

Citation for published version: Gazzola, S, Meng, C & Nagy, JG 2020, 'Krylov Methods for Low-Rank Regularization', SIAM Journal On Matrix Analysis and Applications (SIMAX).

Publication date: 2020

Document Version Peer reviewed version

Link to publication

Publisher Rights CC BY-NC (C) 2020 SIAM

University of Bath

General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

KRYLOV METHODS FOR LOW-RANK REGULARIZATION *

1 2

SILVIA GAZZOLA[†], CHANG MENG[‡], AND JAMES G. NAGY[‡]

3 Abstract. This paper introduces new solvers for the computation of low-rank approximate solutions to large-4 scale linear problems, with a particular focus on the regularization of linear inverse problems. Although Krylov methods incorporating explicit projections onto low-rank subspaces are already used for well-posed systems that arise 5 from discretizing stochastic or time-dependent PDEs, we are mainly concerned with algorithms that solve the so-called 6 nuclear norm regularized problem, where a suitable nuclear norm penalization on the solution is imposed alongside a fit-to-data term expressed in the 2-norm: this has the effect of implicitly enforcing low-rank solutions. By adopting 8 an iteratively reweighted norm approach, the nuclear norm regularized problem is reformulated as a sequence of 9 quadratic problems, which can then be efficiently solved using Krylov methods, giving rise to an inner-outer iteration 10 11 scheme. Our approach differs from the other solvers available in the literature in that: (a) Kronecker product 12properties are exploited to define the reweighted 2-norm penalization terms; (b) efficient preconditioned Krylov 13methods replace gradient (projection) methods; (c) the regularization parameter can be efficiently and adaptively set along the iterations. Furthermore, we reformulate within the framework of flexible Krylov methods both the 14 new inner-outer methods for nuclear norm regularization and some of the existing Krylov methods incorporating 15low-rank projections. This results in an even more computationally efficient (but heuristic) strategy, that does not 16 17rely on an inner-outer iteration scheme. Numerical experiments including image deblurring, computed tomography 18 and inpainting show that our new solvers are competitive with other state-of-the-art solvers for low-rank problems, and deliver reconstructions of increased quality with respect to other classical Krylov methods. 19

20 **Key words.** low-rank solver, nuclear norm regularization, Krylov methods, flexible Krylov methods, Kronecker 21 product, imaging problems

22 AMS subject classifications. 65F20, 65F30

1. Introduction. Consider the following linear system

24 (1.1)
$$Ax = b$$
, where $A \in \mathbb{R}^{M \times N}$, $x \in \mathbb{R}^N$, $b = b^{ex} + \eta \in \mathbb{R}^M$.

We are mainly interested in large-scale linear systems (1.1) arising from inverse problems, where Ais a discretization of the linear forward operator, x is a quantity of interest, and b is the observed perturbed data ($b^{\text{ex}} = Ax^{\text{ex}}$ being the ideally exact data, and η being unknown Gaussian white noise). Our focus is on two-dimensional imaging problems, where the unknown vector $x \in \mathbb{R}^N$ is obtained by stacking the columns of an unknown true image X of size $n \times n$, with $n = \sqrt{N}$ (this operation and its inverse are denoted by x = vec(X) and $X = \text{vec}^{-1}(x)$, respectively).

31 Discrete inverse problems are ill-posed in nature [13] and, because of the presence of noise in (1.1), regularization needs to be applied so that the solution of (1.1) is a meaningful approxima-32 tion to $\boldsymbol{x}^{\text{ex}}$. One typically achieves regularization by replacing the original problem (1.1) with a 33 closely related one that is less sensitive to perturbations: effective regularization methods do so by 34 incorporating known or desired properties of x into the solution process. In imaging applications, 35 Tikhonov (ℓ_2) regularization, ℓ_1 regularization and total variation are typical techniques to be ex-36 plotted, see, for example, [4, 7, 8, 9, 17]. In the field of geophysics, ℓ_0 regularization (also called 37 compact regularization) is sometimes considered (e.g., [21, 34]). 38

^{*}Submitted to the editors DATE.

