Scalable splitting algorithms for big-data interferometric imaging in the SKA era

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ABSTRACT

In the context of next generation radio telescopes, like the Square Kilometre Array, the efficient processing of large-scale datasets is extremely important. Convex optimisation tasks under the compressive sensing framework have recently emerged and provide both enhanced image reconstruction quality and scalability to increasingly larger data sets. We focus herein mainly on scalability and propose two new convex optimisation algorithmic structures able to solve the convex optimisation tasks arising in radio-interferometric imaging. They rely on proximal splitting and forward-backward iterations and can be seen, by analogy with the \textit{clean} major-minor cycle, as running sophisticated \textit{clean}-like iterations in parallel in multiple data, prior, and image spaces. Both methods support any convex regularisation function, in particular the well studied \textit{$\ell_1$} priors promoting image sparsity in an adequate domain. Tailored for big-data, they employ parallel and distributed computations to achieve scalability, in terms of memory and computational requirements. One of them also exploits randomisation, over data blocks at each iteration, offering further flexibility. We present simulation results showing the feasibility of the proposed methods as well as their advantages compared to state-of-the-art algorithmic solvers. Our Matlab code is available online on GitHub.

Key words: techniques: image processing – techniques: interferometric

1 INTRODUCTION

Radio-interferometry (RI) allows the observation of radio emissions with great sensitivity and angular resolution. The technique has been extensively investigated and provides valuable data driving many research directions in astronomy, cosmology or astrophysics (Thompson et al. 2001). Next-generation radio telescopes, such as the L\textsuperscript{O}w Frequency ARray (LOFAR) (van Haarlem et al. 2013) and the future Square Kilometre Array (SKA) (Dewdney et al. 2009), are envisaged to produce giga-pixel images and achieve a dynamic range of six or seven orders of magnitude. This will be an improvement over current instruments by around two orders of magnitude, in terms of both resolution and sensitivity. The amount of data acquired will be massive and the methods solving the inverse problems associated with the image reconstruction need to be fast and to scale well with the number of measurements. Such challenges provided motivation for vigorous research to reformulate imaging and calibration techniques for RI (Wijnholds et al. 2014).

The construction of the first phase of SKA is scheduled to start in 2018. It will consist of two subsystems: a low frequency aperture array, the SKA1-low, operating in the 50-350 MHz frequency range and containing approximately 131,000 antenna elements; a mid frequency array of reflector dishes, the SKA1-mid, operating above 350 MHz, consisting of 197 dishes (Dewdney et al. 2009; Broekema et al. 2015). Both subsystems are planned to operate on the order of 65,000 frequency bands. Data rate estimates in this first phase are around five terabits per second for each subsystem (Broekema et al. 2015) and will present a great challenge for the infrastructure and signal processing. The celebrated \textit{clean} algorithm (Högbom 1974) and its variants do not scale well given the large dimension of the problem. They rely on \textit{local} greedy iterative procedures and are slow.
compared to modern convex optimisation techniques, which are guaranteed to converge towards a global optimal solution. Moreover, they are not designed for large-scale parallelisation or distributed computing (Carrillo et al. 2014).

In the past few years, sparse models and convex optimisation techniques have been applied to RI imaging, showing the potential to outperform state-of-the-art imaging algorithms in the field (Wiaux et al. 2009a; Rau et al. 2009; Li et al. 2011; Carrillo et al. 2012, 2013, 2014; Garsden et al. 2015). These methods typically solve the imaging problem by minimising an objective function defined as a sum of a data term, dependent on the measured visibilities, and several regularisation terms, usually promoting sparsity and positivity. Scalable algorithms, specifically tailored for large-scale problems using parallel and distributed schemes, are just now beginning to gain attention in the context of imaging (Carrillo et al. 2014; Ferrari et al. 2014) and calibration (Yatawatta 2015) for next-generation radio telescopes.

In this context, proximal splitting methods are very popular due to their ability to decompose the original problem into several simpler, easier to solve, sub-problems, each associated with one term of the objective function (Combettes & Pesquet 2011). Another class of algorithms currently gaining traction for large-scale problems in optimisation is based on primal-dual (PD) methods (Komodakis & Pesquet 2015). Such methods efficiently split the optimisation problem and, at the same time, maintain a highly parallelisable structure by solving concomitantly for a dual formulation of the original problem. Building on such tools, the simultaneous direction method of multipliers (SDMM) was recently proposed in the context of RI imaging by Carrillo et al. (2014). It achieves the complete splitting of the functions defining the minimisation task. In the big-data context, SDMM scales well with the number of measurements, however, an expensive matrix inversion is necessary when updating the solution, which limits the suitability of the method for the recovery of very large images.

The scope of this article is to propose two new algorithmic structures for RI imaging. We study their computational performance and parallelisation capabilities by solving the sparsity averaging optimisation problem proposed in the SARA algorithm (Carrillo et al. 2012), previously shown to outperform the standard CLEAN methods. The application of the two algorithms is not limited to the SARA prior, any other convex prior functions being supported. We assume a known model for the measured data such that there is no need for calibration. We use SDMM, solving the same minimisation problem, to compare the computational burden and parallelisation possibilities. Theoretical results ensure convergence, all algorithms reaching the same solution. We also showcase the reconstruction performance of the two algorithms coupled with the SARA prior in comparison with CS-CLEAN (Schwab 1984) and MORESANE (Dabbech et al. 2015).

The first algorithmic solver is a sub-iterative version of the well-known alternating direction method of multipliers (ADMM). The second is based on the PD method and uses forward-backward (FB) iterations, typically alternating between gradient (forward) steps and projection (backward) steps. Such steps can be seen as interlaced CLEAN-like updates. Both algorithms are highly parallelisable and allow for an efficient distributed implementation. ADMM however offers only partial splitting of the objective function leading to a sub-iterative algorithmic structure. The PD method offers the full splitting for both operators and functions. It does not need sub-iterations or any matrix inversion. Additionally, it can attain increased scalability by using randomised updates. It works by selecting only a fraction of the visibilities at each iteration, thus achieving great flexibility in terms of memory requirements and computational load per iteration, at the cost of requiring more iterations to converge. Our simulations suggest no significant increase in the total computation cost.

The remainder of this article is organised as follows. Section 2 introduces the RI imaging problem and describes the state-of-the-art image reconstruction techniques used in radio astronomy. In Section 3 we review some of the main tools from convex optimisation needed for RI imaging. Section 4 formulates the optimisation problem for RI imaging given the large-scale data scenario and presents the proposed algorithms, ADMM and PD, respectively. We discuss implementation details and their computational complexity in Section 5. Numerical experiments evaluating the performance of the algorithms are reported in Section 6. Finally, we briefly present the main contributions and envisaged future research directions in Section 7.

2 RADIO-INTERFEROMETRIC IMAGING

Radio-interferometric data, the visibilities, are produced by an array of antenna pairs that measure radio emissions from a given area of the sky. The projected baseline components, in units of the wavelength of observation, are commonly denoted \((u, v, w)\), where \(w\) identifies the component in the line of sight and \(u = (u, v)\) the components in the orthogonal plane. The sky brightness distribution \(x\) is described in the same coordinate system, with components \(l, m, n\) and with \(l = (l, m)\) and \(n(l) = \sqrt{1 - l^2 - m^2}\). The general measurement equation for non-polarised monochromatic RI imaging can be stated as

\[
y(u) = \int D(l, u) x(l) e^{-2\pi u \cdot l} dl,
\]

with \(D(l, u) = 1/n(l) D(l, u)\) quantifying all the DDEs. Some dominant DDEs can be modelled analytically, like the \(w\) component which is expressed as \(D_w(l, u) = e^{-2\pi u \cdot (n(l) - 1)}\). At high dynamic ranges however, unknown DDEs, related to the primary beam or ionospheric effects, also affect the measurements introducing the need for calibration. Here we work in the absence of DDEs.

The recovery of \(x\) from the visibilities relies on algorithms solving a discretised version of the inverse problem (1). We denote by \(x \in \mathbb{R}^M\) the intensity image of which we take \(M\) visibility measurements \(y \in \mathbb{C}^M\). The measurement model is defined by

\[
y = \Phi x + n,
\]

where the measurement operator \(\Phi \in \mathbb{C}^{M \times N}\) is a linear map from the image domain to the visibility space and \(y\) denotes the vector of measured visibilities corrupted by the additive noise \(n\). Due to limitations in the visibility sampling scheme, equation (2) defines an ill-posed inverse problem. Furthermore, the large number of the data points, \(M \gg N\), introduces additional challenges related to the computational
and memory requirements for finding the solution. In what follows, we assume the operator $\Phi$ to be known is advance such that no calibration step is needed to estimate it.

Due to the highly iterative nature of the reconstruction algorithms, a fast implementation of all operators involved in the image reconstruction is essential, for both regularisation and data terms. To this purpose, the measurement operator is modelled as the product between a matrix $\mathbf{G} \in \mathbb{C}^{M \times n_u}$ and an $n_u$-oversampled Fourier operator,

$$\mathbf{G} = \mathbf{F} \mathbf{Z}.$$  

(3)

The matrix $\mathbf{Z} \in \mathbb{R}^{n_u \times N}$ accounts for the oversampling and the scaling of the image to pre-compensate for possible imperfections in the interpolation (Fessler & Sutton 2003). In the absence of DDEs, $\mathbf{G}$ only contains compact support kernels that enable the computation of the continuous Fourier samples from the discrete Fourier coefficients provided by $\mathbf{F}$. Alternatively, seen as a transform from the $u$–$v$ space to the discrete Fourier space, $\mathbf{G}^\dagger$, the adjoint operator of $\mathbf{G}$, grids the continuous measurements onto a uniformly sampled Fourier space associated with the oversampled discrete Fourier coefficients provided by $\mathbf{F} \mathbf{Z}$. This representation of the measurement operator enables a fast implementation thanks to the use of the fast Fourier transform for generalised compact support in the Fourier domain, which leads to a sparse matrix $\mathbf{G}^\dagger$.

2.1 Classical imaging algorithms

Various methods have been proposed for solving the inverse problem defined by (2). The standard imaging algorithms belong to the CLEAN family and perform a greedy non-linear deconvolution based on local iterative beam removal (Högbom 1974; Schwarz 1978; Schwab 1984; Thompson et al. 2001). A sparsity prior on the solution is implicitly introduced since the method reconstructs the image pixel by pixel. Thus, CLEAN is very similar to the matching pursuit (MP) algorithm (Mallat & Zhang 1993). It may also be seen as a regularised gradient descent method. It minimises the residual norm $\|\mathbf{y} - \mathbf{G} \mathbf{x}\|$ via a gradient descent subject to an implicit sparsity constraint on $\mathbf{x}$. An update of the solution takes the following form

$$\mathbf{x}^{(t)} = \mathbf{x}^{(t-1)} + \mathbf{T} \left( \mathbf{F}^\dagger (\mathbf{y} - \mathbf{G} \mathbf{x}^{(t-1)}) \right),$$  

(4)

where $\mathbf{F}^\dagger$ is the adjoint of the linear operator $\mathbf{F}$. In the astronomy community, the computation of the residual image $\mathbf{F}^\dagger ((\mathbf{y} - \mathbf{G} \mathbf{x}^{(t-1)})$), which represents a gradient step of the residual norm, is being referred to as the major cycle while the deconvolution performed by the operator $\mathbf{T}$ is named the minor cycle. All proposed versions of CLEAN use variations of these major and minor cycles (Rau et al. 2009). CLEAN builds the solution image iteratively by searching for atoms associated with the largest magnitude pixel from the residual image. A loop gain factor controls how aggressive is the update step, by only allowing a fraction of the chosen atoms to be used.

