.5% flux calibration uncertainty we expect the 68%/90%/95% CL upper limits to increase by approximately 7%/12%/14% compared to the case of no flux calibration uncertainty. Similarly, for a 6.3% flux calibration uncertainty we expect the 68%/90%/95% CL upper limits to increase by approximately 13%/22%/27% compared to the case of no flux calibration uncertainty. After fully simulating the effect of the flux calibration uncertainty on our upper limits, we find results that are comparable to the predictions given above (see Table 1).

We have computed confidence intervals for two different SZE spectra: a flat spectrum, $(S_{\nu}^{\text{SZE}})^2 = 2\pi/(\ell + 1)$, for $\ell = 2\pi|\vec{v}|$ and the analytic spectrum calculated by Komatsu & Seljak (2002). The results for both of these spectra are similar, which is reasonable since the analytic spectrum is nearly flat at the scales to which we are most sensitive (4000 $\lesssim \ell \lesssim$ 7000; see Table 1). In addition to the analytic spectrum calculated by Komatsu & Seljak, several SZE power spectra have been determined via hydrodynamic simulations using either moving-mesh hydrodynamic (MMH) or smoothed particle hydrodynamic (SPH) algorithms. Examples of MMH simulations can be found in Zhang et al. (2002), Seljak et al. (2001), Refregier et al. (2000), and Refregier & Teysier (2002). Examples of SPH simulations can be found in da Silva et al. (2001) and Springel et al. (2001). Since most of the simulated SZE spectra are approximately flat at the angular scales we are most sensitive to, we have not determined confidence levels using any of these spectra (see Figure 1 in Komatsu & Seljak 2002).

Komatsu and Seljak determined that the amplitude of the SZE-induced CMB anisotropies scales according to $\sigma_8^2(\Omega_{\text{m}}h)^2$ and is relatively insensitive to all other cosmological parameters (Komatsu & Seljak 2002). Using the results from the WMAP five year data, the best-fit values for $\sigma_8$, $\Omega_m$, and $h$ are 0.796, 0.0440, and 0.719 (Dunkley et al. 2008). These values produce a maximum SZE anisotropy amplitude of less than 10 $\mu K_{\text{MB}}^2$ at our band center of 143 GHz for the analytic Komatsu & Seljak spectrum. For comparison, the 90% confidence level upper limit on the average value of the analytic spectrum weighted by the Bolocam transfer function is 950 $\mu K_{\text{MB}}^2$, including our flux calibration error (see Table 1). Based on this upper limit, assuming the scaling relation given by Komatsu & Seljak and holding all other parameters fixed, the corresponding 90% confidence level upper limit on the three cosmological parameters is $\sigma_8^2(\Omega_{\text{m}}h)^2 < 2.13$. Individually, the best constraint can be placed on $\sigma_8$ since the amplitude depends most strongly on this parameter, with $\sigma_8 < 1.55$ at a confidence level of 90%.

However, this upper limit has been derived by assuming the SZE-induced anisotropy signal is Gaussian, which is a poor assumption. To account for the non-Gaussianity of the signal, we have used a method similar to the one described by Goldstein et al. (2003) to analyze data collected with ACBAR. Based on the results of numerical simulations by White et al. (2002) and Zhang et al. (2002), along with calculations of the trispectrum term from Cooray (2001) and Komatsu & Seljak (2002), they determined that the sample variance of the SZE-induced anisotropy signal should be a factor of 3 larger than the Gaussian equivalent for the $\ell$-range that ACBAR is most sensitive to. For our data, at $\ell \simeq 6000$, the sample variance is approximately four times larger than the expectation for a Gaussian. When we account for this increased sample variance our 68%, 90%, and 95% confidence level upper limits for the average amplitude of the Komatsu & Seljak spectrum are 790, 1060, and 1080 $\mu K_{\text{MB}}^2$, which are approximately 10% higher than the upper limits obtained from assuming the fluctuations in the SZE anisotropy signal are Gaussian. The changes in the upper limits we determine are relatively minor because our uncertainty is dominated by Gaussian instrument noise rather than sample variance on the anisotropy signal. When we convert these upper limits on the anisotropy signal to an upper limit on $\sigma_8$, we find $\sigma_8 < 1.57$ at a 90% confidence level.