Funding: This work was funded by the National Science Foundation under grant no. DMS-1819042. and partially supported by the EPSRC under grant no. EP/T001593/1.

[†]Department of Mathematical Sciences, University of Bath, United Kingdom. (s.gazzola@bath.ac.uk).

[‡]Department of Mathematics, Emory University, Atlanta, GA, USA. (chang.meng@emory.edu, jnagy@emory.edu)

In this paper we consider regularization methods that compute a low-rank approximate solution $X = \text{vec}^{-1}(x)$ of (1.1): this is generally meaningful when the unknown x encodes a high-dimensional quantity and, in particular, in the case of a two-dimensional image. Indeed, two-dimensional images are often assumed to have low-rank or to be well-approximated by low-rank two-dimensional arrays (see [27] and the references therein).

Numerical linear algebra solvers for the estimation of low-rank solutions to linear systems 44 have been developed in the literature, mainly targeting well-posed linear discrete problems, such as 45 those arising when considering the numerical solution of stochastic PDEs (see [22] and the references 46 47 therein). In particular, the authors of [22] devise a restarted GMRES-like method (RS-LR-GMRES) that involves low-rank projections of the basis vectors of the solution subspace, as well as a low-rank 48 projection of the current solution at the end of each cycle. Since, in general, the basic operations 49involved in standard GMRES (such as matrix-vector products and vector sums) increase the ranks 50of the computed quantities, low-rank projections are needed to assure that the computed solution 51is low-rank. In the framework of compressive sensing, the authors of [2] consider a modified version 53 of the conjugate gradient method that incorporates appropriate rank-truncation operations. All the methods mentioned so far employ, often in a heuristic way, Krylov subspace methods together 54with rank-reduction operations (e.g., projections onto a chosen set of low-rank matrices). Since 55many Krylov subspace methods are iterative regularization methods for (1.1), this brings us to the 56question of how incorporating rank-reduction operations would affect the solution of the discrete 57 58 inverse problem (1.1), with a particular focus on imaging applications.

Low-rank matrix estimation can be naturally formulated as a nonconvex optimization problem having either: (i) a least-squares data fitting term as objective function and a rank constraint; (ii) the rank of $\mathbf{X} = \text{vec}^{-1}(\mathbf{x})$ as objective function and a constraint on the least-squares data fitting term. The last instance is commonly referred to as *affine rank minimization problem*, and both formulations are in general NP-hard [27]. In this paper we consider the unconstrained and convex optimization problem

65 (1.2)
$$\min_{x} \|Ax - b\|_{2}^{2} + \lambda \|\operatorname{vec}^{-1}(x)\|_{*}$$

66 where $\lambda > 0$ is a regularization parameter and $\|\cdot\|_*$ denotes the nuclear norm of $\operatorname{vec}^{-1}(\boldsymbol{x}) = \boldsymbol{X}$, 67 defined as the sum of the singular values of \boldsymbol{X} . Indeed, if the singular value decomposition (SVD) 68 of \boldsymbol{X} is given by $\boldsymbol{X} = \boldsymbol{U}_{\boldsymbol{X}} \boldsymbol{\Sigma}_{\boldsymbol{X}} \boldsymbol{V}_{\boldsymbol{X}}^T$, where $\boldsymbol{U}_{\boldsymbol{X}}, \boldsymbol{V}_{\boldsymbol{X}} \in \mathbb{R}^{n \times n}$ are orthogonal matrices, and $\boldsymbol{\Sigma}_{\boldsymbol{X}} \in \mathbb{R}^{n \times n}$ 69 is the diagonal matrix whose diagonal entries are $\sigma_1(\boldsymbol{X}) \ge \cdots \ge \sigma_n(\boldsymbol{X}) \ge 0$, then