Multiple improvements of CLEAN have been suggested. In the multi-scale version (Cornwell 2008) the sparsity model is augmented through a multi-scale decomposition. An adaptive scale variant was proposed by Bhattachar & Cornwell (2004) and can be seen as MP with over-complete dictionaries since it models the image as a superposition of atoms over a redundant dictionary. Another class of solvers, the maximum entropy method (Ables 1974; Gull & Daniell 1978; Cornwell & Evans 1985) solves a regularised global optimisation problem through a general entropy prior. In practice however, CLEAN and its variants have been preferred even though they are slow and require empirically chosen configuration parameters. Furthermore, these methods also lack scalability required for working with huge, SKA-like data.

2.2 Compressed sensing in radio-interferometry

Imaging algorithms based on convex optimisation and using sparsity-aware models have also been proposed, especially under the theoretical framework of compressed sensing (CS), reporting superior reconstruction quality with respect to CLEAN and its multi-scale versions. CS proposes both the optimisation of the acquisition framework, going beyond the traditional Nyquist sampling paradigm, and the use of non-linear iterative algorithms for signal reconstruction, regularising the ill-posed inverse problem through a low dimensional signal model (Donoho 2006; Candès 2006). The key premise in CS is that the underlying signal has a sparse representation, $\mathbf{x} = \mathbf{W} \alpha$ with $\alpha \in \mathbb{C}^{D}$ containing only a few nonzero elements (Fornasier & Rauhut 2011), in a dictionary $\Psi \in \mathbb{C}^{N \times D}$, e.g. a collection of wavelet bases or, more generally, an over-complete frame.

The first study of CS applied to RI was done by Wiaux et al. (2009a), who demonstrated the versatility of convex optimisation methods and their superiority relative to standard interferometric imaging techniques. A CS approach was developed by Wiaux et al. (2010) to recover the signal induced by cosmic strings in the cosmic microwave background. McEwen & Wiaux (2011) generalised the CS imaging techniques to wide field-of-view observations. Non-coplanar effects and the optimisation of the acquisition process were studied by Wiaux et al. (2009b) and Wolz et al. (2013). All the aforementioned works solve a synthesis-based problem defined by

$$\min_{\alpha} \|\alpha\|_1 \text{ subject to } \|\mathbf{y} - \mathbf{G} \Psi \alpha\|_2 \leq \epsilon,$$  

(5)

where $\epsilon$ is a bound on the $\ell_2$ norm of the noise $\mathbf{n}$. Synthesis-based problems recover the image representation $\alpha$ with the final image obtained from the synthesis relation $\mathbf{x} = \Psi \alpha$. Here, the best model for the sparsity, the non-convex $\ell_0$ norm, is replaced with its closest convex relaxation, the $\ell_1$ norm, to allow the use of efficient convex optimisation solvers. Re-weighting schemes are generally employed to
approximate the ℓ₀ norm from its ℓ₁ relaxation (Candès et al. 2008; Daubechies et al. 2010). Imaging approaches based on unconstrained versions of (5) have also been studied (Wenger et al. 2010; Li et al. 2011; Hardy 2013; Garsden et al. 2015). For example, Garsden et al. (2015) applied a synthesis-based reconstruction method to LOFAR data.

As opposed to synthesis-based problems, analysis-based approaches recover the signal itself, solving
\[
\min_{x} \|\Psi^* x\|_1 \text{ subject to } \|y - \Phi x\|_2 \leq \epsilon. \tag{6}
\]
The sparsity averaging reweighed analysis (SARA), based on the analysis approach and an average sparsity model, was introduced by Carrillo et al. (2012). Carrillo et al. (2014) proposed a scalable algorithm, based on SDMM, to solve (6). For such large-scale problems, the use of sparsity operators \(\Psi\) that allow for a fast implementation is fundamental. Hybrid analysis-by-synthesis greedy approaches have also been proposed by Dabbech et al. (2015).

To provide an analogy between CLEAN and the FB iterations employed herein, we can consider one of the basic approaches, the unconstrained version of the minimisation problem (6), namely \(\min_x \|\Psi x\|_1 + \beta \|y - \Phi x\|_2^2\) with \(\beta\) a free parameter. To solve it, modern approaches using FB iterations perform a gradient step together with a soft-thresholding operation in the given basis \(\Psi^*\) (Combettes & Pesquet 2007b). This FB iterative structure is conceptually extremely close to the major-minor cycle structure of CLEAN. At a given iteration, the forward (gradient) step consists in doing a step in the opposite direction to the gradient of the ℓ₂ norm of the residual. It is essentially equivalent to a major cycle of CLEAN. The backward (soft-thresholding) step consists in decreasing the absolute values of all the coefficients of \(\Psi^* x\) that are above a certain threshold by the thresholding factor, and setting to zero those below the threshold. This step is very similar to the minor cycle of CLEAN, with the soft-thresholding value being an analogous to the loop gain factor. The soft-thresholding intuitively works by removing small and insignificant coefficients, globally, on all signal locations simultaneously while keeping big and important coefficients.

3 CONVEX OPTIMISATION

Optimisation techniques play a central role in solving the large-scale inverse problem (2) from RI. Some of the main methods from convex optimisation (Bauschke & Combettes 2011) are presented in what follows.

Proximal splitting techniques are very attractive due to their flexibility and ability to produce scalable algorithmic structures. Examples of proximal splitting algorithms include the Douglas-Rachford method (Combettes & Pesquet 2007a; Bregman proximal algorithm (Baumgartner 1988)), the iterative thresholding algorithm (Daubechies et al. 2004; Beck & Teboulle 2009), the alternating direction method of multipliers (Boyd et al. 2011) or the simultaneous direction method of multipliers (Setzer et al. 2010). All proximal splitting methods solve optimisation problems like
\[
\min_z g_1(z) + \cdots + g_n(z), \tag{7}
\]
with \(g_i, i \in \{1, \ldots, n\}\), proper, lower-semicontinuous, convex functions. No assumptions are required about the smoothness, each non-differentiable function being incorporated into the minimisation through its proximity operator (C1). Constrained problems are reformulated to fit (7) through the use of the indicator function (C2) of the convex set \(C\) defined by the constraints. As a general framework, proximal splitting methods minimise (7) iteratively by handling each function \(g_i\), possibly non smooth, through its proximity operator. A good review of the main proximal splitting algorithms and some of their applications to signal and image processing is presented by Combettes & Pesquet (2011).

Primal-dual methods (Komodakis & Pesquet 2015) introduce another framework over the proximal splitting approaches and are able to achieve full splitting. All the operators involved, not only the gradient or proximity operators, but also the linear operators, can be used separately. Due to this, no inversion of operators is required, which gives important computational advantages when compared to other splitting schemes (Combettes & Pesquet 2012). The methods solve optimisation tasks of the form
\[
\min_z g_1(z) + g_2(Lz), \tag{8}
\]
with \(g_1\) and \(g_2\) proper, lower semicontinuous convex functions and \(L\) a linear operator. They are easily extended to problems, similar to (7), involving multiple functions. The minimisation (8), usually referred to as the primal problem, accepts a dual problem (Bauschke & Combettes 2011),
\[
\min_v g_1^*(-L^*v) + g_2^*(v), \tag{9}
\]
where \(L^*\) is the adjoint of the linear operator \(L\) and \(g_2^*\) is the Legendre-Fenchel conjugate function of \(g_2\), defined in (C3). Under our assumptions for \(g_1\) and \(g_2\) and, if a solution to (8) exists, efficient algorithms for solving together the primal and dual problems can be devised (Condat 2013; Vilius 2013; Combettes & Pesquet 2012). Such PD approaches are able to produce highly scalable algorithms that are well suited for solving inverse problems similar to (2). They are flexible and offer a broad class of methods ranging from distributed computing to randomised or block coordinate approaches (Pesquet & Repetti 2015; Combettes & Pesquet 2015).

Augmented Lagrangian (AL) methods (Bertsekas 1982) have been traditionally used for solving constrained optimisation problems through an equivalent unconstrained minimisation. In our context, the methods can be applied for finding the solution to a constrained optimisation task equivalent to (8),
\[
\min_z g_1(z) + g_2(r), \text{ subject to } r = Lz, \tag{10}
\]
by the introduction of the slack variable \(r\). The solution is found by searching for a saddle point of the augmented Lagrange function associated with (10),
\[
\max_{s, r} \min_{z, v} g_1(z) + g_2(r) + \frac{s}{\mu} (Lz - r) + \frac{1}{2\mu} \|Lz - r\|^2. \tag{11}
\]
The vector $s$ and parameter $\mu$, correspond to the Lagrange multipliers. No explicit assumption is required on the smoothness of the functions $g_1$ and $g_2$. Several algorithms working in this framework have been proposed. The alternating direction method of multipliers (ADMM) (Boyd et al. 2011; Yang & Zhang 2011) is directly applicable to the minimisation (10). A generalisation of the method, solving (7), is the simultaneous direction method of multipliers (SDMM)(Setzer et al. 2010). It finds the solution to an extended augmented Lagrangian, defined for multiple functions $g_i$. Both methods can also be characterised from the PD perspective (Boyd et al. 2011; Komodakis & Pesquet 2015). Algorithmically, they split the minimisation step by alternating between the minimisation over each of the primal variables $z$ and $r$, followed by a maximisation with respect to the multipliers $s$, performed via a gradient ascent.

4 LARGE-SCALE OPTIMISATION

The next generation telescopes will be able to produce a huge amount of visibility data. To this regard, there is much interest in the development of fast and well performing reconstruction algorithms (Carrillo et al. 2014; McEwen & Wiaux 2011). Highly scalable algorithms, distributed or parallelised, are just beginning to gather traction (Carrillo et al. 2014; McEwen & Wiaux 2011). Highly scalable algorithms, distributed or parallelised, are just beginning to gather traction (Carrillo et al. 2014; McEwen & Wiaux 2011). Highly scalable algorithms, distributed or parallelised, are just beginning to gather traction (Carrillo et al. 2014; McEwen & Wiaux 2011).

