We describe redMaPPer, a new red sequence cluster finder specifically designed to make optimal use of ongoing and near-future large photometric surveys. The algorithm has multiple attractive features: (1) it can iteratively self-train the red sequence model based on a minimal spectroscopic training sample, an important feature for high-redshift surveys. (2) It can handle complex masks with varying depth. (3) It produces cluster-appropriate random points to enable large-scale structure studies. (4) All clusters are assigned a full redshift probability distribution $P(z)$. (5) Similarly, clusters can have multiple candidate central galaxies, each with corresponding centering probabilities. (6) The algorithm is parallel and numerically efficient: it can run a Dark Energy Survey-like catalog in $\sim 500$ CPU hours. (7) The algorithm exhibits excellent photometric redshift performance, the richness estimates are tightly correlated with external mass proxies, and the completeness and purity of the corresponding catalogs are superb. We apply the redMaPPer algorithm to $\sim 10,000$ deg$^2$ of SDSS DR8 data and present the resulting catalog of $\sim 25,000$ clusters over the redshift range $z \in [0.08, 0.55]$. The redMaPPer photometric redshifts are nearly Gaussian, with a scatter $\sigma_z \approx 0.006$ at $z \approx 0.1$, increasing to $\sigma_z \approx 0.02$ at $z \approx 0.5$ due to increased photometric noise near the survey limit. The median value for $|\Delta z|/(1 + z)$ for the full sample is 0.006. The incidence of projection effects is low ($\lesssim 5\%$). Detailed performance comparisons of the redMaPPer DR8 cluster catalog to X-ray and Sunyaev–Zel’dovich catalogs are presented in a companion paper.

**Key word:** galaxies: clusters: general

**Online-only material:** color figures, machine-readable tables

1. INTRODUCTION

Over the past several years, galaxy clusters have been recognized as powerful cosmological probes (e.g., Henry et al. 2009; Vikhlinin et al. 2009; Mantz et al. 2010; Rozo et al. 2010; Clerc et al. 2012; Benson et al. 2013; Hasselfield et al. 2013). Galaxy clusters are one of the key probes of dark energy for ongoing and upcoming photometric surveys such as the Dark Energy Survey (DES; The DES Collaboration 2005), Pan-STARRS (Kaiser et al. 2002), the Hyper-Suprime Camera (HSC),12 and the Large Synoptic Survey Telescope (LSST; LSST Dark Energy Science Collaboration 2012).

Because galaxies are obviously clustered on the sky, rich galaxy clusters have been detected as far back as the 1800s (Biviano 2000), with the first large catalogs generated from galaxy overdensities on photographic plates created 50 years ago (e.g., Abell 1958; Zwicky et al. 1968; Abell et al. 1989). More recently, the advent of multi-band data has led to a proliferation of optical cluster-finding algorithms. These algorithms use various techniques for measuring clustering in angular position plus color/redshift space, ranging from simple matched filters to more complicated Voronoi tessellations. These cluster finders can be divided into roughly two classes: those based on photometric redshifts (e.g., Kepner et al. 1999; van Breukelen & Clewley 2009; Milkeraitis et al. 2010; Durret et al. 2011; Szabo et al. 2011; Soares-Santos et al. 2011; Wen et al. 2012), and those utilizing the cluster red sequence (e.g., Annis et al. 1999; Gladders & Yee 2000; Koester et al. 2007a; Gladders et al. 2007; Gal et al. 2009; Thanjavur et al. 2009; Hao et al. 2010; Murphy et al. 2012). However, relatively few of these optical catalogs have been utilized for cosmological parameter constraints (e.g., Rozo et al. 2007, 2010; Mana et al. 2013).

Given the above landscape, it is a fair question to ask whether the world really needs yet another photometric cluster-finding algorithm. As we describe below, we believe that the answer to this question is yes. In particular, there are a variety of important features that any reasonable optical cluster finder must have in order to properly exploit the photometric data that will become available with ongoing or near-future photometric surveys such as the DES or LSST.

What must we require of current photometric cluster finders? The key features are as follows.

1. The algorithm must be able to smoothly detect galaxy clusters in a consistent way across a broad redshift range. This can be a challenge for photometric redshift (“photo-z”) and red-sequence-based algorithms alike. For photometric-redshift-based algorithms, one must be cautious because...
The Astrophysical Journal, 785:104 (33pp), 2014 April 20

Rykoff et al.

biases and scatter in reported photo $z$ increase at fainter magnitudes where spectroscopic training and validation samples can be highly incomplete. For red-sequence-based cluster finders, one must confront the fact that the 4000 Å break characteristic of the early-type galaxy spectra moves across filters. While $g-r$ is an ideal color for cluster detection at low redshift, one should rely primarily on $r-i$ at intermediate redshifts and $i-z$ at higher redshifts (and we note that this will also affect photo-$z$-based finders). By $z \approx 1$, near-infrared (NIR) photometry is required. Being able to smoothly transition from one color to the next—or better yet, to always use all available photometric data—is paramount.

2. To the extent possible, the algorithm should self-train to the available data. For instance, algorithms reliant on a priori parameterizations of the red sequence could easily result in systematic biases if the a priori parameterization differs from reality. Note that this also impacts photo-$z$-based cluster finders, since there can be unknown and difficult-to-calibrate biases in the photometric redshifts of cluster galaxies.

3. The algorithm should be numerically efficient, capable of running on extremely large data sets within reasonable time frames with modest computational resources.

4. The algorithm must be able to properly account for complex survey masks, including varying depth.

5. The algorithm must allow the construction of proper cluster-random points that adequately characterize the effective survey volume for cluster detection in order to enable large-scale structure studies. In particular, it is worth emphasizing that because galaxy clusters are extended objects on the sky, the galaxy mask used to construct the cluster catalog is not the appropriate mask characterizing the angular and redshift selection of galaxy clusters for any particular cluster finder.

6. The algorithm should produce a full $P(z)$ distribution for every cluster. Similarly, given that the center of a galaxy cluster can be observationally uncertain, there should be a corresponding centering distribution in the plane of the sky $P(\hat{\mathbf{r}})$. Note that if one adopts the prior that a galaxy resides at the center of a galaxy cluster, then the probability $P(\hat{\mathbf{r}})$ collapses to the probability that any given cluster galaxy is the correct cluster center. Our expectation is that just as $P(z)$ allows one to adequately recover the redshift distribution of galaxy clusters in a statistical sense, so too will centering probabilities for cluster galaxies allow one to statistically recover the angular distribution of clusters in the sky, a point that is of critical importance for large-scale structure studies.

7. In order to maximize the cosmological utility of the derived cluster samples, the richness estimators should be fully optimized for the purpose of minimizing the scatter in the richness–mass relation.

The red-sequence Matched-filter Probabilistic Percolation (redMaPPer) cluster-finding algorithm is our solution to the above list of must-haves. Concerning the last point in particular, over the past several years we have empirically explored what works and what does not work in estimating cluster richness (Rozo et al. 2009, henceforth R09, 2011; Rykoff et al. 2012, henceforth R12). For instance, we have demonstrated that estimating membership probabilities for every galaxy is very effective, while using hard color cuts to derive cluster membership can lead to large biases. We have fully optimized the optical detection radii, as well as the luminosity cuts employed when counting galaxies. We have also investigated whether total galaxy counts or total cluster luminosity is a better mass proxy, and whether or not trying to add blue galaxies into richness estimates results in improvements. All of these lessons have gone into the creation of the redMaPPer cluster finder.

There is, however, one feature of redMaPPer that represents more of a personal bias as opposed to an empirically driven choice, namely, the fact that redMaPPer is a red sequence cluster finder. Indeed, operationally, redMaPPer can be easily adapted to work in photo-$z$ space rather than working directly in color space. However, we are wary of reliance on photometric redshifts, which become increasingly difficult to characterize for faint galaxies due to a lack of spectroscopic training and validation samples. Furthermore, cluster galaxies are a very particular population, and photo-$z$ estimates tailored for clusters should be derived separately from the total galaxy population. Though there have been some studies comparing different cluster finders (e.g., Goto et al. 2002; Bahcall et al. 2003; Lopes et al. 2004; Rozo & Rykoff 2010), we have not seen any conclusive evidence for photo-$z$-based algorithms outperforming red sequence methods or vice versa. Here we have opted to rely on a red sequence method when developing redMaPPer. Of course, at high enough redshift, as the red sequence begins to disappear, it is obvious that photometric redshift methods must necessarily perform better (e.g., Eisenhardt et al. 2008; Brodwin et al. 2011). We do not, however, expect this to be a problem for near-future large photometric surveys. Note that while it is true that redMaPPer also relies on spectroscopic training samples, an important advantage of our novel red sequence modeling algorithm is that we do not require a locally representative training sample: our spectroscopic training galaxies can be the brightest cluster galaxies at all redshifts, with no degradation in the performance of our photometric redshift estimates.

The redMaPPer algorithm is designed to handle an arbitrary photometric galaxy catalog, with an arbitrary number of photometric bands ($\geq 3$), and will perform well provided that the photometric bands span the 4000 Å break over the redshift range of interest. It is thus well suited to current surveys such as the Sloan Digital Sky Survey (SDSS; York et al. 2000) for low- and moderate-redshift clusters ($z \in [0.05, 0.55]$), as well as upcoming surveys such as DES for low- and high-redshift clusters ($z < 1$). As a case study, in this paper we present the redMaPPer catalog as run on 10.400 deg$^2$ of photometric data from the Eighth Data Release (DR8; Aihara et al. 2011) of the SDSS. We will make the full DR8 redMaPPer catalog available after this paper is accepted for publication.

The layout of this paper is as follows. In Section 2, we describe the SDSS data used for this work, followed in Section 3 with an overall outline of the redMaPPer cluster finder. In Section 4, we describe the multicolor richness estimator $\lambda$, which is an update of the single-color richness estimator used in R12. In Section 5, we describe our strategy for dealing with stellar masks and regions of limited depth in the survey. In Section 6, we describe the self-training of the red sequence parameterization used to detect clusters, as well as measure their photometric redshifts, which is described in Section 7. In Section 8, we describe our new probabilistic cluster centering algorithm. Finally, in Section 9, we put all these pieces together into the redMaPPer cluster finder. The resulting SDSS DR8 redMaPPer cluster catalog is described in Section 10. Next, in Section 11, we describe a new, more accurate method of using the survey data to estimate the purity and completeness of the cluster catalog.
and in Section 12, we describe how these methods can be applied to generate a cluster detection mask over the full sky. A summary is presented in Section 13. In the appendices, we present several systematic checks, including Appendix B, which contains an estimate of the minimum number of training spectra required for an accurate red sequence calibration. A full detailed comparison of redMaPPer and other large photometric survey catalogs to X-ray cluster catalogs is presented in a companion paper (Rozo & Rykoff 2014, henceforth Paper II). When necessary, distances are estimated assuming a flat $\Lambda$CDM model with $\Omega_m = 0.27$ and $h = 1.0$ Mpc, i.e., all quoted distances are in $h^{-1}$Mpc.

2. DATA

As discussed above, the redMaPPer algorithm is designed to handle an arbitrary photometric galaxy catalog, with an arbitrary number of photometric bands ($\geq 3$). Of course, the quality of the output depends on the quality of the photometry. As a case study, in this paper we run redMaPPer on SDSS DR8 data, due to its large area and uniform coverage.

2.1. SDSS DR8 Photometry

The input galaxy catalog for this work is derived from SDSS DR8 data (Aiha et al. 2011). This data release includes more than 14,000 deg$^2$ of drift-scan imaging in the northern and southern Galactic caps. The survey edge used is the same as that used for Baryon Acoustic Oscillation Survey (BOSS) target selection (Dawson et al. 2013), which reduces the total area to $\approx 10,500$ deg$^2$ with high-quality observations and a well-defined contiguous footprint. Similarly, bad field and bright star masks are based on those used for BOSS.

The BOSS bright star mask is based on the Tycho catalog (Høg et al. 2000). However, this catalog is incomplete at the bright end. Cross-matching Tycho to the Yale Bright Star Catalog (Hoffleit & Jaschek 1991), covering 9000 of the brightest stars in the sky (mostly visible to the naked eye), we have found an extra 70 stars—including very bright stars such as Arcturus and Regulus—that obviously impacted galaxy photometry and detection. We have also found that very large, bright galaxies such as M33 cause significant problems for photometry in the area, including many spurious sources masked as galaxies. To handle these issues, we have visually inspected and masked obviously bad regions around 63 objects brighter than $V < 10$ from the New General Catalogue (NGC; Sinnott 1988) that are in the DR8 footprint, as well as the bright stars mentioned above. In total, an additional 36 deg$^2$ ($\approx 0.3\%$ of the total area) is removed by our combined bright star and galaxy mask. After accounting for all the masked regions, the input galaxy catalog covers 10,400 deg$^2$.

As discussed in R12, the careful selection of a clean input galaxy catalog is required for proper cluster finding and richness estimation. Our input catalog cuts are similar to those from Sheldon et al. (2012) used for BOSS target selection, with some modifications. First, we select galaxies as classified by the default SDSS star/galaxy separator. We further limit our catalog to $i < 21.0$, approximately the 10σ limiting magnitude for the survey.\footnote{Although we note that the limiting magnitude is not precisely constant over the survey.} We then filter all objects with any of the following flags set in the $g$, $r$, or $i$ bands: SATURATED, CENTER, BRIGHT, TOOL_MANY, PEAKS, and (NOT BLENDDED) NODEBLEND. Unlike the BOSS target selection, we have chosen to keep objects flagged with SATURATED, NOTCHECKED, and PEAKCENTER.

Particular care has to be made in avoiding overaggressive flag cuts because of the way that the SDSS photo pipeline handles dense regions such as cluster cores. In these cases, the central galaxy (CG) and many satellites may be originally blended into one object and then deblended. However, if there is a problem with one part of the parent object—such as a cosmic-ray hit that is not properly interpolated—then this bad flag is propagated to all the children. We have found that removing objects marked with SATURATED, NOTCHECKED, and PEAKCENTER often masks out cluster centers, while truly saturated objects such as improperly classified stars are also rejected via the SATUR_CENTER flag cut. Overall, by including these objects we increase the number of galaxies in the input catalog by less than 2%, and our tests have shown no significant effect on the richness measurements except for a few clusters for which the cores were inadvertently masked out when galaxies with the above flags were removed. In total, there are 56.5 million galaxies in the input catalog.

In this work, we use CMLDEL_MAG as our total magnitude in the $i$ band, and MDEL_MAG for $u$, $g$, $r$, $i$, and $z$ when computing colors. We limit our input catalog to galaxies that have $m_i < 21.0$, approximately the 10σ limit for galaxy detection such that the characteristic magnitude error for our faintest objects is $\approx 0.1$. The DR8 uncertainty procedure yields magnitude uniformity on the order of 1% in $griz$ and 2% in $u$. The resulting color scatter introduced by the photometry is significantly narrower than the width of the cluster red sequence. All magnitudes and colors are corrected for Galactic extinction using the dust maps and reddening law of Schlegel et al. (1998, SFD).

2.2. Spectroscopic Catalog

Although our cluster finder uses only photometric data, we require spectroscopic data to calibrate the red sequence and to validate our photometric redshifts. For this purpose we use the SDSS DR9 spectroscopic catalog (Ahn et al. 2012). This spectroscopic catalog has over 1.3 million galaxy spectra, including over 500,000 luminous red galaxies (LRGs) at $z \approx 0.5$ from the CMASS sample. As detailed below, we only use $\approx 20\%$ of the available data in our training and use the remaining data set to validate our photometric redshifts.

3. OUTLINE OF THE CLUSTER FINDER

The redMaPPer algorithm finds optical clusters via the red sequence technique. More specifically, it is built around the optimized richness estimator $\lambda$ developed in R09 and R12. The algorithm is divided into two stages: a calibration stage, where we empirically calibrate the properties of the red sequence as a function of redshift, and a cluster-finding stage, where we utilize our calibrated model to identify galaxy clusters and measure their richness. The algorithm is iterative. First, an initial rough color calibration is used to identify clusters. These clusters are then used to better calibrate the red sequence, which enables a new cluster-finding run (see also Blackburne & Kochanek 2012 for a similar approach within the context of cluster finding with spectroscopic data sets). These two calibration/cluster-finding stages are iterated several times before a final cluster-finding run is made.

The calibration itself is also an iterative procedure described in detail in Section 6. We start with a set of “training clusters” that have a red CG with a spectroscopic redshift to calibrate the red-sequence model. As we show in Appendix B, our
minimal training requirements for unbiased cluster richness and photometric redshift estimation are ≃40 clusters per redshift bin of width ±0.025. In the case of SDSS DR8, the spectroscopic availability greatly surpasses this requirement by many orders of magnitude. However, for upcoming surveys such as DES probing much higher redshifts, this will no longer be the case, and we have developed redMaPPer with these limitations in mind.

For the present work on DR8, we construct our sample of training clusters from red spectroscopic galaxies. These red galaxies are used as “seeds” to look for significant overdensities of galaxies of the same color. The significant overdensities thus become our training clusters that are used to fit a full linear red sequence model (including zero point, tilt, and scatter) to the sample of all high-probability cluster members with a luminosity $L \geq 0.2 L_\ast$. This luminosity threshold is optimal for richness measurements (R12). In this way we effectively transfer the “seed” spectroscopic redshift to all high-probability cluster members, which enables a much more accurate measurement of the red sequence. This is especially true for fainter magnitudes where there is very limited spectroscopic coverage. Note that because the algorithm utilizes all colors simultaneously, the “scatter” is characterized not by a single number but by a covariance matrix.

Given a red sequence model, the cluster finding proceeds as follows (see Section 9 for details). First, we consider all photometric galaxies as candidate cluster centers (thereby assuming that the center of a cluster is located at a galaxy position). We use our red sequence model to calculate a photometric redshift for each galaxy ($z_{\text{red}}$; see Section 7.1) and evaluate the goodness of fit of our red galaxy template. Galaxies that are not a reasonable fit to the red galaxy template at any redshift are not considered as possible CGs for the purpose of cluster ranking. We note that as long as a cluster has at least one galaxy that is a reasonable fit to the template, that cluster will be considered in the first step of the cluster-finding stage. We then use the $z_{\text{red}}$ value of the candidate CG as an initial guess for the cluster redshift. Based on this redshift, we compute the cluster richness $\lambda$ and its corresponding likelihood using a multicolor generalization of the method of R12 (see Section 4). When a significant number of red sequence galaxies ($\geq3$) are detected within a 500 $h^{-1}$ kpc aperture, we reestimate the cluster redshift by performing a simultaneous fit of all the high-probability cluster members to the red-sequence model. This procedure is iterated until convergence is achieved between member selection and cluster photometric redshift ($z_{\text{cl}}$; see Section 7.2). The resulting list of candidate cluster centers is then rank-ordered according to likelihood. Starting with the highest-ranked cluster we measure the richness and membership probabilities. These probabilities are then used to mask out member galaxies for lower-ranked clusters in a process we term “percolation” (see Section 9.3). In this way we prevent double-counting of galaxy clusters.

4. RICHNESS ESTIMATOR $\lambda$.

The redMaPPer richness estimator, $\lambda$, is a multicolor extension of the richness estimator developed in R09 and R12, which we now use to denote $\lambda_{\text{col}}$ to indicate that it is a single-color richness. Here we review how we calculate $\lambda$ and highlight the differences relative to R12.