11. CONCLUSIONS

We have surveyed two science fields totaling 1 deg$^2$ with Bolocam at 2.1 mm to search for secondary CMB anisotropies caused by the SZE. The fields are in the Lynx and Subaru/XMM SDS1 fields. Our survey is sensitive to angular scales with an effective angular multipole of $\ell_{\text{eff}} = 5700$ with FWHM$_{\text{e}} = 2800$ and has an angular resolution of 60 arcsec FWHM. Our data provide no evidence for anisotropy. We are able to constrain the level of total astronomical anisotropy, modeled as a flat-band power in $C_\ell$, with most frequent 68%, 90%, and 95% CL upper limits of 560, 760, and 830 $\mu K_{\text{MB}}^2$. We statistically subtract the known contribution from primary CMB anisotropy, including cosmic variance, to obtain constraints on the SZE anisotropy contribution. Now including flux calibration uncertainty, our most frequent 68%, 90%, and 95% CL upper limits on a flat-band power in $C_\ell$ are 690, 960, and 1000 $\mu K_{\text{MB}}^2$. When we instead employ the analytic spectrum suggested by Komatsu & Seljak (2002), and account for the non-Gaussianity of the SZE anisotropy signal, we obtain upper limits on the average amplitude of their spectrum weighted by our transfer function of 790, 1060, and 1080 $\mu K_{\text{MB}}^2$. We obtain a 90% CL upper limit on $\sigma_8$, which normalizes the power spectrum of density fluctuations, of 1.57. These are the first constraints on anisotropy and $\sigma_8$ from survey data at these angular scales at frequencies near 150 GHz.

To calibrate the observations, beam maps were obtained using Uranus and Neptune. Pointing reconstruction was performed using frequent pointing observations of bright sources near our science fields. The data were flux calibrated using techniques similar to those developed to analyze earlier Bolocam survey data collected at 1.1 mm (Laurent et al. 2005), using Uranus and Neptune as absolute calibrators and a number of other sources as transfer calibrators. Internal uncertainty on the pointing and flux calibration contributes negligible uncertainty to the final result; calibration uncertainty in the final result is dominated by uncertainty in models for the absolute brightness temperatures of Mars, Uranus, and Neptune.

Our time streams are dominated by fluctuations in atmospheric thermal emission (sky noise) and we developed several algorithms to subtract this noise from our data. We made use of our simple yet crosslinked scan strategy to develop a pseudo-least-squares map maker that can be run in moderate amounts of time on a single desktop computer. We used simulations to calibrate the transfer function of our data taking and analysis pipeline and map maker. We determined the expected noise properties of our final maps using jackknife realizations of the data obtained by randomly signed combinations of the ~500 independent observations contributing to each science field map. Our final confidence intervals on the anisotropy level are determined using these jackknife realizations combined with the measured transfer function for anisotropies.
We acknowledge the assistance of Minhee Yun and Anthony D. Turner of NASA’s Jet Propulsion Laboratory, who fabricated the Bolocam science array; Toshiro Hatake of the JPL electronic packaging group, who wire-bonded the array; Marty Gould of Zen Machine and Ricardo Paniagua and the Caltech PMA/GPS Instrument Shop, who fabricated much of the Bolocam hardware; Carole Tucker of Cardiff University, who tested metal-mesh reflective filters used in Bolocam; Ben Knowles of the University of Colorado, who contributed to the software pipeline, the day crew and Hilo staff of the Caltech Submillimeter Observatory, who provided invaluable assistance during commissioning and data-taking for this survey data set; and Kathy Deniston, who provided effective administrative support at Caltech. Bolocam was constructed and commissioned using funds from NSF/AST-9618798, NSF/AST-0098737, NSF/AST-9980846, NSF/AST-0229008, and NSF/AST-0206158. J.S. and G.L. were partially supported by NASA Graduate Student Research Fellowships and S.G. was partially supported by an R. A. Millikan Postdoctoral Fellowship at Caltech.