4.1 Convex optimisation algorithms for radio-interferometry

Under the CS paradigm, we can redefine the inverse problem as the estimation of the image $x \in \mathbb{R}^N$ given the measurements $y \in \mathbb{C}^M$ under the constraint that the image is sparse in an over-complete dictionary $\Psi$. Since the solution of interests is an intensity image, we also require $x$ to be real and positive. The analysis formulation (6) is more tractable since it generally produces a simpler optimisation problem when over-complete dictionaries are used (Elad et al. 2007). Additionally, the constrained formulation offers an easy way of defining the minimisation given accurate noise estimates.

Thus, we state the reconstruction task as the convex minimisation problem (Carrillo et al. 2013, 2014)

$$\min_x f(x) + l(\Psi^\dagger x) + h(\Phi x)$$

(12)

with the functions involved including all the aforementioned constraints,

$$l = \|\cdot\|_1,$$

$$f = \mathcal{L}, \quad \mathcal{C} = \mathbb{R}^N_+,$$

$$h(z) = 1_{\mathcal{B}}(z), \quad \mathcal{B} = \{z \in \mathbb{C}^M : \|z - y\|_2 \leq \epsilon\}.$$

(13)

The function $f$ introduces the reality and positivity requirement for the recovered solution, $l$ represents the sparsity prior in the given dictionary $\Psi$ and $h$ is the term that ensures data fidelity constraining the residual to be situated in an $l_2$ ball defined by the noise level $\epsilon$.

We set the operator $\Psi \in \mathbb{C}^{N \times n_1 N}$ to be a collection of $n_1$ sparsity inducing bases (Carrillo et al. 2014). The SARA wavelet bases (Carrillo et al. 2012) are a good candidate but problem (12) is not restricted to them. A re-weighted $l_1$ approach (Candès et al. 2008) may also be used by implicitly imposing weights on the operator $\Psi$ but it is not specifically dealt with herein since it does not change the algorithmic structure. This would serve to approximate the $l_0$ pseudo norm, $\|\Psi^\dagger x\|_0$, by iteratively re-solving the same problem as in (12) with refined weights based on the inverse of the solution coefficients from the previous re-weighted problem.

An efficient parallel implementation can be achieved from (2) by splitting of the data into multiple blocks

$$y = \begin{bmatrix} y_1 \\ \vdots \\ y_{n_d} \end{bmatrix}, \quad \Phi = \begin{bmatrix} \Phi_1 \\ \vdots \\ \Phi_{nd} \end{bmatrix}, \quad C = \begin{bmatrix} G_1 M_1 \\ \vdots \\ G_{nd} M_{nd} \end{bmatrix} FZ. \quad (14)$$

Since $G_i \in \mathbb{C}^{M_i \times n_i N_j}$ is composed of compact support kernels, the matrices $M_i \in \mathbb{R}^{n_i N_j \times n_i N}$ can be introduced to select only the parts of the discrete Fourier plane involved in computations for block $j$, masking everything else. The selected, $n_i N_j$, $N_j \leq N$, frequency points are directly linked to the continuous $u-v$ coordinates associated with each of the visibility measurements from block $y_j$. Thus, for a compact grouping of the visibilities in the $u-v$ space, each block only deals with a limited frequency interval. These frequency ranges are not disjoint since a discrete frequency point is generally used for multiple visibilities due to the interpolation kernels and DDEs modelled through the operator $G_i$. Since both have a compact support in frequency domain, without any loss of generality, we consider for each block $j$ an overlap of $n_i$, such points.

We rewrite (2) for each data block as

$$y_j = \Phi_j x + n_j,$$

(15)

with $n_j$ being the noise associated with the measurements $y_j$. Thus, we can redefine the minimisation problem (12) as

$$\min_x f(x) + \sum_{i=1}^{n_1} l_i(\Psi_i^\dagger x) + \sum_{j=1}^{n_d} h_j(\Phi_j x)$$

(16)

where, similarly to (13), we have

$$l_i = \|\cdot\|_1,$$

$$h_j(z) = 1_{\mathcal{B}_j}(z), \quad \mathcal{B}_j = \{z \in \mathbb{C}^{M_i} : \|z - y_j\|_2 \leq \epsilon_j\}.$$

(17)

Here, $\epsilon_j$ represents the bound on the noise for each block. For the sparsity priors, the $l_1$ norm is additively separable and the splitting of the bases used,

$$\Psi = \begin{bmatrix} \Psi_1 & \ldots & \Psi_{n_i} \end{bmatrix},$$

(18)

with $\Psi_i \in \mathbb{C}^{M_i \times n_i}$ for $i \in \{1, \ldots, n_i\}$, is immediate. The new formulation involving the $l_1$ terms remains equivalent to the original one. Note that there are no restrictions on the number of blocks $\Psi$ is split into. However, a different splitting strategy may not allow for the use of fast algorithms for the computation of the operator.

Hereafter we focus on the block minimisation problem defined in (16) and we describe two main algorithmic structures for finding the solution. The first class of methods uses a proximal ADMM and details the preliminary work of Carrillo et al. (2015). The second is based on the PD framework and introduces to RI, a new algorithm able to achieve the full
splitting previously mentioned. These methods have a much lighter computational burden than the SDMM solver previously proposed by Carrillo et al. (2014). They are still able to achieve a similar level of parallelism, either through an efficient implementation in the case of ADMM or, in the case of PD, by making use of the inherent parallelisable structure of the algorithm. The main bottleneck of SDMM, which the proposed algorithms avoid, is the need to compute the solution of a linear system of equations, at each iteration. Such operation can be prohibitively slow for the large RI data sets and makes the method less attractive. The structure of SDMM is presented in Appendix B, Algorithm 3. For its complete description in the RI context we direct the reader to Carrillo et al. (2014), the following presentation being focused on the ADMM and PD algorithms.

4.2 Dual forward-backward based alternating direction method of multipliers

The ADMM is only applicable to the minimisation of a sum of two functions and does not exhibit any intrinsic parallelisation structure. However, by rewriting the minimisation problem from (16) as

\[
\min_x \tilde{f}(x) + h(\Phi x),
\]

an efficient parallel implementation may be achieved. We define the two functions involved in as

\[
\tilde{f}(x) = f(x) + \sum_{i=1}^{m} \ell_i(\psi_i^T x), \quad h(\Phi x) = \sum_{j=1}^{np} \lambda_j(\phi_j(x)).
\]

Furthermore, since \( \tilde{h} \) is a sum of indicator functions \( \lambda_j(\phi_j(x)) \), we can redefine it as \( \tilde{h}(\Phi x) = \lambda_B(\Phi x) \), with \( B = B_1 \times B_2 \times \ldots \times B_{np} \).

ADMM iteratively searches for the solution to an augmented Lagrangian function similar to (11). The computations are performed in a serial fashion and explicit parallelisation may only be introduced inside each of its three algorithmic steps. Thus, at each iteration, ADMM alternates between the minimisation

\[
\min_x \mu \tilde{f}(x) + \frac{1}{2} \| \Phi x + s - r \|_2^2
\]

over the variable of interest \( x \) and the minimisation involving the slack variable \( r \),

\[
\min_r \mu \tilde{h}(r) + \frac{1}{2} \| r - \Phi x - s \|_2^2.
\]

These are followed by a gradient ascent with a step \( \rho \) performed for the Lagrange multiplier variable \( s \). Given the definition of the function \( h(\Phi x) \), the minimisation involving \( r \) can be split into \( np \) independent sub-problems

\[
\min_{r_j} \mu \lambda_j(r_j) + \frac{1}{2} \| r_j - \Phi_j x - s_j \|_2^2, \quad j \in \{1, \ldots, np\}.
\]

This minimisation amounts to computing the proximity operator of \( \mu \lambda_j \) at \( \Phi_j x + s_j \), which, given the definition of the function \( \lambda_j \), reduces to a projection operation. The method imposes that every \( r_j \) approaches \( \Phi_j x \) while \( x \) converges towards the solution. The convergence speed is governed by the Lagrange multiplier \( \mu \) and by the ascent step \( \rho \) associated with the maximisation over the Lagrange multiplier variable \( s \).

A proximal version of ADMM deals with the non-smooth functions from (21) and (23) by approximating the solution via proximal splitting. Algorithm 1 presents the details. In Figure 1 we present a diagram of the algorithm to provide further insight into its parallelisation and distribution capabilities. It can also be used to understand the algorithm from the CLEAN perspective, performing FB CLEAN-like updates in multiple data, prior and image spaces. Data fidelity is enforced through the slack variables \( r_j \), by minimising (23) and thus constraining the residual to belong to the \( \ell_2 \) balls \( B_j \). This accepts a closed form solution and, for each ball \( j \), represents the projection,

\[
P_{B_j}(z) \triangleq \begin{cases} 
\epsilon_j \|z - y_j\|_2 + y_j & \|z - y_j\|_2 > \epsilon_j \\
z & \|z - y_j\|_2 \leq \epsilon_j
\end{cases}
\]

onto the feasible regions defined by it. Given the structure of the function \( \tilde{h} \), this is implemented in parallel with distributed computations and presented in Algorithm 1, step 8, together with the update of the Lagrange variables \( s^{(k)} \), in step 9. The variables \( \tilde{b}^{(k)} \in \mathbb{C}^{n_\psi} \), computed in steps 3 to 6, are required in the computations and need to be transmitted to the different processing nodes. The nodes compute the solution updates \( \hat{q}^{(k)} \in \mathbb{C}^{n_\psi} \) in step 10 after which they are centralised and used to revise the previous solution estimate \( x^{(k-1)} \) and to compute \( x^{(k)} \). Thus, by carefully defining the minimisation problem, a high degree of parallelism is achieved. Note that this step can easily incorporate all types of weighting of the data specific to RI.