Let $x$ be a vector describing the observable properties of a galaxy (e.g., multiple galaxy colors, $i$-band magnitude, and position). We model the projected distribution within and around clusters as a sum $S(x) = \lambda u(x | \lambda) + b(x)$, where $\lambda$ is the number of cluster galaxies, $u(x | \lambda)$ is the density profile of the cluster normalized to unity, and $b(x)$ is the density of background (i.e., nonmember) galaxies. The probability that a galaxy found near a cluster is actually a cluster member is simply

$$p_{\text{mem}} = p(x) = \frac{\lambda u(x | \lambda)}{\lambda u(x | \lambda) + b(x)}. \tag{1}$$

We note that in Section 9.3, the definition of the membership probability will be modified to allow for proper percolation of the cluster finder. This modification will only impact clusters that are close to each other along the line of sight and at comparable redshifts. Regardless, the total number of cluster galaxies $\lambda$ must satisfy the constraint equation

$$\lambda = \sum p(x | \lambda) = \sum_{R < R_c(\lambda)} \frac{\lambda u(x | \lambda)}{\lambda u(x | \lambda) + b(x)}. \tag{2}$$

The corresponding statistical uncertainty in $\lambda$ is given by

$$\text{Var}(\lambda) = \sum p(x | \lambda) [1 - p(x | \lambda)]. \tag{3}$$

In principle, these sums should extend over all galaxies. In practice, one needs to define a cutoff radius $R_c$ and a luminosity cut $L_{\text{cut}}$. In R12 and Rozo et al. (2011) we showed that the scatter in the mass–richness relation is expected to be minimized when $L_{\text{cut}} = 0.2 L_\ast$, while the optical radial cut scales with richness via

$$R_c(\lambda) = R_0(\lambda/100.0)^{\beta}, \tag{4}$$

where $R_0 = 1.0 h^{-1}$ Mpc and $\beta = 0.2$. We adopt these parameters in redMaPPer.

To determine the cluster richness of a galaxy cluster, note that $\lambda$ is the only unknown in Equations (2) and (4). Therefore, we can numerically solve Equation (2) for $\lambda$ using a zero-finding algorithm. The solution to Equation (2) defines $\lambda$ and naturally produces a cluster radius estimate $R_c$ via Equation (4). We emphasize that this cluster radius is not a proxy for any sort of standard overdensity radius such as $R_{200c}$ or $R_{200m}$.

As in R12, we consider three observable properties of galaxies for our filter function $u(x); R_c$, the projected distance from the cluster center; $m_i$, the galaxy $i$-band magnitude; and a color variable. Ideally, our color variable would be the full color vector (e.g., $c = (u - g, g - r, r - i, i - z)$ in the case of SDSS data). However, practical considerations forced us to reduce this information to a single $\chi^2$ value that gives the goodness of fit of our red sequence template. In doing so, we effectively compress the information contained in the multidimensional color vector into a single number that measures the “distance” in color space between the galaxy of interest and our red-sequence model. This is described in more detail below.

We adopt a separable filter function

$$u(x) = [2\pi R\Sigma(R)]\phi(m_i)\rho(\chi^2), \tag{5}$$

where $\Sigma(R)$ is the two-dimensional cluster galaxy density profile, $\phi(m)$ is the cluster luminosity function (expressed in apparent magnitudes), and $\rho(\chi^2)$ is the $\chi^2$ distribution with $\nu$ degrees of freedom. The pre-factor $2\pi R$ in front of $\Sigma(R)$ accounts for the fact that given $\Sigma(R)$, the radial probability density distribution is $2\pi R \Sigma(R)$. We summarize below the filters used in redMaPPer.

$\text{R}_{200c}$ ($\text{R}_{200m}$) is the radius enclosing an overdensity of 500 (200) times the critical (mean) density of the universe.
4.1. The $\chi^2$ Filter

Assume we have a multicolor red sequence model for which we have $(c_i[z, m_i])$, the mean color of the red sequence galaxies for any given redshift $z$ and $i$-band magnitude $m_i$. Furthermore, assume we have a corresponding covariance matrix $C_{\text{int}}(z)$ to describe the intrinsic scatter and correlations of galaxy colors about the mean.

When comparing a given galaxy with color vector $c$ to the model color, with the assumption of Gaussian errors the distribution of galaxies will be represented by a $\chi^2$ distribution:

$$\chi^2(z) = (c - (c[z, m_i]))(C_{\text{int}}(z) + C_{\text{err}})^{-1}(c - (c[z, m_i])), \quad (6)$$

where $c$ is the color vector of the galaxy under consideration, $m_i$ is the observed galaxy magnitude, $(c[z, m_i])$ is the model color, and $C_{\text{int}}(z)$ is the corresponding covariance matrix, which itself depends on redshift. The matrix $C_{\text{err}}$ describes the photometric error of the galaxy under consideration.

For red sequence cluster members, $\chi^2$ will be distributed according to the $\chi^2$ distribution with $v$ degrees of freedom, $v$ is the number of colors employed when estimating $\chi^2$. Note that for $v = 1$ the $\chi^2$ filter does not reduce to the single color filter of R12. This is because our distance measurement $\chi^2$ does not distinguish between galaxies that are too red and galaxies that are too blue, so there is some loss of information when moving from color space to $\chi^2$. While a full $v$-dimensional Gaussian color filter would work better than our $\chi^2$ filter—and would exactly reduce to the single color $\chi^2$ filter in R12 when $v = 1$—the problem of background estimation for such a filter is much more difficult. In particular, in the case of DR8, it requires one to estimate the galaxy density in a five-dimensional space: $[m_i, u - g, g - r, r - i, i - z]$. We found these background estimates to be very noisy, so we compressed the color information to a single variable $\chi^2$. In this way, at any given redshift the background depends only on $m_i$ and $\chi^2$.

4.2. The Radial and Luminosity Filters

For the radial filter, we follow R09 and R12 and adopt a projected NFW profile (Navarro et al. 1995), which is a good description of the dark matter profile in N-body simulations. In addition, it has been found to be a good description of the radial distribution of cluster galaxies (Lin & Mohr 2004; Hansen et al. 2005; Popesso et al. 2007). In R12 it was shown that in order to minimize the scatter in the mass–richness relation the NFW filter works as well as or better than other possible radial profiles. Therefore, we refer readers to Section 3.1 of R12 for details on the form of the radial filter.

For the luminosity filter, we similarly follow R09 and R12 and adopt a Schechter function (e.g., Hansen et al. 2009), written as

$$\phi(m_i) \propto 10^{-0.4(m_i - m_*)(\alpha + 1)} \exp(-10^{-0.4(m_i - m_*)}). \quad (8)$$

In an update from R12, we have set $\alpha = 1.0$ independent of redshift, which provides a better description of the data. The characteristic magnitude, $m_*$, is the same as used in R12, calculated for a $k$-corrected passively evolving stellar population (Koester et al. 2007b). In the redshift range $0.05 < z < 0.7$, appropriate for DR8, $m_*(z)$ is well approximated ($\delta < 0.02$ mag) by the following polynomials:

$$m_*(z) = \begin{cases} 
22.44 + 3.36 \ln(z) + 0.273 \ln(z)^2 & \text{if } z \leq 0.5, \\
-0.0616 \ln(z)^3 - 0.0227 \ln(z)^4 & \text{if } z > 0.5, \\
-27.11 \ln(z)^3 - 18.02 \ln(z)^4 
\end{cases} \quad (9)$$

For each cluster, $m_*$ is taken at the appropriate redshift and the luminosity filter is normalized to unity at the appropriate magnitude cutoff. As with R12, this is taken to be $0.2 L_*$, or $m_* + 1.75$ mag. Although in the current version of redMaPPer both $\alpha$ and $m_*$ are fixed as described above, in future releases we will replace these parameters with those directly measured from calibration clusters. We emphasize, however, that modest changes to the shape of the luminosity filter result in insignificant changes to the recovered richness. Of course, changes to the magnitude limit above which one counts galaxies have an obvious systematic impact on the richness as one moves up and down the luminosity function, although we have found that these modest shifts do not significantly impact the mass–richness scatter (see R12).

4.3. Background Estimation

As in R12, we assume that the background density is uniform, such that $b(x|z) = 2\pi R_0 \Sigma_p(m_i, \chi^2|z)$, where $\Sigma_p(m_i, \chi^2|z)$ is the galaxy density as a function of galaxy $i$-band magnitude and $\chi^2$, where $\chi^2$ is evaluated using the red sequence model at redshift $z$. In this way, the effective background for every cluster is different and depends on the cluster redshift. Note that since clusters reside in overdense regions, this mean background is likely an underestimate of the true background, but the incurred bias is small (see R12 and Rozo et al. 2011) and is irrelevant for cosmological purposes, as it is completely degenerate with the amplitude of the richness–mass relation. Variability of the background (which is expected) is more problematic and can lead to catastrophic projections. These are expected to be small (Rozo et al. 2011) and will be addressed in a future paper (E. Rozo et al., in preparation).

To calculate the mean galaxy density, we first calculate the $\chi^2$ value for all galaxies in a grid of redshifts with spacing 0.02. For computation purposes, we only calculate the $\chi^2$ for galaxies that are brighter than 0.1 $L_*$ at a given redshift bin. This implies that the magnitude range being sampled is different for each redshift bin. At each redshift we bin the full galaxy catalog in $\chi^2$ and magnitude using a cloud-in-cells (CIC) algorithm (e.g., Hockney & Eastwood 1981) and divide by the survey area. For our cells, we use $\chi^2 \in [0, 20]$ with a bin size of 0.5, and $i \in [12, m_{\text{lim}}]$ with a bin size of 0.2. The $\chi^2 < 20$ cut can be justified by the fact that the sum total of cluster membership probability in the redMaPPer catalog for galaxies with $\chi^2 \in [15, 20]$ is only 0.7%. That is, our $\chi^2 < 20$ cut impacts our results at well below the 1% level. The final galaxy number density is normalized by the width of each color and magnitude bin. To evaluate the background at an arbitrary redshift, we linearly interpolate between the backgrounds computed along our redshift grid. As noted in R12, because the background is measured per square degree, the average number of background galaxies as a function of $\chi^2$,
magnitude, and redshift is automatically accounted for as the angular size of the clusters changes with redshift.

5. HANDLING MASKED REGIONS AND LIMITED DEPTH

In an ideal world, our survey would have uniform depth, be deep enough to reach 0.2 L∗ at all redshifts of interest, and there would be no missing and/or masked regions, e.g., due to bright stars. Most previous optical cluster finders make this simple assumption.16 Here, we describe how we can properly correct for these effects. Our approach is conceptually straightforward. Given a cluster model and a geometric and magnitude mask, we can effectively calculate the fraction of cluster galaxies that we expect to be masked out. This correction factor is then applied to the “raw” richness to compensate for the masked region. In practice, this correction can be self-consistently incorporated into the richness estimation, as described below. Our method is simple to implement with any geometric mask, including those that describe variations in depth. However, we do not take into account masks that contain one or more missing bands.

5.1. The Correction Term

Looking back at Equation (2), we have

\[ 1 = \sum_i \frac{u(x_i)}{\lambda u(x_i) + b(x_i)}. \]  

(10)

where \( x_i \) describes the radial position, color (via \( \chi^2 \), and luminosity (via \( m_i \)) of each galaxy. This formulation works if we can see all galaxies, but in reality we cannot. Let us then pixelize all observable space \( x \) into infinitesimal pixels, and let \( N_i \) be the number of galaxies in pixel \( i \). Most pixels have \( N_i = 0 \), but a few have \( N_i = 1 \). Thus, the sum over all galaxies can be rewritten in terms of a sum over all pixels via

\[ 1 = \sum_i N_i \frac{u(x_i)}{\lambda u(x_i) + b(x_i)}. \]  

(11)

In the case of masking, we can only observe the galaxies that are inside the mask, so we can split this sum into

\[ 1 = \sum_{\text{in}} N_i \frac{u(x_i)}{\lambda u(x_i) + b(x_i)} + \sum_{\text{out}} N_i \frac{u(x_i)}{\lambda u(x_i) + b(x_i)}. \]  

(12)

The “in” term is the raw \( \lambda \) that we usually compute, and it can be replaced by the standard sum over all observed galaxies. The “out” term is now a correction to the standard expression, call it \( C \),

\[ C = \sum_{\text{out}} N_i \frac{u(x_i)}{\lambda u(x_i) + b(x_i)}. \]  

(13)

In terms of \( C \), Equation (2) can be rewritten as

\[ 1 - C = \sum_{\text{gals}} \frac{u(x)}{\lambda u(x) + b(x)}. \]  

(14)

Now, while \( C \) is unknown (we cannot see the masked region), we can compute its expected value for a cluster of richness \( \lambda \). Using the fact that

\[ \langle N_i \rangle = [\lambda u(x_i) + b(x_i)] \Delta x_i, \]  

(15)

we see that the expectation value of \( C \) is given by

\[ \langle C|\lambda \rangle = \int_{\text{out}} dx \ u(x). \]  

(16)

In the above equation, we have made explicit the fact that \( C \) depends on \( \lambda \), both via the cutoff radius used in the sum over galaxies and because the radial filter depends on \( \lambda \). Thus, in the presence of missing data, our richness estimate is given by the solution to

\[ 1 - \langle C|\lambda \rangle = \sum_{\text{gals}} \frac{u(x)}{\lambda u(x) + b(x)}. \]  

(17)

Note, however, that because \( C \) is an unknown, there must also be additional measurement error associated with this unknown correction. To calculate the variance of \( C \), we note that \( \text{Var}(N_i) = \langle N_i \rangle \). To compute \( \text{Cov}(N_i, N_j) = \langle N_i N_j \rangle - \langle N_i \rangle \langle N_j \rangle \), we first compute \( \langle N_i N_j \rangle \). For infinitesimal pixels, \( N_i N_j = 1 \) implies that both pixels \( i \) and \( j \) contain cluster galaxies, or that one pixel contains a cluster galaxy while the other pixel contains a background galaxy, or that both pixels contain a background galaxy. Consequently,

\[ P(N_i, N_j = 1) = \Delta x^2 \lambda(\lambda - 1)u_i u_j + \lambda(\lambda - 1)(u_i b_j + u_j b_i + b_i b_j). \]  

(18)

Since \( \langle N_i N_j \rangle = P(N_i, N_j = 1) \), subtracting off \( \langle N_i \rangle \langle N_j \rangle \), we arrive at

\[ \text{Cov}(N_i, N_j) = -\Delta x^2 u_i u_j \]  

(19)

for \( i \neq j \). Putting it all together, we arrive at

\[ \text{Var}(C) = \sum_i \langle N_i \rangle p_i - \sum_i \sum_{j \neq i} \text{Cov}(N_i, N_j) p_i p_j \]  

(20)

\[ = \int_{\text{out}} dx \ u(x) p(x) - \left[ \int_{\text{out}} dx \ u(x) p(x) \right]^2. \]  

(21)

To propagate the error in \( C \) into the error in \( \lambda \), we set

\[ \sigma_\lambda = \left. \frac{d\lambda}{dC} \right|_{\lambda(\lambda)} \sigma_C, \]  

(22)

where \( \sigma_C = [\text{Var}(C)]^{1/2} \). The derivative of \( \lambda \) with respect to \( C \) is evaluated numerically about the expectation value of \( \langle C|\lambda \rangle \), where \( \lambda \) is the solution to Equation (17).

For future reference, it will be useful to define the “scale factor”

\[ S(z) = \frac{1}{1 - \langle C|z \rangle} \]  

(23)

for the case in which the only source of masking is due to limited depth. With this definition, a cluster with richness \( \lambda \) has a total of \( \lambda / S(z) \) galaxies above the limiting magnitude of the survey.

In addition, it is useful to calculate the fraction of the effective cluster area that is masked solely by geometrical factors such as bright stars, bad fields, and survey edges. This is complementary to \( S(z) \) defined above in that it contains all the masking except the magnitude limit. The cluster mask fraction is then

\[ f_{\text{mask}} = \frac{\int_{\text{out}} dx \ u(x)}{\int dx \ u(x)}. \]  

(24)

This quantity is very useful because clusters that are strongly affected by edges are more likely to be catastrophically miscentered and to have poor richness estimates. Consequently, when defining our cluster catalog, we will apply a cut in \( f_{\text{mask}} \).
5.2. Evaluating the Mask Correction

Evaluation of the mask correction and its associated error on $\lambda$ can be difficult. Fortunately, this problem is well suited to Monte Carlo integration. First, define the selection function $S(x)$, so that $S(x) = 1$ if the galaxy is in a region where it is detected, and $S(x) = 0$ if the position $x$ is masked out. Then, for any function $f(x)$,

$$
\int_{\text{out}}^{} f(x) = \int_{\text{cluster}}^{} [1 - S(x)] f(x) dx,
$$

(25)

where the integral on the right-hand side is over the full cluster region, i.e., $R \in [0, R(\lambda)]$, $L \geq L_{\text{min}}$, and over all colors.

Applying this to Equation (16), we find

$$
\langle C \rangle = \int_{\text{cluster}}^{} [1 - S(x)] u(x) dx.
$$

(26)

Since $u(x)$ is the probability distribution for $x$, we can evaluate the above integral using Monte Carlo integration by randomly sampling $N$ sets of model galaxy parameters from the $u(x)$ filter function and then computing the sample mean of the function $1 - S(x)$. That is, $\langle C \rangle$ is simply the fraction of random draws that fall in the masked region. Similarly, we can evaluate the integrals defining $\text{Var}(C)$ via

$$
\text{Var}(C) = \frac{1}{N} \sum_{i=1}^{N} p(x_i) - \left( \frac{1}{N} \sum_{i=1}^{N} p(x_i) \right)^2,
$$

(27)

where $N$ is again the total number of random draws. For simplicity, we have replaced the $1 - S(x)$ terms with a summation over all galaxies that are outside the detectable region due to the mask, as these are the only galaxies that contribute to the summation.

One of the slowest part of this process is drawing random realizations of $x_i$. However, these random draws do not need to be independent from cluster to cluster. In practice, we generate a template distribution of 5000 galaxies and scale the radius and magnitude to the appropriate values for each galaxy cluster. We find that this number of galaxies gives accurate results for the recovered richnesses and richness errors, except for galaxy clusters that are largely masked out. Consequently, our final cluster selection criteria include the requirement that the cluster mask fraction, $f_{\text{mask}}$, is less than 20%. In Appendix E, we demonstrate with DR8 data that given our filter function this formalism accurately corrects for masked regions and limited depth.

6. CALIBRATION OF THE RED SEQUENCE

6.1. Outline

Suppose we have a complete sample of red galaxies with spectroscopic redshifts down to the limiting magnitude of the survey. One can then directly fit a red sequence model to these galaxies to calibrate the color as a function of redshift for these galaxies. The question then becomes, how does one get a sample of red spectroscopic galaxies? Note that in order to calibrate the tilt of the red sequence, it is important to include a significant number of less luminous galaxies, which are more difficult to come by.

Our solution is to simply use the cluster members themselves. If we know the spectroscopic redshift of a cluster, then all the cluster members can be tagged with the spectroscopic redshift of their CG, leveraging one spectroscopic redshift into many. Of course, from photometric data one can only identify likely cluster members, so the fit of the red sequence model must account for contamination by non-cluster members.

In order to fit the red-sequence model, all that is required is a sample of galaxy clusters with known spectroscopic redshifts. As discussed in Section 3, we only require a limited set of these training clusters. These training clusters can be derived from external X-ray and Sunyaev–Zel’dovich (SZ) catalogs, or from spectroscopic follow-up of likely centrals in dense regions by running redMaPPer with an ad hoc red sequence model.

In the specific case of SDSS DR8, we construct this training cluster sample based on existing spectroscopic galaxy samples. Each spectroscopic galaxy is used as a “seed” around which we look for galaxy clusters by identifying nearby overdensities of galaxies with the same color as the seed galaxy. The details of these steps are described below. We emphasize that our calibration is performed using only 2000 deg$^2$ of SDSS data. As we show in Appendix B, while we have the full wealth of SDSS spectroscopic data at our disposal, an equivalent red-sequence model may be derived from only 400 CGs from $z \in [0.05, 0.6]$.