Facilities: CSO.

APPENDIX
APPENDIX MATERIAL

The goal of our analysis is to determine the amplitude of the power spectrum due to emission from astronomical sources by measuring an excess noise in the maps of the science fields. This excess noise is the difference between the actual noise of the map in the absence of the desired astronomical signal. For a flat-band power spectrum due to emission from astronomical sources by \(\vec{\nu} T\), the spatial power spectral profile of the expected signal spectrum. Therefore, we need measurements of the following quantities:

1. \(X_{\vec{v}}\). The measured PSD of the science field map at pixel \(\vec{v}\) in units of \(\mu K_{\text{CMB}}^2\). \(\vec{v}\) is a two-dimensional value, \(\vec{v} = (v_{R.A.}, v_{\text{decl}})\), describing a location in the spatial Fourier transform of the map, and has units of 1 rad\(^{-1}\).
2. \(P_{\vec{v}}\). The predicted PSD of the science field map at pixel \(\vec{v}\) in the absence of the desired astronomical signal. \(P_{\vec{v}}\) is estimated from jackknife realizations, along with the PSDs of unwanted astronomical sources in the map (i.e., primary CMB anisotropies in our case).
3. \(S^2_{\vec{v}}\). The spatial power spectral profile of the expected astronomical signal. For a flat-band power \(S^2_{\vec{v}} = 2\pi/(\ell(\ell + 1))\), where the angular multipole \(\ell\) is described by \(\ell = 2\pi |\vec{v}|\).
4. \(B^2_{\vec{v}}\). The peak-normalized square of the \(\vec{v}\)-space Bolocam beam profile. Since astronomical signals are attenuated by the beam, \(B^2_{\vec{v}}\) acts like a transfer function or a filter. Note that the broadening of the beam in the map space due to our pointing uncertainty is included in \(B^2_{\vec{v}}\).
5. \(T_{\vec{v}}\). The effective transfer function, or window function, of the data processing applied to the time-stream data. Analogous to \(B^2_{\vec{v}}\), \(T_{\vec{v}}\) describes how much astronomical signal is attenuated.

With this convention, the expected PSD of the map can be described by

\[
(X_{\vec{v}}) = P_{\vec{v}} + AS^2_{\vec{v}}B^2_{\vec{v}}T_{\vec{v}},
\]

where \(A\) is the amplitude of the excess anisotropy power, in \(\mu K_{\text{CMB}}^2\).

The anisotropy amplitude can be estimated by determining what value of \(A\) maximizes the likelihood of the measured map PSD, \(X_{\vec{v}}\). Therefore, we need to determine the probability density function (PDF) describing \(X_{\vec{v}}\), given \(A\). First, note that

\[
X_{\vec{v}} = |\alpha + i\beta|^2,
\]

where \(\alpha\) is the real part of the Fourier transform of the science field map and \(\beta\) is the imaginary part of the Fourier transform of the science field map. If we assume that the noise properties of the map are Gaussian, then the PDFs for \(\alpha\) and \(\beta\) are the same and are given by

\[
f(\alpha) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\alpha^2/2\sigma^2} \quad \text{and} \quad f(\beta) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\beta^2/2\sigma^2},
\]

(A2)

where \(\sigma^2 = \langle X_{\vec{v}}^2 \rangle/2\). Next, after a change of variables to \(\alpha = r\cos(\theta)\) and \(\beta = r\sin(\theta)\), the PDF in Equation (A2) becomes

\[
f(r, \theta) = \frac{1}{2\pi\sigma^2} r e^{-r^2/2\sigma^2}.
\]