For our specific problem, the minimisation over \( x \) from (21) does not accept a closed form solution. We approximate it by using a FB step. The forward step corresponds to a gradient step and the backward step is an implicit sub-gradient-like step performed through the proximity operator. Thus, in step 12, the solution is updated using the descent step \( \rho \) in the direction of the gradient of the smooth part. This is followed by the iterative dual FB (Combettes et al. 2011) updates necessary to approximate the proximity operator to the non smooth \( \tilde{f} \). Algorithm 1, function DUALFB, details the required sub-iterations. In steps 23 and 20, the method alternates between projections onto the convex set \( \mathcal{C} \), which, component wise, are defined as

\[
P_{\mathcal{C}}(z) \triangleq \begin{cases} 
\mathbb{R}(z_k) & \mathbb{R}(z_k) > 0, \\
\mathbb{R}(z_k) & \mathbb{R}(z_k) \leq 0
\end{cases} \quad \forall k,
\]

and the application of the proximity operator to the sparsity prior functions \( \ell_t \), which is the component wise soft-thresholding operator

\[
S_{\alpha}(z_k) \triangleq \begin{cases} 
\frac{z_k \{z_k - \alpha\}_+}{|z_k|} & |z_k| > 0 \\
0 & |z_k| = 0
\end{cases} \quad \forall k,
\]

with threshold \( \alpha \). The soft threshold resulting for the algorithm is \( \frac{1}{\eta} \mu \). However, since \( \mu \) is a free parameter, we reparameterise the operation to use the soft threshold \( \kappa |\psi|_s \), with \( \kappa \) as a new scale-free parameter, independent of the operator \( \Psi \) used. Here, we denote by \( |\Psi|_s \) the operator norm of the sparsifying transform. The operator \( \{\cdot\}_+ \) from (26) sets the negative values to 0. The parameter \( \eta \) serves as an update step for the sub-problem. In step 20, we have additionally used the Moreau decomposition (C4) to replace

MNRAS 000, 1–23 (2016)
Algorithm 1 Dual forward-backward ADMM.
1: given $x^{(0)}, r^{(0)}, s^{(0)}$; $j = 0, \rho, \theta$
2: repeat for $t = 1, \ldots$
3: $b^{(t)} = \mathbb{F} z^{(t-1)}$
4: $\forall j \in \{1, \ldots, n_d\}$ set
5: $b_{j}^{(t)} = M_{j} b_{j}^{(t)}$
6: $\forall j \in \{1, \ldots, n_d\}$ distribute $b_{j}^{(t)}$ and do in parallel
7: $s_{j}^{(t)} = \mathcal{P}_{S_{j}} \left( G_{j} b_{j}^{(t)} + s_{j}^{(t-1)} \right)$
8: $s_{j}^{(t-1)} = s_{j}^{(t-1)} + \varrho \left( G_{j} b_{j}^{(t)} - r_{j}^{(t)} \right)$
9: $q_{j}^{(t)} = G_{j} \left( G_{j} b_{j}^{(t)} + r_{j}^{(t)} - s_{j}^{(t-1)} \right)$
10: end and gather $q_{j}^{(t)}$
11: $z^{(t)} = x^{(t-1)} - \rho z^{(t)} F^{T} \sum_{j=1}^{n_d} M_{j} q_{j}^{(t)}$
12: $x^{(t)} = \text{DUALFB} \left( z^{(t)}, \kappa \right)$
13: until convergence
14: function DUALFB $z, \kappa$
15: given $d^{(0)}$, $\eta$
16: $\tilde{z}^{(0)} = \mathcal{P}_{C} \left( z \right)$
17: $\tilde{z}^{(0)} = \mathcal{P}_{C} \left( z \right)$
18: $\forall i \in \{1, \ldots, n_b\}$ do in parallel
19: $d_{i}^{(k)} = \frac{\rho}{\kappa} \left( I - \mathcal{S}_{i} \right) \left( \eta d_{i}^{(k-1)} + \Psi_{i} z^{(k-1)} \right)$
20: $d_{i}^{(k)} = \Psi_{i} d_{i}^{(k)}$
21: end
22: $z^{(k)} = \mathcal{P}_{C} \left( z - \sum_{i=1}^{n_b} d_{i}^{(k)} \right)$
23: until convergence
24: return $z^{(k)}$

4.3 Primal-dual algorithms with randomisation

The main advantage that makes the PD algorithms attractive for solving inverse problems is their flexibility and scalability. They are able to deal with both differentiable and non-differentiable functions and are applicable to a broad range of minimisation tasks. The inherent parallelisation on the level of splitting the functions gives a direct approach for solving (16). Another important aspect is given by the use of randomisation, allowing the update for a given component function to be performed less often and thus lowering the computational cost per iteration. Block coordinate computations are also supported but are not explicitly used herein.

We define the minimisation task to be solved using PD methods, similarly to (16), as

$$
\min f(x) + \gamma \sum_{i=1}^{n_b} l_{i}(\Psi_{i} x) + \sum_{j=1}^{n_d} h_{j}(\Phi_{j} x),
$$

where $\gamma$ is an additional tuning parameter. Note that the minimisation problem does not change, regardless of the

![Figure 1](image-url). The diagram of the structure of ADMM, detailed in Algorithm 1, showcasing the parallelism capabilities and overall computation flow. The algorithm performs in parallel proximal and gradient updates (similarly to the CLEAN performing major-minor cycle) for all data fidelity terms. Its structure is sub-iterative and enforces sparsity and positivity through the dual FB algorithm. These updates, performed in parallel for each sparsity basis, can be again seen as analogous to CLEAN. Thus, the whole algorithm can be seen as composed of interlaced CLEAN-like proximal splitting and FB updates running in parallel in multiple data, prior, and image spaces.
form all updates on the dual variables flow. In contrast with ADMM, the PD algorithm is able to perform all updates on the dual variables conjugates. A diagram of the structure is presented in Fig-

Algorithm 2 Randomised forward-backward PD.

1: given $x^{(0)}, u^{(0)}, b^{(0)}, e^{(0)}, \nu^{(0)}, \phi^{(0)}, \kappa, \sigma_i, s_j, \lambda$
2: repeat for $t = 1, \ldots$
3: generate sets $\mathcal{P} \subset \{1, \ldots, n_b\}$ and $\mathcal{D} \subset \{1, \ldots, n_d\}$
4: $\tilde{b}^{(t)} = \text{FZ}[z^{(t-1)}]$ 
5: $\forall j \in \mathcal{D}$ set
6: $b_j^{(t)} = M_j \tilde{b}_j^{(t)}$
7: end
8: run simultaneously 
9: $\forall j \in \mathcal{D}$ distribute $b_j^{(t)}$ and do in parallel
10: $\bar{v}_j^{(t)} = (I - \mathcal{P} b_j) \left( v_j^{(t-1)} + G_j b_j^{(t)} \right)$
11: $v_j^{(t)} = v_j^{(t-1)} + \lambda \left( \bar{v}_j^{(t)} - v_j^{(t-1)} \right)$
12: $\tilde{v}_j^{(t)} = G_j v_j^{(t)}$
13: end and gather $\bar{v}_j^{(t)}$
14: $\forall j \in \{1, \ldots, n_d\} \setminus \mathcal{D}$ set
15: $v_j^{(t)} = v_j^{(t-1)}$
16: $\tilde{v}_j^{(t)} = \hat{v}_j^{(t-1)}$
17: end
18: $\forall i \in \mathcal{P}$ do in parallel
19: $\bar{u}_i^{(t)} = (I - \mathcal{S}_{b | \psi}) \left( u_i^{(t-1)} + \psi_i^{(t-1)} \right)$
20: $u_i^{(t)} = u_i^{(t-1)} + \lambda \left( \bar{u}_i^{(t)} - u_i^{(t-1)} \right)$
21: $\hat{u}_i^{(t)} = \psi_i u_i^{(t)}$
22: end
23: $\forall i \in \{1, \ldots, n_b\} \setminus \mathcal{P}$ set
24: $u_i^{(t)} = u_i^{(t-1)}$
25: $\hat{u}_i^{(t)} = \bar{u}_i^{(t-1)}$
26: end
27: end
28: $\tilde{x}^{(t)} = \text{FZ} \left( x^{(t-1)} - \tau Z^T \mathcal{F} \left( \sum_{i=1}^{n_b} \sum_{j=1}^{n_d} \lambda_j \hat{u}_j^{(t)} + \sum_{i=1}^{n_b} \sigma_i \bar{u}_i^{(t)} \right) \right)$
29: $x^{(t)} = x^{(t-1)} + \lambda \left( \tilde{x}^{(t)} - x^{(t-1)} \right)$
30: $\tilde{x}^{(t)} = 2\tilde{x}^{(t)} - x^{(t-1)}$
31: until convergence

value $\gamma$ takes due to the use of the indicator functions in $f$ and $b_j$ which are invariant to scaling. This fits under the framework introduced by Condat (2013); Vău (2013); Pesquet & Repetti (2015) and we devise a PD algorithm towards finding the solution. The method iteratively alternates between solving the primal problem (27) and the dual problem,

$$
\min_{u_i} f_i^f - \sum_{i=1}^{n_b} \Psi_i u_i - \sum_{j=1}^{n_d} \Phi_j v_j
+ \frac{1}{\gamma} \sum_{i=1}^{n_b} l_i^*(u_i) + \sum_{j=1}^{n_d} h_j^*(v_j)
$$

(28)

essentially converging towards a Kuhn-Tucker point. This produces the algorithmic structure of Algorithm 2 where additionally we have used the Moreau decomposition (C4) to rewrite the proximal operations and replace the function conjugates. A diagram of the structure is presented in Figure 2 further exemplifying the conceptual analogy between the PD algorithm and CLEAN. The algorithm allows the full split of the operations and performs all the updates on the dual variables in parallel. The update of the primal variable, the image of interest $x^{(t)}$, requires the contribution of all dual variables $u^{(t)}$ and $v^{(t)}$. The algorithm uses the update steps $\tau$, $\sigma_i$, and $s_j$ to iteratively revise the solution and allows for a relaxation with the factor $\lambda$. FB iterations, consisting of a gradient descent step coupled with a proximal update, are used to update both the primal and the dual variables. These FB updates can be seen as CLEAN-like steps performed in the multiple signal spaces associated with the primal and the dual variables. In the deterministic case, the active sets $\mathcal{P}$ and $\mathcal{D}$ are fixed such that all the dual variables are used. The randomisation capabilities of the algorithm are presented later on, given a probabilistic construction of the active sets.

When applied in conjunction with the functions from (17), the primal update from step 28 is performed through the projection (25) onto the positive orthant defined by $C$. The dual variables are updated in steps 10 and 19 using the proximity operators for $h_j$ and $l_i$, which become the projection onto an $\ell_2$ ball $B_2$ defined by (24) and the component wise soft-thresholding operator (26). We use the Moreau decomposition (C4) to replace the proximity operator of the
conjugate functions \( l^*_i \) and \( h^*_j \) with that of the function \( l_i \) and \( h_j \), respectively. The identity operator is denoted by \( I \).

Step 19 also contains a re-parametrisation similar to the one performed for ADMM. We replace the implicit algorithmic soft-threshold size \( \gamma \), with \( \kappa |\mathbf{u}|_p \) by appropriately choosing the free parameter \( \gamma \). This ensures that we are left with the scale-free parameter \( \kappa \) independent to the operator \( \mathbf{\Psi} \).

Steps 11, 20 and 29 represent the relaxation of the application of the updates. To make use of the parallelisation, the application of the operators \( \mathbf{G}_j \) and \( \mathbf{\Psi} \), is also performed in parallel, in steps 12 and 21. Note that the splitting of the operators is presented in (14), more specifically \( \Phi_j = \mathbf{G}_j \mathbf{M}_j \mathbf{F}_j \), \forall j \in \{1, \ldots, n_d\}. These operations are given in steps 4 to 7.