6.2. Selecting Seed Galaxies and the Initial Color Model

The initial calibration of the red sequence relies on spectroscopic “seed” galaxies. This may simply be a set of training clusters with spectroscopy (see Appendix B), or in the case of DR8, a broad spectroscopic sample that contains a sufficient number of red galaxies in galaxy clusters. For SDSS spectroscopy, the first step is to identify the subsample of spectroscopic galaxies that are red. This is achieved by using a single color that samples the 4000 Å break for early-type galaxies. With SDSS data, we use $g - r$ for $z < 0.35$ and $r - i$ for $z > 0.35$. Because we wish to have a relatively clean selection of red-sequence galaxies as our seeds, we approach the problem of selecting these galaxies in several steps. We emphasize that some of these steps are only necessary for cutting the full list of SDSS spectra to an appropriate red galaxy sample.

Step 1: Perform an approximate red galaxy selection. To make this selection, we bin the galaxies in redshift bins of width ±0.025. We then use the error-corrected Gaussian Mixture Method (Hao et al. 2009) to estimate the mean and intrinsic width $\sigma_{\text{int}}(z)$ of the red-sequence galaxies. Those galaxies within 2$\sigma$ of the expected mean for red galaxies are considered approximately red.

Step 2: Use this approximate red galaxy selection to measure the mean color of the red galaxies as a function of redshift. Given our approximate red galaxy sample, we refine our initial estimate of the mean color–redshift relation by minimizing the function

$$
s = \sum_{i} |c_i - \tilde{c}(z_i)|,
$$

(28)

where $\tilde{c}(z)$ is our model for the color as a function of redshift. The function $\tilde{c}(z)$ is defined via spline interpolation, and the values of the spline nodes are the parameters with respect to which $s$ is minimized. The spline nodes are placed on a redshift grid with spacing of 0.1. Our use of $\tilde{c}(z)$ indicates that this is our early calibration model, which is distinct from the full model color $c(z)$ that is derived at the end of the calibration procedure. In defining the function $s$, we rely on the sum of absolute values rather than the sum of the squares to make the resulting minimization more robust to gross outliers.
The function $s$ is minimized using the downhill-simplex method of Nelder & Mead (1965) as implemented in the IDL AMOEBA function.

**Step 3:** Use the mean color–redshift relation to estimate the width of the color–redshift relation. We can now improve upon our initial estimate of the width of the color–redshift relation by minimizing the function

$$s = \sum_i |c_i - \tilde{c}(z_i)| - \text{MAD},$$

where $\tilde{c}(z) = 1.4826 \times \text{MAD}$, where MAD is the median absolute deviation of the sample about the median, and the factor of 1.4826 relates the MAD to the standard deviation for a Gaussian distribution. The value MAD is again defined via spline interpolation, with the free parameters being the values of the function at the nodes.

**Step 4:** Generate a final sample of seed galaxies. Finally, with the full red spectroscopic galaxy model in hand ($\tilde{c}(z), \tilde{\sigma}_{\text{int}}(z)$), we can cleanly select our seed sample. We select all galaxies within $2\tilde{\sigma}_{\text{int}}(z)$ of the model color at the spectroscopic redshift of the galaxy, using $g - r$ at $z < 0.35$ and $r - i$ at $z > 0.35$.

In Figure 1, we show the final seed spectroscopic galaxy selection for the $g - r$ and $r - i$ colors. The large red points show the median colors at the node positions, and the dashed red lines show the cubic spline interpolation. Note that the single-color selection leaves a small number of outliers in the complementary color, as well as a small number of large outliers with anomalously large photometric errors.

(A color version of this figure is available in the online journal.)

![Figure 1. Sample of red spectroscopic galaxies selected for training in 2000 deg$^{-2}$ of DR8. Top panel shows $g - r$ color and bottom panel $r - i$ color. The red galaxy selection is done in $g - r (r - i) < 0.35 (z > 0.35)$, selecting all galaxies within $2\tilde{\sigma}_{\text{int}}(z)$ of the spectroscopic redshift of the galaxy. Note that this selection leaves a small number of outliers in the complementary color, as well as a small number of large outliers with anomalously large photometric errors.](http://example.com/figure1.png)

6.3. Single Color Member Selection

Having selected our seed galaxies and calibrated a rough initial color–redshift relation, we now proceed to find likely cluster members around each of our seed galaxies. For this first iteration, we rely on single-color-based membership. Specifically, in R12, we demonstrated that for moderately rich ($\lambda \gtrsim 20$) clusters, one can reliably estimate the red sequence directly from the data as follows. First, we select all galaxies within a color window around the seed galaxy. Next, we fit for the amplitude and tilt of the red sequence of that galaxy cluster directly from the galaxy data. However, in extending this algorithm to high redshift, we found that large photometric errors can introduce an unacceptable amount of noise in the initial color box selection of galaxies. Therefore, rather than drawing a color box around the color of the CG for the initial fit, we draw the color box around the model color $\tilde{c}(z)$ calibrated in the previous section. In detail, we perform the following.

1. Take a red galaxy of known spectroscopic redshift (the "seed").
2. Select all galaxies within $500 h^{-1}$ kpc of the spectroscopic galaxy, as well as $2\sigma$ of the model color determined in Section 6.2. For the model color, we use $g - r$ at $z \lesssim 0.35$ and $r - i$ at $z \gtrsim 0.35$. The width $\sigma$ of the color box is set to 0.05 and 0.03, respectively, which we expect to be the approximate red sequence width (e.g., R12).
3. Fit the red sequence (slope and intercept) of these galaxies.
4. Measure the single-color $\lambda_{\text{col}}$ using the method of R12 and a fixed aperture of $500 h^{-1}$ kpc.
5. For all overdensities with $\lambda_{\text{col}} > 10$, take the galaxies with nonzero $p_{\text{mem}}$ and assign them the spectroscopic redshift of the initial seed galaxy. In practice, we limit our analysis to those galaxies with $p_{\text{mem}} > 0.7$.

At this point, we have leveraged the spectroscopic seed galaxies to generate a set of red galaxies as faint as $0.2 L_*$ over the redshift range of interest. Although not all of these galaxies are true cluster members, we have an estimate $p_{\text{mem}}$ of the probability that each such galaxy is indeed a red-sequence cluster member, as in Equation (1). Consequently, we can model the contamination of non-red-sequence galaxies in our sample, as shown below.

We emphasize that it is essential that we leverage our spectroscopic redshifts to fainter magnitudes to properly model the red sequence. In the case of DR8, our initial seed galaxy sample is composed of 42,000 galaxies associated with $\lambda > 5$ clusters, almost all of which are preferentially bright. By contrast, our final calibration sample (see Section 6.5) is composed of over 600,000 red sequence galaxies that extend to much fainter magnitudes. This is illustrated in Figure 2. The magnitude distribution of our seed galaxies in the redshift slice $z \in [0.24, 0.26]$ (solid black histogram) is contrasted with the membership-weighted magnitude distribution of our final photometrically selected training sample (red dashed histogram). We see that the gain in the effective number of red sequence training galaxies is enormous, allowing for an accurate calibration of the red sequence (amplitude, tilt, and scatter) as a function of redshift. We have explicitly verified that modest changes to the cuts applied in this section do not impact our final calibration of the red sequence resulting from the subsequent analysis described in the next section.

6.4. Modeling the Red Sequence

Given a list of galaxies with multidimensional color ($c$), redshift ($z$), taken to be the cluster redshift), and membership probability ($p_{\text{mem}}$), we can now proceed to calibrate the full red sequence model. Our model is well motivated by observations of galaxy clusters, in that the red sequence at any given redshift in a given color can be described by a simple linear relation between color and $i$-band magnitude $m_i$ with intrinsic scatter $\sigma_{\text{int}}$. For example, Figure 3 shows the composite red sequence...
at $z = 0.25$ for both $g-r$ and $r-i$ colors, for all galaxies selected in the final calibration iteration with $p_{\text{mem}} > 0.9$.

Our red-sequence model is defined in terms of smoothly evolving functions of redshift characterizing the amplitude and slope of the mean color–redshift relation and the corresponding covariance matrices. We have opted to use a cubic spline interpolation to parameterize these functions. Given the large number of colors (four for SDSS) and broad redshift range, our model necessarily contains a large number of free parameters. For instance, in our SDSS DR8 implementation, we required a total of 118 parameters to fully characterize the red sequence model. In principle, we would like to fit the full red sequence model simultaneously. However, to make the problem more tractable, we fit the red-sequence parameters governing the mean relation and the diagonal elements of the covariance matrix one color at a time. Once these terms are in place, we fit the off-diagonal terms of the covariance matrix. We are also cautious that our model does not have too many free parameters given the training data such that overfitting becomes possible. As shown in Appendix B, this can be a problem in the case of very sparse training data.

An additional complication comes from the fact that our selection of red-sequence galaxies is not entirely clean. Our fit of the red sequence must take into account the background density of nonmember galaxies, as described below. In addition, we also have to contend with blue cluster galaxies that are not taken into account by a global background term. These blue galaxies will tend to have two effects. First, as the blue fraction increases at lower luminosities, they will tend to steepen the apparent red sequence tilt. Second, the blend of red and blue galaxies will tend to broaden the apparent intrinsic width of the red sequence.

In order to deal with both of these effects of blue cluster galaxies, we have taken a pragmatic approach. When fitting the red sequence for a given color, we first perform a sharp color cut to concentrate on the core of the red galaxy distribution.

---

**Figure 2.** Magnitude distribution of the red spectroscopic seed galaxies (black histogram) and photometrically selected cluster galaxies with $p_{\text{mem}} > 0.7$ (red dashed histogram) in the redshift slice $z \in [0.24, 0.26]$. The “cluster member” histogram weights each galaxy by its membership probability. The gain in the training sample of faint red-sequence galaxies through our photometric selection is of critical importance for an accurate calibration of the red sequence as a function of redshift, particularly at the faint end.

(A color version of this figure is available in the online journal.)

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**Figure 3.** Composite red sequence at $0.25 < z < 0.26$ for color-selected galaxies with $p_{\text{mem}} > 0.9$. A linear model (red dashed line) with roughly constant intrinsic scatter is a good representation of the red sequence in both $g-r$ and $r-i$. We note that the $p_{\text{mem}} > 0.9$ cut is employed for illustration purposes only.

(A color version of this figure is available in the online journal.)

Naively, this cut would introduce biases in the recovered red-sequence model, leading to underestimates of the scatter. We avoid this difficulty by explicitly modeling such a color cut into our likelihood function. All that remains is to specify the color cut. Here we apply a color cut of $1.5\sigma$ about the median color of the high-probability member galaxies, where $\sigma$ is the median absolute deviation of the color about the median.

### 6.4.1. Measuring the Model Mean and Color Scatter

As noted above, we begin by measuring the model color $\langle c|m_i,z \rangle$ as a function of galaxy magnitude $m_i$ and cluster redshift $z$ for each color, one at a time. The first step in this process is to define the pivot point $\vec{m}_i(z)$ used to calibrate the amplitude and tilt of the mean red-sequence relation at redshift $z$. We write

$$\langle c|z, m_i \rangle = \vec{c}(z) + \vec{s}(z)[m_i - \vec{m}_i(z)].$$  

(30)

We wish to select a pivot point that is characteristic of most cluster members. To do so, starting from our full members list, we apply a $p_{\text{mem}} > 0.7$ cut. Using this subsample, we minimize the cost function $E$, where

$$E = \sum |m_i - \vec{m}_i(z)|,$$

(31)

where $\vec{m}_i(z)$ is defined via spline interpolation, and the model parameters are the value of $\vec{m}_i(z)$ at the nodes.

Having defined our pivot point as a function of redshift, we turn to calibrating the amplitude and slope of the mean relation, i.e., $\vec{c}(z)$ and $\vec{s}(z)$ in Equation (30). As a first step, we do a rough estimate of the amplitude and scatter, which we will use to isolate the core of the color distribution of member galaxies. These rough estimates for the amplitude and scatter are denoted $\vec{c}(z)$ and $\vec{s}(z)$ and are obtained by selecting galaxies with $p_{\text{mem}} > 0.7$ and then fitting for these functions as was done in Section 6.2. Specifically, the functions are spline interpolated, with model parameters being the value of these functions at the nodes. The best-fit parameters are found by minimizing Equation (28), and $\vec{s}(z)$ is defined by minimizing Equation (29). The primary
difference between these new color estimates and scatter relative to those derived in Section 6.2 is that these parameters are now appropriate to the full red sequence rather than simply the (brightest) spectroscopic galaxies.

We now turn to measuring the actual model parameters defining the amplitude $\tilde{c}(z)$, slope $\tilde{s}(z)$, and scatter $C_{m}^{\text{int}}(z)$. As before, we use a cubic spline interpolation to parameterize these smoothly evolving functions of redshift. For DR8, we have chosen to use a node spacing of 0.05 for $\tilde{c}(z)$, 0.1 for $\tilde{s}(z)$, and 0.15 for $C_{m}^{\text{int}}(z)$. We have found that a relatively tight spacing is required for $\tilde{c}(z)$, as this function can change relatively rapidly at filter transitions. Fortunately, $\tilde{c}(z)$ is the most robust parameter and thus is amenable to smaller node spacings. The slope and scatter are not expected to vary as rapidly and are also noisier to estimate, so we have chosen wider node spacings. Overall, the calibration is not very sensitive to the node spacings chosen provided that there are sufficient calibration galaxies (though see Appendix B).

Starting from the photometrically selected galaxy training set from the previous section, we first apply a color cut $|c - \bar{c}(z)| < 1.5\tilde{c}(z)$, which ensures that the red-sequence parameters are based on the core of the red galaxy distribution and are therefore less likely to be biased by blue galaxies. In our model, the probability that a red-sequence cluster galaxy has a color $c$ is given by a truncated Gaussian distribution,

$$G(c) = \frac{1}{\sqrt{2\pi}\sigma} e^{-(c-\bar{c}(z))^2/2\sigma^2} \text{erf}(\frac{1.5\tilde{c}(z)}{\sqrt{2}\sigma}),$$

where the expectation value $\langle c|m_{i}, z \rangle$ is defined in terms of our model functions $\bar{c}(z)$ and $\tilde{s}(z)$ as per Equation (30), and the scatter $\sigma$ is the sum in quadrature of the intrinsic scatter and the photometric error of the galaxy,

$$\sigma = \sqrt{\sigma_{z}^{2} + \sigma_{\text{int}}^{2}},$$

where $\sigma_{\text{int}} = \sqrt{C_{z}^{\text{int}}}$. The erfterm in the denominator accounts for the fact that $G(c)$ is truncated at $\bar{c}(z) \pm \tilde{c}(z)$, under the approximation $\tilde{s}(z) = \tilde{c}(z)$. This approximation is only used in the overall normalization of the distribution.

The total probability distribution for all of our calibration galaxies must account for the fact that some of our galaxies are in fact background galaxies, so the full color distribution is given by

$$P(c) = p_{\text{mem}}G(c) + (1 - p_{\text{mem}})b(c, m_{i}),$$

where $b(c, m_{i})$ is the distribution in color and magnitude of galaxies about random points. The shape of the background function is obtained by binning all galaxies in color and magnitude bins and using a CIC algorithm as in Section 4.3.

In the end, our task is to calculate the set of $\bar{c}(z)$, $\tilde{s}(z)$, and $C_{m}^{\text{int}}(z)$ values at the given cubic spline nodes that maximizes the total likelihood given by

$$\ln \mathcal{L} = \sum_{i} \ln P_{i}.$$  

As above, we accomplish this maximization by making use of the downhill-simplex method. The maximum likelihood point defines the model functions $\bar{c}(z)$, $\tilde{s}(z)$, and $C_{m}^{\text{int}}(z)$. We emphasize that the likelihood is explicitly truncated as the data are, so that the recovered scatter is unbiased relative to the full population of cluster member galaxies, as we have confirmed with simple mock red sequences and blue clouds.

In Figure 4, we show the color evolution of red-sequence galaxies with $p_{\text{mem}} > 0.9$. The red points indicate the $\tilde{c}(z)$ values at the spline node positions and the long-dashed lines are the spline interpolation. The short-dashed red lines indicate the $3\sigma_{\text{int}}$ range. Note that the colors in the figure are not corrected for red sequence tilt. The cyan dash-dotted line shows the color model for the bright spectroscopic sample from Figure 1, which tend to be brighter and redder than the full population. We caution that the intrinsic width of the red sequence can be wider than the $p_{\text{mem}} > 0.9$ subsample of galaxies in this illustration suggests, since high-probability membership requires the galaxy to fall close to the expected average color. Conversely, the larger number of outliers in $g - r$ above reflects the fact that the photometric errors in $g - r$ at high redshift are larger than the intrinsic width of the red sequence.

(A color version of this figure is available in the online journal.)

6.4.2. Measuring $C_{m}^{\text{int}}(z)$

With the intercept and slope of the red sequence in hand, as well as the diagonal elements of the covariance matrix, we now estimate the off-diagonal elements of the covariance matrix, $C_{jk}^{\text{int}}(z)$. Once again, we use a cubic spline interpolation, with the same 0.15 node spacing as used for $C_{jj}^{\text{int}}(z)$.

In order to make the calculation tractable, to constrain the off-diagonal elements of the covariance matrix, we consider the problem piecewise, tackling two colors at a time. Each individual piece of the covariance matrix constrained in this way will be positive-definite and thus a valid covariance matrix. Unfortunately, due to noise in the estimation of the parameters, this method does not guarantee that the total covariance matrix, $C_{\text{int}}(z)$, will also be positive-definite.

To ensure that $C_{\text{int}}(z)$ is positive-definite, we constrain the parameters for pairs of colors in a specific priority order, ensuring that the best-constrained colors have precedence. In the case of DR8 data in the redshift range $z \in [0.05, 0.6]$, these
are \(g - r\) and \(r - i\). Then, at each step in the downhill-simplex estimation described below we do not allow any terms in \(C^\text{int}(z)\) that result in a minimum eigenvalue in the total covariance matrix \(C_\text{int}(z)\) that is less than 0.012. In this way, the first color pair to be constrained \((g - r, r - i)\) is essentially free, while the final (and noisiest) color pair to be constrained \((u - g, i - z)\) will not result in a non-invertible covariance matrix \(C_\text{int}(z)\).

To perform the pairwise constraints on the off-diagonal elements, let us consider the residuals in two colors \(x_j\) and \(x_k\). We start with Equation (30),

\[
x = c - \{c|m_i, z\} = (\tilde{c}(z) + \bar{s}(z))[m_i - \tilde{m}_i(z)].
\]  

(36)

The probability density function (pdf) is again a Gaussian, though this time we explicitly leave the covariance matrix in the equation

\[
G(x) = \frac{1}{\sqrt{2\pi}|C|^{1/2}} \exp \left[-\frac{1}{2}xC^{-1}x\right],
\]  

(37)

where \(x = \{x_j, x_k\}\) is the vector of residuals, and the total covariance matrix \(C\) is

\[
C = C_\text{int}(z) + C_\text{err}(z).
\]  

(38)

Here \(C_\text{int}(z)\) and \(C_\text{err}(z)\) are the covariance matrices characterizing the intrinsic scatter and photometric error, respectively. The intrinsic scatter is simply

\[
C_\text{int} = \begin{pmatrix}
\sigma^2_{\text{int},j} & r\sigma_{\text{int},j}\sigma_{\text{int},k} \\
r\sigma_{\text{int},j}\sigma_{\text{int},k} & \sigma^2_{\text{int},k}
\end{pmatrix},
\]  

(39)

where \(\sigma_j\) and \(\sigma_k\) are known from the previous section, and \(r\) is the only unknown. The covariance matrix \(C_\text{err}(z)\) is derived from the photometric error in each band. Given two colors \(c_j = m_\gamma - m_\beta\) and \(c_k = m_\gamma - m_\delta\), the covariance matrix characterizing the photometric error is given by

\[
C_\text{err}(z) = \begin{pmatrix}
\sigma^2_a + \sigma^2_\beta & \eta \\
\eta & \sigma^2_\gamma + \sigma^2_\delta
\end{pmatrix}
\]  

(40)

and

\[
\eta = \begin{cases}
-\sigma^2_\beta & \text{if } \gamma = \beta \\
0 & \text{otherwise}.
\end{cases}
\]  

(41)

Here we are assuming that neighboring colors are of the form \(c_{\alpha\beta}\) and \(c_{\gamma\delta}\), i.e., that the “shared” magnitude is \(m_\beta = m_\gamma\). The covariance between photometric errors arises precisely because, for example, the neighboring colors \(g - r\) and \(r - i\) are both derived from the same \(r\)-band magnitude.