Since the \(\theta\) dependence of \(f(r, \theta)\) is trivial, we can reduce the above PDF to \(f(r) = 2\pi f(r, \theta)\), with

\[
f(r) = \frac{r}{\sigma^2} e^{-r^2/2\sigma^2}.
\]

Finally, after one more change of variables using the relation \(X_{\vec{v}} = r^2\), we find that the PDF for \(X_{\vec{v}}\) is equal to

\[
f(X_{\vec{v}}) = \frac{1}{2\sigma^2} e^{-X_{\vec{v}}/2\sigma^2},
\]

(A3)

where the factor of \(r\) has been replaced by one half due to the change in the differential element. Equation (A3) can be written in terms of our measured parameters as

\[
f(X_{\vec{v}}|A) = \frac{1}{P_{\vec{v}} + AS^2_{\vec{v}}B^2_{\vec{v}}T_{\vec{v}}} \exp\left(-\frac{X_{\vec{v}}}{P_{\vec{v}} + AS^2_{\vec{v}}B^2_{\vec{v}}T_{\vec{v}}}\right)
\]

(A4)

using Equation (A1). Note that we have made use of the fact that \(2\sigma^2 = \langle X_{\vec{v}}^2 \rangle = P_{\vec{v}} + AS^2_{\vec{v}}B^2_{\vec{v}}T_{\vec{v}}\) to go from Equation (A3) to Equation (A4).

The next step is to calculate a likelihood function, \(L\), from Equation (A4) by multiplying \(f(X_{\vec{v}}|A)\) over all of the \(\vec{v}\)-space pixels. This product can be turned into a sum by taking the logarithm of \(L\), with

\[
\log(L) = \sum_{\vec{v}} \left(-\log\left(P_{\vec{v}} + AS^2_{\vec{v}}B^2_{\vec{v}}T_{\vec{v}}\right) - \frac{X_{\vec{v}}}{P_{\vec{v}} + AS^2_{\vec{v}}B^2_{\vec{v}}T_{\vec{v}}}\right).
\]

(A5)

Note that half of the \(\vec{v}\)-space is discarded from the sum in Equation (A5) since our maps are real (\(X_{\vec{v}} = X_{-\vec{v}}\)).

Then, the most probable value of the anisotropy amplitude for our measured map PSD can be determined by maximizing \(\log(L)\) with respect to \(A\). In practice, we maximize Equation (A5) by evaluating \(\log(L)\) at a range of values for \(A\). Since the number of \(\vec{v}\)-space pixels is \(\lesssim 10,000\), the computational time required to evaluate \(\log(L)\) at each value of \(A\) is minimal, which means that we can determine the best-fit value of \(A\) to almost any desired precision using this numerical method.

\[29\] This is an extremely good assumption (see Figure 11). Although the anisotropy signal may not follow a Gaussian distribution, \(AS^2_{\vec{v}}B^2_{\vec{v}}T_{\vec{v}} \ll P_{\vec{v}}\) for a single \(\vec{v}\)-space pixel, so the underlying distribution function for \(X_{\vec{v}}\) will still be well approximated by a Gaussian.
However, it is also instructive to analytically approximate the value of $A$ that maximizes Equation (A5). To start, we take the derivative of $\log(L)$ with respect to $A$, yielding

$$
\frac{\partial \log(L)}{\partial A} \bigg|_{A=\hat{A}} = \sum_P \Theta_P \left( \frac{\hat{X}_P - \hat{P}_P - A \Theta_P}{\hat{P}_P} \right)^2 \left( X_P - P_P - A \Theta_P \right) = 0,
$$

where $\Theta_P = S^2 \mathcal{B}^2 T_3$ and $\hat{A}$ is the best-fit value of $A$. For any given $\vec{v}$-space pixel, $\hat{P}_P \gg A \Theta_P$ for any physically reasonable value of $A$. Therefore, we can simplify Equation (A6) to

$$
\sum_P \Theta_P \left( 1 - \frac{2 A \Theta_P}{\hat{P}_P} + \mathcal{O} \left( \frac{A^2 \Theta_P^2}{\hat{P}_P^2} \right) \right) \left( X_P - P_P - A \Theta_P \right) \bigg|_{A=\hat{A}} \approx 0.
$$