The computation of the dual variables \( \mathbf{u}^{(t)}_i \) associated with the sparsity priors requires the current solution estimate. This solution estimate is then revised with the updates \( \hat{\mathbf{u}}^{(t)}_i \) computed from the dual variables. Both \( \mathbf{x}^{(t)} \) and \( \hat{\mathbf{u}}^{(t)}_i \) are of size \( N \) and their communication might not be desirable in a loosely distributed system. In such case all computations involving \( \hat{\mathbf{u}}^{(t)}_i \) can be performed in parallel but not in a distributed fashion. The dual variables \( \mathbf{v}^{(t)}_i \), associated with the data fidelity functions, should be computed over a distributed computing network. They only require the communication of the updates \( \mathbf{b}^{(t)} \in \mathbb{C}^{n \times N} \) and dual updates \( \tilde{\mathbf{v}}^{(t)}_j \in \mathbb{C}^{n \times N} \) which remains feasible.

The main challenge associated with the inverse problem defined by (2) is linked with the dimensionality of the data. The large data size is a limiting factor not only from the computational perspective but also from that of memory availability. A randomisation of the computations following the same PD framework (Pesquet & Repetti 2015) is much more flexible at balancing memory and computational requirements. By selectively deciding which data fidelity and sparsity prior functions are active at each iterations, full control over the memory requirements and computational cost per iteration can be achieved. In Algorithm 2, this is controlled by changing the sets \( \mathcal{P} \), containing the active sparsity prior dual variables, and \( \mathcal{D} \), which governs the selection of the data fidelity dual variables. At each iteration, each dual variable has a given probability of being selected, \( p_P \), for the sparsity prior, and \( p_D \), for the data fidelity, respectively. These probabilities are independent of each other. Note that the algorithm has inertia still performing the primal updates using all dual variables even though some dual variables remain unchanged.

5 IMPLEMENTATION DETAILS AND COMPUTATIONAL COMPLEXITY

An efficient implementation of the ADMM and the PD algorithms takes advantage of the data split and of the implicit parallelisation from the definition of the minimisation problem. For presentation simplicity, we consider the processing to be split between a central meta-node, a single processing unit or possibly a collection of nodes, centralising the update on the desired solution \( \mathbf{x}^{(t)} \) and performing the computations associated with the sparsity priors, and a number of data fidelity nodes dealing with the constraints involving the balls \( \mathcal{B}_j \). The computation of the sparsity prior terms can be easily parallelised however, the distribution of the data can be too costly. In this case, a shared memory architecture might be more appropriate than distributed processing. For the data nodes, the communication cost is low and a distributed approach is feasible. We have assumed these two different strategies for dealing with the different terms in the presentation of Algorithms 1 and 2.

Most of the operations to be performed are proportional with \( N \) since the main variable of interest \( \mathbf{x}^{(t)} \) is the image to be recovered. The most demanding operation performed on \( \mathbf{x}^{(t)} \) is the application of the oversampled Fourier operators. When computed with a fast Fourier algorithm (FFT) (Cooley & Tukey 1965), the computational cost of the transforms \( \mathbf{F} \) and \( \mathbf{F}^\dagger \) applied to \( n_o \)-oversampled data scales as \( O(n_o N \log n_o N) \). It should be noted that the FFT implementation can be sped up by using multiple processing cores or nodes. The wavelet operators \( \mathbf{\Psi} \) and \( \mathbf{\Psi}^\dagger \) are applied to the image \( \mathbf{x}^{(t)} \) as well. The Discrete Wavelet Transform (DWT) can be performed with fast wavelet implementations using lifting schemes or filter banks (Cohen et al. 1993; Daubechies & Sweldens 1998; Mallat 2008) and achieves a linear complexity of \( O(N) \) for compactly supported wavelets. A distributed processing of the operations involved in the application of each sparsity basis \( \mathbf{\Psi} \), may be used. However, this requires the communication of the current solution estimate, which might not be feasible. We consider that these computations are performed locally, on the central meta-node.

For the data nodes, a manageable computational load and an efficient communication can be achieved by both algorithms by adopting a balanced and compact split of the data; splitting the data into blocks of similar size having a compact frequency range as proposed in (14). An overlap of size \( n_o \) between discrete frequency ranges is necessary for an efficient interpolation (Fessler & Sutton 2003) to the uniform frequency grid which allows fast Fourier computations or to include DDEs (Wolz et al. 2013). Besides this overlap, each block only deals with a limited frequency range reducing the communication performed. In such case, the matrices \( \mathbf{M}_j \) mask out the frequencies outside the range associated with the blocks \( \mathbf{y}_j \). Furthermore, the use of compact support interpolation kernels and DDEs with compact support in the Fourier domain makes \( \mathbf{G}_j \) sparse, which lowers the computational load significantly. We consider it has a generic sparsity percentage \( n_o \).

Details on the levels of parallelisation and the scaling to multiple nodes for both methods are presented below. As mentioned earlier, the main computational difficulties arise from working with large images and data sets, thus making important the way the complexity of the algorithms scales with \( N \) and \( M \). An overview of the complexity requirements is presented in Table 1.

5.1 Alternating direction method of multipliers

The efficient implementation of ADMM for the problem defined by (19) offloads the data fidelity computations to the data nodes. As can be seen from Figure 1 and Table 1, the basic structure of the algorithm is serial and the processing is just accelerated by parallelising each serial step.

The iterative updates follow the operations presented in Algorithm 1. The central node computes an estimate \( \tilde{\mathbf{z}}^{(t)} \) of the solution and iteratively updates it to enforce sparsity and positivity. The update from step 12 requires
$O(n_oN \log n_oN)$ operations for the computation of the oversampled FFT. Given a compact partitioning of the matrix $\mathbf{G}$, the sum involving the updates $\mathbf{b}_j^{(i)}$ requires computations of the order $O(n_oN) + O(n_by)$. Note that it may be accelerated by using the data node network, however since generally $n_y$ is not large, the gain remains small. The computation of the Fourier coefficients from step 3 also incurs a complexity $O(n_oN \log n_oN).

For the approximation of the proximal operator of the function $f$, the algorithm essentially remains serial and requires a number $n_f$ of iterations. In this case, the complexity of each update performed for the sparsity prior is dominated by the application of the operators $\mathbf{Ψ}$ and $\mathbf{Ψ}^\dagger$, which, given an efficient implementation of the DWT requires $O(N)$ operations. The updates $\mathbf{d}_j^{(k)}$ and $\mathbf{d}_j^{(k)}$ from step 20 and 21 may be computed in parallel. Given a serial processing however this would need $O(n_oN)$ computations. Note that although in this case the complexity scales linearly with $N$, the scaling constants can make the computations to be of the same level as the FFT.

The data fidelity nodes perform steps 8 to 10 in parallel using the Fourier coefficients $\mathbf{b}_j^{(i)}$ precomputed in step 5. The computations are heavier due to the linear operators $\mathbf{G}_j$. As mentioned earlier, the operator has a very sparse structure. This reduces the computation cost for applying $\mathbf{G}_j$ to $O(n_oM_jn_oN_j)$, where $n_oN_j$ is the number of uniformly gridded, frequency points associated with each visibility block $y_j$. The remaining operations only involve vectors of size $M_j$. The overall resulting complexity per node is $O(n_oM_jn_oN_j) + O(M_j)$. Under the assumption that the blocks contain an equal number of visibilities, this further reduces to $O(n_oM_jn_oN_j) + O(M_j)$. The communication required between the central and the data fidelity nodes is of order $n_oN_j$, the size of frequency range of each data block.

5.2 Primal-dual algorithm

An implementation of the PD algorithms benefits from the full split achieved by the methods which allows for the computation of all the dual variables to be completed in parallel.

The processing is performed in two synchronous alternating serial steps to update the primal and dual variables, respectively. Each step is however highly parallelisable. The central node uses the current estimate of the solution $\mathbf{z}^{(t-1)}$ and distributes the oversampled Fourier transform coefficients $\mathbf{b}_j^{(i)}$ to the data fidelity nodes. The data fidelity and central nodes compute simultaneously the dual variables and provide the updates $\mathbf{b}_j^{(i)}$ and $\mathbf{u}_j^{(i)}$ to be centralised and included in the next solution estimate on the central node. Such a strategy requires at each step the propagation of variables of size $n_oN_j$, between the central and data fidelity nodes. As suggested in Algorithms 2, the computation of the sparsity prior dual variables is also highly parallelisable. However, the communication of the current image estimate is required, limiting the possibility to distribute the data due to its large size. We leave the computation to be performed by the central node, without an explicit exploitation of the possible parallelism.

All dual variables can be computed simultaneously as can be seen in Figure 2. The data fidelity nodes need to apply the linear operators $\mathbf{G}_j$ as in steps 10 and 12. Similarly to ADMM, this incurs the heaviest computational burden. Given the very sparse structure of the matrix $\mathbf{G}_j$, this accounts for a complexity of $O(n_oM_jn_oN_j)$ with $n_oN_j$ being the previously mentioned number of, uniformly gridded, frequency points for the visibilities $y_j$. The remaining operations only involve vectors of size $M_j$ and thus the overall resulting complexity is $O(2n_oM_jn_oN_j) + O(2M_j)$. The wavelet decomposition from steps 19 and 21 achieves a linear complexity of $O(N)$ for compactly supported wavelets. The other operations from steps 19 and 20 are of order $O(N)$ resulting in a load for the sparsity prior nodes that scales linearly with $N$.

In step 28 of Algorithm 2, the summing of the sparsity prior updates requires $O(n_oN)$ operations. For the $\ell_2$ data fidelity terms, given a compact partitioning in frequency for the matrix $\mathbf{G}$, the computation requires $O(n_oN) + O(n_oN_y)$ operations. The computational cost of the transforms $\mathbf{F}$ and $\mathbf{F}^\dagger$, steps 4 and 28, scales as $O(n_oN \log n_oN)$ since this requires the FFT computation of the $n_o$-oversampled image. The remaining operations, including the projection,
are $O(N)$, giving the complexity of the primal update step $O(n_b N \log n_b N) + O((n_b + n_a) N) + O(N) + O(n_b n_a)$. We kept the terms separate to give insight on how the algorithms scales for different configurations. Similarly to ADMM, the sums may be performed over the network in a distributed fashion, further reducing the complexity and leaving the primal update step dominated by the Fourier computations.

The randomised primal-dual algorithm introduces an even more scalable implementation. To achieve a low computational burden per data node, the number of nodes has to be very large in order to reduce the size of $M_j$ and $N_j$ for each block. The randomised algorithms achieve greater flexibility by allowing some of the updates for the sparsity prior or data fidelity dual variables, to be skipped at the current iteration. Given a limited computing infrastructure, by carefully choosing the probabilities we can ensure that data fit into memory and that all available nodes are processing parts of it. The average computational burden per iteration is lowered proportionally to the probability of selection, $p_{D_i}$ and $p_{D_j}$. In practice this also produces an increase in the number of iterations needed to achieve convergence, requiring a balanced choice for the probabilities.

5.3 Splitting the data

As reported earlier, the modality in which the data are split can have a big impact in the scalability of the algorithms. Ideally, each data node should process an identical number of visibilities for the computation to be spread evenly. If the visibilities used by one node are however spread over the whole $u$–$v$ plane, their processing requires all the discrete Fourier points. Due to this, a compact grouping in frequency domain is also important since it determines the size of the data to be communicated. Ideally, the splitting should be performed taking into account the computing infrastructure and should balance the communication and computation loads which are linked to the size of $N_j$ and $M_j$.