70
$$\|\boldsymbol{X}\|_{*} = \sum_{i=1}^{n} \sigma_{i}(\boldsymbol{X}).$$

Problem (1.2) is referred to as a nuclear norm regularized (NNR) problem. In particular, the nuclear 71 norm is a convex function that has been proven to be the best convex lower approximation of the 72 rank function over the set of matrices X such that $||X||_2 \leq 1$ (see [27] and the references therein). 73 The nuclear norm has been used in many applications, such as low-rank matrix completion and 74 compressed sensing; see, e.g., [3, 10, 16, 24, 27], where the constrained formulation of problem (1.2) 75has also been considered (note that, for a proper choice of $\lambda > 0$, constrained and unconstrained 76 formulations are equivalent; see, e.g., [29]). In the framework of compressive sensing, under the 77 assumption that the matrix A satisfies a certain null-space property, recovery guarantees for the 78

LOW-RANK KRYLOV METHODS

⁷⁹ affine rank minimization problem are proven in [5, 25]. We also consider the following formulation

80 (1.3)
$$\min_{\boldsymbol{x}} \|\boldsymbol{A}\boldsymbol{x} - \boldsymbol{b}\|_{2}^{2} + \lambda \|\operatorname{vec}^{-1}(\boldsymbol{x})\|_{*,p}, \quad \text{where} \quad \|\boldsymbol{X}\|_{*,p} = \sum_{i=1}^{n} (\sigma_{i}(\boldsymbol{X}))^{p}, \quad 0$$

Problem (1.3) is referred to as *NNRp problem*, and it generalizes problem (1.2) (which is obtained taking p = 1 in (1.3)). The constrained version of (1.3) is already considered in [25], where the authors empirically show an improved recovery performance of the constrained formulation of problem (1.3) with p < 1 with respect to p = 1. Note, however, that the choice p < 1 in (1.3) results in a nonconvex minimization problem.

Many different optimization methods, such as singular value thresholding (i.e., projected gradient descent) and continuation methods [10], have been proposed for the solution of problem (1.2) or its constrained counterpart. In particular, the so-called IRLS(-p) (i.e., iteratively reweighted least squares) family of methods has recently attracted a lot of attention [5, 25, 27]. IRLS(-p) solves the affine rank minimization problem by solving a sequence of problems whose objective function only involves an iteratively updated weighted 2-norm term. The authors of [23] apply the IRLS(-p) framework to the unconstrained problem (1.3), requiring the solution of a sequence of sub-problems

93 (1.4)
$$\min_{\boldsymbol{x}} \|\boldsymbol{A}\boldsymbol{x} - \boldsymbol{b}\|_{2}^{2} + \lambda \|\boldsymbol{W}_{k} \operatorname{vec}^{-1}(\boldsymbol{x})\|_{F}^{2}$$

where W_k is an appropriate weight matrix to be employed to solve the kth sub-problem, and $\|\cdot\|_F$ 94 denotes the Frobenius norm of a matrix. A gradient (projection) algorithm is typically used to 95solve each sub-problem (1.4). Since an IRLS(-p) approach is also commonly applied to objective 96 functions involving a quadratic fit-to-data term and a general p-norm penalization on \boldsymbol{x} , and since 97 efficient strategies based on Krylov methods have been devised to solve each quadratic sub-problem 98 in the IRLS(-p) sequence [28, 30], this brings us to the question of how Krylov methods can be best 99 employed to solve each problem (1.4) (recall that $\|vec^{-1}(\boldsymbol{x})\|_{*,p}$ can be regarded as a *p*-norm of the 100 vector whose entries are the singular values of $X = \text{vec}^{-1}(x)$. 101