The color distribution function of the full galaxy population is again given by Equation (34), noting that now the background term \(b(c_j, c_k, m_i)\) is given by a three-dimensional binning in two colors and \(r\)-band magnitude. In addition, we implement a prior on \(r\) with 0 mean and width 0.45 for each of the nodes. We find that this prior reduces the noise in the parameter constraints, which is especially important at high redshift where the photometric errors dominate and the covariance matrix is largely unconstrained. At the same time, this prior allows high correlations \((r \sim 0.9)\) if strongly favored by the data. Our total likelihood is now given by

\[
\ln L = \sum_i \ln P_i - \sum_n \frac{(r_n/0.45)^2}{2},
\]  

(42)

where \(\sum_n\) is a sum over all the nodes, and \(r_n\) is the correlation coefficient at that node. That is, the prior is placed at each of the nodes. Maximization of the likelihood function defines the final values for the correlation coefficients that characterize the intrinsic scatter covariance matrix.

6.5. Iterating the Red-sequence Model

We emphasize that the estimation of the red-sequence parameters in the previous section depends on the membership probabilities \(P_{\text{mem}}\) of the red-sequence galaxies. Of course, the membership probabilities themselves depend on the red-sequence model. In order to obtain a red-sequence model that is consistent with the membership probabilities, we take an iterative approach.

After we calibrate the red-sequence parameters based on single-color membership probabilities, we run the cluster finder on the training data, as described in Section 9. During these calibration runs, we restrict ourselves to finding clusters associated with our seed galaxies so that we can affirmatively associate a spectroscopic redshift with each cluster. We note that our cluster finder starts with the photometric redshift estimate \(z_{\text{red}}\) of each cluster galaxy, so spectroscopic galaxies whose colors are incompatible with the red sequence at the spectroscopic redshift never result in galaxy clusters. Thus, our training sample at this point results in robust clusters with spectroscopic redshifts. Further, failures in the photo-\(z\) of the galaxies for red-sequence galaxies are rare (see Figure 7), so any such failures simply slightly reduce the sample of training clusters, without otherwise adversely affecting our training sample. The resulting cluster catalog includes cluster member lists and new membership probability estimates \(P_{\text{mem}}\) based on the full color model. With these in hand we can re-estimate the red-sequence model as described in Section 6.4.

As we iterate, the largest shifts in the model occur between the first and second iteration, reflecting the shift from estimating membership probabilities based on a single color to estimating membership probabilities with the full multicolor data. Figure 5 shows the red-sequence parameters \((\bar{c}(z), \bar{s}(z), \text{and } \sigma_{\text{int}}^m)\) for each of the first three iterations of our red-sequence calibration. For illustration, only the figures for the \(g - r\) color are shown. The color at the reference magnitude \(m_\gamma(z)\) and slopes characterizing the average color of red-sequence galaxies converge quickly and are generally well measured, except for \(u - g\) at high redshift, where the large photometric errors in \(u\) make our model estimates noisy. The scatter model, on the other hand, converges slowly, particularly at high redshift, where the intrinsic scatter is often subdominant to photometric errors. As we now show, however, by the third iteration our model is well converged.

We define convergence of the red-sequence model in terms of the relevant quantity for our purposes, i.e., the cluster richness \(\lambda\). That is, we require that cluster richness estimates be insensitive to further iterations. To this end, we have run the calibration through 10 iterations. Given the red-sequence model for each of these 10 iterations, we estimate the photometric redshift and cluster richness of a standard set of galaxy clusters while fixing the CG of these systems. Let then \(\lambda_i\) and \(z_i\) denote the richness and redshift estimates, respectively, from iteration \(i\). We bin the clusters in narrow redshift slices \((\pm 0.01)\), and we calculate (1) the median ratio \(\lambda_i/\lambda_3\) and (2) the median offset \((\lambda_i - \lambda_3)/\sigma_3\), where \(\sigma_3\) is the error estimate in the richness as estimated from iteration 3.

In Figure 6, we show the results of these iteration checks for the first six iterations in the DR8 training region. Even for
7. PHOTOMETRIC REDSHIFT ESTIMATION

At the end of our calibration we have a complete red sequence model as a function of redshift. Note, however, that in order to estimate the richness of a photometric cluster we need to know the cluster redshift. If we have some initial, reasonably accurate estimate, and we iterate this procedure through convergence.

We then simultaneously fit our red sequence model to all high-richness and determine the high-probability cluster members. We now describe this full procedure in detail, including the construction of our initial cluster redshift guess $z_{\text{init}}$.

7.1. Redshift Initialization: $z_{\text{red}}$

For the full SDSS DR8 survey, we have multiple photometric redshifts based on large training sets (e.g., Csabai et al. 2007; Sheldon et al. 2012). However, these methods have certain limitations. First, they require training sets that span a broad range of magnitudes, which, although abundant at $z \lesssim 0.5$ for SDSS data, will be much sparser at higher redshifts for large surveys such as DES. Second, these methods—in particular $p(z)$ methods such as that of Sheldon et al. (2012)—are very good at estimating the ensemble of redshifts for a broad class of galaxies. However, our needs are much more specific: we wish to have a good initial single-value estimate of the redshift of the CG of galaxy clusters to initialize our cluster photometric redshift estimation procedure. To that end, we have developed our own photometric redshift estimator $z_{\text{red}}$, which is specifically designed to work on red sequence galaxies.

Given a red-sequence galaxy at redshift $z$ with $i$-band magnitude $m_i$, color vector $c$, and photometric error $C_{\text{err}}(z)$, the probability distribution of its color is simply

$$P(c) \propto \exp \left( -\frac{1}{2} \chi^2 \right),$$

(43)

where $\chi^2$ is given by Equation (6), i.e.,

$$\chi^2 = (c - \langle c | z, m_i \rangle) (C_{\text{err}}(z) + C_{\text{err}}(z))^{-1} (c - \langle c | z, m_i \rangle).$$

(44)

The corresponding log-likelihood is therefore simply $\ln \mathcal{L} = -0.5 \chi^2$. In practice, we also include an additional volume prior that accounts for the fact that there is more volume at higher redshifts. Assuming that the luminosity function does not evolve over the redshift uncertainties, the probability that a galaxy of a given luminosity is at redshift $z$ is

$$P_0(z) \propto \frac{dV}{dz} = (1 + z)^2 D_N^2(z) cH^{-1}(z),$$

(45)

which leads us to the likelihood

$$\ln \mathcal{L}_{\text{red}} = -\frac{\chi^2}{2} + \ln \left| \frac{dV}{dz} \right|.$$ 

(46)

The redshift estimator $z_{\text{red}}$ is that which maximizes the above likelihood. We use the “red” subscript to indicate that the redshift estimator assumes a red-sequence galaxy model.
use parabolic interpolation to find the correct maximum. This search is restricted to galaxies with $m_i < m_\ast(z) + 2.5$, since galaxies fainter than this fall well below the luminosity threshold used to define cluster richness (recall that $m_\ast(z)$ is defined in Section 4.2). The error estimate for $z_{\text{red}}$ is estimated as the standard deviation of the redshift over its posterior, i.e.,

$$\sigma^2_{z_{\text{red}}} = \langle z^2 \rangle - \langle z \rangle^2,$$

where

$$\langle z^n \rangle = \frac{\int dz L_{\text{red}}(z) z^n}{\int dz L_{\text{red}}(z)}.$$

We could, of course, store the posterior of the redshift distribution, but we have chosen not to do so since the only use of $z_{\text{red}}$ in the redMaPPer algorithm is that of providing an initial redshift estimate for galaxy clusters.

The top left panel of Figure 7 shows $z_{\text{red}}$ for DR8 cluster training galaxies with $p_{\text{mem}} \geq 0.9$ versus the spectroscopic redshift of the corresponding CG $z_{\text{CG}}$. We see that $z_{\text{red}}$ performs very well, with low bias and scatter, and very few gross outliers. The “flare-up” of the points around $z_{\text{CG}} \sim 0.35$ is due to the 4000 Å break moving from the $g-r$ to the $r-i$ color.

The performance of $z_{\text{red}}$ is better illustrated in the bottom left panel of the same figure. The black triangles show the mean offset $z_{\text{red}} - z_{\text{CG}}$ in redshift bins, the blue dashed line shows the average error in $z_{\text{red}}$ as estimated above, while the red dashed line shows the observed rms of the redshift offset in each of the redshift bins. The magenta dotted line shows the fraction of $4\sigma$ outliers. It is clear from the figure that our errors are somewhat overestimated, and that there is a small redshift bias in $z_{\text{red}}$.

We correct for the deficiencies revealed in the left panel of Figure 7 by applying an afterburner. Specifically, for the above cluster sample we define the mean redshift offset as a function of redshift,

$$dz(z) = \langle z_{\text{red}}^0 - z_{\text{CG}} \rangle_{z_{\text{CG}}},$$

where $z_{\text{red}}^0$ is the original, uncorrected redshift estimate defined above. That is, $dz(z)$ is the curve traced by the black triangles in the bottom left panel of Figure 7. We define a corrected $z_{\text{red}}$ redshift, as the solution to the equation

$$z_{\text{red}} = z_{\text{red}}^0 + dz(z_{\text{red}}).$$

In practice, the above treatment is slightly simplified, since our correction afterburner allows for the redshift bias to be a function of magnitude. For details, we refer the reader to Appendix A.1.

In the right panel of Figure 7, we show the corrected value of $z_{\text{red}}$ as a function of $z_{\text{CG}}$ after applying our afterburner, again for a sample of galaxies with $p_{\text{mem}} > 0.9$. The notation is the same as for the left panel. The biases are improved at high redshift, although there are still some residual issues at $z \sim 0.4$ where $z_{\text{red}}$ is biased by $\sim 0.3\sigma$. The reason the biases are not completely removed is due to the asymmetric and non-Gaussian nature of the scatter at the filter transition. We also note that the afterburner removes residual biases observed as a function of $m_i$ (not shown). The overall small bias and scatter in $z_{\text{red}}$ allow us to use this photometric redshift estimate as a good initial guess with which to initialize our photometric cluster redshift estimator.

7.2. Cluster Redshift Estimation: $z_{\lambda}$

Our approach to computing the cluster photometric redshift $z_{\lambda}$ is essentially an iterative extension of $z_{\text{red}}$. Specifically, given a CG candidate, we perform the following steps.

1. Start with a cluster redshift $z_{\lambda,i}$, where $i$ indexes the iteration. In the first iteration, we set $z_{\lambda,0} = z_{\text{red}}$.
2. Calculate the richness $\lambda$ around the candidate CG setting $z_{\text{cluster}} = z_{\lambda,i}$, and get the associated set of membership probabilities $p_{\text{mem}}$.
3. Select high membership probability galaxies to estimate a new redshift $z_{\lambda,i+1}$ by maximizing the likelihood function given by Equation (52) below.
4. Repeat from step 2 until convergence, such that $|z_{\lambda,i+1} - z_{\lambda,i}| < 0.0002$.

All that remains then is the definition of a suitable likelihood function. To begin, let us assume that we have a sample of known cluster member galaxies. Then, the log-likelihood of the observed colors for these galaxies would be

$$\ln L = \sum \frac{-\chi_i^2}{2} - \frac{\ln |C|}{2}. \quad (51)$$

In Equation (44) we take into account the log of the determinant of the covariance matrix, $\ln |C|$. We have found that, unlike the case of $z_{\text{true}}$, including this term improves the performance of $z_{\lambda}$ when the intrinsic scatter is varying rapidly. This makes sense, given that when utilizing multiple galaxies, one can directly probe the scatter in the red sequence, which is an observable that is inaccessible when estimating single-galaxy photo $z$s.

Of course, in practice, we do not have a list of known members, but rather a list of likely members with membership probabilities. One might be inclined to adopt a sharp cut $p_{\text{mem}} \geq p_{\text{min}}$ in order to define a likelihood that can be used to compute the cluster redshift. However, we find that a sharp cut in $p_{\text{mem}}$ leads to numerical instabilities in the iterative process because galaxies can scatter in and out of the sample in the course of the iteration.

To overcome this problem, we adopt instead a soft cut and define a new likelihood

$$\ln L = \sum \left[ -\frac{\chi_i^2}{2} + \ln |C| \right], \quad (52)$$

where each galaxy contributes a weight $w$ that smoothly varies from $w = 1$ at $p_{\text{mem}} = 1$ to $w = 0$ at $p_{\text{mem}} = 0$.

The assignment of these weights is somewhat ad hoc. We assume that $w(p_{\text{mem}})$ follows a Fermi-Dirac distribution. The transition from $w = 0$ to $w = 1$ occurs at $p_{70}$, which is the probability threshold that accounts for 70% of the total richness, i.e.,

$$0.7\lambda = \sum_{p_{\text{mem}} > p_{70}} p_{\text{mem}}. \quad (53)$$

The advantage of defining the probability threshold in this way—as opposed to a redshift independent threshold $p_{\text{cut}}$—is that $p_{70}$ varies with cluster redshift in such a way that one always uses the same fraction of cluster galaxies when estimating redshifts. Were we to take a constant $p_{\text{mem}}$ cut, the number of galaxies contributing to $z_{\lambda}$ would decrease with increasing redshift, since galaxy $p_{\text{mem}}$ values decrease as the photometry becomes noisier. The width of the distribution is set to 0.04, which we found is sufficient to regularize the iterative process. Thus, our galaxy weights are defined via

$$w(p_{\text{mem}}) = \frac{1}{\exp\left[((p_{70} - p_{\text{mem}})/0.04) + 1\right]}. \quad (54)$$

In Figure 8, we illustrate how the iterative process in our redshift estimate works. Fundamentally, each loop in the iteration takes a $z_{\text{in}}$ value for the redshift and produces a redshift $z_{\text{out}}$, and we wish to find the stable point where $z_{\text{out}} = z_{\text{in}}$. In the figure, we show $z_{\text{out}}(z_{\text{in}})$ for three sample clusters. For the two typical clusters denoted with red short-dashed lines, this function is well behaved, and we quickly achieve convergence. However, there are also $\sim 1$%–2% of clusters that have convergence curves like the blue long-dashed line. These appear to be projection effects between multiple nearby structures.

(A color version of this figure is available in the online journal.)

Figure 8. Redshift difference $z_{\text{out}} - z_{\text{in}}$ for one loop of our iterative photometric redshift estimator, as a function of the input redshift $z_{\text{in}}$. Two typical, well-behaved clusters are shown with red short-dashed lines. However, $\sim 1$%–2% of clusters have convergence curves like the blue long-dashed line. These appear to be projection effects between multiple nearby structures.
problem in a future paper. We see too that there is a feature at 0.35 ≲ z ≲ 0.45, in both the bias and scatter, reflecting the additional difficulties introduced by the fact that the 4000 Å break goes from being sampled by g − r to r − i. This is also the redshift range where we start running into the limit of the DR8 photometry, which further aggravates these failures. Indeed, these features are greatly reduced when redMaPPer is run on deeper data (e.g., SDSS Stripe 82 co-adds; Annis et al. 2011, not shown).

One interesting thing to note about the top panel in Figure 9 is that the “large” (Δz ∼ 0.1) redshift offsets in this plot do not reflect errors in the cluster redshift estimates, but rather cluster miscentering. That is, when we compare zλ to the redshift of the CG, large offsets are primarily due to our selection of a CG that is not, in fact, a cluster member. To demonstrate this, we have created a “clean” sample of clusters where we demand that there be at least two spectroscopic cluster members with p_{mem} > 0.8 within 1000 km s^{-1} of the spectroscopic redshift of the CG, thereby ensuring that the CG is in fact a cluster member. Of the 13,178 redMaPPer clusters in DR8 with spectroscopic redshifts, 1829 (or 14%) meet this criterion. The vast majority of outliers are due to errors in cluster centering, i.e., the offset zλ − zCG is large not because zλ is incorrect, but rather because the chosen CG is not actually a cluster member. To demonstrate this, we have created a “true” sample of clusters where we demand that there be at least two spectroscopic cluster members with p_{true} ≥ 0.8 within 1000 km s^{-1} of the spectroscopic redshift of the CG, shown with a black solid histogram. We show the fraction of outliers, as labeled. These are computed using the full redMaPPer cluster sample, with no additional spectroscopic requirements on member galaxies (unlike in Figure 10).

We can get a better sense of the fraction of gross redshift outliers from Figure 11, where we show the fraction of 3σ, 4σ, and 5σ outliers. A cluster is considered an Nσ outlier if |zλ − zCG| ≥ Nσzλ. To estimate the fraction of outliers as a function of redshift, for each redshift z we collect all clusters with redshift z_{mem} ∈ [z − 0.025, z + 0.025] and directly measure the fraction of Nσ outliers. By moving the window [z − 0.025, z + 0.025], we recover the outlier fraction as a function of redshift. We see that ≈1% of our galaxy clusters are 4σ redshift outliers. We note that the outlier fraction is considerably larger than expected if the errors were simply Gaussian. We emphasize that this fraction is measured using the full cluster sample, not the cleaned version used to produce Figure 10.

Finally, in Figure 12 we test whether the redMaPPer estimates for the cluster redshift probability distributions P(z_{true}|z_{obs}) are accurate. First, we select all clusters with spectroscopic CGs to create a “true” N(z_{CG}), shown with a black solid histogram. We note that this is not representative of the full cluster population...
8. CLUSTER CENTERING

The issue of galaxy cluster centering is very important for constraining cosmology with photometric surveys. In particular, miscentered clusters are a leading source of systematic error in stacked weak-lensing mass estimates (e.g., Johnston et al. 2007; Mandelbaum et al. 2008; Rozo et al. 2010), as well as mean velocity dispersions (e.g., Becker et al. 2007). In addition, the cluster richness estimates themselves depend on the choice of center (Lopes et al. 2006). Thus, a well-characterized centering model is essential for precision cosmology.

We assume that every galaxy cluster halo has a bright, dominant galaxy residing at its center (e.g., Matthews et al. 1964; Oemler 1976; Schombert 1986; von der Linden et al. 2007, 2014; Menanteau et al. 2013; Mahdavi et al. 2013; Song et al. 2012b; Stott et al. 2012; see also Paper II). In our current implementation, we also assume that the CG is red, which is the case for the vast majority of massive clusters. The exceptions are strong cool-core clusters such as A1835 (Allen 1995), where there is enough star formation for the broadband color of the CG to no longer be consistent with that of a red-sequence galaxy (e.g., McNamara et al. 2006). Although blue CGs are more common (although still rare) at the group scale (e.g., George et al. 2011, 2012; More et al. 2011; Tinker et al. 2012), the redMaPPer clusters are much more massive than the scale at which this is an issue.

Miscentering of galaxy clusters wherein the CG is undergoing strong star formation is a known failure of the redMaPPer centering algorithm (see Section 8.4 and Paper II). Simply removing the requirement that CGs be consistent with the red sequence—i.e., relying solely on luminosity and proximity—can fix some of these clusters, but at the expense of miscentering ∼10% of the clusters on foreground galaxies. Likewise, our tests have shown that both galaxy centroids and luminosity-weighted galaxy centroids result in worse centering properties than the algorithm currently implemented below (e.g., see also George et al. 2012). Thus, centering on red galaxies is, as far as we can tell, the “least bad” option. In its current implementation, the centering success rate is ≈85% (see Paper II). We intend to continue working on improving our centering model for future data releases, as this is currently the dominant source of systematic failures in the redMaPPer cluster catalog.

