Clustered calibration: an improvement to radio interferometric direction-dependent self-calibration

S. Kazemi, 1 S. Yatawatta 2 and S. Zaroubi 1

1 Kapteyn Astronomical Institute, University of Groningen, PO Box 800, NL-9700 AV Groningen, the Netherlands
2 ASTRON, Postbus 2, NL-7990 AA Dwingeloo, the Netherlands

ABSTRACT

The new generation of radio synthesis arrays, such as Low Frequency Array and Square Kilometre Array, have been designed to surpass existing arrays in terms of sensitivity, angular resolution and frequency coverage. This evolution has led to the development of advanced calibration techniques that ensure the delivery of accurate results at the lowest possible computational cost. However, the performance of such calibration techniques is still limited by the compact, bright sources in the sky, used as calibrators. It is important to have a bright enough source that is well distinguished from the background noise level in order to achieve satisfactory results in calibration. This paper presents ‘clustered calibration’ as a modification to traditional radio interferometric calibration, in order to accommodate faint sources that are almost below the background noise level into the calibration process. The main idea is to employ the information of the bright sources’ measured signals as an aid to calibrate fainter sources that are nearby the bright sources. In the case where we do not have bright enough sources, a source cluster could act as a bright source that can be distinguished from background noise. For this purpose, we construct a number of source clusters assuming that the signals of the sources belonging to a single cluster are corrupted by almost the same errors. Under this assumption, each cluster is calibrated as a single source, using the combined coherencies of its sources simultaneously. This upgrades the power of an individual faint source by the effective power of its cluster. The solutions thus obtained for every cluster are assigned to each individual source in the cluster. We give performance analysis of clustered calibration to show the superiority of this approach compared to the traditional unclustered calibration. We also provide analytical criteria to choose the optimum number of clusters for a given observation in an efficient manner.

Key words: methods: numerical – methods: statistical – techniques: interferometric.

1 INTRODUCTION

Low-frequency radio astronomy is undergoing a revolution as a new generation of radio interferometers with increased sensitivity and resolution, such as the LOw Frequency ARray (LOFAR) 1, the Murchison Wide-field Array (MWA) 2 and the Square Kilometre Array (SKA) 3 are being devised and some are already becoming operational. These arrays form a large effective aperture by the combination of a large number of antennas using aperture synthesis (Thompson, Moran & Swenson 2001). In order to achieve the full potential of such an interferometer, it is essential that the observed data are properly calibrated before any imaging is done. Calibration of radio interferometers refers to the estimation and reduction of errors introduced by the atmosphere and also by the receiver hardware, before imaging. We also consider the removal of bright sources from the observed data part of calibration that enable imaging the weak background sources. For low-frequency observations with a wide field of view, calibration needs to be done along multiple directions in the sky. Proper calibration across the full field of view is required to achieve the interferometer’s desired precision and sensitivity giving us high dynamic range images.

Early radio astronomy used external (classical) calibration which estimates the instrumental unknown parameters using a radio source with known properties. This method has a relatively low computational cost and a fast execution time, but it generates results strongly affected by the accuracy with which the source properties are known. In addition, existence of an isolated bright source, which is the most important requirement of external calibration, in a very

E-mail: kazemi@astro.rug.nl

1 http://www.lofar.org
2 http://www.mwatelescope.org
3 http://www.skatelescope.org

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Published by Oxford University Press on behalf of the Royal Astronomical Society
wide field of view is almost impractical, and even when it does, external calibration would only give information along the direction of the calibrator. The external calibration was then improved by self-calibration (Pearson & Readhead 1984). Self-calibration has the advantage of estimating both the source and instrumental unknowns, without the need of having a prior knowledge of the sky, only utilizing the instrument’s observed data. Moreover, it can achieve a superior precision by iterating between the sky and the instrumental parameters.

The accuracy of any calibration scheme, regardless of the used technique, is determined by the level of signal-to-noise ratio (SNR). This limits the feasibility of any calibration scheme to only bright sources that have a high enough SNR to be distinguished from the background noise (Liu, tegmark & Zaldarriaga 2009; Bernardi et al. 2011; Pindor et al. 2010). Note that interferometric images are produced using the data observed during the total observation (integration) time. However, calibration is done at shorter time intervals of that total duration. This increases the noise level of the data for which calibration is executed compared to the one in the images. In other words, there are some faint sources that appear well above the noise in the images while they are well below the noise level during calibration. It has been a great challenge to calibrate such faint sources having a very low SNR. In this paper, we present clustered self-calibration, introduced in Kazemi, Yatawatta & Zaroubi (2011a), and emphasize its better performance compared to unclustered calibration below the calibration noise level. Existing unclustered calibration approaches (Intema et al. 2009; Smirnov 2011) can only handle a handful of directions where strong enough sources are present and we improve this, in particular for subtraction of thousands of sources over hundreds of directions in the sky, as in the case for LOFAR (Bregman 2012).

The implementation of clustered calibration is performed by clustering the sources in the sky, assuming that all the sources in a single cluster have the same corruptions, and calibrating each cluster as a single source. At the end, the obtained calibration solutions for every source cluster is assigned to all the sources in that cluster. This procedure improves the information used by calibration by incorporating the total of signals observed at each cluster instead of each individual source’s signal. Therefore, when SNR is very low, it provides a considerably better result compared to unclustered calibration. Moreover, calibrating for a set of source clusters instead of for all the individual sources in the sky reduces the number of directions along which calibration has to be performed, thus considerably cutting down the computational cost. However, there is one drawback in clustered calibration: the corruptions of each individual source in one given cluster will almost surely be slightly different from the corruptions estimated for the whole cluster by calibration. We call this additional error as ‘clustering error’ and in order to get the best performance in clustered calibration, we should find the right balance between the improvement in SNR as opposed to degradation by clustering error.

Recently, clustering methods have gained a lot of popularity in dealing with large data sets (Kaufman & Rousseuw 1990). There are various clustering approaches that could be applied to calibration based on their specific characteristics. An overview of different clustering methods is given in Xu & Wunsch (2008). Clustering of radio sources should take into account (i) their physical distance from each other and (ii) their individual intensity. The smaller the angular separations of sources, the higher the likelihood that they share the same corruptions in their radiated signals. Moreover, in order to get the best accuracy in the calibration results, there should be a balance between effective intensities of different clusters. Thus, in the clustering procedure, every source should be weighted suitably according to its brightness intensity.

The brightness distribution of radio source in the sky is a power law and the spatial distribution is Poisson. Therefore, clustering the sources via probabilistic clustering approaches is computationally complex. Hierarchical clustering (Johnson 1967) is a well-known clustering approach with a straightforward implementation suitable for our case. But, its computational cost grows exponentially with the size of its target data set which can be a disadvantage when the number of sources is huge. Weighted $K$-means clustering (Mac-Queen 1967; Kerdprasop, Kerdprasop & Sattayatham 2005) is also one of the most used clustering schemes applicable in clustered calibration. The advantage of this clustering technique is its low computational cost, which is proportional to the number of clusters and the size of the target data set. Nonetheless, we emphasize that the computational time taken by any of the aforementioned clustering algorithms is negligible compared with the computational time taken by the actual calibration. Therefore, we pursue all clustering approaches in this paper. However, the use of fuzzy $C$-means clustering (Bezdek 1981) in clustered calibration requires major changes in the calibration data model and will be explored in future work.

This paper is organized as follows. First, in Section 2, we present the general data model used in clustered calibration. In Section 3, we present modified weighted $K$-means and divisive hierarchical clustering for clustering sources in the sky. Next, in Section 4, we focus on analysing the performance of clustered calibration, with an a priori clustered sky model, and compare the results with unclustered calibration. In clustered calibration, there is contention between the improvement of SNR by clustering sources and the errors introduced by the clustering of sources itself. Thus, we relate the clustered calibration’s performance to the effective signal-to-interference plus noise ratio (SNIR) obtained for each cluster. For this purpose, we use statistical estimation theory and the Cramer–Rao Lower Bounds (CRB) (Kay 1993). In Section 5, we derive criteria for finding the optimum number of clusters for a given sky. We use the SNIR analysis and adopt Akaike’s information criterion (AIC; Akaike 1973) and the likelihood ratio test (LRT; Graves 1978) to estimate the optimum number of clusters. We present simulation results in Section 6 to show the superiority of clustered calibration to unclustered calibration and the performance of the presented criteria in detecting the optimum number of clusters. Finally, we draw our conclusions in Section 7. Through this paper, calibration is executed by the Space Alternating Generalized Expectation maximization (SAGE) (Fessler & Hero 1994; Yatawatta et al. 2009; Kazemi et al. 2011b) algorithm. Moreover, in our simulations, radio sources are considered to be uniformly distributed in the sky and their flux intensities follow Raleigh distribution, which is the worst case scenario. In real sky models, there usually exist only a few (two or three) number of bright sources which dominate the emission. In the presence of these sources and the background noise, it is impractical to solve for the other faint sources in the field of view individually. Therefore, obtaining a better calibration performance via the clustered calibration approach, compared to the unclustered one, is guaranteed. Therefore, in Section 6, we illustrate this using simulations in which the brightness distribution of sources is a power law with a very steep slope. On top of that, Kazemi, Yatawatta & Zaroubi (2012) and Yatawatta et al. (2012) also present the performance of clustered calibration on real observations using LOFAR.