The goal of this paper is to propose new efficient Krylov methods for the estimation of low-rank 102103 solutions to (1.1). We will mainly consider an IRLS(-p) approach to problem (1.3) (rather than incorporating low-rank projections into a linear solver for (1.1), the upside being that low-rank is 104implicitly enforced into the solution by penalizing the p-norm of the singular values for a suitable 105 choice of λ . Our main contributions are the new IRN-GMRES-NNRp and IRN-LSQR-NNRp meth-106 ods for (1.3), where automatic strategies for choosing a suitable λ are naturally incorporated. Here 107 and in the following, the IRN acronym indicates an iteratively reweighted norm (rather than an 108 iteratively reweighted least squares problem, [30]). One of the key points in deriving the new meth-109 ods is expressing in matrix form the invertible linear operator mapping x to the reweighted 2-norm 110 of the singular values of $\mathbf{X} = \text{vec}^{-1}(\mathbf{x})$: this can be achieved in a computationally affordable way by 111 exploiting Kronecker product properties. Each iteratively reweighted quadratic sub-problem of the 112form (1.4) can then be expressed as a Tikhonov regularization problem in general form, which can 113 be straightforwardly transformed into standard form. In this way, the inverse of the linear operator 114 mapping x into the reweighted 2-norm of the singular values of $X = \text{vec}^{-1}(x)$ formally acts as a 115 preconditioner for A, and the so-called hybrid methods [26] based on the preconditioned Arnoldi (if 116 117 A is square) or Golub-Kahan bidiagonalization algorithms can be used to efficiently approximate the solution of each problem of the form (1.4). Once a hybrid method is adopted, many automatic, 118 adaptive, and efficient parameter choice strategies can be employed to choose a suitable λ ; see [18] 119for an overview. Therefore, contrarily to many existing methods for (1.3), IRN-GMRES-NNRp and 120

121 IRN-LSQR-NNRp have the advantage of not requiring a regularization parameter (either λ or the 122 desired rank of the solution) to be available in advance of the iterations, nor the repeated solution 123 of (1.3) for different regularization parameters.

Although inherently efficient, both the IRN-GMRES-NNRp and IRN-LSQR-NNRp methods 124are inner-outer iteration schemes, where each outer iteration requires running a "preconditioned" 125Krylov subspace method until convergence (inner iteration) before updating the weights (and there-126127 fore the "preconditioner") in the next outer iteration. In order to avoid inner-outer iterations and with the aim of generating only one approximation subspace for the solution of (1.3), where a new 128129"preconditioner" is incorporated as soon as a new approximate solution becomes available (i.e., at each iteration), we propose to solve (1.3) using flexible Krylov subspace methods, such as those based 130 on the flexible Arnoldi [31] and Golub-Kahan [4] algorithms. The use of flexible Krylov methods for p-norm regularization of inverse problems was already proposed in [4, 7]; however, differently from the available solvers, our new approach involves iteratively defining both weights and transform 133 matrices (i.e., the linear operator mapping $vec^{-1}(x)$ into its singular values). Switching from IRN-134GMRES-NNRp and IRN-LSQR-NNRp to their flexible counterparts (dubbed FGMRES-NNRp and 135FLSQR-NNRp, respectively) allows for savings in computations and, although FGMRES-NNRp 136and FLSQR-NNRp are purely heuristic, it leads to approximate solutions whose accuracy on many 137test problems is comparable to the ones of other well established solvers for (1.2). Motivated by 138 the same idea of avoiding inner-outer iteration cycles while adaptively incorporating (low-rank) 139information into the approximation subspace for the solution, we also propose a flexible version 140of the projected and restarted Krylov subspace methods (such as RS-LR-GMRES, [22]) that were 141 originally devised for square matrices, considering also extensions to rectangular matrices A.