8.1. Basic Framework

We introduce a fundamentally new way of thinking about identifying the CG of a cluster: rather than specifying a unique cluster center, redMaPPer estimates the probability that a given galaxy is the CG of the cluster. Some clusters have well-defined cluster centers, exhibiting a single galaxy with a centering probability $P_{\text{cen}} \approx 1$, whereas others can have two or more reasonable central candidates, with the most likely center having $P_{\text{cen}} \approx 50\%$. We note that these centering probabilities are the angular position equivalent of the standard photo-z distributions $P(z)$. That is, just as a cluster has an uncertain redshift position characterized by a redshift probability distribution, so too does the cluster have an uncertain angular position on the sky, characterized by the probability of any given galaxy of being the correct cluster center. The importance of this new way of treating cluster centering is that it opens up the possibility of a statistical treatment of cluster centering akin to the statistical treatment of photometric redshifts, allowing us to improve our estimates of the cluster richness functions and cluster correlation functions. A detailed description of this framework will be presented and tested in a future work.

The key insight that allows us to estimate centering probabilities is that there are three different types of galaxies in a cluster: a central galaxy (“CG”), satellite galaxies, and unassociated foreground and background galaxies. Let $x$ be an observable vector for a galaxy, e.g., color (via $z_{\text{red}}$), luminosity ($m$), and position of each galaxy. We define $u_{\text{cen}}$, $u_{\text{sat}}$, and $u_{\text{fg}}$ as the distribution of $x$ for central, satellite, and background galaxies, respectively. The $u_{\text{cen}}$ and $u_{\text{sat}}$ filters are assumed to depend on cluster redshift and richness, while $u_{\text{fg}}$ depends only on cluster redshift (via $z_{\text{red}}$). We use the subscript “fg” as we expect that foreground galaxies will be more likely to be misidentified as CGs. Given a galaxy with observable $x$, the probability that it is the CG of a cluster is

$$p_{\text{cen}}(x|\lambda, z_{\text{red}}) = p_{\text{free}} \frac{u_{\text{cen}}}{u_{\text{cen}} + \lambda_{\text{sat}}u_{\text{sat}} + u_{\text{fg}}}.$$  (57)

where $p_{\text{free}}$ is the probability that a galaxy has not been partially masked by a higher ranked cluster (as described in Section 9.3; typically $p_{\text{free}} \approx 1$), and $\lambda_{\text{sat}} = \lambda - 1$ is the total number of
solar satellite galaxies. This formula can be thought of as the simple
definition of probabilities, or it can be interpreted as a Bayesian
classification algorithm.

Note, however, that the probability $P_{cen}$ is not the same thing
as the probability $P_{cen}$ that the galaxy is the unique CG of
the cluster. By assumption, there can be only one CG, so if
galaxy $i$ is the CG, then every other galaxy $j \neq i$ must not be a
central. Consequently, the probability that galaxy $i$ is the CG of a
cluster is

$$\text{The Astrophysical Journal}$$

$$P_{cen} \propto p_{cen}(x_i) \prod_{j \neq i} (1 - p_{cen}(x_j)). \quad (58)$$

The proportionality constant is set by the condition that there is
just one CG in the cluster,

$$1 = \sum_i P_{cen}(x_i). \quad (59)$$

In addition to the CG probability, we can also calculate
the probability that a cluster is centered on a satellite galaxy, given by

$$P_{sat} = (1 - P_{cen}) \frac{\ln u_{sat}}{\ln u_{sat} + u_{fg}}. \quad (60)$$

All that remains for us to be able to estimate centering probabilities is the definition of the filters $u_{cen}$, $u_{sat}$, and $u_{fg}$.

### 8.2. Centering Filters

With the basic formalism laid out, we need to specify the observable $x$ and the corresponding filters. There are three observables that we use to select the CG: the galaxy $i$-band magnitude $m_i$, the red sequence photometric redshift $z_{red}$ of the galaxy, and a weight $w$ that characterizes the local cluster galaxy density around the proposed CG. We also explored replacing our photometric redshift $z_{red}$ with $\chi^2$, the “distance” in color space to the red sequence. However, we have found empirically that $z_{red}$ works better for estimating central probabilities, in that small amounts of star formation and/or small color errors due to deblending have a much smaller impact on $z_{red}$ than they do on $\chi^2$. We consider each of the filters in turn.

#### 8.2.1. Luminosity Filter: $\phi_{cen}$

The magnitude of the CG is correlated with both richness and
redshift, so we define the CG magnitude filter as

$$\phi_{cen}(m_i | \bar{m}_i, \sigma_m) = \frac{1}{\sqrt{2\pi}\sigma_m} \exp\left(-\frac{(m_i - \bar{m}_i)^2}{2\sigma_m^2}\right). \quad (61)$$

where in principle both $\bar{m}_i$ and $\sigma_m$ depend on richness
and redshift. In practice, we expect $\sigma_m$ to be roughly redshift
independent, whereas $\bar{m}_i$ obviously depends on redshift. We
assume that $\bar{m}_i$ traces $m_{cen}(z_i)$, so that the full richness- and
redshift-dependent parameterization of $\bar{m}_i$ is

$$\bar{m}_i(z_{cen}, \lambda) = m_{cen}(z_i) + \Delta_0 + \Delta_1 \ln \frac{\lambda}{\lambda_p}, \quad (62)$$

where $\Delta_0$ and $\Delta_1$ are redshift-independent constants, and $\lambda_p$ is
the median richness of the sample. Our algorithm for fitting for
$\Delta_0$ and $\Delta_1$ is detailed below.

#### 8.2.2. Photometric Redshift Filter: $G_{cen}(z_{red})$

For the photometric redshift filter, we use the red-sequence
photometric redshift $z_{red}$ for each galaxy in the field. We model
this as a Gaussian function, with the form

$$G_{cen}(z_{cen}) = \frac{1}{\sqrt{2\pi}\sigma_{z_{red}}} \exp\left(-\frac{(z_{red} - z_{cen})^2}{2\sigma_{z_{red}}^2}\right). \quad (63)$$

As the error in the single galaxy photometric redshift dominates
that from the cluster photometric redshift, we have set the scatter
in $G_{cen}(z_{red})$ to that of the individual galaxy. In addition to
the photometric redshift filter, we employ a hard cut such that
$\chi^2(z_{red}) < 100$ (with 4 degrees of freedom). Investigations
of DR8 spectroscopic galaxies have shown that galaxies with
$\chi^2 > 100$ are all catastrophic outliers in $z_{red}$, which is not
surprising considering the bad fit to the red-sequence template.
By allowing galaxies with $\chi^2 < 100$, we allow some flexibility
for galaxies that have slightly offset colors to still be considered
as CGs. This is especially an issue for SDSS DR8 for bright,
neighboring CGs that may have color shifts caused by deblending
problems.

#### 8.2.3. Local Galaxy Density Filter: $f_{cen}(w)$

The motivation behind the local galaxy density filter is to define
an observable $w$ that is a pseudo-gravitational potential
connecting each galaxy to every other cluster member. The weight $w$ assigned to a given central candidate is

$$w = \ln \left[ \sum_i (\rho_{cen}(x_i) L_i / R_c(\lambda/S)^{1/2}) \right], \quad (64)$$

where the sum is over all galaxies within the scale radius $R_c(\lambda)$
around the candidate central, $r_c = 50$ $h^{-1}$ kpc is a core radius
used to soften the $1/r$ dependence, $L_i$ is each galaxy’s $i$-band
luminosity, and $\rho_{cen}$ are the usual $\lambda$ membership probabilities.
The denominator is chosen to make the argument of the natural
log dimensionless and to remove the obvious dependence of the
numerator of $w$ on the total number of terms in the sum.
Normalized in this fashion, we expect that $w$ does not scale
with cluster richness or redshift.

We assume that for CGs, $w$ follows a lognormal distribution
$f_{cen}(w)$,

$$f_{cen}(w) = \frac{1}{\sqrt{2\pi}\sigma_w} \exp\left(-\frac{(\ln(w) - \bar{w}_{cen})^2}{2\sigma_w^2}\right). \quad (65)$$

As noted above, we expect $\bar{w}_{cen}$ to be richness and redshift
independent. On the other hand, $\sigma_w$ will certainly depend on
richness. The noise in $w$ should scale with raw galaxy counts
$(\lambda/S)_{1/2}$, where $S(z)$ is the redshift-dependent factor that relates
the raw galaxy counts to a richness estimate when the survey is
not sufficiently deep to reach $0.2 L_c$ at the redshift of the cluster
(see Equation (23)). For Poisson noise, we set

$$\sigma_w = \sigma_{w, cen} \left(\frac{\lambda}{S\lambda_p}\right)^{-1/2}, \quad (66)$$

where $\sigma_{w, cen}$ is a constant that we fit for. As above, the pivot
point $\lambda_p$ should be chosen to match the median richness of the
sample.
With these definitions, the product
\[ u_{\text{cen}} = \phi_{\text{cen}}(m_1 | z_\ell, \lambda) G_{\text{cen}}(z_{\text{red}}) f_{\text{cen}}(w | z_\ell, \lambda) \] (67)
is the filter characterizing the distribution of CGs.

8.2.4. Satellite Filter: \( u_{\text{sat}} \)

Satellite galaxies on the red sequence can be described by a filter function analogous to Equation (67), except with parameters appropriate for the satellite galaxies, \( \phi_{\text{sat}} \) and \( \sigma_{w, \text{sat}} \). The satellite luminosity function, \( \phi_{\text{sat}} \), is a Schechter function as described in Equation (8). The redshift filter \( G_{\text{sat}}(z_{\text{red}}) \) is identical to \( G_{\text{cen}}(z_{\text{red}}) \).

8.2.5. Foreground Filter: \( u_{\text{fg}} \)

The foreground filter is defined as the expected number of unassociated galaxies within the cluster radius \( R_c(\lambda) \),
\[ u_{\text{fg}} = \sum_{g,z} (m_1 | z_{\text{red}}) f_{\text{fg}}(w) \pi R_c^2(d_A^2), \] (69)
where \( \sum_{g,z} \) is the background density per deg\(^2\) per \( m_1 \) per \( z_{\text{red}} \), calculated in a similar fashion as the red sequence background described in Section 4.3. In addition, the area subtended by the cluster in Mpc\(^2\) must be converted to deg\(^2\) via the angular diameter distance \( d_A \), with \( d_A \) measured in Mpc deg\(^{-1}\). Finally, the \( f_{\text{fg}} \) filter describes the local galaxy density filter from Equation (65), with parameters appropriate for random points (\( \bar{w}_{\text{fg}} \) and \( \sigma_{w,\text{fg}} \)) as described below.

8.3. Implementation

Implementing this formalism requires that we calculate the parameters that describe the filters for CGs, satellite galaxies, and foreground/background galaxies. Of course, calibrating these parameters depends on having a training sample to start with. As usual, we approach this problem in an iterative fashion, where the centering model is constrained at the same time as the redMaPPer red-sequence model. In the first iteration, we generate a catalog with roughly correct centering and use this to provide an initial calibration of the filters. In subsequent iterations we make use of the centering filters and use the output to recalibrate. This procedure is iterated until convergence.

8.3.1. First Iteration and Initial Filter Calibration

First, we implement a rough centering algorithm: for every cluster, we simply select the brightest high-probability (\( p_{\text{mem}} > 0.8 \)) member galaxy as the central galaxy of the cluster. In this fashion, we obtain a full training catalog with a set of CGs that should be roughly correct.

We now use this first iteration of a CG catalog to determine the filter parameters that we will use in subsequent iterations. Note that because the initial CG catalog contains some miscenterings, when calibrating the CG filters it is important to account for this contamination.

If a cluster is improperly centered on a satellite galaxy, it is most often centered on the brightest satellite. Consequently, the luminosity distribution of satellites that are mistaken as central is not simply a Schechter function (\( \phi_{\text{sat}} \)). Rather, the luminosity distribution of satellite galaxies in the CG catalog is given by \( \phi_{\text{sat}}(m_1 | \lambda, m_\star) \), the magnitude distribution of the brightest satellite in clusters of richness \( \lambda \). The expected magnitude and \( z_{\text{red}} \) distribution of the galaxies in our CG catalog is
\[ \rho(m_1, z_{\text{red}} | z) = P_{\text{cen}} \phi_{\text{cen}}(m_1 | G(z_{\text{red}} | z) + P_{\text{sat}} \phi_{\text{sat}}(m_1 | G(z_{\text{red}} | z) + (1 - P_{\text{cen}} - P_{\text{sat}}) \sum_{g,z} \frac{\pi R_c^2}{d_A^2}, \] (70)
where \( P_{\text{cen}} \) is the probability that the galaxy in question is the central galaxy, as in Equation (58), and \( P_{\text{sat}} \) is the probability that the cluster is centered on the brightest satellite galaxy. The redshift, \( z \), is the spectroscopic redshift of the “seed” galaxy used in the training step.

Our primary goal is to constrain the parameters \( \Delta_0, \Delta_1 \), and \( \sigma_m \). However, we also have the parameters \( P_{\text{cen}} \) and \( P_{\text{sat}} \), which are unknown in the first iteration. For these parameters, we have found that setting them in the first iteration at any reasonable initial estimate (\( P_{\text{cen}} \in [0.7, 0.9] \); \( P_{\text{sat}} \in [0.05, 0.2] \)) has no marked effect on the final calibration of the filter parameters. Therefore, for simplicity we set \( P_{\text{cen}} = 0.9 \) and \( P_{\text{sat}} = 0.05 \) for each individual cluster in this first iteration.

Before we can continue, we must estimate the parameters for \( \phi_{\text{sat}}(m_1 | \lambda, m_\star) \). This is modeled as a Gaussian distribution with central value
\[ m_\star = m_\star(z) + \sigma_{m_\star} \ln(\lambda/\lambda_p), \] (71)
where \( m_\star(z) \) is obtained from Equation (9) and \( \lambda_p \) is the median richness of the sample. The width of the distribution is similarly modeled as
\[ \sigma_m = c_{m_\star} + s_{m_\star} \ln(\lambda/\lambda_p). \] (72)
The central value must scale with richness because as we sample more galaxies from the luminosity function, we are more likely to find a very bright galaxy. In order to obtain these parameters, we run a simple Monte Carlo with the luminosity function parameters from Section 4.2. For \( \alpha = -1.0 \) with \( \lambda_p = 30 \), we find that \( c_{m_\star} = -0.95, s_{m_\star} = -0.32, c_{\sigma_m} = 0.40, \) and \( s_{\sigma_m} = -0.09 \). We note that these parameters depend only on \( \alpha \) and \( \lambda_p \) and thus do not need to be updated in subsequent iterations.

Finally, in order to constrain the parameters \( \Delta_0, \Delta_1 \), and \( \sigma_m \), we define our likelihood based on Equation (70) as
\[ \ln L = \sum_i \ln \rho(m_1, z_{\text{red}} | z). \] (73)
By maximizing this likelihood with respect to \( \Delta_0 \) and \( \Delta_1 \), we can use our training clusters to estimate the \( u_{\text{cen}} \) filter parameters.

8.3.2. Calibrating the \( w \) Filters

We now turn to calibrating the \( w \) filters. We begin by calibrating the foreground and satellite filters, \( f_{\text{fg}}(w) \) and \( f_{\text{sat}}(w) \). For these purposes we assume that all satellites follow the same spatial profile independent of brightness. Both \( f_{\text{fg}}(w) \) and \( f_{\text{sat}}(w) \) can be calibrated in a Monte Carlo fashion. We note that foreground galaxies will uniformly sample the cluster area \( \pi R_c^2(\lambda) \); thus to evaluate \( f_{\text{fg}}(w) \), we draw random points uniformly within the disk of every training cluster and compute the corresponding \( f_{\text{fg}}(w) \) parameters, \( \bar{w}_{\text{fg}} \) and \( \sigma_{w,\text{fg}} \).
The satellite filter \( f_{\text{sat}}(w) \) is computed in a similar fashion. First, for every training cluster we randomly select a cluster member with a probability \( p \) that is proportional to the membership probability. For this randomly selected member we compute \( w \) at the location of that satellite. After computing \( w \) for all the training clusters, we compute the corresponding \( f_{\text{sat}}(w) \) parameters, \( \bar{w}_{\text{sat}} \) and \( \sigma_{w,\text{sat}} \).

We can now turn our attention to the distribution \( f_{\text{cen}}(w) \). Consider now the distribution \( f(w) \) for all CGs. This total distribution is then

\[
f(w) = P_{\text{cen}} f_{\text{cen}}(w) + P_{\text{sat}} f_{\text{sat}}(w) + (1 - P_{\text{cen}} - P_{\text{sat}}) f_{\text{fg}}(w).
\]

The only unknowns in this equation are the mean \( \bar{w}_{\text{cen}} \) and rms \( \sigma_{w,\text{cen}} \) for the central filter, so we write the likelihood

\[
\ln L(\bar{w}_{\text{cen}}, \sigma_{w,\text{cen}}) = \sum_i \ln f(w),
\]

where the sum is over all training clusters. We then maximize this likelihood to find \( \bar{w}_{\text{cen}} \) and rms \( \sigma_{w,\text{cen}} \).

In Figure 13, we show the central, satellite, and foreground \( f(w) \) filters for the final training iteration. It is clear that there is some power here to differentiate between CGs and satellites and foregrounds, but it is far from perfect. In particular, satellites are only slightly less well connected than CGs.

### 8.3.3. Subsequent Iterations

As noted above, in our first iteration our catalog of CGs is constructed using a simple centering algorithm: i.e., select the brightest high-probability member as the cluster center. In subsequent iterations of the cluster finder calibration we use the probabilistic centering algorithm described in Section 8.1. After application of our centering algorithm, we have the important advantage that each cluster now comes tagged with \( P_{\text{cen}} \) and \( P_{\text{sat}} \). Therefore, we can now repeat the calibrations from Sections 8.3.1 and 8.3.2 while using the correct \( P_{\text{cen}} \) and \( P_{\text{sat}} \) for each individual galaxy. In this way we continuously improve our centering model with multiple iterations of the cluster-finding algorithm.

### 8.4. A Note on Blue Central Galaxies

At the beginning of this section, we noted that CGs undergoing strong star formation pose a particular problem for the redMaPPer centering algorithm. In Figure 14 we show an SDSS image of A1835, a well-known strong cool-core cluster with star formation in the CG. The true center is denoted with a red circle, while the top two candidate redMaPPer centers are indicated in blue. As expected, when the color of the CG is far from the red sequence, our centering algorithm instead chooses one of the bright red satellites as a possible center. However, we emphasize that the cluster will not be missing, it will simply be miscentered.

We have also made use of SDSS spectra to get an initial estimate on the rate of bad centers due to galaxies that do not agree with our color model (both very blue and very red). We have taken every cluster with a spectroscopic central and a second brighter spectroscopic galaxy that is within \( 0.167 \, h^{-1} \) Mpc for the cluster, it is considered a possible catastrophic center. In all, \( 4.6\% \) of the clusters fit this category. Visual inspection of the richest clusters shows that the majority of these galaxies are in the outskirts of the cluster and are not centrals, confirming that the redMaPPer centering algorithm is not simply taking the brightest member as the center. Alternatively, we can look at the subset of these galaxies that are not in the members list and thus were not considered as candidate centrals at all. Only \( 0.7\% \) of clusters contain such galaxies. From these estimates, we expect that the rate of miscentering due to galaxies that are too blue to be selected as CGs is \( \lesssim 2\% \).
9. THE CLUSTER FINDER

We have now described in detail all the ingredients that go into the redMaPPer cluster finder. Here, we focus on how these ingredients are blended within the context of the cluster finder to produce a catalog. In particular, we discuss how clusters are ultimately defined and percolated to ensure that every cluster is found once and only once. From a practical perspective, the cluster finding is broken into three stages. First, we look for overdensities around each individual galaxy using $z_{\text{red}}$ as an estimate of the cluster redshift. Second, we calculate the cluster likelihoods for each of the galaxies that have a sufficient overdensity. Third, after sorting by cluster likelihood, we percolate through the full catalog while probabilistically masking out cluster members. A process flowchart for reference in this section is shown in Figure 15.