The following notations are used in this paper: bold, lowercase letters refer to column vectors, e.g. $\mathbf{y}$. Upper case bold letters refer
to matrices, e.g. \( \mathbf{C} \). All parameters are complex numbers, unless stated otherwise. The inverse, transpose, Hermitian transpose and conjugation of a matrix are represented by \((\cdot)^{-1}, (\cdot)^T, (\cdot)^H\) and \((\cdot)^\ast\), respectively. The statistical expectation operator is referred to as \(E[\cdot] \). The matrix Kronecker product and the proper (strict) subset of a matrix are denoted by \(\otimes\) and \(\mathcal{S}\), respectively. \(\mathbf{I}_n\) is the \(n \times n\) identity matrix and \(\emptyset\) is the empty set. The Kronecker delta is presented by \(\delta_{ij}\). \(\mathbb{R}\) and \(\mathbb{C}\) are the sets of real and complex numbers, respectively. The Frobenius norm is shown by \(||\cdot||\). Estimated parameters are denoted by a hat, \(\hat{\cdot}\). All logarithmic calculations are to the base \(e\). The multivariate Gaussian and Complex Gaussian distributions are denoted by \(\mathcal{N}\) and \(\mathcal{CN}\), respectively.

## 2 Clustered Self-Calibration Data Model

In this section, we present the measurement equation of a polarimetric clustered calibration in detail (Hamaker, Bregman & Sault 1996; Hamaker 2006). Suppose we have a radio interferometer consisting of \(N\) polarimetric antennas where each antenna is composed of two orthogonal dual-polarization feeds that observe \(K\) compact sources in the sky. Every \(i\)th source signal, \(i \in \{1, 2, \ldots, K\}\), causes an induced voltage of \(\tilde{v}_{pi} = [v_{p1i} v_{p2i}]^T\) at \(X\) and \(Y\) dipoles of every \(p\)th antenna, \(p \in \{1, 2, \ldots, N\}\). In practice, the signal \(v_{pi}\), obtained at every antenna \(p\) is a linear superposition of the \(K\) sources corrupted signals \(\tilde{v}_{pi}\) where \(i \in \{1, 2, \ldots, K\}\), plus the antenna’s thermal noise. The multitude of ignored fainter sources also contributes to this additive noise.

The voltages collected at the instrument antennas get corrected for geometric delays, based on the location of their antennas, and some instrumental effects, like the antenna clock phases and electronic gains. Then, they are correlated in the array’s correlator to generate visibilities (Hamaker et al. 1996). The visibility matrix of the baseline \(p - q\), \(\mathbf{E}(\mathbf{v}_{pq} \otimes \mathbf{v}_{pq}^\ast)\), is given by

\[
\mathbf{V}_{pq} = \mathbf{G}_p \sum_{i=1}^{K} \mathbf{J}_{pi}(\theta) \mathbf{C}_{i [pq]} \mathbf{J}_{pi}(\theta)^\ast \mathbf{G}_q^\ast + \mathbf{N}_{pq}. \tag{2}
\]

In equation (2), \(\theta \in \mathbb{C}^P\), \(P = 4KN\), is the unknown instrumental and sky parameter vector, \(\mathbf{N}_{pq}\) is the additive \(2 \times 2\) noise matrix of the baseline \(p - q\), and \(\mathbf{C}_{i [pq]}\) is the Fourier transform of the \(i\)th source coherency matrix \(\mathbf{C}_i = \mathbf{E}(\mathbf{v}_i \otimes \mathbf{v}_i^\ast)\) (Born & Wolf 1999; Hamaker et al. 1996). If the \(i\)th source radiation intensity is \(I_i\), then \(\mathbf{C}_i = \frac{1}{2} I_i \mathbf{I}_2\). Considering this source to have equatorial coordinates (right ascension \(\alpha\), declination \(\delta\)), equal to \((\alpha_i, \delta_i)\), and the geometric components of baseline \(p - q\) to be \((u, v, w)\), then

\[
\mathbf{C}_{i [pq]} = e^{-2\pi j (u \sin \delta + v \cos \delta)} C_{i}, \tag{3}
\]

where \(j^2 = -1\) and,

\[
l = \sin(\alpha_i - \alpha_q) \cos(\delta_i), \quad m = \cos(\delta_i) \sin(\delta_i) - \cos(\alpha_i - \alpha_q) \cos(\delta_i) \sin(\delta_0)
\]

are the source direction components corresponding to the observation phase reference of \((\alpha_0, \delta_0)\) (Thompson et al. 2001). The errors common to all directions, such as the receiver delay and amplitude errors, are given by \(\mathbf{G}_p\) and \(\mathbf{G}_q\). Initial calibration at a finer time and frequency resolution is performed to estimate and correct for \(\mathbf{G}_p\)’s and the corrected visibilities are obtained as:

\[
\tilde{\mathbf{V}}_{pq} = \mathbf{G}_p^{-1} \mathbf{V}_{pq} \mathbf{G}_q^{-\ast}. \tag{4}
\]

The remaining errors are unique to a given direction, but residual errors in \(\mathbf{G}_p\)’s are also absorbed into these errors, which are denoted by \(\mathbf{J}_{pi}\) in the usual notation.

Vectorizing equation (4), the final visibility vector of the baseline \(p - q\) is given by

\[
\mathbf{v}_{pq} = \sum_{i=1}^{K} \mathbf{J}_{pi}(\theta) \otimes \mathbf{J}_{pi}(\theta)^\ast \mathbf{vec}(\mathbf{C}_{i [pq]}) + \mathbf{n}_{pq}. \tag{5}
\]

Stacking up all the cross-correlations (measured visibilities) and noise vectors in \(\mathbf{y}\) and \(\mathbf{n}\), respectively, the unclustered self-calibration measurement equation is given by

\[
\mathbf{y} = \sum_{i=1}^{K} \mathbf{s}_i(\theta) + \mathbf{n}. \tag{6}
\]

In equation (6), \(\mathbf{y}, \mathbf{n} \in \mathbb{C}^M\), \(M = 2N(N - 1)\), the noise vector is considered to have a zero mean Gaussian distribution with covariance \(\mathbf{P}, \mathbf{n} \sim \mathcal{N}(0, \mathbf{P}_{\emptyset} \otimes \mathbf{M})\), and the non-linear function \(\mathbf{s}_i(\theta)\) is defined as:

\[
\mathbf{s}_i(\theta) = \left[ \begin{array}{c} \mathbf{J}^T_{pi}(\theta) \otimes \mathbf{J}_{i1}(\theta)^\ast \mathbf{vec}(\mathbf{C}_{i [pq]}) \\ \vdots \\ \mathbf{J}^T_{pi}(\theta) \otimes \mathbf{J}_{i(N-1)}(\theta)^\ast \mathbf{vec}(\mathbf{C}_{i [pq]}) \end{array} \right]. \tag{7}
\]

Calibration is essentially the maximum likelihood (ML) estimation of the unknown parameters \(\theta\) (P complex values or 2P real values), or of the Jones matrices \(\mathbf{J}(\theta)\), from equation (6) and removal of the \(K\) sources. Note that calibration methods could also be applied to the uncorrected visibilities of equation (4) to estimate \(\mathbf{G}_p\), and \(\mathbf{G}_q\) errors as well.

The Jones matrix \(\mathbf{J}_{pi}\), for every \(i\)th direction and at every \(p\)th antenna, is given as:

\[
\mathbf{J}_{pi} = \mathbf{E}_{pi} \mathbf{Z}_{pi} \mathbf{F}_{pi}. \tag{8}
\]

In equation (8), \(\mathbf{E}_{pi}\), \(\mathbf{Z}_{pi}\) and \(\mathbf{F}_{pi}\) are the antenna’s voltage pattern, ionospheric phase fluctuation and Faraday Rotation Jones matrices, respectively. In practice, the \(\mathbf{E}, \mathbf{Z}\) and \(\mathbf{F}\) Jones matrices obtained for nearby directions and for a given antenna are almost the same. Thus, for every antenna \(p\), if the \(i\)th and \(j\)th sources have a small angular separation from each other, we may consider

\[
\mathbf{J}_{pi} \approx \mathbf{J}_{pj}. \tag{9}
\]

This is the underlying assumption for clustered calibration.

Clustered calibration first assigns source clusters, \(L_i\) for \(i \in \{1, 2, \ldots, Q\}\), where \(Q \ll K\), on which the sky variation is considered to be uniform. Then, it assumes that there exists a unique \(\mathbf{J}_{pi}\) which is shared by all the sources of the \(i\)th cluster \(L_i\), \(i \in \{1, 2, \ldots, Q\}\), at receiver \(p\), \(p \in \{1, 2, \ldots, N\}\). Based on that, the visibility at every baseline \(p - q\), given by equation (4), is reformulated as:

\[
\tilde{\mathbf{V}}_{pq} = \sum_{i=1}^{Q} \mathbf{J}_{pi}(\theta) \left\{ \sum_{l \in L_i} \mathbf{C}_l \right\} \mathbf{J}_{pi}(\theta)^\ast + \mathbf{N}_{pq}. \tag{10}
\]
In equation (10), $\tilde{N}_{pq}$ is the clustered calibration’s effective noise at baseline $p - q$ which will be explicitly discussed at Section 4. Note that clustered calibration estimates the new unknown parameter $\theta \in \mathbb{C}^p$, where $P = 4QN$. Denoting the effective signal of every $i$th cluster at baseline $p - q$ by

$$\tilde{C}_{i\mid(pq)} \equiv \sum_{j \in \mathcal{C}_i\mid(pq)} C_{i\mid(pq)},$$

(11)

the clustered calibration visibility vector at this baseline (vectorized form of equation 10) is

$$v_{pq} = \sum_{i=1}^{Q} \tilde{J}_{i\mid(pq)}(\tilde{\theta}) \otimes J_{i\mid(pq)}(\tilde{\theta})\text{vec}(\tilde{C}_{i\mid(pq)}) + \tilde{n}_{pq}.$$  

(12)

Finally, stacking up the visibilities of all the instrument’s baselines in vector $y$, clustered calibration’s general measurement equation is resulted as:

$$y = \sum_{i=1}^{Q} \tilde{s}_{i}(\tilde{\theta}) + \tilde{n}.$$  

(13)

In equation (13), $\tilde{s}_i$ is defined similar to $s_i$ in equation (7) where $J$ and $C$ are replaced by $\tilde{J}$ and $\tilde{C}$, respectively.