This paper is organized as follows. In Section 2 we review the available low-rank Krylov methods 143 for square linear systems and, after surveying the available flexible Krylov solvers, we formulate 144 new low-rank flexible Krylov solvers for both square and rectangular problems, where the basis 145vectors for the approximation subspace are truncated to low-rank. In Section 3 we derive the new 146 iteratively reweighted methods for (1.3) as fixed-point methods, and we describe how to efficiently 147solve each reweighted problem of the form (1.4) using preconditioned Krylov methods: this leads 148 to the IRN-GMRES-NNRp and IRN-LSQR-NNRp methods; their flexible counterparts (FGMRES-149150NNRp and FLSQR-NNRp, respectively) are also derived. Some implementation details, such as stopping criteria and regularization parameter choice strategies for the new methods, are unfolded 151in Section 4. Numerical results on image deblurring, computed tomography and inpainting are 152presented in Section 5, including comparisons between the proposed methods, low-rank projection 153methods, projected gradient methods, and standard Krylov subspace methods. Conclusions are 154drawn in Section 6.

Definitions and notations. Matching lower and upper case letters are used to denote the 156"vectorized" and "matricized" versions of a given quantity, respectively; e.g., c = vec(C) and 157 $C = \text{vec}^{-1}(c)$. We denote the *i*th entry of a vector c by $[c]_i$, and the (i, j)th entry of a matrix C158by $[C]_{ij}$ or, using MATLAB-like notations, $[c]_i = c(i), [C]_{ij} = C(i, j)$. Using again MATLAB-like notations, d = diag(C) defines a vector d whose entries are the diagonal elements of a matrix 160 C. $\operatorname{Tr}(C)$ denotes the trace of a matrix C. $\mathcal{R}(C)$ denotes the range (or column space) of a 161 matrix C, and $\mathcal{K}_m(A, b)$ denotes the m-dimensional Krylov subspace defined by A and b, i.e., 162 $\mathcal{K}_m(\mathbf{A}, \mathbf{b}) = \operatorname{span} \{ \mathbf{b}, \mathbf{A}\mathbf{b}, \mathbf{A}^2\mathbf{b}, \dots, \mathbf{A}^{m-1}\mathbf{b} \}.$ We denote by $\mathbf{I} \in \mathbb{R}^{d \times d}$ the identity matrix of order 163d, and by e_i the *i*th canonical basis vector of \mathbb{R}^d , where d should be clear from the context. Note 164that, in the following, we will quite often interchange x and X and, with a slight abuse of notations, we will denote the action of a linear operator on x or X by $\mathcal{A}(X) = \mathcal{A}X = Ax$, and the action of 166

LOW-RANK KRYLOV METHODS

167 the adjoint operator by $\mathcal{A}^*(\mathbf{Y}) = \mathcal{A}^*\mathbf{Y} = A^T \operatorname{vec}(\mathbf{Y}).$

2. Low-rank projection methods: classical and new approaches. As recalled in Section 1, when solving square well-posed linear systems coming from the discretization of some instances of stochastic or time-dependent PDEs, a suitable rearrangement of the solution is expected to be low-rank: for this reason, schemes that incorporate low-rank projections within the basis vectors and the approximate solution obtained by a Krylov method have been proposed in the literature. In the following we summarize the working ideas underlying the so-called restarted low-rank-projected GMRES (RS-LR-GMRES) method proposed in [22].

The starting points for the derivation of RS-LR-GMRES are the basic properties and relations underlying GMRES. Indeed, one can define GMRES for the solution of (1.1) with a square $A \in \mathbb{R}^{N \times N}$ and initial guess $x_0 = 0$ by generating a matrix $V_m = [v_1, \ldots, v_m] \in \mathbb{R}^{N \times m}$ with orthonormal columns, such that $\mathcal{R}(V_m) = \mathcal{K}_m(A, b)$, and imposing that the residual $r_m = b - Ax_m$ is orthogonal to $U_m = AV_m$. In practice, at the *k*th iteration of GMRES, one computes

180 (2.1)
$$\boldsymbol{u}_{k} = \boldsymbol{A}\boldsymbol{v}_{k-1} \text{ and } \|\boldsymbol{v}_{k}\|_{2}\boldsymbol{v}_{k} = (\boldsymbol{I} - \boldsymbol{V}_{k-1}(\underbrace{\boldsymbol{V}_{k-1}^{T}\boldsymbol{V}_{k-1}}_{=\boldsymbol{I}})^{-1}\boldsymbol{V}_{k-1}^{T})\boldsymbol{u}_{k},$$