9.1. First Pass

In the first pass, we wish to identify galaxies that are credible centers of galaxy clusters. This task involves a lot of data handling, and so we wish to make it as efficient as possible.

We begin by taking every galaxy in the input catalog with $\chi^2(z_{\text{red}}) < 20$ and brighter than $0.2\, L_*$ at the red-sequence photometric redshift $z_{\text{red}}$. These are very generous cuts, yet they reduce the input DR8 catalog from 56 million galaxies to 23 million possible cluster centers in the redshift region $0.05 < z_{\text{red}} < 0.6$. Next, we take all galaxies within $0.5\, h^{-1}\, \text{Mpc}$ of a candidate center and measure $\lambda$, setting the cluster redshift to $z_{\text{red}}$. Candidate centers with $\lambda/S < 3$ are rejected. The scale value $S = 1/(1 - C)$ (from Section 5.1) is used to ensure that we have detected at least three red galaxies above the magnitude limit. This cut rejects a further $\sim 60\%$ of the catalog of candidate centrals. Finally, for all centers that pass these cuts we calculate $z_\lambda$, as described in Section 7.2 to better refine the redshift of the possible cluster.

9.2. Likelihood Sorting

Given our list of possible clusters from the first pass, we now calculate the cluster likelihood for each of these clusters. The total likelihood is a combination of the $\lambda$ likelihood and the centering likelihood. To calculate the $\lambda$ likelihood, we first calculate the richness $\lambda$ using the optimized radial scale parameters with $R_0 = 1.0\, h^{-1}\, \text{Mpc}$ and $\beta = 0.2$ as described in Section 4. The $\lambda$ likelihood is then given by

$$ \ln L_\lambda = -\frac{\lambda}{S} - \sum \ln(1 - p_{\text{mem}}), $$

where $\lambda$ is evaluated at the cluster photometric redshift $z_\lambda$.

Next, following Section 8 and Equation (67), the centering likelihood is given by

$$ \ln L_{\text{cen}} = \ln[\phi_{\text{cen}}(m_i|z_\lambda, \lambda)G_{\text{cen}}(z_{\text{red}})f_{\text{cen}}(w|z_\lambda, \lambda)], $$

where we combine the luminosity, $z_{\text{red}}$, and local galaxy density $w$ of each galaxy. The total likelihood used in the ranking of possible cluster centers is

$$ \ln L = \ln L_\lambda + \ln L_{\text{cen}}. $$

\[^{18}\text{In the case of the training runs, we take every “seed” galaxy at the spectroscopic redshift.}\]
9.3. Percolation

Having rank-ordered the cluster candidates according to likelihood, we now need to percolate the list to assign galaxies to clusters and ensure that no cluster is counted multiple times. The basic outline of the percolation proceeds as follows.

1. Given cluster number \(i\) in the list, recompute \(\lambda\) and \(z_\perp\) based on the percolated galaxy catalog. At the beginning of the percolation, the percolated galaxy catalog is simply the input galaxy catalog.
2. Determine the cluster center and centering probability via the method outlined in Section 8.
3. Perform a final calculation of \(\lambda\) and \(z_\perp\) with respect to the new CG.
4. Update the percolated galaxy catalog by masking out galaxies based on their membership probabilities.
5. Remove all lower-ranked possible centers that have a membership probability \(p_{\text{mem}} > 0.5\) of being a member of cluster \(i\). Note that these galaxies are still allowed to provide membership weight to lower-ranked clusters as part of the percolated galaxy catalog.
6. Repeat step 1 for the next cluster galaxy in the ranked list.

9.3.1. Masking Galaxies

Masking galaxies based on their membership probabilities is the “probabilistic percolation” step of the redMaPPer algorithm. To perform this step, we keep track of the “total probability” that a galaxy belongs to a cluster, which we call \(p_{\text{taken}}\). The probability \(p_{\text{free}} = 1 - p_{\text{taken}}\) is the probability that the galaxy does not belong to any cluster. Initially, one has \(p_{\text{taken}} = 0\) and \(p_{\text{free}} = 1\) for all galaxies. Upon finding a galaxy cluster, the entire galaxy catalog is percolated by updating the probability \(p_{\text{taken}}\) via

\[
p_{\text{taken},i+1} = p_{\text{taken},i} + p_{\text{free},i} p_{\text{mem}},
\]

where \(p_{\text{mem}}\) is given by Equation (1).

Now, when we reestimate the richness of cluster \(i + 1\), we must take into account the fact that some of the galaxies have a nonzero probability of belonging to a cluster \(j < i + 1\). We do so by modifying the richness calculation from Equation (2) via

\[
\lambda = \sum p_{\text{free}} p_{\text{mem}}(x|\lambda).
\]

The first factor above is simply the probability that a galaxy is “free” to belong to the new cluster, and \(p_{\text{mem}}\) is the standard membership probability from Equation (1). For instance, suppose a galaxy has a probability \(p_{\text{mem}} = 0.3\) of belonging to the first cluster in the rank-ordered list. In this case, the galaxy still has 70% of its probability to give to a cluster lower in the list. In practice, when quoting cluster membership probabilities \(p_{\text{mem}}\) we report not the raw \(p_{\text{mem}}\) value as given by Equation (1), but rather the product \(p_{\text{free}} p_{\text{mem}}\) for that galaxy–cluster pair. That is, the reported value is the correct probability that the given galaxy belongs to the cluster under consideration. For galaxy clusters that are sparse in the sky (e.g., at high richness) these corrections are negligible.

9.3.2. Extent of Clusters and Percolation Radius

As noted above, cluster richness is measured within a radius \(R_c(\lambda)\) that optimizes the signal-to-noise ratio of the richness measurements (R12), but is not in any way chosen to be related to standard definitions of the extent of a halo, say, \(R_{200c}\). For cosmological purposes, it is useful to differentiate between the radius \(R_c(\lambda)\), which defines the richness measurement, and the percolation radius that is used to mask out cluster members and blend or deblend nearby systems. In particular, ideally one selects the percolation radius so that it matches as best as possible the percolation radii employed in the halo definition used to calibrate the corresponding halo mass function.

In the Appendix of R12 we used maxBCG clusters to obtain an approximate scaling of mass to \(\lambda\) richness. We found that the slope of the mass–\(\lambda\) relation is consistent with 1, and that \(R_{200c} \approx 1.5 R_c(\lambda)\). Consequently, we have adopted \(1.5 R_c(\lambda)\) as our default percolation radius. That is, galaxies are masked out to this radius. Note that while galaxies outside the \(R_c(\lambda)\) radius are not used in the summand in Equation (2), we can still estimate \(p_{\text{mem}}\) in exactly the same way as we do with all other galaxies out to an arbitrary radius, which is how we implement the large percolation radius above.

In practice, for the \(\lambda/S > 20\) richness threshold we have employed, changing the mask radius by \(\pm 50\%\) has a very small impact on the resulting cluster catalog. Only a small number of clusters (\(~\%5\%)—primarily satellites of the richest \(\lambda > 100\) clusters—are affected at all by making this change. We expect to return to the question of what the optimal masking radius is in future work, particularly within the context of cosmological constraints from galaxy clusters.

9.4. A Sample Cluster

At this point, it would be useful to investigate a sample rich cluster to explore the distribution of cluster members. We have selected RM J164019.8+464241.5 (A2219), one of the richest clusters in SDSS, at a redshift of \(z = 0.23\). In Figure 16 we show four aspects of this cluster. In the top left panel, we show the SDSS image of the cluster. In the top right panel, we show the distribution of cluster members with \(p_{\text{mem}} > 0.05\), with the symbol size proportional to the membership probability. As can be seen in the histogram of \(p_{\text{mem}}\) in the bottom-right panel, this distribution is strongly peaked near \(p_{\text{mem}} \approx 0.9\). We note that we do not show another, larger peak at \(p_{\text{mem}} = 0\), for the vast majority of galaxies in the field that are not red sequence cluster members. Finally, in the bottom-left panel we show the distribution of cluster members as a function of radius. The red dashed line shows the expected distribution of galaxies for an NFW profile with \(r_s = 0.5 h^{-1}\) Mpc, thus showing that the radial distribution of cluster galaxies is broadly consistent with the NFW model.

10. THE redMaPPer SDSS DR8 CLUSTER CATALOG

We have run the redMaPPer cluster-finding algorithm in the SDSS DR8 photometric catalog described in Section 2. The full cluster finder run contains all clusters with \(\lambda \geq 5 S(z_\perp)\) and \(z_\perp \in [0.05, 0.6]\). However, we have chosen to apply very conservative cuts to our catalog. The cuts we apply are as follows.

\(^{19}\) As shown in Appendix F, \(\lambda_{\text{col}}\) used in R12 is within \(~10\%\) of the multicolor \(\lambda\) used in this work.
Figure 16. Top left: SDSS image of RM J164019.8+464241.5 (A2219). Top right: locations of cluster members with $p_{\text{mem}} > 0.05$, with the symbol size proportional to the membership probability. Bottom left: distribution of cluster members as a function of radius. The red dashed line shows the expected distribution of galaxies for an NFW profile with $r_s = 0.5 h^{-1}$ Mpc. Bottom right: distribution of $p_{\text{mem}}$ for cluster members. This is strongly peaked near $p_{\text{mem}} > 0.9$. We note that we do not show another, larger peak at $p_{\text{mem}} = 0$, for the vast majority of galaxies in the field that are not red sequence cluster members.

(A color version of this figure is available in the online journal.)

1. The richness is cut to $\lambda \geq 20 S(z_\lambda)$. Roughly speaking, this requires that every cluster have at least 20 galaxy counts above the flux limit of the survey or $0.2 L_*$ at the cluster redshift, whichever is higher. From R12, we estimate that this results in an effective mass cut of $M_{200} \gtrsim 10^{14} M_\odot$.

2. The redshift range is cut to $z_\lambda \in [0.08, 0.55]$, so as to minimize edge effects from the training sample.

3. Only clusters with $f_{\text{mask}} < 0.2$ are included (see Equation (24)), ensuring that clusters are not overly compromised by bad fields, bright stars, and survey edges.

The resulting cluster catalog contains 25,236 systems. In Figure 17 we show the full footprint of the catalog. The color scale shows the density contrast relative to the mean cluster density, where red regions are denser than average and blue regions are less dense, as estimated using all $\lambda \geq 5$, $z_\lambda \in [0.1, 0.3]$ clusters so as to give a better sense of the large-scale structure in the survey. The image was produced by binning the catalog into a Mangle simple pixelization scheme of depth 7 (Swanson et al. 2008).

In Figure 18, we show the comoving density of redMaPPer clusters with $\lambda/S(z) > 20$ over the full redshift range of interest. The comoving density is roughly constant at $z_\lambda \lesssim 0.35$ where...
is roughly constant at $z_{\lambda} < 0.35$, where the catalog is volume limited (denoted by the vertical dashed line). Above this redshift the comoving density falls off rapidly as the detection threshold rapidly increases.

(A color version of this figure is available in the online journal.)

Figure 18. Comoving density of redMaPPer DR8 clusters as a function of photometric redshift ($z_{\lambda}$) for clusters with $\lambda/S(z) > 10, 20, 40$. All densities have been computed by taking the sum of cluster $\rho(z)$. The comoving density is roughly constant at $z_{\lambda} < 0.35$, where the catalog is volume limited (denoted by the vertical dashed line). Above this redshift the comoving density falls off rapidly as the detection threshold rapidly increases.

(A color version of this figure is available in the online journal.)

the catalog is volume limited. At $z_{\lambda} \sim 0.35$ the richness and redshift scatter are significantly boosted by both the 4000 Å break filter transition and the magnitude limit of the survey reaching 0.2 $L_\odot$. Therefore, the comoving density is boosted by low richness clusters scattering up into our sample. A full accounting for this scatter must be made in order to precisely calculate the redMaPPer abundance function, which we leave to future work. Above this redshift the magnitude limit starts to kick in (via the scale factor $S(z)$), and we only observe the most massive clusters. As an illustration of this effect, in Figure 19, we plot the richness $\lambda$ versus the photometric redshift $z_{\lambda}$ for the final redMaPPer catalog. The red dashed line shows the redshift-dependent richness cut $\lambda > 20S(z)$.

Finally, we show a sample redMaPPer cluster. In Figure 20, we show REDM J003208.2+180625.3, the richest redMaPPer cluster not found within the MCXC cluster catalog (Piffaretti et al. 2011), a system with $\lambda = 236 \pm 12$ at redshift $z_{\lambda} = 0.396 \pm 0.013$. We note that this cluster is associated with a source in the ROSAT Bright Source Catalog (Voges et al. 1999). The specific data available for each of the clusters and members are described in Appendix G, and a detailed comparison of the redMaPPer clusters to X-ray and SZ catalogs is presented in Paper II.

11. PURITY AND COMPLETENESS

Purity and completeness can mean many different things depending on the context. There is a tendency to think of purity as the probability that a cluster in the catalog is a real cluster, and to think of completeness as the probability that a real cluster is in the catalog. However, it is often incorrect to think of these quantities as calibrating failure rates of the algorithm. Here, we adopt specific definitions of purity and completeness and discuss their implications. For an alternative definition of purity and completeness by comparing redMaPPer-detected clusters to X-ray catalogs, we refer the reader to Paper II.

For cluster cosmology, the relevant quantity is the probability of detecting a halo of mass $M$ with richness $\lambda_{\text{obs}}$, which can be decomposed into a convolution of two components,

$$P(\lambda_{\text{obs}}|M) = \int d\lambda_{\text{true}} P(\lambda_{\text{obs}}|\lambda_{\text{true}}) P(\lambda_{\text{true}}|M).$$  \hspace{1cm} (81)

The probability $P(\lambda_{\text{true}}|M)$ is a feature of the universe and must be properly marginalized over in any cosmological study that relies on the cluster number function. Constraining this probability distribution can also be supplemented by utilizing realistic mock catalogs (e.g., Song et al. 2012a; R. H. Wechsler et al., in preparation), which we return to in future work.
On the other hand, \( P(\lambda_{\text{obs}}|\lambda_{\text{true}}) \) is a feature of the cluster-finding algorithm itself. This probability fully contains all of the information associated with measurement error in our catalog. In the present work, we define completeness and purity as specific integrals over this distribution.

Purity and completeness, used as a simple parameterization of \( P(\lambda_{\text{obs}}|\lambda_{\text{true}}) \), can be estimated in several ways. Perhaps the simplest consists of removing galaxy clusters, randomizing galaxy positions, and then reinserting galaxy clusters. The cluster-finding algorithm can then be rerun, and one can determine which clusters are detected, and how many “false” clusters are detected (e.g., Goto et al. 2002; Koester et al. 2007b; Hao et al. 2010). However, as shown in the literature (Lopes et al. 2004; Rozo et al. 2011), such an algorithm is fundamentally flawed because background galaxies are not uniformly distributed. Consequently, we take a somewhat different approach, as described below.

1. Generate random points. We generate a list of random points uniformly sampling the input survey mask. These points are to be the centers of mock clusters that will be inserted into the dataset. This procedure ensures that we sample all the systematics in the survey, as well as the effect of masked regions. We note that these locations are not sampled from the DR8 galaxy positions.

2. Sample “true” cluster richness and redshift. Using the full cluster catalog (\( \lambda > 5 \)), we randomly sample galaxy clusters to generate pairs of parameters (\( \lambda_{\text{true}}, z_{\text{true}} \)). This ensures that our model cluster distribution has the same richness and redshift distribution as the final catalog, including covariances.

3. For each pair of sampled values (\( \lambda_{\text{true}}, z_{\text{true}} \)), assign them a spatial location using a random point from step 1, and sample galaxies using the cluster model. Using the same method as in Section 5.2, we use Monte Carlo sampling to generate 5000 galaxies with the model radial and luminosity profiles. From this sample of 5000 cluster galaxies, we randomly sample \( \lambda_{\text{true}} \) galaxies from within \( R_c(\lambda_{\text{true}}) \), as well as \( k \lambda_{\text{true}} \) galaxies from \( R_c(\lambda_{\text{true}} < r < R_c(2\lambda_{\text{true}}) \), where \( k \) scales with \( \lambda_{\text{true}} \) as appropriate for the radial profile. This ensures that our fake clusters do not have artificial hard edges.

4. Measure \( \lambda_{\text{obs}} \) for the generated fake cluster at the random location, and repeat 100 times. When measuring \( \lambda_{\text{obs}} \), we mask out galaxies according to the bright stars and edges in the survey mask, as well as applying any necessary magnitude limits. We do not, however, make corrections for higher-order effects such as blending of galaxies.

In this way, we generate a map over the full sky of the detectability of clusters as a function of redshift and richness, while taking into account the large-scale structure that is already imprinted on the galaxy catalog. This is a more stringent test than a random-background test, but still does not capture the additional effect that correlated large-scale structure can have on the galaxy clusters. However, we expect that these additional effects are subdominant: while the correlation length of galaxy clusters can be as large as \( \sim 20 \) Mpc, the typical length scale over which projections are effective is \( \sim 100 \) Mpc or more, so for most of this volume \( 1 + \xi \approx 1.0 \). Roughly speaking, we would expect no more than 20% corrections to our estimated impurity from these effects, so, for example, if 5% of our clusters suffer from projection effects in this analysis, it is likely that this fraction is underestimated by \( \sim 0.2 \times 0.05 = 1\% \). A more detailed treatment of projection effects will be presented in a future work (see also the discussion on the impact of the background on \( \lambda \) in Rozo et al. 2011).

One additional concern with regard to the above test is that the clusters that we input into our data set are produced with the same filters that we used to find galaxy clusters. As discussed in Rykoff et al. (2012), the choice of luminosity and radial filters has negligible impact on the richness measurements. Rozo et al. (2011) also demonstrated that cluster ellipticity is largely irrelevant. Thus, we expect our results to be robust to changes to the cluster model used to generate the inserted clusters. We have verified this by inserting real redMaPPer clusters as above, where the cluster galaxies are sampled from the members list according to their membership probabilities. Our results are nearly identical to those derived from the model clusters above.

In Figure 21, we illustrate how we use the above outputs to define purity and completeness. The figure shows the expectation value (\( \langle \lambda_{\text{obs}} \rangle \)) for simulated clusters in the narrow redshift range \( 0.2 < z < 0.22 \). Note that although \( \lambda_{\text{true}} \) is a fixed value, each cluster has a distribution of \( \lambda_{\text{obs}} \) and we have plotted the mean value. To measure completeness, we consider the subsample of clusters in a richness bin in \( \lambda_{\text{true}} \), defined by the vertical red short-dashed lines. While most of the cluster sample falls within a tight locus around \( \lambda_{\text{obs}} \approx \lambda_{\text{true}} \), there are some clusters that fall above the \( 4\sigma \) contours defined by the diagonal red short-dashed lines. These outliers are projection effects, where we placed a fake cluster atop an existing richer structure, and are counted toward incompleteness. For reference, the one-to-one line is shown with the magenta dotted line. To measure purity, we consider the subsample of clusters in a richness bin in \( \lambda_{\text{obs}} \), defined by the blue long-dashed lines. The clusters that are significant outliers with low \( \lambda_{\text{true}} \) are impurities where the measured cluster is a result of a projection effect of multiple systems. The gray region denotes “unphysical” projections where \( \lambda_{\text{obs}} \geq 2 \lambda_{\text{true}} \), and as such the fake cluster with richness \( \lambda_{\text{true}} \) is the secondary rather than the primary halo.