Because of the similarity between the clustered and the unclustered calibration’s measurement equations presented by equations (13) and (6), respectively, they could utilize the same calibration techniques. Thus, the only difference between these two types of calibration is that clustered calibration solves for clusters of sources instead of for the individual ones. That upgrades the signals which should be calibrated for from equation (3) to equation (11).

### 3 CLUSTERING ALGORITHMS

Clustering is grouping a set of data so that the members of the same group (cluster) have some similarities (Kauffman & Rousseau 1990). This similarity is defined based on the application of the clustering method.

We need to define clustering schemes in which two radio sources merge into a single cluster based on the similarity in their direction-dependent gain errors (equation 8). Radiations of sources that are close enough to each other in the sky are assumed to be affected by almost the same corruptions (equation 9). Based on that, we aim to design source clusters with small angular diameters. On the other hand, every cluster’s intensity is the sum of the intensities of its members (equation 11). In order to keep a balance between different clusters’ intensities, we intend to apply weighted clustering techniques in which the sources are weighted proportional to their intensities.

Suppose that the $K$ sources, $x_1, x_2, \ldots, x_K$ have $(\alpha_1, \delta_1), (\alpha_2, \delta_2), \ldots, (\alpha_K, \delta_K)$ equatorial coordinates, respectively. The aim is to provide $Q$ clusters so that the objective function $f = \sum_{q=1}^{Q} D(L_q)$ is minimized. $D(L_q)$ is the angular diameter of cluster $L_q$, for $q \in \{1, 2, \ldots, Q\}$, defined as:

$$D(L_q) \equiv \max \{d(x_i, x_j) | x_i, x_j \in L_q\},$$

(14)

d$(x_i, x_j)$ is the angular separation between any two points on the celestial sphere. Having two radio sources $a$ and $b$ with equatorial coordinates $(\alpha_a, \delta_a)$ and $(\alpha_b, \delta_b)$, respectively, the angular separation $d(\alpha, \delta)$, in radians, is obtained by

$$\tan^{-1} \frac{\sqrt{\cos^2 \delta_a \sin^2 \alpha + \cos \delta_a \sin \alpha - \sin \delta_b \cos \delta_a \cos \Delta \alpha}}{\sin \delta_a \sin \delta_b + \cos \delta_a \cos \delta_b \cos \Delta \alpha},$$

(15)

where $\Delta \alpha = \alpha_b - \alpha_a$.

For defining the centroids, we associate a weight to every source $x_i$, as:

$$w_i = w(x_i) = \frac{I_i}{T}, \text{ for } i \in \{1, 2, \ldots, K\},$$

(16)

where $I_i$ is the source’s intensity and $T = \min \{I_1, I_2, \ldots, I_K\}$. Applying the weights to the clustering procedure, the centroids of the clusters lean mostly towards the brightest sources. That causes a tendency in faint sources to gather with brighter sources close to them into one cluster. Thus, their weak signals are promoted being added up with some brighter sources’ signals. Moreover, very strong sources will be isolated such that their signals are calibrated individually, without being affected by the other faint sources.

We cluster radio sources using weighted K-means (Kerdprasop et al. 2005) and divisive hierarchical clustering (Johnson 1967) algorithms. Since the source clustering for calibration is performed offline, its computational complexity is negligible compared with the calibration procedure itself.

#### 3.1 Weighted K-means clustering

**Step 1.** Select the $Q$ brightest sources, $x_1, x_2, \ldots, x_Q$, and initialize the centroids of $Q$ clusters by their locations:

$$c_q = [\alpha_q^*, \delta_q^*], \text{ for } q \in \{1, 2, \ldots, Q\}, q^* \in \{1^*, 2^*, \ldots, Q^*\}.$$  

(17)

**Step 2.** Assign each source to the cluster with the closest centroid, defining the membership function

$$m_{L_q}(x_i) = \begin{cases} 1, & \text{if } d(x_i, c_j) = \min[d(x_i, c_j)] \text{ for } j = 1, \ldots, Q \\ 0, & \text{Otherwise.} \end{cases}$$

(18)

**Step 3.** Update the centroids by

$$c_q = \frac{\sum_{i=1}^{K} m_{L_q}(x_i) w_i x_i}{\sum_{i=1}^{K} m_{L_q}(x_i) w_i}, \text{ for } q \in \{1, 2, \ldots, Q\}.$$  

Repeat steps 2 and 3 until there are no reassignments of sources to clusters.

#### 3.2 Divisive hierarchical clustering

**Step 1.** Initialize the cluster counter $Q^*$ to 1, assign all the $K$ sources to a single cluster $L_1$ and $\varnothing$ to a set of null clusters $\mathcal{A}$.

**Step 2.** Choose cluster $L_{Q^*}$, for $q^* \in \{1, 2, \ldots, Q\} - A$, with the largest angular diameter

$$D(L_{Q^*}) = \max[D(L_q) | q \in \{1, 2, \ldots, Q\} - A].$$

(19)

**Step 3.** Apply the presented weighted K-means clustering technique to split $L_{Q^*}$ into two clusters, $L_{Q^*}'$ and $L_{Q^*}''$.

**Step 4.** If $D(L_{Q^*}') + D(L_{Q^*}'') < D(L_{Q^*})$, then set $Q' = Q^* + 1$, $L_{Q'} \equiv L_{Q^*}'$, $L_{Q''} \equiv L_{Q^*}''$, and $\mathcal{A} = \varnothing$; otherwise set $A = A \cup \{q^*\}$.

Repeat steps 2, 3 and 4 until $Q = Q^*$. 

#### 3.3 Comparison of clustering methods

Hierarchical clustering method tends to design clusters with almost the same angular diameters, whereas, the K-means clustering method tends to keep the same level of intensity at all its clusters. In practice, since hierarchical clustering method makes less errors in dedicating the same solutions to sources in small clusters, it performs better than weighted K-means clustering in a clustered
Figure 1. A simulated 8° by 8° sky of 50 point sources with intensities below 3 Jy. The source positions and their brightness are following uniform and Rayleigh distributions, respectively. The marker sizes are proportional to sources intensities.

calibration procedure. But, when the number of sources (and clusters) is very large ($Q \geq 100$), its prohibitive computational costs makes the fast $K$-means clustering method preferable.

3.3.1 Example 1: weighted $K$-means and hierarchical clustering

We simulate an 8° by 8° sky with 50 point sources with intensities below 3Jy. The source positions and their brightness follow uniform and Rayleigh distributions, respectively. The result is shown by Fig. 1 in which the symbol sizes are proportional to intensities of sources. Weighted $K$-means and divisive hierarchical clustering methods are applied to cluster the 50 sources into 10 source clusters. The results are presented in Figs 2 and 3, respectively. Fig. 2 shows that the weighted $K$-means clustering could design source clusters with considerably large angular diameters. Assigning the same calibration solutions to the sources of these large clusters could cause significant errors. However, as Fig. 3 shows, this is not the case for the hierarchical clustering and it constructs clusters with almost the same angular diameters.

Since the number of sources in this simulation is not that large ($K = 50$), the difference between execution time of the two clustering methods is not significant. Hence, in such a case, the use of hierarchical clustering method, rather than the weighted $K$-means, is advised. However, this is not the case when we have a large number of sources, and subsequently a large number of source clusters, in the sky. To demonstrate this, we use the two clustering techniques for clustering thousand of sources ($K = 1000$) into $Q$ source clusters, $Q \in \{3, 4, \ldots, 100\}$. The methods’ computational times versus the number of clusters are plotted in Fig. 4. As Fig. 4 shows, for large $Q$s, the computational cost of weighted $K$-means is much lower.

Figure 2. Fifty point sources are clustered into 10 source clusters by weighted $K$-means clustering technique. There is not a good balance between different clusters angular diameters.

Figure 3. Fifty point sources are clustered into 10 source clusters via hierarchical clustering method. Different clusters have almost the same angular diameters.

Figure 4. Weighted $K$-means and divisive hierarchical clustering methods computational costs. For small number of source clusters, there is no difference between execution times of the two clustering methods. But, when the number of source clusters is large, the computational cost of Weighted $K$-means becomes much cheaper than the one of the hierarchical clustering.
cheaper than the one of the hierarchical clustering. That can make the weighted K-means clustering method more suitable than the hierarchical clustering for such a case.

4 PERFORMANCE ANALYSIS

In this section, we explain the reasons for clustered calibration’s better performance, compared to unclustered calibration, at a low SNR (Kazemi et al. 2012). This superiority is in the sense of achieving an unprecedented precision in solutions with a considerably lower computational complexity, given the optimum clustering scheme. In the next section, we present different criteria for finding the optimum number of clusters at which the clustered calibration performs the best.

4.1 Cramer–Rao lower bounds

The most fundamental assumption in clustered calibration is that the sources at the same cluster have exactly the same corrections in their radiated signals. This assumption is of course incorrect; nonetheless, it provides us with a stronger signal, the sum of the signals in the whole cluster. We present an analytic comparison of clustered and unclustered calibration where we use the CRLB (Kay 1993) as a tool to measure the performance of the calibration.