6 SIMULATIONS AND RESULTS

We study the performance of the algorithms developed herein for different configuration parameters and compare the reconstruction performance against CS-CLEAN (Schwab 1984) and MORESANE (Dabbech et al. 2015). We denote the methods as follows: SDMM, the method introduced by Carrillo et al. (2014); ADMM, the approach described in Algorithm 1; PD and PD-R, the algorithms presented in Algorithm 2 without and with randomisation, respectively; MORESANE, the algorithm$^2$ from Dabbech et al. (2015); CS-CLEAN, the Cotton-Schwab clean (Schwab 1984) algorithm$^3$. For both MORESANE and CS-CLEAN we perform tests for three types of weighting: natural weighting denoted by -N, uniform weighting denoted by -U and Briggs weighting with the robustness parameter set to 1 denoted by -B.

2 We have used the MORESANE implementation from ws-clean (Offringa et al. 2014), https://sourceforge.net/p/wsclean/wiki/Home/.

3 We have used the CS-CLEAN implementation of LWImager from Casacore, https://github.com/casacore/.

Figure 3. The test images, from left to right, top to bottom, a 256 × 256 image of the M31 galaxy, a 512 × 512 galaxy cluster image, a 477 × 1025 image of Cygnus A and a 1024 × 1024 image of the W28 supernova remnant, all shown in log$_{10}$ scale.

The reconstruction performance is assessed in terms of the signal to noise ratio,

$$\text{SNR} = 20 \log_{10} \left( \frac{\|x^o\|_2}{\|x^o - \hat{x}^{(i)}\|_2} \right),$$

(29)

where $x^o$ is the original image and $\hat{x}^{(i)}$ is the reconstructed estimate of the original, averaged over 10 simulations performed for different noise realisations. For the tests involving the comparison with CS-CLEAN and MORESANE on the VLA and SKA coverages we do not perform this averaging. In the latter case we also report the dynamic range

$$\text{DR} = \frac{\sqrt{N} \| \Phi \|_2^2}{\| \Phi (y - \Phi x) \|_2} \max_{k,l} |x_{k,l}|,$$

(30)

obtained by all algorithms.

6.1 Simulation setup

In the first part of the simulations, we evaluate the influence of the different configuration parameters for PD, PD-R, and ADMM. Here, we also validate their performance against SDMM, a previously proposed solver (Carrillo et al. 2014) for the same optimisation task. The test images, as shown in Figure 3, represent a small 256 × 256 image of the Hi region of the M31 galaxy, a 512 × 512 high dynamic range.
range image of a galaxy cluster with faint extended emissions, and a $477 \times 1025$ image of the Cygnus A radio galaxy, respectively. The galaxy cluster image was produced using the \textsc{faraday} tool (Murgia et al. 2004). We reconstruct them from simulated visibility data. We use a $u-v$ coverage generated randomly through Gaussian sampling, with zero mean and variance of 0.25 of the maximum frequency, creating a concentration of visibility data in the centre of the plane, for low frequencies. We introduce holes in the coverage with an inverse Gaussian profile, placing the missing spectrum information predominantly in high frequency. This generates very generic profiles and allows us to study the algorithm performance with a large number of different coverages. A typical $u-v$ coverage is presented in Figure 4.

The second part of the simulations involves testing the algorithm reconstruction using simulated VLA and SKA coverages\textsuperscript{4} corresponding to 5 and 9 hours of observations, respectively. The coverages are presented in Figure 4. For the tests we use an additional large $1024 \times 1024$ image, also presented in Figure 3, representing the W28 supernova remnant\textsuperscript{5}. We showcase the reconstruction quality and speed of convergence for PD and ADMM without performing any re-weighting\textsuperscript{6} and compare the results with those produced by CS-CLEAN and MORESANE.

In both cases, we have normalised the frequencies to the interval $[-\pi, \pi]$. The visibilities are corrupted by zero mean complex Gaussian noise producing a signal to noise level of 20 dB. The bound $\epsilon_j$, for the ball $B_j$ defined by (17), can be therefore estimated based on the noise variance $\sigma_j^2$ of the real and imaginary parts of the noise, the residual norm being distributed according to a $\chi^2$ distribution with $2M_j$ degrees of freedom. Thus, we impose that the square of the global bound $\epsilon_j^2$ is 2 standard deviations above the mean of the $\chi^2$ distribution, $\epsilon_j^2 = (2M_j + 2\sqrt{4M_j}) \sigma_j^2$. The resulting block constraints must satisfy $\sum_{j=1}^{n_d} \epsilon_j^2 = \epsilon^2$. When all blocks have the same size, this results in $\epsilon_j^2 = (2M_j + \frac{2\sqrt{4M_j}}{n_d}) \sigma_j^2$.

We work with pre-calibrated measurements. For simplicity we assume, without loss of generality, the absence of DDEs and a small field of view, the measurement operator reducing to a Fourier matrix sampled at the $M$ frequencies that characterise the visibility points. We have used an oversampled Fourier transform $F$ with $n_o = 4$ and a matrix $G$ that performs an interpolation of the frequency data, linking the visibilities to the uniformly sampled frequency space. The $8 \times 8$ interpolation kernels (Fessler & Sutton 2003) average nearby uniformly distributed frequency values to estimate the value at the frequencies associated with each visibility. A scaling is also introduced in image space to pre-compensate for imperfections in the interpolation. This allows for an efficient implementation of the operator.

To detail the behaviour of the algorithms, we vary the number of blocks $n_d$ used for the data fidelity term. Tests are performed for 4, 16 and 64 blocks. In each case, the blocks are generated such that they have an equal number of visibility points, which cover a compact region in the $u-v$ space. An example of the grouping for the 16 blocks is overlaid on the randomly generated coverage from Figure 4. The figure also contains, marked with dashed lines, an example of the discrete frequency points required to model the visibilities for two of the blocks, under our previous assumptions, for the M31 image. The number of discrete frequency points required for each block would only grow slightly in the presence of DDEs due to their, possible larger, compact support. The overall structure from Figure 4 would remain similar. For the SKA and VLA coverages, the data are also split into blocks of equal size. The resulting block structure is also presented in Figure 4. As sparsity prior, we use the SARA collection of wavelets (Carrillo et al. 2012), namely a concatenation of a Dirac basis with the first eight Daubechies wavelets. We split the collection of bases into $m_b = 9$ individual basis.
6.2 Choice of parameters

The ADMM, PD, and PD-R algorithms converge given that (D1) and (D3), respectively, are satisfied. To ensure this we set for PD $\sigma = 1/\sqrt{w_j^2}$, $\varsigma = 1/\sqrt{\|\Phi x\|^2}$ and $\tilde{r} = 0.49$. The relaxation parameter is set to 1. For the ADMM algorithm we set $\rho = 1/\sqrt{w_j^2}$ and $\eta = 1/\sqrt{\|\Phi x\|^2}$. The ascent step is set $q = 0.9$. The maximum number of sub-iterations is set to $n_j = 100$. We consider the convergence achieved, using a criterion similar to (31), when the relative solution variation for $\tilde{x}^{(k)}$ is below $10^{-3}$. The norms of the operators are computed a priori using the power iterative method. They act as a normalisation of the updates, enabling the algorithm to deal with different data or image scales.

We leave the normalised soft-threshold values $\kappa$ as a configuration parameter for both PD and ADMM. SDMM has a similar parameter $\kappa$. It influences the convergence speed which is of interest since, given the scale of the problem, we want to minimise the computational burden which is inherently linked to the number of iterations performed. We aim at providing a general prescription for this tuning parameter, similarly to the standard choices for the loop gain factor used by CLEAN. Intuitively, this soft-thresholding parameter can be seen as analogous to this factor, deciding how aggressive we are in enforcing the sparsity requirements. The stopping parameter $\delta$, essentially linked to the accuracy of the solution given a certain convergence speed, is also configurable. For simplicity we also set equal probabilities for PD-R, namely $p_{\rho v} = p_{\rho}$, $\forall v$ and $p_{\rho v_j} = p_{\rho}$, $\forall j$ and we show how the different choices affect the performance. We choose to randomise only over the data fidelity terms since the SARA sparsity prior is light from the computational perspective when compared to the data fidelity term, thus $p_{\rho} = 1$ for all tests performed. Different strategies for the choice of probabilities, with values different for each block, are also possible. For example setting a higher probability for the blocks containing low frequency data will recover faster a coarse image. The details are incorporated into the solution through the lower probability updates of the high frequency data. An overview of all the parameters used for defining the optimisation task and for configuring both ADMM and PD algorithms is presented in Appendix A, Table A1 and Table A2, respectively.

We ran MORESANE with a 5 major loops and a major loop gain 0.9. The loop gain inside MORESANE was set to 0.1. We use the model image to compare against the other methods. CS-CLEAN was run with two loop gain factors, $l_\rho = 0.1$ and $l_\rho = 0.001$. The results shown are the best of the two. We compare against the model image convolved with a Gaussian kernel associated with the main beam. We scale the resulting image to be closest to the true model image in the least square sense. Additionally, we also present results with the main beam scaled by a factor 6 chosen such that the best SNR is achieved. This introduces a large advantage for CS-CLEAN when compared to the other algorithms. To avoid edge artefacts, both MORESANE and CS-CLEAN were configured to produce a padded double sized image and only the centre was used for comparison.

For PD, PD-R, and ADMM, the stopping criterion for the algorithms is composed of two criteria. We consider the constraints satisfied when the global residual norm is in the vicinity of the bound $\epsilon$ of the global $\ell_2$ ball, namely below a threshold $\epsilon$. This is equivalent to stopping if $\sum_{j=1}^{n_j} \|y_j - \Phi x^{(t)}\| \leq \epsilon^2$. We set $\epsilon^2 = (2M + 3\sqrt{10M}) \sigma_n^2$, namely 3 standard deviations above the mean. The second criterion relates to the relative variation of the solution, measured by

$$
\delta = \frac{\|x^{(t)} - x^{(t-1)}\|_2}{\|x^{(t)}\|_2}.
$$

Figure 5. The evolution of the SNR for PD, ADMM and SDMM as a function of the number of iterations for the M31 test image. The configuration parameter, $\kappa = 10^{-3}$, is the same for ADMM, PD and SDMM. The number of visibilities $M$ used is 10N, 5N and 2N. The input data are split into 4 blocks.

The iterations stop when the $\ell_2$ ball constraints are satisfied and when the relative change in the solution norm is small, $\delta \leq \delta$. The data fidelity requirements are explicitly enforced, ensuring that we are inside or very close to the feasible region. However, this does not guarantee the minimisation of the $\ell_1$ prior function. The algorithms should run until the relative variation of the solution is small between iterations.