(A color version of this figure is available in the online journal.)
outliers fall above the main cloud of points: these are projection effects, where we placed a fake galaxy cluster atop an existing richer structure. It is precisely to characterize these outliers that we do not limit our model clusters to avoid the existing structure in the catalog. The completeness (\(\xi_{\text{true}}\)) is defined as the fraction of the non-outlier points to the total number of clusters in the richness bin, i.e., it is the ratio of the number of clusters within the red dashed parallelogram (\(\xi_{\text{obs}}\) is consistent with \(\xi_{\text{true}}\) within the scatter) to the number of clusters in the \(\xi_{\text{true}}\) bin.

Note that with this definition \(\xi_{\text{true}} \leq 1\) does not imply that we are missing clusters. Instead, it is simply estimating the fraction of clusters at a given \(\xi_{\text{true}}\) that suffer from severe projection effects. Similarly, for clusters with \(\xi_{\text{true}}\) near the detection threshold, a fraction of these clusters will have \(\xi_{\text{obs}}\) less than the detection threshold. Thus, these clusters are only “missing” due to well-understood observational scatter.

Similarly, we can estimate purity by considering clusters in a bin in \(\xi_{\text{obs}}\), e.g., that defined by the blue long-dashed lines in Figure 21. Several of the clusters in this bin are clear outliers compared to their corresponding \(\xi_{\text{true}}\). The fraction of such outliers in the \(\xi_{\text{obs}}\) bin is the impurity. However, we note one additional restriction; that is, we discard all outliers with \(\xi_{\text{obs}} \geq \xi_{\text{true}}/2\), denoted by the gray region in the figure. We note that in any projection effect, \(\xi_{\text{obs}} = \xi_{\text{main}} + \xi_{\text{proj}}\), the richness of the main and projected halo, respectively. By definition, the main halo has a richness \(\xi_{\text{main}} \geq \xi_{\text{obs}}/2\), and we are concerned with calculating the purity of main halos only. Thus, any fake cluster with \(\xi_{\text{true}} \leq \xi_{\text{obs}}/2\) is necessarily a projection on a real, significantly richer cluster in the catalog and should be discarded in this analysis. As with the completeness calculation, we emphasize that the resulting purity is the fraction of galaxy clusters as a function of the observed richness that suffer from projection effects and does not represent an absence of galaxies.

To formalize all of the above discussion, we define the completeness in a bin of richness \(\xi_{\text{true}}\) as

\[
\text{completeness} = \frac{\sum_{i} \xi_{\text{true}} \int_{\xi_{\text{true}}}^{\xi_{i}} d\lambda_{\text{obs}} \xi_{\text{true}} \int_{\xi_{\text{true}}}^{\lambda_{\text{true}}} d\lambda_{\text{obs}} \text{P}(\lambda_{\text{true}}|\lambda_{\text{true}})}{N(\lambda_{\text{true}})},
\]

where the sum is over all clusters in a given bin of \(\xi_{\text{true}}\). We define \(\lambda_{0} = \lambda_{\text{true}} - 4\sigma\), with the restriction that \(\lambda_{0} > 0\); \(\lambda_{1} = \lambda_{\text{true}} + 4\sigma\); \(\sigma^{2} = \sigma_{\text{true}}^{2} + \sigma_{\text{proj}}^{2}\); and \(N(\lambda_{\text{true}})\) is the total number of clusters in the bin. We estimate \(\sigma_{\text{int}}\) directly from the output as the intrinsic scatter in the \(\lambda_{\text{true}}\) distribution, and we have chosen to define an outlier (incomplete) cluster as any cluster that has a measured richness \(\lambda_{\text{obs}}\) that is more than \(4\sigma\) discrepant from its true richness \(\lambda_{\text{true}}\).

By the same token, the purity is defined as

\[
\text{purity} = \frac{\sum_{i} \xi_{\text{true}} \int_{\xi_{\text{true}}}^{\lambda_{i}} d\lambda_{\text{obs}} \xi_{\text{true}} \int_{\lambda_{\text{true}}}^{\lambda_{i}} d\lambda_{\text{obs}} \text{P}(\lambda_{\text{true}}|\lambda_{\text{true}})}{\sum_{i} \xi_{\text{true}} \int_{\xi_{\text{true}}}^{\lambda_{i}} d\lambda_{\text{obs}} \xi_{\text{true}} \int_{\lambda_{\text{true}}}^{\lambda_{i}} d\lambda_{\text{obs}} \text{P}(\lambda_{\text{true}}|\lambda_{\text{true}})},
\]

where now the sums are over all clusters in a given bin of \(\lambda_{\text{obs}}\). Note that we have the additional restriction that the sum is restricted to systems with \(\lambda_{\text{true}} \geq \lambda_{\text{obs}}/2\). We define \(\lambda_{0} = \lambda_{\text{true}} - 4\sigma_{\text{int}}\) and \(\lambda_{1} = \lambda_{\text{true}} + 4\sigma_{\text{int}}\) as before, and \(\lambda_{\text{obs}}, 1\) and \(\lambda_{\text{obs}}, 2\) are the extent of the richness bin in question.

In Figure 22, we show the completeness and purity as a function of richness for five redshift bins for the DR8 galaxy and cluster catalog. At low redshifts (\(z < 0.3\)) the completeness is essentially \(\geq 99\%\) at \(\lambda > 30\), but falls below this threshold due to clusters randomly scattering in and out of the selection threshold. At higher redshift, as we encounter the magnitude limit of the DR8 catalog our richness threshold increases and thus the richness at which these threshold effects come into play also increases.

Our purity is \(\geq 95\%\) for all richness and redshift bins, with the richest systems being less pure. This can be understood very simply: consider a chance superposition of two clusters of richness \(\lambda\) leading to a single detection of richness \(2\lambda\). This factor of two shift has a much more dramatic impact on the overall abundance function at the rich end, simply because the richness function is steeper there, i.e., a constant projection fraction in \(\lambda_{\text{true}}\) translates into a projection fraction that decreases with \(\lambda_{\text{obs}}\). Again, all these “impurities” actually correspond to real clusters; it’s just that the observed richness has been systematically overestimated.

One curious feature of our purity is that it seems to increase with decreasing richness and with increasing redshift. This is a consequence of our definition; at lower richness and higher redshifts, the measurement errors in the richness are larger, so a cluster that is a \(4\sigma\) outlier needs to be more and more of an extreme projection, which makes such \(4\sigma\) outliers rarer. That is, the purity increases not because there are fewer projections, but rather because the projections that do occur become increasingly less important relative to the observational errors in the richness estimates.

In this context, it is important to emphasize that the purity that we have defined here is fundamentally different from the purity that is usually defined for X-ray or SZ cluster finding. Let us take SZ as a specific example: a false detection in SZ occurs when random Gaussian noise in the map produces a fluctuation that can be mistaken for an SZ cluster. In our galaxy catalog, galaxies are detected at \(> 10\sigma\), so a “false detection”...
of a cluster near our threshold ($\lambda = 20$) would require 20 simultaneous $10\sigma$ noise fluctuations collocated in the sky and strongly correlated across all five bands to mimic red sequence colors. If regions with bad photometry have been properly culled out (by application of the BOSS mask), this simply does not happen. Every galaxy overdensity identified by redMaPPer is a true galaxy overdensity: there are no true false detections driven by observational noise. The key, however, is that the galaxy overdensity is a cylindrical galaxy overdensity that may contain more than one massive halo, and it is precisely this rate that we have tried to characterize with our definition of purity.

12. CLUSTER MASKS

One of the great advantages of using model clusters placed randomly on the real sky is that we can use the same output to map the detectability of redMaPPer clusters across the entire survey. In this way, we can directly construct a set of random points directly applicable to the cluster mask, which is not the same as the galaxy mask that defines the survey. An appropriate set of random points is essential for cross-correlation studies for cluster cosmology (e.g., Landy & Szalay 1993).

As an illustration of the difference between the galaxy mask that defines the survey and the cluster mask that defines the redMaPPer catalog, we have run a dense sample of random points in the vicinity of Arcturus using the methods described in Section 11. This very bright star contaminates the SDSS photometry over a large area and thus effectively masks out a region of the sky that is 0.8 in radius. To isolate the effect of the survey mask, all the random points shown are associated with model clusters of the same true richness $\lambda_{\text{true}} = 40$, with a redshift distribution appropriate for redMaPPer.

In Figure 23, we show the map of the detectability of $\lambda_{\text{true}} = 40$ clusters in a $4^\circ \times 4^\circ$ region around Arcturus. Each pixel shows the fraction of time a sample cluster is detected, using Equation (82), where black is 0% and white is 100%. In the low-redshift bin ($0.1 < z < 0.2$, top-left panel) the detectability of $\lambda_{\text{true}} = 40$ clusters is essentially 100% outside the Arcturus mask, except for a few pixels around bright Tycho stars. Note, however, that due to our requirement that the area of a cluster must not be significantly masked out ($f_{\text{mask}} < 0.2$), the edge for the detectability of a cluster at these redshifts is slightly farther from the center of Arcturus than the edge of the galaxy mask (denoted by the red dashed line). At higher redshift these edges drift closer to each other as the angular extent of the clusters decreases. However, in the highest-redshift bin ($0.4 < z < 0.5$, bottom right panel) the cluster is only detected $\sim 60\% \pm 30\%$ of the time (see Figure 22) due to the survey depth. The detectability varies significantly when clusters approach the threshold, and we see a strong dependence on the local depth and structure.

13. SUMMARY

In this paper, we have introduced redMaPPer, a red-sequence cluster finder that is designed to make optimal use of large photometric surveys. As a case study in the implementation of the algorithm, we have run on the SDSS DR8 photometric catalog. We have shown that redMaPPer improves significantly on previous cluster finders (see also Paper II), with many features that will be required to take advantage of upcoming surveys such as DES and LSST. In particular:

1. redMaPPer is based on a multicolor extension of the optimized richness estimator $\lambda$, which has been shown to be a good mass proxy (R12 and Paper II).
2. redMaPPer is self-training, with a modest requirement in the number of training spectra, which can themselves be limited to the brightest cluster galaxies. This makes it particularly well suited to high-redshift surveys. Furthermore, the multicolor red-sequence model makes optimal use of all color data at all redshifts, with no sharp features as the 4000 Å break transitions between filters.
3. redMaPPer can handle complex survey masks. Both mask-corrected richness values can be computed, as well as cluster-appropriate random point catalogs for large-scale structure studies.
4. All clusters are assigned a redshift probability distribution $P(z)$, which enables a more accurate reconstruction of the redshift distribution of the cluster population relative to simple point-redshift estimates.
5. The centering of clusters is fully probabilistic. In this way, the uncertainty in the position of the cluster can be handled in an analogous way to the redshift uncertainty provided by $P(z)$.
6. The algorithm is numerically efficient and can be run on large surveys with modest computing power.

Using the red-sequence model derived in the redMaPPer calibration phase, we have derived two red-sequence-based photometric redshifts. The first, $z_{\text{red}}$, is a red sequence template-based photo $z$, which has been designed to generate a good “first-guess” estimation of the redshift in each cluster. We have...
also shown, in Appendix C, that \( z_{\text{red}} \) compares very well to existing DR8 photometric redshifts for this specific class of galaxies. However, \( z_{\text{red}} \) has the advantage that it requires many fewer spectroscopic training galaxies. Moreover, these galaxies can be the brightest galaxies in the clusters, with no penalty to the performance of \( z_{\text{red}} \) at the faint end of the galaxy sample. The second, \( z_\lambda \), is a very precise photo \( z \) derived from fitting all cluster members simultaneously to the red sequence model. In addition, we derive a \( P(z) \) estimator for \( z_\lambda \), which we show is superior to point-based photometric redshifts for the purposes of estimating the redshift distribution of the galaxy clusters. In DR8, this is especially true in the region of the filter transition at \( z \sim 0.35 \).

As a case study in the implementation of the algorithm, we have run redMaPPer on the 10,400 deg\(^2\) BOSS region from the SDSS DR8 photometric catalog. Using red galaxy spectroscopic redshifts from one-fifth of the total area from \( z \in [0.05, 0.6] \), we are able to constrain a robust red sequence model that defines both the richness and photometric redshift estimators. The photometric redshifts, \( z_\lambda \), have small bias and low scatter, ranging from \( \sigma = 0.06 \) at \( z \sim 0.1 \) to \( \sigma = 0.02 \) at \( z \sim 0.5 \), due to increased photometric noise near the survey limit. The rate of catastrophic outliers is low, with only \( \lesssim 1\% \) of galaxy clusters appearing as \( 4\sigma \) outliers. Note that because of our high photo-\( z \) precision, a cluster at \( z = 0.1 \) with a redshift offset as small as \( \Delta z = 0.025 \) is considered a catastrophic redshift failure. Furthermore, we show that the majority of these outliers are bad centers rather than bad redshifts; when the catalog is cleaned by demanding that central and satellite galaxies with spectroscopy must all be within 1000 km s\(^{-1}\), the failure rate decreases to \( \lesssim 0.2\% \).

After running redMaPPer on the full DR8 photometric catalog, we apply a conservative selection cut of \( \lambda/S(z) > 20 \), for a total of 25,236 clusters in the redshift range of \( z \in [0.08, 0.55] \). As shown in Paper II, the comoving density of redMaPPer clusters satisfying this cut is lower than that of all other SDSS photometric cluster catalogs. The scale factor, \( S(z) \), given by Equation (23), defines the correction factor on the richness caused by the survey depth. The catalog is volume-limited at \( z < 0.35 \), where \( S = 1 \) and the survey depth is brighter than the fiducial luminosity cut of 0.2 \( L_\odot \) used by the \( \lambda \) richness. Because our selection threshold corresponds to a total of 20 galaxy detections, as we lose galaxies at high redshift due to the magnitude limit of the survey, these 20 galaxies must all be due to bright members. Therefore, the corresponding richness threshold of \( 20/S(z) \) is much higher. This increased detection threshold results in fewer galaxy clusters at high redshifts. Our adopted richness threshold of 20 detected red-sequence galaxies is chosen to provide the most robust cluster catalog possible, with a mass threshold of \( M \gtrsim 10^{14} M_\odot \) where our catalog is volume limited at \( z \lesssim 0.35 \) (R12; Paper II). Although the full redMaPPer catalog extends to lower richnesses, we expect that performance will worsen as one moves toward lower and lower richness thresholds.

Finally, we investigate the purity and completeness of our cluster-finding algorithm, focusing on the observationally relevant probability distribution \( P(\lambda_{\text{obs}}|\lambda_{\text{true}}) \). We have defined impurity and incompleteness as the fraction of clusters for which the observed richness \( \lambda_{\text{obs}} \) is significantly different from the true richness \( \lambda_{\text{true}} \). These outliers are caused by projection effects: when two halos are merged together, this manifests itself as incompleteness (a cluster with richness \( \lambda_{\text{true}} \) is up-scattered significantly, so it is “missing” from where it should have been) or impurity (the richness \( \lambda_{\text{obs}} \) of such a cluster is significantly overestimated). We note that while the completeness of redMaPPer is near 100%, the purity is \( \sim 95\% \) at the rich end, increasing at lower richness. This decrease simply reflects larger observational error (in a proportional sense) for lower richness clusters; i.e., “outliers” become more rare not because projection effects are less rare, but because projection effects become subdominant to observational uncertainties. Our estimate of the incidence of projection effects is thus \( \sim 5\% \), similar to what was estimated in Rozo et al. (2011). A more detailed analysis of projection effects for redMaPPer clusters will be presented in a future work.

In Paper II, we present a detailed comparison of the redMaPPer cluster catalog to various X-ray and SZ catalogs with high-quality mass proxies. In all cases, we show that the redMaPPer richness \( \lambda \) is a low scatter mass proxy with high completeness and low impurity compared to these “truth” tables. We also compare the performance of redMaPPer to other photometric cluster finders that have been run on SDSS data and show that redMaPPer outperforms these other algorithms in all metrics (e.g., photo-\( z \) performance, mass scatter, and purity and completeness), though some do perform equally well in subsets of these categories in specific redshift ranges.

While this present work has focused on the application of redMaPPer to the SDSS DR8 catalog, we emphasize that this algorithm was developed specifically for upcoming large photometric surveys such as DES and LSST. In particular, its ability to simultaneously utilize all available photometric data, its smooth handling of the filter transition of the 4000 Å break across filter passes, and its ability to self-calibrate using only minimal spectroscopic training samples of bright cluster galaxies are all specifically designed to enable cluster finding in these new photometric data sets. This will be especially advantageous at \( z \gtrsim 0.7 \) in the Southern Hemisphere, where we do not have the advantages of more than a decade of survey data from the SDSS spectrograph. Thus, in short order we expect that redMaPPer will be capable of producing large, high-quality catalogs of \( \sim 80,000 \) clusters at \( z < 1 \) with DES, opening a new era of high-redshift cluster cosmology.

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A.1. Constraining $z_{\text{red}}$ Correction Parameters

Our approach to constraining the $z_{\text{red}}$ correction parameters $\tilde{c}_z(z)$ and $\tilde{s}_z(z)$ is similar to that employed for the red-sequence calibration (note the $z$ subscript). As before, we have chosen to constrain these parameters at a given node spacing, using cubic spline interpolation between the nodes. The node spacing we have chosen for DR8 is 0.05 for $\tilde{c}_z(z)$ and 0.10 for $\tilde{s}_z(z)$, suited to the characteristic variation scales.

One significant complication that we have to deal with is that we have membership probabilities for all the galaxies. In order to properly make use of the probabilities, as in Equation (34) we need to know the background pdf. Unfortunately, there is no first-principle way of calculating the $z_{\text{red}}$ background as a function of $z_{\text{true}}$. Therefore, we have chosen to assume that the background is a Gaussian function with zero mean and finite width, and to marginalize over this background as a set of nuisance terms. As above, we assume that the background width, $\sigma_b(z)$, is a smoothly interpolated function with a node spacing of 0.10. To ensure that we are calculating the correction factors appropriate for red galaxies, and not blue cluster members and interlopers, we limit ourselves to galaxies with $p_{\text{mem}} > 0.7$.

Given a model correction,

$$\langle c_z | z_{\text{true}}, m_i \rangle = \tilde{c}_z(z_{\text{true}}) + \tilde{s}_z(z_{\text{true}})[m_i - \bar{m}_i(z)](z_{\text{true}}), \quad (A1)$$

then we have a Gaussian pdf for the true galaxies,

$$G_1 = \frac{1}{\sqrt{2\pi}\sigma_{z_{\text{red}}}} \exp \left( -\frac{[z_{\text{red}} - z_{\text{true}}] - \langle c_z | z_{\text{true}}, m_i \rangle)^2}{2\sigma_{z_{\text{red}}}^2} \right), \quad (A2)$$

and for the background,

$$G_2 = \frac{1}{\sqrt{2\pi}\sigma_b} \exp \left( -\frac{[z_{\text{red}} - z_{\text{true}}]^2}{2\sigma_b^2} \right). \quad (A3)$$

The total likelihood is then

$$\mathcal{L} = w[p_{\text{mem}}G_1 + (1 - p_{\text{mem}})G_2], \quad (A4)$$

where we have made the addition of a weight function, $w$, which is a smooth function of $\chi^2$ that de-weights galaxies with large $\chi^2$ and that are possible outliers. The weight $w$ is

$$w = \frac{1}{\exp \left( (\chi^2 - \chi^2_{95})/0.2 \right) + 1}, \quad (A5)$$

where $\chi^2_{95}$ is the 95th percentile of all galaxies with $p_{\text{mem}} > 0.7$.