4.1.1 Estimations of CRLB for two sources at a single cluster

For simplicity, first consider observing two point sources at a single baseline, let us say baseline $p - q$. Based on equation (4), the visibilities are given by

$$\tilde{V}_{pq} = J_{pq} \mathcal{C}_{1(pq)} J_{pq}^\dagger + J_{pq} \mathcal{C}_{2(pq)} J_{pq}^\dagger + N_{pq},$$

(20)

in the unclustered calibration strategy. Vectorizing $V_{pq}$, the visibility vector is

$$\mathbf{y} = \mathbf{J}_{pq}^* \otimes \mathbf{J}_{pq} \mathcal{V}(\mathbf{C}_{1(pq)}) + \mathbf{J}_{pq}^* \otimes \mathbf{J}_{pq} \mathcal{V}(\mathbf{C}_{2(pq)}) + \mathbf{n}_{pq}.$$  

Assuming $\mathbf{n}_{pq} \sim \mathcal{CN}(\mathbf{0}, \sigma^{-2} \mathbf{I}_4)$, we have

$$\mathbf{y} \sim \mathcal{CN}(\mathbf{s}(\theta), \sigma^{-2} \mathbf{I}_4),$$

(21)

where

$$\mathbf{s}(\theta) = \sum_{i=1,2} \mathbf{J}_{pq}^* \otimes \mathbf{J}_{pq} \mathcal{V}(\mathbf{C}_{1(pq)}).$$

(22)

Using equation (21), the log-likelihood function of the visibility vector $\mathbf{y}$ is given by

$$\mathcal{L}(\theta | \mathbf{y}) = -4 \ln \left( \frac{\pi}{\sigma^2} \right) - \sigma^{-2} (\mathbf{y} - \mathbf{s}(\theta))\mathcal{H}(\mathbf{y} - \mathbf{s}(\theta)).$$

(23)

CRLB is a tight lower bound on the error variance of any unbiased parameter estimators (Kay 1993). Based on its definition, if the log-likelihood function of the random vector $\mathbf{y}$, $\mathcal{L}(\theta | \mathbf{y})$, satisfies the ‘regularity’ condition

$$E_\mathbf{y} \left[ \frac{\partial}{\partial \theta} \mathcal{L}(\theta | \mathbf{y}) \right] = 0, \quad \text{for all } \theta,$$

(24)

then for any unbiased estimator of $\theta$, $\hat{\theta}$,

$$\text{var}(\hat{\theta}) \geq \mathbf{I}^{-1}(\theta)|_{\theta = \mathbf{0}}, \quad \text{for } i \in \{1, \ldots, M\},$$

(25)

where $\mathbf{I}(\theta)$ is the Fisher information matrix defined as:

$$\mathbf{I}(\theta) = -E_\mathbf{y} \left[ \frac{\partial^2 \mathcal{L}(\theta | \mathbf{y})}{\partial \theta \partial \theta^*} \right].$$

(26)

In other words, the variance of any unbiased estimator of the unknown parameter vector $\theta$ is bounded from below by the diagonal elements of $[\mathbf{I}(\theta)]^{-1}$. Using equations (23) and (26), the Fisher information matrix of the visibility vector $\mathbf{y}$ is obtained as:

$$\mathbf{F}(\theta) = 2\sigma^{-2} \mathcal{R}(\mathbf{J}_{pq}^\dagger \mathbf{J}_{pq}),$$

(27)

where $\mathbf{J}_{pq}$ is the Jacobian matrix of $\mathbf{s}$ with respect to $\theta$

$$\mathbf{J}_{pq}(\theta) = \sum_{i=1}^2 \frac{\partial}{\partial \theta} [\mathbf{J}_{pq}^* \otimes \mathbf{J}_{pq} \mathcal{V}(\mathbf{C}_{1(pq)})].$$

(28)

Thus, variations of any unbiased estimator of parameter vector $\theta$, let us say $\hat{\theta}$, is bounded from below by the CRLB as:

$$\text{Var}(\hat{\theta}) \geq [2\sigma^{-2} \mathcal{R}(\mathbf{J}_{pq}^\dagger \mathbf{J}_{pq})]^{-1}.$$ 

(29)

Let’s try to bound the error variations of the clustered calibration parameters assuming that the two sources construct a single cluster, called cluster number 1. We reform equation (20) as:

$$\mathbf{V}_{pq} = \mathbf{J}_{pq} \mathcal{C}_{1(pq)} + \mathbf{J}_{pq} \mathcal{C}_{2(pq)} \mathbf{J}_{pq}^\dagger + \mathbf{N}_{pq},$$

(30)

where $\mathbf{C}_{1(pq)}$, referred to as the ‘clustering error’ matrices, are given by

$$\mathbf{C}_{1(pq)} = \mathbf{J}_{pq} \mathcal{C}_{1(pq)} \mathbf{J}_{pq}^\dagger - \mathbf{J}_{pq} \mathcal{C}_{2(pq)} \mathbf{J}_{pq}^\dagger.$$ 

(31)

and $\mathbf{J}_{pq}(\hat{\theta})$ is the clustered calibration solution at receiver $p$. Equation (30) implies that what is considered as the noise matrix $\mathbf{N}_{pq}$ in the clustered calibration data model, equation (10), is in fact

$$\mathbf{N}_{pq} = \mathbf{C}_{1(pq)} + \mathbf{C}_{2(pq)} + \mathbf{N}_{pq}.$$ 

(32)

Vectorizing equation (30), the clustered calibration visibility vector is obtained by

$$\mathbf{y} = \mathbf{J}_{pq}^* \otimes \mathbf{J}_{pq} \mathcal{V}(\mathbf{C}_{1(pq)}) + \mathbf{J}_{pq}^* \otimes \mathbf{J}_{pq} \mathcal{V}(\mathbf{C}_{2(pq)}) + \mathbf{n}_{pq},$$

(33)

where $\mathbf{n}_{pq} = \mathcal{N}(\mathbf{0})$.

We point out that depending on the observation as well as the positions of the two sources on the sky, the clustering error $\mathbf{C}_{1(pq)}$ will have different properties. However, in order to study the performance of the clustered calibration in a statistical sense, and to simplify our analysis, we make the following assumptions.

(i) Consider statistical expectation over different observations and over different sky realizations where the sources are randomly distributed on the sky. In that case, almost surely $\mathbf{E}[\mathbf{J}] \rightarrow \mathbf{E}[\mathbf{J}]$ and consequently,

$$\mathbf{E}[\mathbf{N}_{pq}] \rightarrow \mathbf{0}.$$ 

(34)

In other words, we assume the clustering error to have zero mean over many observations of different parts of the sky.

(ii) We assume that the closer the sources are together in the sky, the smaller the errors introduced by clustering would be. Therefore, given a set of sources, the clustering error will reduce as the number of clusters increase. In fact, this error introduced by clustering vanishes when the number of clusters is equal to the number of sources (each cluster contains only one source). Therefore, given a set of sources, the variance of $\mathbf{C}_{1(pq)}$ will decrease as the number of clusters increase.

Using equation (34) and bearing in mind that $\mathbf{E}[\mathbf{N}_{pq}] = \mathbf{0}$, we can consider $\mathbf{n}_{pq} \sim \mathcal{CN}(\mathbf{0}, \sigma^{-2} \mathbf{I}_4)$, where $\mathbf{E}[\mathbf{n}_{pq}] = \mathbf{0}$. 

$$\mathbf{E}[\mathbf{n}_{pq}] = \mathbf{0}.$$ 

(35)
Therefore,\
\[ y \sim \mathcal{CN}(\tilde{s}, \sigma^2 I_4), \]
\[ \tilde{s} = J_{p1}^2 \otimes J_{p1} \text{vec}(C_{1(pq)} + C_{2(pq)}), \]
and similar to equation (29), we have
\[ \text{Var}(\theta) \geq (2\sigma^2)^{-1} \text{Re}(J_{p1}^2 \Gamma_{p1})^{-1}. \] (35)

We use numerical simulations to compare the unclustered and clustered calibrations performances via their CRLBs which are given by equations (29) and (35), respectively.

### 4.1.2 Example 2: CRLB for two sources and one cluster

We simulate a 12 h observation of two point sources with intensities \( I_1 = 11.25 \) and \( I_2 = 2.01 \) Jy at sky coordinates \((l, m)\) equal to \((-0.014, -0.005)\) and \((-0.011, -0.010)\) rad, respectively. We use the uv coverage of Westerbork Synthesis Radio Telescope (WSRT) with 14 receivers in this simulation.

We consider the Jones matrices \( J \) in equation (20) to be diagonal. Their amplitude and phase elements follow \( U(0.75, 0.95) \) and \( U(0.003, 0.004) \) distributions, respectively. The background noise is \( N \sim \mathcal{CN}(0, 10I) \). Jones matrices of the clustered calibration, \( J_{p1} \) for \( p = 1, 2 \), are obtained as \( J_{p1} = U(0.02, 0.40) e^{i\theta(pq)} \). For 20 realizations of \( J \) matrices, we calculated CRLB of the unclustered and clustered calibrations using equations (29) and (35), respectively. The results are presented in Fig. 5. As shown in this figure, for small enough errors matrices \( \Gamma \) of equation (31), the clustered calibration’s CRLB stands below the unclustered calibration’s CRLB.

On the other hand, with increasing power of error matrices, or the power of effective noise \( N \), the unclustered calibration’s CRLB becomes lower than the clustered calibration’s CRLB.

### 4.1.3 Analysis of CRLB

Generally, if source 1 is considerably brighter than source 2, \( ||C_{1(pq)}|| \gg ||C_{2(pq)}|| \), and if the weak source power is much lower than the noise level, \( ||C_{2(pq)}|| \ll ||N_{pq}|| \), then clustered calibration’s performance is better than unclustered calibration. Note that the worst performance of both calibrations is at the faintest source and we are more concerned with comparing the CRLBs for this source.

The CRLBs obtained for the unclustered and clustered calibrations in equations (29) and (35), respectively, are both almost equal to the inverse of the SINR, \( \text{SINR}^{-1} \). In unclustered calibration, the effective signal for the faintest source is \( C_{2(pq)} \) where the noise is \( N_{pq} \). Therefore, SINR for this source is
\[ \text{SINR}_c = \frac{||C_{2(pq)}||^2}{||N_{pq}||^2}. \] (36)

But, in clustered calibration, the effective signal and noise are \( C_{(pq)} = C_{1(pq)} + C_{2(pq)} \) and \( N_{pq} \), respectively. Thus, SINR for the cluster is
\[ \text{SINR}_c = \frac{||C_{(pq)}||^2}{||N_{pq}||^2}. \] (37)

Clustered calibration has an improved performance when
\[ \text{SINR}_c \gg \text{SINR}_c. \] (38)

Consider the two possible extremes in a clustered calibration procedure.