181 and the approximate solution is computed as

182 (2.2)
$$\boldsymbol{x}_k = \boldsymbol{V}_k \boldsymbol{y}_k$$
, where $(\boldsymbol{U}_k^T \boldsymbol{A} \boldsymbol{V}_k) \boldsymbol{y}_k = \boldsymbol{U}_k^T \boldsymbol{b}$

183 This procedure is mathematically equivalent to the somewhat more standard procedure that, at the

*k*th iteration of GMRES, updates the partial Arnoldi factorization and computes the approximate solution as follows:

186 (2.3)
$$\boldsymbol{A}\boldsymbol{V}_{k} = \boldsymbol{V}_{k+1}\boldsymbol{H}_{k}, \quad \boldsymbol{x}_{k} = \boldsymbol{V}_{k}\boldsymbol{y}_{k}, \text{ where } \boldsymbol{y}_{k} = \arg\min_{\boldsymbol{y} \in \mathbb{R}^{k}} \|\boldsymbol{H}_{k}\boldsymbol{y} - \|\boldsymbol{b}\|_{2}\boldsymbol{e}_{1}\|_{2}.$$

Note that, in particular, the matrix V_k appearing in (2.2) coincides with the matrix V_k appearing 187 in (2.3). However, since matrix-vector products and vector sums of low-rank vectorized matrices 188 189increase the rank of the latter, relations (2.1) and (2.2) obviously do not guarantee that the new basis vectors v_k for the solution nor the new solution x_k are low-rank. To force the basis vector 190for the solution and the approximate solution to be low-rank, a truncation operator should be 191 incorporated into the GMRES algorithm. Given a vectorized matrix $\boldsymbol{c} = \operatorname{vec}(\boldsymbol{C})$, and given a 192desired low-rank κ for C, one can define a truncation operator $\tau_{\kappa}(c)$ by the following standard 193194operations:

195 (2.4)

$$\begin{bmatrix}
1. & \text{Take } \boldsymbol{C} = \text{vec}^{-1}(\boldsymbol{c}); \\
2. & \text{Compute the SVD of } \boldsymbol{C}, \, \boldsymbol{C} = \boldsymbol{U}_{\boldsymbol{C}} \boldsymbol{\Sigma}_{\boldsymbol{C}} \boldsymbol{V}_{\boldsymbol{C}}^{T}; \\
3. & \text{Compute } \boldsymbol{C}_{\kappa} = \boldsymbol{U}_{\boldsymbol{C}}(:, 1:\kappa) \boldsymbol{\Sigma}_{\boldsymbol{C}}(1:\kappa, 1:\kappa) \boldsymbol{V}_{\boldsymbol{C}}(:, 1:\kappa)^{T}; \\
4. & \text{Take } \tau_{\kappa}(\boldsymbol{c}) = \text{vec}(\boldsymbol{C}_{\kappa}).
\end{bmatrix}$$

RS-LR-GMRES is a restarted version of the standard GMRES method where the basis vectors for the solution are truncated at each inner iteration, and the solution itself is truncated at the beginning of each outer iteration. Note that truncating the basis vectors does not guarantee that the solution has low rank (which is the reason we still need to truncate the approximate solution). The reason for truncating the basis vectors is to keep the original solution rank from increasing drastically, since it is computed as a linear combination of basis vectors. More precisely, at the ℓ th

S. GAZZOLA, C. MENG AND J. NAGY

outer iteration of RS-LR-GMRES, one takes $v_1 = r_{\ell-1}/||r_{\ell-1}||_2$, where $r_{\ell-1} = b - Ax_{\ell-1}$, and, at the *k*th inner iteration, one computes