To better understand the behaviour of the algorithms, for most simulations we perform tests over a fixed number of iterations without applying the stopping conditions above.

The stopping criterion for MORESANE and CS-CLEAN was set to be 3 standard deviations above the noise mean. This level was seldom reached by CS-CLEAN after the deconvolution, the algorithm seeming to stop because of the accumulation of false detections leading to the increase of the residual between iterations.

6.3 Results using random coverages

We begin by analysing the evolution of the SNR for the ADMM and PD algorithms in comparison with that produced by the previously proposed SDMM solver. Figure 5 contains the SNR as a function of number of iterations for the three algorithms for the reconstruction of the M31 image from $M = 10N$, $M = 5N$ and $M = 2N$ visibilities. The two newly introduced algorithms have the same convergence rate as SDMM but have a much lower computational burden per iteration, especially the PD method. In these tests, all three method use the parameter $\kappa = 10^{-3}$, suggested also by Carrillo et al. (2014). The reconstruction performance is comparable for the different test cases, the PD and ADMM obtaining the same reconstruction quality. Adding more data improves the reconstruction SNR by 2-3 dB because the noise is better averaged. However, note that the SNR gain stagnates slightly when more visibility data are added mainly because the holes in the frequency plane are still not covered. The problem remains very ill-posed with similar coverage. In a realistic situation, adding
Figure 6. The reconstruction of the M31 image from $M = 10N$ visibilities. The input data are split into 4 blocks. (top) The evolution of the SNR for PD and ADMM as a function of the number of iterations for different values of the parameter $\kappa$. (bottom) The value of $\delta$ for both methods.

Figure 7. The reconstruction of the Cygnus A image from $M = N$ visibilities. The input data are split into 4 blocks. (top) The evolution of the SNR for PD and ADMM as a function of the number of iterations for different values of the parameter $\kappa$. (bottom) The value of $\delta$ for both methods.

Figure 8. The reconstruction of the galaxy cluster image from $M = 2N$ visibilities. The input data are split into 4 blocks. (top) The evolution of the SNR for PD and ADMM as a function of the number of iterations for different values of the parameter $\kappa$. (bottom) The value of $\delta$ for both methods.

more data will also fill the coverage more and the SNR improvement will be larger. Since all three algorithms explicitly solve the same minimisation problem, they should have similar behaviour for any other test case.

We continue by investigating the performance of the PD and ADMM algorithms as a function of the parameter $\kappa$ in Figures 6, 7 and 8 for the reconstruction of the M31, Cygnus A and galaxy cluster test images, respectively. The parameter $\kappa$ serves as a normalised threshold and essentially governs the convergence speed. The values $\kappa = 10^{-3}$ to $\kappa = 10^{-5}$ generally produce good and consistent performance. This behaviour was also observed for similar tests, with smaller $M$. Larger values for $\kappa$ reduce the convergence speed since they emphasise greatly the sparsity prior information at the expense of the data fidelity. The smaller values place less weight on the sparsity prior and, after an initial fast convergence due to the data fidelity term, typically require more iterations to minimise the $\ell_1$ prior. The average variation of the solution norm $\delta$ is also reported since the stopping criterion is based on it. It links the convergence speed with the recovery performance. For the galaxy cluster, the tests exhibit slower convergence speed when compared to the M31 and Cygnus A tests. The values $\kappa = 10^{-3}$ and $\kappa = 10^{-5}$ produce similar behaviour. It should be also noted that the variation of the solution decreases smoothly until convergence and that ADMM shows a larger variability.

The convergence speed of the randomised algorithm, PD-R, is studied in Figures 9, 10 and 11 for the M31, Cygnus A and galaxy cluster test images, with three choices for the data splitting. As expected, the convergence speed decrease when the probability of update $p_D$ is lowered. The number of iterations required for convergence increases greatly for probabilities below 0.25. Similar behaviour is achieved for the reconstruction of the test images from a smaller number of measurements. Again, the convergence speed for the galaxy cluster test image is slower. There is also a very small decrease in the convergence speed for all tests when the data are split into a larger number of blocks. This is due to the fact that, in order to reach the same global $\epsilon$, the resulting bounds imposed per block are more constraining and due to the fact that achieving a consensus between a larger number of blocks is more difficult.

Generally, the convergence speed decreases gradually as the probability $p_D$ gets lower, PD-R remaining competitive and able to achieve good complexity as can be seen in Figure 12. Here, we exemplify the performance in more
Figure 9. The SNR for the reconstruction of the M31 image from $M = 10N$ visibilities for the PD and PD-R algorithms with parameter $\kappa = 10^{-3}$. The algorithms split the input data into: (top) 4 blocks, (middle) 16 blocks, (bottom) 64 blocks.

detail when using the 64 blocks with parameter $\kappa = 10^{-3}$, the stopping threshold $\delta = 10^{-4}$ and the $\ell_2$ ball stopping threshold $\ell^2 = (2M + 3\sqrt{M}) \sigma^2$. Our tests show that the total number of iterations performed is roughly inversely proportional to the probability $p_D$. Additionally, we provide a basic estimate of the overall global complexity given the data from Table 1 and the number of iterations required. We only take into account the computationally heaviest operations, the FFT and the operations involving the data fidelity terms. The computations involving the sparsity priors are performed in parallel with the data fidelity computations and are much lighter. Since the analysis is made up to a scaling factor, for better consistency, we normalised the complexity of PD-R with respect to that of the PD.

The total complexity of PD-R remains similar to that of the non-randomised PD which makes PD-R extremely attractive. Generally, if the main computational bottleneck is due to the data term and not to the FFT computations it is expected that the total complexity of PD-R will remain comparable to that of the non-randomised PD. This is of great importance since, for a very large number of visibilities when the data does not fit in memory on the processing nodes, PD-R may be the only feasible alternative. When a more accurate stopping criterion is used, either with a smaller $\epsilon_3$ or relative variation of the solution $\delta$, the randomised algorithms start to require increasingly more iterations to converge and their relative complexity grows. Randomisation over the sparsity bases is also possible but, due to the low computational burden of the priors we use, it is not of interest herein. However, randomisation over the prior functions can become an important feature when computationally heavier priors are used or when the images to be reconstructed are very large.

6.4 Results with the VLA and SKA coverages

In Figure 13 we present the SNR evolution as a function of the number of iterations for the PD and ADMM algorithms for the reconstruction of the Cygnus A and galaxy cluster images using the VLA coverage, and of the W28 supernova remnant test image using the SKA coverage. The visibilities are split into 64 equal size blocks and the parameter $\kappa = 10^{-5}$. We also overlay on the figures the SNR achieved using CS-CLEAN and MORESANE with the different types of weighting.

The dirty images produced using natural weighting for the same tests are presented in Figure 14. For all three test cases, we showcase the reconstructed images, the reconstruction error images and the dirty residual images in Figures 15, 16, and 17. We present the naturally weighted residual images for all methods even when they perform the deconvolution using a different weighting. Since any other type of weighting essentially biases the data and decreases the sensitivity of the reconstruction, this is the more natural choice of visualising the remaining information in the residual image. Although both CS-CLEAN and MORESANE generally
achieve better reconstruction for other weighting types, we present the naturally weighted dirty residual since it represents an unbiased estimation of the remaining structures.

For the reconstruction of the Cygnus A and galaxy cluster images, the methods developed herein outperform MORESANE, using the best performing type of weighting, by approximately 5 dB. Comparing against CS-CLEAN with the best weighting and beam size $b$, the SNR is around 10 dB in favour of the reconstruction performed by the PD and ADMM methods. Visually, both CS-CLEAN and MORESANE fail to recover properly the jet present in the Cygnus A image while for PD and ADMM it is clearly visible. It should be noted that the residual images show also very little structure for PD and ADMM while CS-CLEAN and MORESANE still allow for a more structured residual image. This is partially due to the biasing of the data when the uniform and Briggs weighting is performed. PD and ADMM also achieve a better reconstruction of the galaxy cluster image. They are able to better estimate the three bright sources in the centre of the image. They are however slower to converge if compared to the recovery of the Cygnus A image. MORESANE-N also performs well for this test image and is able to produce a relatively smoother residual image in comparison to the Cygnus A case. Note also that the performance of both CS-CLEAN and MORESANE is inconsistent and varies greatly with the weighting type.

The last test is performed for the reconstruction of the W28 supernova remnant image using the SKA coverage. In this case, the coverage is dominated by the low frequency points and lowers the convergence speed of both PD and ADMM algorithms. Both PD and ADMM achieve good SNR, again around 5 dB over that reached by MORESANE. CS-CLEAN is 2 dB worse than MORESANE and is only able to recover the brightest sources as can be seen in Figure 17. Again, both of our methods are able to recover more of the faint regions surrounding the bright sources. The dirty residual images show less structure for the methods developed herein since they work directly with the naturally weighted visibilities. Note that in Figure 17, in order to achieve a better visualisation, the scale of the dirty residual images for CS-CLEAN is different than that of the other methods. Also, the performance of both CS-CLEAN and MORESANE is again very inconsistent and varies greatly with the weighting types.

Both PD and ADMM methods show decreased convergence speed for the recovery of the galaxy cluster and W28 supernova remnant images. A future study should, possibly by using generalised proximity operators (Pesquet & Repetti 2015), address the acceleration of the convergence which is influenced by the relative distribution of the visibilities in frequency. Coverages dominated by low frequency points, like the SKA one, generally produce slower convergence speed. Furthermore, if a faster convergence is achieved, a reweighing $\ell_1$ approach becomes more attractive and should increase the reconstruction quality significantly.

7 CONCLUSIONS

We proposed two algorithmic frameworks based on ADMM and PD approaches for solving the RI imaging problem. Both methods are highly parallelisable and allow for an efficient distributed implementation which is fundamental.
in the context of the high dimensionality problems associated with the future SKA radio telescope. The structure of ADMM is sub-iterative, which for much heavier priors than the ones used herein may become a bottleneck. The PD algorithm achieves greater flexibility, in terms of memory requirements and computational burden per iteration, by using full splitting and randomised updates. Through the analogy between the CLEAN major-minor loop and a FB iteration, both methods can be understood as being composed of sophisticated CLEAN-like iterations running in parallel in multiple data, prior, and image spaces.

The reconstruction quality for both ADMM and PD methods is similar to that of SDMM. The computational burden is much lower. Experimental results with realistic coverages show impressive performance in terms of parallelisation and distribution, suggesting scalability to extremely large data sets. We give insight into the performance as a function of the configuration parameters and provide a parameter setup, with the normalised soft-thresholding values between $10^{-3}$ and $10^{-5}$, that produce consistently stable results for a broad range of tests. The solution to the optimisation problem solved herein was shown to greatly outperform the standard methods in RL which further motivates the use of our methods. Our tests also confirm the reconstruction quality in the high dynamic range regime.