(i) Clustering many sources in a large field of view to a very small number of clusters. In this case, the angular diameter of a cluster is probably too large for the assumption of uniform corruptions to apply. Subsequently, dedicating a single solution to all the sources of every cluster by clustered calibration introduces clustering error matrices \( \Gamma \) with a large variance (see equation 31). Having high interference power, the clustered calibration effective noise \( N \) of equation (32) becomes very large. Therefore, clustered calibration SINR will be very low and it does not produce high quality results.

(ii) Clustering sources in a small field of view to a very large number of clusters. In this case, the variances of \( \Gamma \) matrices are almost zero while the signal powers of source clusters are almost as low as of the individual sources. Therefore, the SINR of clustered calibration is almost equal to the unclustered calibration SINR and the calibration performance is expected to be almost the same as well.

Thus, the best efficiency of clustered calibration is obtained at the smallest number of clusters for which equation (38) is satisfied. We use the SINR of equation (38) as an efficient criterion for detecting the optimum number of clusters.

### 4.1.4 Generalization to many sources and many clusters

For the visibility vector \( y \) of unclustered calibration’s general data model, presented by equation (6), we have
\[ y \sim \mathcal{CN} \left( \sum_{i=1}^{K} s_i(\theta), \Pi \right). \] (39)

Therefore, the CRLB of unclustered calibration is
\[ \text{var}(\theta) \geq \left[ 2 \text{Re} \left\{ \left( \sum_{i=1}^{K} J_i(\theta) \right)^\dagger \Pi^{-1} \left( \sum_{i=1}^{K} J_i(\theta) \right) \right\} \right]^{-1}, \]
where \( J_i \) is the Jacobian matrix of \( s_i \) with respect to \( \theta \).

Computing the exact CRLB is more complicated when we have clustered calibration. In the clustered calibration measurement equation, given by equation (13), we have
\[ \tilde{n} = \sum_{i=1}^{K} \Gamma_i + n. \] (40)
where \( \mathbf{n} \) is the unclustered calibration’s noise vector,

\[
\mathbf{\Gamma}_i = \left[ \text{vec}(\mathbf{I}_{(12)})^T \cdots \text{vec}(\mathbf{I}_{(N-1,N)})^T \right]^T,
\]

and \( \mathbf{\Gamma}_{(pq)} \) is given by equation (31). Due to the existence of the nuisance parameters \( \mathbf{\Gamma}_i \) in the clustered calibration data model, calculation of its conventional CRLB is impractical. This leads us to the use of Cramer–Rao like bounds devised in the presence of the nuisance parameters (Gini & Reggiannini 2000). We apply the modified CRLB (MCRLB; Gini, Reggiannini & Mengali 1998) to the performance of clustered calibration.

The MCRLB for estimating the errors of \( \hat{\mathbf{\theta}} \) in the presence of the nuisance parameters \( \mathbf{\Gamma} \) (clustering error) is defined as:

\[
\text{var}(\hat{\mathbf{\theta}}) \geq \left[ E_{\mathbf{\theta},\mathbf{\Gamma}} \left\{ -E_{\mathbf{\theta},\mathbf{\Gamma}} \left[ \frac{\partial}{\partial \mathbf{\theta}} \frac{\partial}{\partial \hat{\mathbf{\theta}}} \ln P(\mathbf{y}|\hat{\mathbf{\theta}}) \right] \right\} \right]^{-1},
\]

where \( P(\mathbf{y}|\hat{\mathbf{\theta}}) \) is the probability density function of the visibility vector \( \mathbf{y} \) assuming that the matrices \( \mathbf{\Gamma} \) of equation (41) are a priori known. Since \( \mathbf{n} \sim \mathcal{CN}(0, \mathbf{\Pi}) \), from equation (13) we have

\[
\mathbf{y}|\mathbf{\Gamma} \sim \mathcal{CN} \left( \sum_{i=1}^{Q} \mathbf{s}_i \mathbf{\Gamma}_i, \mathbf{\Pi} \right),
\]

and therefore in equation (42), \( -E_{\mathbf{\theta},\mathbf{\Gamma}} \left[ \frac{\partial}{\partial \mathbf{\theta}} \frac{\partial}{\partial \hat{\mathbf{\theta}}} \ln P(\mathbf{y}|\hat{\mathbf{\theta}}) \right] \), which is called the modified Fisher information, is equal to

\[
2\pi \left\{ \sum_{i=1}^{Q} J_k(\mathbf{\theta}) + \sum_{i=1}^{K} J_{\mathbf{\Gamma}_i}(\hat{\mathbf{\theta}}) \right\} \mathbf{\Pi}^{-1} \left( \sum_{i=1}^{Q} J_k(\mathbf{\theta}) + \sum_{i=1}^{K} J_{\mathbf{\Gamma}_i}(\hat{\mathbf{\theta}}) \right).
\]

Note that \( E_{\mathbf{\theta},\mathbf{\Gamma}} \) in equation (42) could be estimated by Monte Carlo method.

As a rule of thumb, reducing the heavy computational cost of MCRLB, one can interpret the SINR test of equation (38) as follows. If in average the effective SINR of clustered calibration, \( \text{SINR}_c \), gets higher than the effective SINR of unclustered calibration obtained for the weakest observed signal, \( \text{SINR}_u \),

\[
\text{E} [\text{SINR}_c] \geq \text{E} [\text{SINR}_u],
\]

then clustered calibration can achieve a better results. In equation (44), the expectation is taken with respect to the noise \( \mathbf{N} \), error matrices \( \mathbf{\Gamma} \), and all the baselines.

4.1.5 Example 3: MCRLB and SINR estimations

We simulate WSRT including \( N = 14 \) receivers which observe 50 sources with intensities below 15 Jy. The source positions and their brightness follow uniform and Rayleigh distributions, respectively. The background noise is \( \mathbf{N} \sim \mathcal{CN}(0, 15 \mathbf{I}_{14}) \), where \( M = 2N(N-1) = 364 \). We cluster sources using divisive hierarchical clustering, into \( Q \) number of clusters, where \( Q \in \{3, 4, \ldots, 50\} \). Clustered calibration’s Jones matrices, \( \mathbf{J} \), are generated as \( \mathbf{J}(0.9, 1.1)e^{i\Delta(0.02)} \).

Since for smaller number of clusters, we expect larger interference (errors) in clustered calibration’s solutions, for every \( Q \), we consider \( \sum_{i=1}^{50} \mathbf{\Gamma}_i \sim \mathcal{CN}(0, 1 \mathbf{I}_{14}) \). The choice of the complex Gaussian distribution for the error matrices \( \mathbf{\Gamma} \) is due to the central limit theorem and the assumptions made in Section 4.1.1.

We proceed to calculate the clustered calibration’s MCRLB, given by equation (42), and \( \text{E} [\text{SINR}_c] \), utilizing the Monte Carlo method. Jacobian matrices for MCRLB are calculated numerically and in computation of \( \text{E} [\text{SINR}_c] \), signal power of every cluster is obtained only using the cluster’s brightest and faintest sources. The estimated results of MCRLB and \( \text{E} [\text{SINR}_c] \) are presented in Figs 6 and 7, respectively. As we can see in Fig. 6, for very small \( Q \), the effect of interference is large, MCRLB is high. By increasing the number of clusters, MCRLB decreases and reaches its minimum where the best performance of the clustered calibration is expected. After that, due to the dominant effect of the background noise, MCRLB starts to increase until it reaches the unclustered calibration CRLB.

As Fig. 7 shows, \( \text{E} [\text{SINR}_c] \) is low for very small \( Q \), when the interference is large. By increasing the number of clusters the SINR increases and gets its highest level for which the best performance of the clustered calibration is expected. After that, it decreases due to the dominant effect of the background noise, and converges to the unclustered calibration SINR.
4.2 Computational cost

In the measurement equation of unclustered calibration, presented in equation (6), we have \( M = 2N(N-1) \) constraints given by the visibility vector \( y \), and need to solve for \( P = 4KN \) unknown parameters \( \theta \). If \( P > M \), then equation (6) will be an underdetermined non-linear system. This clarifies the need of having a small enough \( N \) (number of antennas) and a large enough \( K \) (number of sources) for estimating \( \theta \). However, clustered calibration, equation (13), has the advantage of decreasing the number of directions, \( K \), relative to the number of source clusters, \( Q \ll K \). This considerably cuts down the number of unknown parameters \( P \) that needs to be calibrated, thus reducing the computational cost of calibration.

5 SELECTION OF NUMBER OF CLUSTERS

Consider a clustered calibration procedure with a predefined clustering scheme. There is no guarantee that the calibration results for \( Q \) number of clusters, where \( Q \in \{1, 2, \ldots, K\} \) is randomly chosen, is the most accurate. Thus, we seek the optimum number of clusters at which the clustered calibration performs the best. In this section, we describe the use of: (i) AIC (Akaike 1973), as well as (ii) LRT (Graves 1978), in finding this optimum \( Q \) for a given observation. Some other alternative criteria could also be found in Wax & Kailath (1985). Note that for different clustering schemes the optimum \( Q \) is not necessarily the same.