If we rearrange some terms, and again keep only the lowest order terms in $A \Theta_P / \hat{P}_P$, then we find

$$
\hat{A} \simeq \frac{\sum_P \Theta_P \left( X_P - P_P \right)}{\sum_P \Theta_P \left( 2X_P - P_P \right)}.
$$

Finally, because $A \Theta_P \ll \hat{P}_P$, we can make the approximation that $\langle X_P \rangle \simeq \hat{P}_P$, which means that $\langle 2X_P - P_P \rangle \simeq \hat{P}_P$. With this approximation we find

$$
\hat{A} \simeq \frac{\sum_P \Theta_P \left( X_P - P_P \right)}{\sum_P \Theta_P \hat{P}_P}.
$$

(A7)

To understand this result, consider that for a single $\vec{v}$-space pixel the best estimate of $A$ is $(X_P - P_P) / \Theta_P$. Therefore, Equation (A7) determines the weighted mean of $A$ over all pixels, assuming that the uncertainty on the value of $A$ for each $\vec{v}$-space pixel is proportional to $\hat{P}_P / \Theta_P$, which is a reasonable assumption. This means that the variance on $\hat{A}$ implied by Equation (A7) is proportional to

$$
\sigma^2 \hat{A} \propto \frac{1}{\sum_P \Theta_P \hat{P}_P^2}.
$$

(A8)

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However, it is also instructive to analytically approximate the value of $A$ that maximizes Equation (A5). To start, we take the derivative of $\log(L)$ with respect to $A$, yielding

$$
\frac{\partial \log(L)}{\partial A} \bigg|_{A=\hat{A}} = \sum_P \Theta_P \left( \frac{\hat{X}_P - \hat{P}_P - A \Theta_P}{\hat{P}_P} \right)^2 \left( X_P - P_P - A \Theta_P \right) = 0,
$$

where $\Theta_P = S^2 \mathcal{B}^2 T_3$ and $\hat{A}$ is the best-fit value of $A$. For any given $\vec{v}$-space pixel, $\hat{P}_P \gg A \Theta_P$ for any physically reasonable value of $A$. Therefore, we can simplify Equation (A6) to

$$
\sum_P \Theta_P \left( 1 - \frac{2 A \Theta_P}{\hat{P}_P} + \mathcal{O} \left( \frac{A^2 \Theta_P^2}{\hat{P}_P^2} \right) \right) \left( X_P - P_P - A \Theta_P \right) \bigg|_{A=\hat{A}} \approx 0.
$$

Finally, because $A \Theta_P \ll \hat{P}_P$, we can make the approximation that $\langle X_P \rangle \simeq \hat{P}_P$, which means that $\langle 2X_P - P_P \rangle \simeq \hat{P}_P$. With this approximation we find

$$
\hat{A} \simeq \frac{\sum_P \Theta_P \left( X_P - P_P \right)}{\sum_P \Theta_P \hat{P}_P}.
$$

(A7)

To understand this result, consider that for a single $\vec{v}$-space pixel the best estimate of $A$ is $(X_P - P_P) / \Theta_P$. Therefore, Equation (A7) determines the weighted mean of $A$ over all pixels, assuming that the uncertainty on the value of $A$ for each $\vec{v}$-space pixel is proportional to $\hat{P}_P / \Theta_P$, which is a reasonable assumption. This means that the variance on $\hat{A}$ implied by Equation (A7) is proportional to

$$
\sigma^2 \hat{A} \propto \frac{1}{\sum_P \Theta_P \hat{P}_P^2}.
$$

(A8)