Our Matlab code is available online on GitHub, http://baasp-group.github.io/pd-and-admm-for-ri/. In the near future, we intend to provide an efficient implementation, using the MPI communication library, for a distributed computing infrastructure. This will be included in the PURYF C++ package, which currently only implements a sequential version of SDMM. The acceleration of the algorithms for coverages dominated by low frequency points will also be investigated, by leveraging a generalised proximal operator. Additionally, recent results suggest that the conditions for convergence for the randomised PD can be relaxed, which would accelerate the convergence speed making these methods to be even more competitive. We also envisage to use the same type of framework to image in the presence of DDEs, such as the $w$ component, as well as to jointly solve the calibration and image reconstruction problems.

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Figure 15. (top 4 images) Log scale reconstructed images; (middle 4 images) log scale of the absolute value of the estimation errors; (bottom 4 images) log scale of the absolute value of the naturally weighted residual images, for the 477 × 1025 Cygnus A test image using the VLA coverage. For each group, the algorithms are: (top left) PD with the reconstruction SNR = 30.51 dB and the corresponding DR = 108620; (top right) ADMM with the reconstruction SNR = 30.52 dB and DR = 107050; (bottom left) CS-CLEAN-N with $l_g = 0.001$ and $b = 0.53$ with the reconstruction SNR = 19.95 dB and DR = 10773; (bottom right) MORESANE-U with the reconstruction SNR = 25.82 dB and DR = 11661. The images correspond to the best results obtained by all algorithms as presented in Figure 13.
Figure 16. (left to right) The reconstructed images, absolute value of the estimation errors, and absolute value of the naturally weighted residual images, all in log scale, for the $512 \times 512$ galaxy cluster test image using the VLA coverage. The algorithms are: (from top to bottom) PD having the reconstruction SNR = 30.98 dB and the corresponding DR = 475300; ADMM having the reconstruction SNR = 31.08 dB and DR = 432070; CS-CLEAN-N with $l_p = 0.001$ and $b = 0.32$ having the reconstruction SNR = 18.03 dB and DR = 21884; MORESANE-N having the reconstruction SNR = 24.96 dB and DR = 351850. The images correspond to the best results obtained by all algorithms as presented in Figure 13.
Figure 17. (left to right) The reconstructed images, absolute value of the estimation errors, and absolute value of the naturally weighted residual images, all in log scale, for the $1024 \times 1024$ W28 test image using the SKA coverage. The algorithms are: (from top to bottom) PD having the reconstruction SNR = 21.86 dB and the corresponding DR = 737720; ADMM having the reconstruction SNR = 21.99 dB and DR = 735620; CS-CLEAN-U with $l_g = 0.1$ and $b = 1.1$ having the reconstruction SNR = 14.14 dB and DR = 515; MORESANE-B having the reconstruction SNR = 15.89 dB and DR = 10990. The images correspond to the best results obtained by all algorithms as presented in Figure 13. Note that the scale for the residual image of CS-CLEAN-U is in the same range as the dirty image presented in Figure 14 while for PD, ADMM and MORESANE the scale of the residual image is below that.
APPENDIX A: PARAMETER OVERVIEW

An overview of the parameters used to define the minimisation problems is presented in Table A1. The configuration parameters for the algorithms are presented in Table A2.

APPENDIX B: SDMM ALGORITHM

The structure of SDMM, solving the specific RI problem (16), is presented for completeness in Algorithm 3.

APPENDIX C: CONVEX OPTIMISATION TOOLS

Definition 1. The proximity operator (Moreau 1965) applied to any lower-semicontinuous and proper convex function $g$ is defined as

$$\pro{x}{g}(z) \triangleq \arg\min_{x} g(x) + \frac{1}{2} \|x - z\|^2_2.$$  \hfill (C1)

Definition 2. The indicator function $\mathcal{I}$ of any set $\mathcal{C}$ is defined as

$$\mathcal{I}(z) \triangleq \begin{cases} 0 & z \in \mathcal{C} \\ +\infty & z \notin \mathcal{C}. \end{cases}$$ \hfill (C2)

In convex optimisation, it allows the use of an equivalent formulation for constrained problems by replacing the explicit convex constraints with the indicator function of the convex
The Legendre-Fenchel conjugate function solvers.

\[ \Psi(x) \] the \( n_b \) wavelet bases in which the signal is considered sparse; other priors can be incorporated as well by redefining the functions \( l_i \) and their associated proximity operators

\( n_b \) the number of data blocks generally linked to the computing infrastructure

\( B_j \) the \( \ell_2 \) balls imposing data fidelity; they are linked to the modality in which the data are split into blocks \( y_j \)

\( \epsilon_j \) the size of the \( \ell_2 \) balls defining the data fidelity; they are linked to the statistics of the noise; herein \( \epsilon_j \) are set based the \( \chi^2 \) distribution associated with the noise.

Table A1. Overview of the parameters for defining the optimisation problem (12).

| Parameter | Description |
|-----------|-------------|
| \( \Psi \) | Wavelet bases |
| \( n_b \) | Number of data blocks |
| \( B_j \) | \( \ell_2 \) balls |
| \( \epsilon_j \) | \( \ell_2 \) ball sizes |

Algorithm 1 (ADMM)

| Parameter | Description |
|-----------|-------------|
| \( \kappa > 0 \) | Configurable; influences the convergence speed |
| \( \delta \leq 10^{-3} \) | Configurable; stopping criteria; linked to the accuracy of the desired solution |
| \( \epsilon_j \) | Stop-iteration stopping criteria; linked to the accuracy of the desired solution |
| \( \rho_g > 0 \) | Configurable; randomisation probabilities; linked to the computing infrastructure |
| \( \tau = 0.49 \) | Configurable; stopping criteria; need to satisfy (D3) |
| \( \epsilon \) | Fixed; algorithm convergence parameter |

Algorithm 2 (PD)

| Parameter | Description |
|-----------|-------------|
| \( \kappa > 0 \) | Configurable; influences the convergence speed |
| \( \delta \leq 10^{-3} \) | Configurable; stopping criteria; linked to the accuracy of the desired solution |
| \( \rho_g > 0 \) | Configurable; randomisation probabilities; linked to the computing infrastructure |
| \( \tau = 0.49 \) | Fixed; algorithm convergence parameter |

Table A2. The configuration parameters for the ADMM (top) and PD (bottom) algorithms.

Property 1. (Moreau decomposition) The Moreau decomposition links the proximity operator of a lower-semicontinuous and proper convex function \( g \) to that of its Legendre-Fenchel conjugate \( g^* \) as

\[
(\forall v) \quad g^*(v) = \sup_z z^Tv - g(z).
\] (C3)

Definition 3. The Legendre-Fenchel conjugate function \( g^* \) of a function \( g \) is

\[ g^*(v) = \sup_z z^Tv - g(z). \] (C3)

APPENDIX D: ALGORITHM CONVERGENCE

D1 Alternating Direction Method of Multipliers

The convergence of Algorithm 1 is achieved through a careful choice of the parameters \( \rho \) and \( \eta \). The algorithm converges for any choice of the Lagrange parameter \( \mu \) satisfying \( \mu > 0 \). This imposes the same constraint on \( \kappa \). For the convergence of the dual FB sub-iterations, the update parameter \( \eta \) should satisfy \( 0 < \eta < 2/\|\Psi\|_S^2 \).

Assuming that the measurement operator \( \Phi \) is full column rank and that convergence has been reached with the dual FB sub-iterations, the convergence for the whole algorithm is achieved in terms of both objective function \( f(x) + h(\Phi x) \) and iterates \( x^{(l)} \), \( r_i^{(l)} \) and, \( s_i^{(l)} \) (Komodakis & Pesquet 2015; Boyd et al. 2011). It requires that

\[
\rho \|\Phi\|_S^2 + \beta < 2,
\] (D1)

with \( \|\Phi\|_S \) being the spectral norm of the measurement operator and the parameters \( \rho \) and \( \beta \) being the update step used for the proximal splitting and the gradient ascent step, respectively.

In practice however, the RI imaging problem is very ill-conditioned and the operator \( \Phi \) is typically not full rank. Under these relaxed conditions, the convergence is guaran-
mented only with respect to the objective function and the multipliers $s_j^{(t)}$, without any guarantees for the iterates $x^{(t)}$ and $r^{(t)}$ (Boyd et al. 2011). A possible way to improve this is to replace $\bar{h}$ with an augmented function $\tilde{h}$,

$$\tilde{h} \left( \begin{bmatrix} \Phi \\ \Gamma \end{bmatrix} x \right) = \bar{h}(\Phi x) + 0(\Gamma x),$$  \tag{D2}

where $0$ represents the null function, zero for any $x$. Such a trick (Pesquet et al. 2012) replaces the measurement operator $\Phi$ with the augmented operator representing the concatenation of both $\Phi$ and $\Gamma$. The new resulting operator is full rank for a proper choice of the matrix $\Gamma$. In practice Algorithm 1 produces reliable performance and we did not employ such a trick herein.

### D2 Primal-Dual Algorithm

The variables $x^{(t)}$, $v_i^{(t)}$, and $u_j^{(t)}$, $\forall i, j$, are guaranteed to converge to the solution of the PD problem (27)-(28) for a proper set of configuration parameters. The convergence, defined given two general preconditioning matrices $U$ and $W$, requires (Pesquet & Repetti 2015, Lemma 4.3) that

$$\|U^{1/2}LW^{1/2}\|_2 < 1, \tag{D3}$$

with the linear operator $L$ being a concatenation of all the used operators, in our case a concatenation of both $\Psi^\dagger$ and $\Phi$. By choosing diagonal preconditioning matrices, with the config parameters $\tau, \sigma_i = \sigma, \varsigma_i = \varsigma, \forall i, j$, on the adequate diagonal locations, the conditions from (D3) can be restated explicitly for Algorithm 2 as

$$\left\| \begin{bmatrix} \sigma I & 0 \\ 0 & \varsigma I \end{bmatrix}^{1/2} \begin{bmatrix} \Psi^\dagger \\ \Phi \end{bmatrix} \begin{bmatrix} \tau I \end{bmatrix}^{1/2} \right\|_2^2 \leq \tau \sigma \|\Psi^\dagger\|_2^2 + \tau \varsigma \|\Phi\|_2^2 < 1, \tag{D4}$$

with the use of the triangle and Cauchy-Schwarz inequalities and with the diagonal matrices $I$ of a proper dimension. It should be noted that this formulation does not limit the use to only two parameters $\sigma$ and $\varsigma$. However, having more independent update steps scales poorly due to the increasing difference between the resulting bound, computed similarly to (D4), and the requirements (D3). This translates to having increasingly small values for the update steps, the more independent parameters we employ, with the convergence speed slowing down considerably in such situation. It is also required that the relaxation parameter is chosen such that $0 < \lambda \leq 1$. The additional parameter $\gamma > 0$ imposes that $\kappa > 0$ as well.

For the randomised setup, the same parameters satisfying (D3) suffice, granted that the probabilities of update $p_P$ and $p_D$ are nonzero and the activated variables are drawn in an independent and identical manner along the iterations.

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