5.1 Akaike’s Information Criterion

We utilize AIC to find the optimum \( Q \) for clustered calibration. Consider having \( \tilde{n} \sim \mathcal{CN}(0, \sigma^2 I_{2N}) \) in the general data model of clustered calibration, equation (13). Then, the log-likelihood of the visibility vector \( y \) is given by

\[
\mathcal{L}(\theta|y) = -M \log \pi - M \log \tilde{\sigma}^2 \]

\[
\quad -\frac{1}{\tilde{\sigma}^2} (y - \sum_{i=1}^{Q} \tilde{s}_i(\tilde{\theta}))^H (y - \sum_{i=1}^{Q} \tilde{s}_i(\tilde{\theta})) .
\]  

(45)

The ML estimation of the noise variance \( \tilde{\sigma}^2 \) is

\[
\tilde{\sigma}^2 = \frac{1}{M} \left( y - \sum_{i=1}^{Q} \tilde{s}_i(\tilde{\theta}) \right)^H \left( y - \sum_{i=1}^{Q} \tilde{s}_i(\tilde{\theta}) \right).
\]

(46)

Substituting equation (46) in equation (45), we arrive at the ML estimation of \( \tilde{\theta} \),

\[
\mathcal{L}(\tilde{\theta}|y) = -M \log \pi - M \log \tilde{\sigma}^2
\]

\[-M \log \frac{1}{M} \left( y - \sum_{i=1}^{Q} \tilde{s}_i(\tilde{\theta}) \right)^H \left( y - \sum_{i=1}^{Q} \tilde{s}_i(\tilde{\theta}) \right) .
\]

(47)

Using equation (47), the AIC is given by

\[
\text{AIC}(Q) = -2 \mathcal{L}(\tilde{\theta}|y) + 2(2P).
\]

(48)

The optimum \( Q \) is selected as the one that minimizes \( \text{AIC}(Q) \).

5.2 Likelihood-ratio test

Errors in clustered calibration originate from the system (sky and instrumental) noise, ‘clustering errors’ introduced in Section 4.1.1, and ‘solver noise’ which is referred to as errors produced by the calibration algorithm itself. We assume that the true Jones matrices along different directions (clusters) at the same antenna are statistically uncorrelated. Note that this assumption is only made for the LRT to produce reasonable results and this assumption is not needed for clustered calibration to work. Therefore, if such correlations exist, they are caused by the aforementioned errors. Consequently, the more accurate the clustered calibration solutions are, the smaller their statistical similarities would be. Based on this general statement, the best number of clusters in a clustered calibration procedure is the one which provides us with the minimum correlations in the calibrated solutions. Note that for a fixed measurement, the correlation due to the system noise is fixed. Therefore, differences in the statistical similarities of solutions obtained by different clustering schemes are only due to ‘clustering errors’ and ‘solver noise’.

To investigate the statistical interaction between the gain solutions we apply the LRT.

Consider the clustered calibration solution \( \tilde{J}_{pl}(\tilde{\theta}) \) for directions \( i, i \in \{1, 2, \ldots, Q\} \), at antennas \( p \), where \( p \in \{1, 2, \ldots, N\} \).

\[
\tilde{J}_{pl} = \begin{bmatrix}
\tilde{J}_{11,p} & \tilde{J}_{12,p} \\
\tilde{J}_{21,p} & \tilde{J}_{22,p}
\end{bmatrix}
\]

(49)

Then, the parameter vector \( \tilde{\theta}_{pl} \) (corresponding to the \( i \)th direction and \( p \)th antenna) is obtained by

\[
\tilde{\theta}_{pl} = \left[ \text{Re}(\tilde{J}_{11,p}) \text{Im}(\tilde{J}_{11,p}) \ldots \text{Re}(\tilde{J}_{22,p}) \text{Im}(\tilde{J}_{22,p}) \right]^T.
\]

(50)

Let us define for each antenna \( p \) and each pair of directions \( k \) and \( l \), where \( k \) and \( l \) are belong to \( \{1, 2, \ldots, Q\} \), a vector \( z_{pl} \) as:

\[
z_{pl} = \left[ \theta_{pk}^T \theta_{pl}^T \right]^T.
\]

(51)

In fact, we are concatenating the solutions of the same antenna for two different directions (clusters) together.

We define the null \( H_0 \) model as:

\[
H_0 : z_{pl} \sim \mathcal{N}(m, \Sigma_0).
\]

(52)

where

\[
m = \left[ \bar{m}(\tilde{\theta}_{pk})^T \bar{m}(\tilde{\theta}_{pl})^T \right]^T,
\]

(53)

and

\[
\Sigma_0 = \begin{bmatrix}
s^2(\tilde{\theta}_{pk}) & 0 \\
0 & s^2(\tilde{\theta}_{pl})
\end{bmatrix}.
\]

(54)

In equations (53) and (54), \( \bar{m} \) and \( s^2 \) are denoting sample mean and sample variance, respectively. Note that having a large number of samples in hand, the assumption of having a Gaussian distribution for solutions is justified according to the central limit theorem. The structure of the variance matrix \( \Sigma_0 \) tells us that the statistical correlation between the components of the random vector \( z_{pl} \), or between the solutions \( \theta_{pk} \) and \( \theta_{pl} \), is zero. This is the ideal case in which there are no estimation errors.

To investigate the validity of the null model compared with the case in which there exists some correlation between the solutions, we define the alternative \( H_1 \) model as:

\[
H_1 : z_{pl} \sim \mathcal{N}(m, \Sigma_1),
\]

(55)

where the variance matrix \( \Sigma_1 \) is given by

\[
\Sigma_1 = \begin{bmatrix}
s^2(\tilde{\theta}_{pk}) \text{Cov}(\tilde{\theta}_{pk}, \tilde{\theta}_{pl}) \\
\text{Cov}(\tilde{\theta}_{pk}, \tilde{\theta}_{pl})^T & s^2(\tilde{\theta}_{pl})
\end{bmatrix}.
\]

(56)

\text{Cov}(\tilde{\theta}_{pk}, \tilde{\theta}_{pl}) \) in equation (56) is the 8 × 8 sample covariance matrix.
Using the above models, the likelihood-ratio is defined as
\[ \Lambda = -2 \ln \left( \frac{\text{Likelihood for null model}}{\text{Likelihood for alternative model}} \right), \]
which has a \( \chi^2 \) distribution with 64 degrees of freedom. As \( \Lambda \) becomes smaller, the null model, in which the statistical correlation between the solutions is zero, becomes more acceptable rather than the alternative model. Therefore, the smaller the \( \Lambda \) is, the less the clustered calibration's errors are, and vice versa.

6 SIMULATION STUDIES

We use simulations to compare the performance of unclustered and clustered calibration. Working with simulations has the advantage of having the true solutions available, which is not the case in real observations. That makes the comparison much more objective. Nevertheless, the better performance of the clustered calibration in comparison with the unclustered ones in calibrating for real observations of LOFAR is also shown by Kazemi et al. (2011a) and Yatawatta et al. (2012).

We simulate an east–west radio synthesis array including 14 antennas (similar to WSRT) and an 8\(^\circ\) by 8\(^\circ\) sky with 50 sources with low intensities, below 3 Jy. The source positions and their brightness follow uniform and Rayleigh distributions, respectively. The single channel simulated observation at 355 MHz is shown in Fig. 8.

We proceed to add gain errors, multiplying source coherencies by the Jones matrices, as it is shown in equation (2), to our simulation. The amplitude and phase of the Jones matrices' elements are generated using linear combination of sine and cosine functions. We aim at simulating a sky with almost uniform variations on small angular scales. In other words, we provide very similar Jones matrices for sources with small angular separations. To accomplish this goal, for every antenna, we first choose a single direction as a reference and simulate its Jones matrix as it is explained before. Then, for the remaining 49 sources, at that antenna, the Jones matrices (amplitude and phase terms) are that initial Jones matrix multiplied by the inverse of their corresponding angular distances from that reference direction. The result of adding such gain errors to our simulation is shown in Fig. 9.

6.1 Performance comparison of the clustered and unclustered calibrations at SNR = 2

We add noise \( n \sim \text{CN}(0, \sigma^2 I) \) with \( \sigma^2 = 28 \), as it is shown in equation (6), to our simulation. The result has an SNR = 2 and is presented in Fig. 10. We have chosen to present the case of SNR = 2 since for this particular simulated observation both the divisive hierarchical and the weighted \( K \)-means clustered calibrations achieve their best performances at the same number of clusters, as will be shown later in this section.

We apply unclustered and clustered calibration on the simulation to compare their efficiencies. The 50 sources are grouped into \( Q \in \{3, 4, \ldots, 49\} \) number of clusters, using the proposed divisive hierarchical and weighted \( K \)-means clustering algorithms. Self-calibration is implemented via Space Alternating Generalized Expectation Maximization (SAGE) algorithm (Fessler & Hero 1994; Yatawatta et al. 2009; Kazemi et al. 2011b) with nine iterations. Plots of the averaged Frobenius distance between the simulated (true) Jones matrices and the obtained solutions is shown in Fig. 11. As we can see in Fig. 11, for both clustering schemes, increasing the number of clusters decreases this distance and the minimum is reached at approximately 33 clusters (\( Q = 33 \)). Beyond this number of clusters, it increases until the 50 individual sources become individual clusters. This shows that the best performance of both the divisive hierarchical and the weighted \( K \)-means clustered calibrations is approximately at 33 clusters and is superior to that of the unclustered calibration.

The Frobenius distance curves in Fig. 11, the MCRLB curve in Fig. 6, and the SINR curve in Fig. 7 illustrate that clustered calibration with an extremely low number of clusters does not necessarily
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Figure 10. Simulated image of sky, corrupted by gain errors and by noise, as in equation (6). The simulated noise vector $n$ has zero mean complex Gaussian distribution and the SNR is equal to 2.