As before, we find the $\tilde{c}_z(z)$, $\tilde{s}_z(z)$, and $\sigma_b(z)$ parameters by maximizing $\Sigma z$ using the downhill-simplex method.

A.2. Constraining $z_{\lambda}$ Correction Parameters

Our approach to constraining the $z_{\lambda}$ correction parameters is analogous to that used for the $z_{\text{true}}$ parameters in Appendix A.1. However, our job is a little easier because we are applying corrections such that $\langle c_{z_{\lambda}} | z_{\lambda} \rangle$ is unbiased rather than the converse. Therefore, the correction term can be a function of $z_{\lambda}$. For DR8, we use a cubic spline interpolation with node spacing of 0.04 for $\langle c_{z_{\lambda}} | z_{\lambda} \rangle$. In addition, we allow an additional variance term as we find that our raw $z_{\lambda}$ errors are underestimated. For $\sigma_{z_{\lambda}}$, we use a smooth function with a node spacing of 0.10. To ensure that we are using well-measured clusters, we limit ourselves to calibration clusters that have $\lambda/S(z) > 10$, where $S$ is the scale factor defined in Equation (23). Essentially, this limits us to clusters with at least 10 red galaxies above the luminosity threshold or magnitude limit.

Given a model correction $\langle c_{z_{\lambda}} | z_{\lambda} \rangle$ and intrinsic scatter correction $\sigma_{z_{\lambda}}$, we have a Gaussian pdf for the clusters,

$$G = \frac{1}{\sqrt{2\pi}\sigma_{\text{tot}}} \exp \left( -\frac{[\langle z_{\lambda} \rangle - z_{\lambda} - \langle c_{z_{\lambda}} | z_{\lambda} \rangle]^2}{2\sigma_{\text{tot}}^2} \right), \quad (A6)$$

where $\sigma_{\text{tot}} = \sqrt{\sigma_{z_{\lambda}}^2 + \sigma_{z_{\lambda}}^2}$. The total likelihood is then given by $\ln \mathcal{L} = \sum \ln G$. As before, we find $\langle c_{z_{\lambda}} | z_{\lambda} \rangle$ and $\sigma_{z_{\lambda}}^2$ by maximizing this likelihood using the downhill-simplex method.

With this parameterization in hand, we can calculate the corrected $z_{\lambda}$ and error as $z_{\lambda} = z_{\lambda,\text{raw}} + \langle c_{z_{\lambda}} | z_{\lambda} \rangle$, and $\sigma_{z_{\lambda}}^2 = \sigma_{z_{\lambda,\text{raw}}}^2 + \sigma_{z_{\lambda}}^2$. However, we find that after applying these corrections there may still be small residuals in the training sample. Therefore, we iterate on this solution two further times to obtain a final corrected redshift $z_{\lambda}$.

After the calibration is complete, we must also apply these corrections to the $P(z_{\lambda}|z_{\lambda})$ estimation for each cluster. To replicate the $z_{\lambda}$ offset represented by $\langle c_{z_{\lambda}} | z_{\lambda} \rangle$, we first offset the central value of the $P(z)$ distribution. Next, to replicate the increased scatter we “expand the space” between the $P(z)$ bins, so that a Gaussian fit to $P(z)$ will measure the same width as the corrected $z_{\lambda}$ value. We find that this does an adequate job of maintaining asymmetries in the $P(z)$ distribution that show up near the filter transitions.

APPENDIX B

HOW MANY TRAINING CLUSTERS?

When calibrating the red sequence in Section 6 on DR8 data, we make use of all the spectroscopy available in our 2000 deg$^2$ training region. However, much of this is superfluous. First, most of the spectroscopic galaxies—even the LRG samples—are not in massive clusters. Second, our strategy of leveraging CG spectroscopy to all the galaxies in a cluster means that we do not require thousands of clusters to perform the calibration. In this section, we investigate how many training clusters—each represented by a single spectroscopic redshift for the CG—are required to create an accurate and unbiased richness and redshift estimate.

To test the number of required training clusters, we follow the method of Section 6.5 to measure the bias in the recovered richness and photometric redshift values on a predetermined set of test clusters. For our test suite, we select the {5, 10, 20, 40, 80} richest clusters per redshift bin of ±0.025 in the training region. The redshift binning is used to ensure that we have a relatively uniform coverage over the redshift range of interest. In practice, of course, the training clusters need not be so uniformly sampled.
For each of these spectroscopic training sets we recalibrate the red sequence and measure the richness \(\lambda\) and redshift \(z_\lambda\) for each of the test clusters from Section 6.5.

In Figure 24, we show the results of these test runs. The left panel shows the richness bias and significance of the bias as a function of redshift for the various training samples. Although we can get a reasonable calibration of the red sequence with as few as five spectra per ±0.025 redshift bin, the resulting richnesses are significantly biased (∼1σ) at the transition redshift \(z \sim 0.35\). In order to achieve unbiased richness estimates (<0.3σ), we require ~40 clusters per redshift bin. We assume that any residual biases are due to the noise in estimating the off-diagonal elements of the covariance matrix. This results in a total of ~400 spectra to achieve essentially the same fidelity of calibration as we can achieve with millions of SDSS spectra. The right panel shows the photometric redshift bias and significance, similar to the left panel. For accurate photo-z estimation, we require even fewer training spectra: ~20 per redshift bin, or a total of 200.

For upcoming photometric surveys such as DES, we can obtain these spectra by first running a crude run with an approximate red sequence model. After selecting bright CGs, these can easily be followed up spectroscopically, as they are the most luminous galaxies at any redshift. For example, over 85% of the training spectra required for DR8 training are brighter than \(m_1 < 18.5\). Thus, our method allows for an incredibly efficient use of limited spectroscopic resources to enable science in large photometric surveys.

APPENDIX C

COMPARISON OF \(z_{\text{red}}\) TO SDSS DR8 PHOTO \(z_{\text{photo}}\)

We consider two sets of photometric redshift estimates available for all of DR8. The first, \("z_{\text{photo}}\)\), uses an updated method of Sheldon et al. (2012). In this section, we make use of high-probability cluster member galaxies to compare these photometric redshifts to \(z_{\text{red}}\) at both bright magnitudes (where training galaxies are plentiful) and fainter magnitudes.

For our “pseudo-spectroscopic” test sample, we start with all clusters with \(\lambda > 5\) and a CG with spectroscopic redshift \(z_{\text{CG}}\). We then select all members with \(p_{\text{mem}} > 0.9\). We thus expect a contamination rate of up to 10%, although the real rate should be smaller than this. By assigning each high-probability member to the spectroscopic redshift of the CG, we can leverage the red sequence to obtain spectroscopic quality redshifts to much fainter magnitudes than available in the SDSS main or LRG spectroscopic samples.

Figure 25 shows the density map of the photometric redshift biases as a function of magnitude for \(z_{\text{red}}, z_{\text{photo}}, \) and \(p(z)\) for a narrow redshift slice of \(0.195 < z_{\text{CG}} < 0.205\). For \(z_{\text{red}}\) and \(z_{\text{photo}}\), we have assumed a pdf that is Gaussian with mean \(z_{\text{red}}(z_{\text{photo}})\) and width \(\sigma_{\text{red}}(\sigma_{\text{photo}})\). For the \(p(z)\) values, we use a spline interpolation to smooth the pdf and normalize the area to unity. On the right-hand side are projected histograms from the density field. The dotted red lines show the \(z_{\text{red}}\) distribution for comparison. Note that the density plot clearly shows the separation in magnitude between central and satellite galaxies. Both \(z_{\text{red}}\) and \(z_{\text{photo}}\) perform well down to the 0.2 \(L_\odot\) limit of the redMaPPer richness estimation, while the \(p(z)\) values have a broader distribution at the faint end. There is also obvious structure in the photo-z bias as a function of magnitude.

Figure 26 shows the same map for a narrow redshift slice of \(0.395 < z_{\text{CG}} < 0.405\). While all the photometric redshifts handle the luminous galaxies very well, there appear to be slight biases at the faint end in the case of the DR8 \(z_{\text{photo}}\), and a bifurcation of the distribution for the \(p(z)\) redshifts. The evolution of the bias in the \(p(z)\) estimates is due to a combination of effects. First, the \(r\)-band magnitude was used as an input to the photo-z estimator. For a field galaxy, a fainter magnitude correlates with a higher redshift. For cluster galaxies, however, galaxies of a wide range of luminosity occupy the same cluster.
As a result, when using magnitude-based photo-$z$ estimators on galaxies in clusters, one should expect an increasing bias with magnitude, which is simply a manifestation of the intra-cluster luminosity function. The large width of the error distributions relative to the other estimators is due primarily to the lack of training set galaxies in that range. As discussed in Sheldon et al. (2012), the main focus was on recovering the full $r<21.8$ galaxy sample. To avoid biases induced by training set selection, the authors did not include the most recent BOSS LRG samples in that work and deferred LRG-optimized $p(z)$ estimates to a future paper. It is also worth pointing out that, despite the extra width of the error distributions obtained when using $p(z)$, the recovered redshift distributions obtained by summing the $p(z)$ of Sheldon et al. (2012) are still superior to the distributions estimated using the DR8 $z_{\text{photo}}$ or single-point $z_{\text{red}}$ estimates.

There are two important messages from this comparison. First, the performance of state-of-the-art photo-$z$ estimators appears to be sufficiently accurate for bright galaxies that we would likely be able to use these in the initialization phase of redMaPPer without any loss. Second, $z_{\text{red}}$ appears to be at least as good—if not better—than what is currently achieved, with much smaller spectroscopic training samples. As shown in Appendix B, we can achieve this redshift performance with only $\approx 400$ of the brightest CG spectra. With the technique of assigning the spectroscopic redshift of the CGs to the members, we effectively increase the faint end of our training sample. This is very useful for future surveys because of the high cost of obtaining spectroscopic redshifts of faint galaxies.

As a color version of this figure is available in the online journal.

**Figure 25.** Top: density of total $p(z_{\text{red}} - z_{\text{CG}})$ as a function of $i$-band magnitude $m_i$ for all cluster members with $p_{\text{num}} > 0.9$ and $0.195 < z_{\text{CG}} < 0.205$ in clusters with $\lambda > 5$. The right panel shows the total pdf at all magnitudes. There is a small bias in $z_{\text{red}}$, though it is constant with magnitude. Middle: same as top panel, with $z_{\text{photo}}$ calculated with the algorithm of Csabai et al. (2007). The performance is good down to $0.2 L_\ast$. The right panel compares the distribution for $z_{\text{photo}}$ (black line) to $z_{\text{red}}$ (red dotted line). Bottom: same as top panel, with $p(z)$ values from Sheldon et al. (2012). While the bright galaxy performance is good, there are biases at the faint end and the distribution is significantly wider. (A color version of this figure is available in the online journal.)

**Figure 26.** Same as Figure 25, with members selected from clusters with $0.395 < z_{\text{CG}} < 0.405$. (A color version of this figure is available in the online journal.)

**APPENDIX D**

**COMPUTING PERFORMANCE BENCHMARKS**

The redMaPPer algorithm has been designed to be fast, efficient, flexible, and trivially parallelized. As there are two parts to running redMaPPer, the calibration and cluster-finding stages, we split the performance benchmarks into two parts.

For the calibration phase, the runtime depends on the number of training spectra and clusters. For the DR8 training sample on 2000 deg$^2$, the full calibration takes $\sim 30$ CPU hr on a three-year-old 2.8 GHz AMD Opteron 8389. Current Intel processors can run the calibration roughly twice as fast. For the minimal training sample of 40 clusters per redshift bin (see Appendix B) calibration takes $\sim 13$ CPU hr.

The cluster-finding stage is designed to be split into chunks of arbitrary size on the sky. For these purposes, we use the Mangle simple pixelization scheme (Swanson et al. 2008), although any pixelization scheme will work. As long as the overlap region between pixels is wider than twice the largest size of any cluster in the catalog, then the percolation of clusters within each cell is guaranteed to be unique. For the DR8 catalog, this corresponds to a border region of 1.5$'$, corresponding to twice the size of a cluster of richness $\sim 300$ at $z = 0.05$, given the mask radius parameters. In total, running the cluster finder on the full DR8 catalog requires $\sim 500$ CPU hr including all galaxy mask corrections. On a modestly sized compute cluster this can be run in much less than one day.

**APPENDIX E**

**VALIDATING THE CORRECTION C**

In Section 5, we laid out our methodology for correcting the richness for survey holes and a magnitude limit that is brighter than 0.2 $L_\ast$. In order to validate the calculation of the correction term $C$ described in that section, we have taken
a subsample of clusters with $0.15 < z < 0.3$ and simulated a more restrictive magnitude limit. We have chosen a magnitude limit of $m_i < 19.6$, which is $0.2 L_*$ at $z = 0.2$, so that all clusters at higher redshift will have their richness corrected according to our formalism. The average correction for the $z = 0.3$ clusters is similar to that for the highest redshift clusters in our catalog, so this test will sample the full range of corrections employed.

In Figure 27 we show the results of our test. In the top panel we show $\lambda_{19.6}$ versus $\lambda_{0.2}$ for all clusters with $0.2 < z < 0.3$, where $\lambda_{19.6}$ is the richness calculated with a magnitude limit of $m_i < 19.6$ and $\lambda_{0.2}$ is the standard $\lambda$ with a $0.2 L_*$ cut. When calculating $\lambda_{19.6}$, we have re-fit the photometric redshift $z_1$ to ensure that our comparison is as fair as possible. It is clear in the top panel that the correction richness scales with uncorrected richness, with some scatter as expected. In the bottom panel we show the richness scale value ($S = \frac{\lambda_{19.6}}{\lambda_{0.2}} = \frac{\lambda_{\text{scaled}}}{\lambda_{\text{raw}}}$) as a function of redshift. The black squares show the median estimated value of $S$ derived from Equation (16), while the black error bars represent the median error in $S$ as derived from Equation (22). The red diamonds show the median measured value of $S$, and the red error bars represent the observed width in the distribution of $S$. Our predicted correction factor does scale with redshift as expected. However, our errors are slightly overestimated for the largest corrections.

APPENDIX F
COMPARING $\lambda$ TO $\lambda_{\text{col}}$

We now explore how the richness estimate used in this work, $\lambda$, compares to the single-color richness $\lambda_{\text{col}}$ used in R12. As detailed in Section 4, the primary difference in richness estimators is the replacement of the Gaussian color filter with a multicolor $\chi^2$ filter. However, we emphasize that there is also a subtle difference in the background model, as described in Section 4.1. That is, the $\chi^2$ filter does not distinguish between galaxies that are too red or too blue relative to the model, and while the red sequence model is symmetric, the background model is not.

To make our comparisons, we have started with all redMaPPer clusters with $\lambda > 20$. We then calculate $\lambda_{r-i}$ and $\lambda_{\text{col}}$ using the appropriate color model from the red sequence parameterization. Our expectation is that $\lambda_{r-i}$ should trace $\lambda$ at low redshift where the dominant signal is from the $g-r$ color, and $\lambda_{\text{col}}$ should trace $\lambda$ at high redshift.

In Figure 28, we show the statistics from comparing $\lambda_{r-i}$ and $\lambda_{\text{col}}$ to $\lambda$. In the top panel we show the median ratio as a function of redshift. At all redshifts the bias between the appropriate $\lambda_{\text{col}}$ and $\lambda$ is $\lesssim 10\%$. In the bottom panel, we show the median normalized deviation, which is $\sim 1\sigma$ at low redshift and less so at high redshift where the richness errors are much larger due to the magnitude limit. We attribute this bias at low redshift to the different background model employed, as galaxies that are redder than the red sequence are down-weighted in the $\lambda$ model compared to the $\lambda_{\text{col}}$ model. These biases are not large, but they are significant and thus show the importance of using the same color model and consistent survey data to achieve the best richness estimation.

The middle panel of Figure 28 shows the width of the $\lambda_{\text{col}}/\lambda$ distribution as a function of redshift. The scatter is $\lesssim 15\%$ for the appropriate color except at $z \sim 0.35$, where the 4000 Å break...
is transitioning from $g$ to $r$. It is in this transition region that a single-color richness estimator does especially poorly and we have the biggest advantage of using a multicolor estimator.

**APPENDIX G**

**DESCRIPTION OF COLUMNS IN THE DR8 CLUSTER CATALOG**

The full redMaPPer DR8 cluster and member catalogs are available at [http://risa.stanford.edu/redmapper/](http://risa.stanford.edu/redmapper/) in FITS format, and from the online journal in machine-readable formats. A summary of the cluster catalog information is given in Table 1. A summary of the member information is given in Table 2.

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**Table 1**

| Column | Name | Format | Description |
|--------|------|--------|-------------|
| 1.0    | ID   | I7     | redMaPPer cluster identification number |
| 2.0    | NAME | A20    | redMaPPer cluster name |
| 3.0    | RA   | F12.7  | Right ascension in decimal degrees (J2000) |
| 4.0    | DEC  | F12.7  | Declination in decimal degrees (J2000) |
| 5.0    | Z_LAMBDA | F6.4 | Cluster photo-$z_{\lambda}$ |
| 6.0    | Z_LAMBDA_ERR | F6.4 | Gaussian error estimate for $z_{\lambda}$ |
| 7.0    | LAMBDA | F6.2  | Richness estimate $\lambda$ |
| 8.0    | LAMBDA_ERR | F6.2 | Gaussian error estimate for $\lambda$ |
| 9.0    | S    | F6.3   | Richness scale factor (see Equation 23) |
| 10.0   | Z_SPEC | F8.5  | SDSS spectroscopic redshift for most likely center ($-1.0$ if not available) |

(This table is available in its entirety in a machine-readable form in the online journal. A portion is shown here for guidance regarding its form and content.)

**Table 2**

| Column | Name | Format | Description |
|--------|------|--------|-------------|
| 1.0    | ID   | I7     | redMaPPer cluster identification number |
| 2.0    | RA   | F12.7  | Right ascension in decimal degrees (J2000) |
| 3.0    | DEC  | F12.7  | Declination in decimal degrees (J2000) |
| 4.0    | R    | F5.3   | Distance from cluster center ($h^{-1}$Mpc) |
| 5.0    | P_MEM | F5.3  | Membership probability $p_{mem}$ |
| 6.0    | IMAG | F6.3   | $i$-band cmodel magnitude (dereddened) |
| 7.0    | IMAG_ERR | F6.3 | Error on $i$-band cmodel magnitude |
| 8.0    | MODEL_MAG_U | F6.3 | $u$ model magnitude (dereddened) |
| 9.0    | MODEL_MAGERR_U | F6.3 | Error on $u$ model magnitude |
| 10.0   | MODEL_MAG_G | F6.3  | $g$ model magnitude (dereddened) |
| 11.0   | MODEL_MAGERR_G | F6.3 | Error on $g$ model magnitude |
| 12.0   | MODEL_MAG_R | F6.3  | $r$ model magnitude (dereddened) |
| 13.0   | MODEL_MAGERR_R | F6.3 | Error on $r$ model magnitude |
| 14.0   | MODEL_MAG_I | F6.3  | $i$ model magnitude (dereddened) |
| 15.0   | MODEL_MAGERR_I | F6.3 | Error on $i$ model magnitude |
| 16.0   | MODEL_MAG_Z | F6.3  | $z$ model magnitude (dereddened) |
| 17.0   | MODEL_MAGERR_Z | F6.3 | Error on $z$ model magnitude |
| 18.0   | Z_SPEC | F8.5  | SDSS spectroscopic redshift ($-1.0$ if not available) |
| 19.0   | OBJID | I20    | SDSS DR8 CAS object identifier |

(This table is available in its entirety in a machine-readable form in the online journal. A portion is shown here for guidance regarding its form and content.)