Figure 11. The average Frobenius distance between the simulated (true) Jones matrices and the solutions of clustered and unclustered calibrations. The two curves represent clustered calibration via divisive hierarchical and weighted $K$-means clustering algorithms. By increasing the number of clusters, for both clustering methods, this distance is decreased and gets its minimum approximately at 33 clusters. After that, it is increased till the 50 individual sources. That shows that the best performance of the clustered calibration is at around 33 clusters.

perform better than the unclustered calibration. The reason is that when there are only a small number of clusters, the interference, or the so-called clustering errors introduced in Section 4.1.1, is relatively large. Therefore, the effect of this interference dominates the clustering of signals. On the other hand, we reach the theoretical performance limit approximately after 25 number of clusters and therefore increasing the number beyond this point gives highly variable results, mainly because we are limited by the number of constraints as opposed to the number of parameters that we need to solve for. But, this is not the case for the plots in Figs 6 and 7. The reason is that the MCRLB results of Fig. 6 as well as the SINR results of Fig. 7 are obtained by the Monte Carlo method with iterations over 50 different sky and noise realizations. However, Fig. 11 is limited to the presented specific simulation with only one sky and one noise realization.

The residual images of the unclustered calibration as well as the divisive hierarchical and weighted $K$-means clustered calibrations for $Q = 33$ are shown by Figs 12 and 13, respectively. As it is shown in Fig. 12, in the result of unclustered calibration, the sources are almost not subtracted at all and there is a significant residual error remaining. The residuals have asymmetric Gaussian distribution with variance $\sigma^2 = 82.29$ which is much larger than the true noise variance $\sigma^2 = 10.85$. On the other hand, the sources have been perfectly subtracted in the case of clustered calibration, Fig. 13, and the residuals converge to the simulated background noise distribution. The residuals of the divisive hierarchical and weighted $K$-means clustered calibrations follow symmetric zero mean Gaussian distributions with $\sigma^2 = 20.17$ and $\sigma^2 = 18.76$, respectively. These variances are closer to the simulated one $\sigma^2 = 10.85$ and this indicates the promising performance of clustered calibration. As we can see, hierarchical clustered calibration provides a slightly better result compared to the $K$-means one. This is due to the fact that hierarchical clustering constructs clusters of smaller angular diameters and thus it assigns the same calibration solutions to the sources with smaller angular separations.

We also calculate the root-mean-squared error of prediction (RMSEP) to assess the performance of clustered and unclustered calibrations’ non-linear regressions. The results of log(RMSEP), presented by Fig. 14, also justify that the best efficiencies of the hierarchical and $K$-means clustered calibrations are obtained at 33 clusters. But, note that there is a difference between the behaviour of log(RMSEP) plot of Fig. 14 and the plots of MCRLB, SINR and Frobenius distance between the simulated Jones matrices and solutions in Figs 6, 7 and 11, respectively. In Fig. 14, log(RMSEP) of clustered calibration is less than that of unclustered calibration, even for extremely low number of clusters. This means that even with those low number of clusters, clustered calibration performs...
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Figure 13. The residual images of the clustered calibration using hierarchical (right) and weighted $K$-means (left) clustering methods with 33 source clusters. Calibration is implemented by SAGE algorithm. The sources are subtracted perfectly and the residuals converge to the simulated background noise distribution. The hierarchical clustering and weighted $K$-means residuals follow symmetric zero mean Gaussian distributions with $\sigma^2 = 20.17$ and $\sigma^2 = 18.76$, respectively, where the simulated noise distribution is $CN(0, 10.85)$. 

Figure 14. The RMSEP for clustered and unclustered calibrations. The results are obtained using a base 10 logarithmic scale. The two curves are corresponding to clustered calibration via divisive hierarchical and weighted $K$-means clustering algorithms. By increasing the number of clusters, the results are decreased and the minimum result is obtained at around 33 clusters. After that, the results are increased till the 50 individual sources. That shows the superior performance of the clustered calibration compared to the unclustered one. The best performance of the clustered calibration for both of the applied clustering methods is at around 33 clusters. 

After executing a calibration for the above data model, there is a distance between the target parameters $\hat{\theta}$ and the estimated solutions $\hat{\tilde{s}}(\hat{\theta})$. This is the so-called solver noise, mentioned in Section 5.2. Thus, the residuals are given as:

$$y = \sum_{i=1}^{Q} \hat{s}_i(\hat{\theta}) = \sum_{i=1}^{Q} [\hat{s}_i(\hat{\theta}) - \hat{s}_i(\hat{\theta})] + \sum_{i=1}^{K} \Gamma_i + n. \quad (59)$$

From equation (59), we immediately see that the background noise $n$ is fixed and the ‘clustering errors’ are calculated for all the sources as $\sum_{i=1}^{K} \Gamma_i$. However, since we solve only for $Q$ directions and not for all the $K$ sources individually, the ‘solver noise’ part, $\sum_{i=1}^{Q} \hat{s}_i(\hat{\theta}) - \hat{s}_i(\hat{\theta})$, is also calculated only for $Q$ clusters and not for all the $K$ sources. It is clear that for a very small $Q$, this term could be much less than for $Q = K$. Therefore, in Fig. 11, the result of RMSEP at a very low number of clusters is still less than the ones at $Q = K$. When $Q$ is not at the two extremes of being very small or very large (almost equal to the number of individual sources), then the result of RMSEP is promising. Moreover, applying more accurate calibration methods or increasing the number of iterations, the ‘solver noise’ will decrease and subsequently, we expect the RMSEP curve to have the same behaviour as the curves of Figs 7 and 11.

6.2 Optimum number of clusters for SNR = 2

We utilize AIC and LRT to select the optimum number of clusters for which clustered calibration achieves its best performance. The methods are applied to our simulation for the case of SNR = 2. The AIC and LRT results are shown in Figs 15 and 16, respectively. They both agree on $Q = 33$ as the optimum number of clusters for the divisive hierarchical and weighted $K$-means clustered calibrations. Likelihood-ratio plot of Fig. 16 has almost the same behaviour as the plot of Frobenius distance between the simulated Jones matrices.
Figure 15. AIC plot for clustered and unclustered calibrations. Both the weighted $K$-means and divisive hierarchical clustered calibrations get their minimum AIC at about 33 clusters. This illustrates that their best performances are obtained at this number of clusters. Also, their AIC results at 33 clusters is lower than the unclustered calibration’s AIC, which shows their better performances compared to the unclustered calibration.

Figure 16. Likelihood-ratio of the gain solutions obtained by clustered and unclustered calibrations. In the both cases of weighted $K$-means and divisive hierarchical clustered calibrations, the minimum likelihood-ratio values belong to approximately 33 clusters. These minimums are also lower than the unclustered calibration’s likelihood-ratio result. Therefore, clustered calibration via both the clustering methods performs better than the unclustered calibration and it achieves the best accuracy in its solutions at 33 clusters.

and the obtained solutions presented in Fig. 11. The reason is that the results of the both plots are obtained using the solutions themselves as the input data. However, since AIC results are computed using the residual errors as inputs, which is also the case for obtaining the RMSEP curves of Fig. 14, AIC curves of Fig. 15 are slightly steeper than the Frobenius distance between the simulated Jones matrices and solutions and the likelihood-ratio curves of Figs 11 and 16, respectively.

6.3 Clustered calibration’s efficiency at different SNRs

We start changing the noise in our simulation to see how it affects the clustered calibration’s efficiency. We simulate the cases for which SNR $\in \{1, 2, \ldots, 15\}$ and apply clustered calibration on them. Since the sky model does not change, the clusters obtained by divisive hierarchical and weighted $K$-means methods for the case of SNR = 2 remain the same. Fig. 17 shows the optimum number of clusters, on which the best performances of clustered calibration are obtained, at different SNRs. For low SNRs, the efficiency of the clustered calibration is superior to the unclustered calibration. As the SNR gets higher, clustered calibrations achieve their best solutions utilizing a higher number of clusters. Finally, when the SNR is high enough, the performance of unclustered calibration becomes better than the clustered one.

Figure 17. The optimum number of clusters, on which the best performance of clustered calibration is obtained, at different SNRs. For low SNRs, the efficiency of the clustered calibration is superior to the unclustered calibration. As the SNR gets higher, clustered calibrations achieve their best solutions utilizing a higher number of clusters. Finally, when the SNR is high enough, the performance of unclustered calibration becomes better than the clustered one.

6.3.1 Empirical estimation of SINR

Having the results of Fig. 17 in hand, we could find an empirical model for estimating the optimum number of clusters for various SNRs. Note that by changing the observation and the instrument characteristics, this model will also be changed.

As it is explained in Section 4.1.3, the best performance of clustered calibration is obtained when the SINR is at its highest level. Fig. 17 shows the number of clusters on which the maximum SINR was obtained, where the signal (sky) and the noise powers are known a priori. Thus, the only unknown for estimating the SINR is the interference, or the ‘clustering errors’, for which we need to have a prediction model. After estimating the SINR using this model, finding the optimum $Q$ will be straightforward.

Consider the definition of ‘clustering errors’ given by equation (31). It is logical that for every source, the difference between its true Jones matrix and the clustered calibration solution, $||J - \tilde{J}||$, is a function of the angular distance between the source and the centroid of the cluster that it belongs to. Based on this and using equations (31) and (34), for the interference of the $i$th cluster at baseline $p - q$ we assume that

$$\sum_{l \in L_i} \Gamma_{l|pq} \sim \mathcal{CN}(0, \eta ||\tilde{C}_{l|pq}||^2 (D(L_i))^\nu L_j).$$

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$$\sum_{l \in L_i} \Gamma_{l|pq} \sim \mathcal{CN}(0, \eta ||\tilde{C}_{l|pq}||^2 (D(L_i))^\nu L_j).$$
where \( \eta \) and \( \nu \) are unknowns. Equation (60), in fact, considers an interference power (variance) of \( \eta ||C_{pq}||^2 (D(L_i))^\nu \) for every \( i \)th cluster, \( i \in \{1, 2, \ldots, Q\} \), at baseline \( p - q \). Assuming the interferences of different clusters to be statistically independent from each other, and bearing in mind that the baseline’s additive noise \( N_{pq} \) has also a complex Gaussian distribution independent from those interferences, then the noise plus interference power for the \( i \)th cluster at baseline \( p - q \) is obtained by

\[
\eta ||C_{pq}||^2 (D(L_i))^\nu + ||N_{pq}||^2.
\] (61)

Fitting suitable \( \eta \) and \( \nu \) to equation (61), the SINR for the \( i \)th cluster at baseline \( p - q \) is equal to the cluster’s signal power, \( ||C_{pq}||^2 \), divided by the result of equation (61). Subsequently, estimation of \( E\{\text{SINR}_c\} \) will be straightforward where the expectation is calculated with respect to all the source clusters and all the baselines. Note that simulation provides us with the true noise power, \( ||N_{pq}||^2 \). In the case of having a real observation, this power could be estimated by equation (46).

Fig. 18 shows the number of clusters on which divisive hierarchical and weighted K-means clustered calibrations achieve their maximum estimated \( E\{\text{SINR}_c\} \). The results are calculated for SNR \( \in \{1, 2, \ldots, 15\} \). For the hierarchical clustering \( \eta = 1550 \) and \( \nu = 0.3 \), and for the K-means clustering \( \eta = 2500 \) and \( \nu = 0.003 \). As we can see, for both the clustering methods, these maximum

\[ E\{\text{SINR}_c\} \]'s are mostly obtained at the true optimum number of clusters for which clustered calibration performed the best. Introducing more refined models compared to equation (60) could even improve the current result.

### 6.4 Realistic sky models

So far, we have limited our studies to sky models in which the brightness and position of the radio sources follow Rayleigh and uniform distributions, respectively. These characteristics provide us with a smooth and uniform variation of flux intensities in our simulated skies. In such a case, the effects of the background noise on the faintest and the strongest signals are almost the same. Therefore, if clustered calibration performs better than the unclustered calibration that would be only based on upgrading the signals against the noise. Although, in nature, we mostly deal with the sky models in which the distribution of the flux intensities is a power law, with a steep slope, and the spatial distribution is Poisson. Hence, there exist a few number of very bright sources, whose signals are considerably stronger than the others, and they are sparse in the field of view. The corruptions of the background noise plus the interferences of the strong signals of those few bright sources make the calibration of the other faint point sources impractical. Thus, there is the need for utilizing the clustered calibration which applies the solutions of the bright sources to their closed by fainter ones or solves for upgraded signals obtained by adding up a group of faint signals together. This has been shown by Kazemi et al. (2011a) and Yatawatta et al. (2012), when comparing the efficiency of the clustered and unclustered calibrations on LOFAR real observations.

In this section, using simulations, we also reveal the superiority of clustered calibration compared to the unclustered calibration for such sky models.

We simulate a sky of 52 radio point sources which are obtained by modified Jelić et al. (2008) foreground model. The brightness distribution of the point sources follows the source count function obtained at 151 MHz (Willett et al. 2001), while the angular clustering of the sources are characterized by a typical two-point correlation function,

\[
\rho(d) = Ad^{-0.8}.
\] (62)

In (62), \( \rho \) is the two-point correlation function, \( d \) is the angular separation and \( A \) is the normalization amplitude of \( \rho \). The flux cut-off is 0.1 Jy.

We corrupt the signals with gain errors which are linear combinations of sin and cos functions, as in our previous simulations. At the end, a zero mean Gaussian thermal noise with a standard deviation of 3 mJy is added to the simulated data. The result is shown in Fig. 19. In Fig. 19, all the bright sources are concentrated on the right-hand side of the image, rather than being uniformly distributed in the field of view, and the rest of the sources are so faint that are almost invisible.

We apply the clustered and unclustered calibrations on \( Q \in \{3, 4, \ldots, 51\} \) number of clusters and \( K = 52 \) number of individual sources, respectively. The clustering method used is the divisive hierarchical and the calibrations are executed via SAGE algorithm with nine number of iterations. The residual noise variances obtained are demonstrated in Fig. 20. As Fig. 20 shows, the level of the residual noise obtained by the clustered calibration for \( Q \in \{15, 16, \ldots, 45\} \) number of clusters is always below the result of the unclustered calibration. This proves the better performance of the clustered calibration. The best result of the clustered calibration,
The noise variances of the residual images, obtained by clustered and unclustered calibrations, in mJy. The level of the residual noise obtained by the clustered calibration for $Q \in \{15, 16, \ldots 45\}$ number of clusters stands below the result of the unclustered calibration. That reveals the superior performance of the clustered calibration in comparison with the unclustered one. The best result of the clustered calibration which achieves the minimum noise level is at $Q = 27$ number of clusters.

with the minimum noise level, is achieved for $Q = 27$ number of clusters.

The residual images of the clustered calibration with $Q = 27$ number of source clusters, and the unclustered calibration for $K = 52$ individual sources are shown in Fig. 21. On the right-hand side of the residual image of the unclustered calibration there exist artificial stripes caused by over and under estimating the brightest sources of the field of view. That shows the problematic performance of the unclustered calibration. However, clustered calibration has generated much less artificial effects after subtracting these sources. On top of that, the zoomed-in window on the left-hand side of the images of Fig. 21 show that the faint sources are not removed by the unclustered calibration at all, while being almost perfectly subtracted by the clustered calibration. Moreover, the residual noise of the clustered calibration follows a symmetric zero mean Gaussian distributions with a standard deviation of 4.2 mJy, while the one from the unclustered calibration has an asymmetric Gaussian distribution with mean and standard deviation equal to $-1.2$ and $5.3$ mJy, respectively. Taking to account that the simulated noise distribution is a zero mean Gaussian distribution with a variance of 3 mJy; the superior performance of the clustered calibration compared to the unclustered one is evident.

As the final conclusion of this simulation, calibrating below the noise level, clustered calibration always performs better than the unclustered calibration. This is regardless of the sky model and is only based on the fact that solving for individual sources with very poor signals is impractical. Nevertheless, when some sources are very close to each other, the sky corruption on their signals would be exactly the same and there is no point in solving for every of them individually.

Figure 21. Simulated observation of 52 sources which are obtained by modified Jelić et al. (2008) foreground model. The corrupting gain errors are generated as linear combinations of sin and cos functions. The image size is $8' \times 8'$ and the additive thermal noise is a zero mean Gaussian noise with a standard deviation of 3 mJy.

Figure 20. The noise variances of the residual images, obtained by clustered and unclustered calibrations, in mJy. The level of the residual noise obtained by the clustered calibration for $Q \in \{15, 16, \ldots 45\}$ number of clusters stands below the result of the unclustered calibration. That reveals the superior performance of the clustered calibration in comparison with the unclustered one. The best result of the clustered calibration which achieves the minimum noise level is at $Q = 27$ number of clusters.

7 CONCLUSIONS

In this paper, we demonstrate the superior performance of ‘clustered calibration’ compared to unclustered calibration especially in calibrating sources that are below the calibration noise level. The superiority is in the sense of having more accurate results by the enhancement of SNR as well as by the improvement of computational efficiency by reducing the number of directions along which calibration has to be performed.

In a ‘clustered calibration’ procedure, sky sources are grouped into some clusters and every cluster is calibrated as a single source. That replaces the coherencies of individual sources by the total coherency of the cluster. Clustered calibration is applied to these new coherencies that carry a higher level of information compared with the individual ones. Thus, for the calibration of sources below the noise level it has a considerably better performance compared to unclustered calibration. Another way of looking at clustering is to consider the distribution of source flux densities, i.e. the number of sources versus the source flux density curve. Regardless of the intrinsic flux density distribution, clustering makes the number of clusters versus the cluster flux density curve more uniform, thus yielding superior performance. An analytical proof of this superiority, for an arbitrary sky model, is presented using MCRLB and SINR analysis.

KLD and LRT are utilized to detect the optimum number of clusters, for which the clustered calibration accomplishes its best performance. A model for estimating SINR of clustered calibration is also presented by which we could find the optimum number of clusters at low computational cost.

Divisive hierarchical as well as weighted $K$-means clustering methods are used to exploit the spatial proximity of the sources. Simulation studies reveal clustered calibration’s improved performance at a low SNR, utilizing these clustering algorithms. Both the clustering methods are hard clustering techniques which divide data into distinct clusters. However, we expect more accurate results using fuzzy (soft) clustering, which constructs overlapping clusters with uncertain boundaries. Application and performance of this type of clustering for clustered calibration will be explored in future work.
Figure 21. The residual images of the clustered calibration for $Q = 27$ number of clusters (right) and the unclustered calibration for $K = 52$ (left). Calibration is implemented by SAGE algorithm with nine number of iterations and the clustering method applied is divisive hierarchical. On the right-hand side of the residual image of the unclustered calibration, we see stripes of over and under estimations due to problematic performance of the unclustered calibration in subtracting the brightest sources. The zoomed in window on the left-hand side of the image also shows that the faint sources are not removed at all. However, clustered calibration could remove all the faint sources almost perfectly and has generated much less artefacts after subtracting for the brightest sources on the right-hand side of the field of view. Moreover, the residual noise of the clustered calibration follows a symmetric zero mean Gaussian distribution with a standard deviation of 4.2 mJy, while the one from the unclustered calibration has an asymmetric Gaussian distribution with mean and standard deviation equal to $-1.2$ and $5.3$ mJy, respectively. Taking to account that the simulated noise distribution is a zero mean Gaussian distribution with a standard deviation of $3$ mJy, the better performance of the clustered calibration compared to the unclustered one is evident.

ACKNOWLEDGMENTS

The first author would like to gratefully acknowledge NWO grant 436040 and to thank N. Monshizadeh for the useful discussions. The authors also thank the referee for a careful review and valuable comments that helped enhance this paper.

REFERENCES

Graves S., 1978, British J. Phil. Sci., 29, 1
Johnson S. C., 1967, Psychometrika, 32, 241

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