204 (2.5)
$$\boldsymbol{u}_{k} = \boldsymbol{A}\boldsymbol{v}_{k-1}$$
 and $\|\boldsymbol{v}_{k}\|_{2}\boldsymbol{v}_{k} = \tau_{\kappa}\left((\boldsymbol{I} - \boldsymbol{V}_{k-1}(\boldsymbol{V}_{k-1}^{T}\boldsymbol{V}_{k-1})^{-1}\boldsymbol{V}_{k-1}^{T})\boldsymbol{u}_{k}\right)$

Once m inner iterations are performed, the approximate solution at the ℓ th outer iteration is computed as

207 (2.6)
$$\boldsymbol{x}_{\ell} = \tau_{\kappa} \left(\boldsymbol{x}_{\ell-1} + \boldsymbol{V}_m \boldsymbol{y}_m \right), \text{ where } \left(\boldsymbol{U}_m^T \boldsymbol{A} \boldsymbol{V}_m \right) \boldsymbol{y}_m = \boldsymbol{U}_m^T \boldsymbol{r}_{\ell-1}.$$

The operations in (2.5) and (2.6) heavily depend on the value κ of the truncated rank, which 208 eventually coincides with the rank of the approximate solution. In the framework of stochastic 209 PDEs, a suitable estimate for κ can be obtained by first performing coarse-grid computations (see 210[22] for details, and [19, 33] for similar approaches). Comparing (2.5) and (2.1) one can see that, 211 as in standard GMRES, RS-LR-GMRES computes a new basis vector for the solution by applying 212213the linear operator A to the previous basis vector v_{k-1} and orthogonalizing it against the previous basis vectors v_i , $i = 1, \ldots, k - 1$. However, since the basis vectors are truncated to low rank, 214 the matrix V_k does not have orthonormal columns anymore, and $\mathcal{R}(V_m)$ is not a Krylov subspace 215anymore. This remark leads us to the derivation of alternative low-rank projection solvers, which 216can be (re)casted into the framework of flexible Krylov methods and can work with both square 217 and rectangular systems (1.1). 218

Low-rank flexible GMRES (LR-FGMRES) and low-rank flexible LSQR (LR-FLSQR). Flexible 219Krylov methods are a class of linear solvers that can handle iteration-dependent preconditioners: 220 they were originally introduced in [31] for FGMRES, where a preconditioner for GMRES was al-221 222 lowed to change from one iteration to the next (either because at each iteration the preconditioner is implicitly defined by applying an iterative linear solver, or because the preconditioner can be 223 updated with newly-computed information; see [32] for an overview). In the framework of regular-224izing linear solvers, flexible Krylov methods were proposed in [4, 7, 9], where the iteration-dependent 225"preconditioner" was associated to an iteratively reweighted norm approach to Tikhonov-like reg-226ularized problems involving penalization terms expressed in some p-norm, 0 (and, indeed,these "preconditioners" have the effect of enforcing specific regularity into the approximation sub-228space for the solution, rather than accelerating the convergence of the iterative solvers). Leveraging 229flexible Krylov subspaces in this setting comes with the upside of avoiding restarts of the itera-230tive solver, which is the approach commonly used when adopting an iteratively reweighted norm 231 method. When considering low-rank projections of the basis vectors within RS-LR-GMRES, we 232 enforce the basis vectors to have low-rank, so to better reproduce available information about the 233solution of (1.1) (i.e., the solution should be low-rank). It is therefore natural to consider flexible 234Krylov methods that involve truncation of the basis vectors at each iteration, as a computationally 235cheaper alternative to RS-LR-GMRES that does not involve restarts. 236

Considering first the case of a square $\mathbf{A} \in \mathbb{R}^{N \times N}$, we can use the flexible Arnoldi algorithm [31] to naturally incorporate low-rank basis vectors for the solution of (1.1). In general, starting with $\mathbf{x}_0 = \mathbf{0}$, at the *k*th iteration, FGMRES updates a partial flexible Arnoldi factorization and computes the *k*th approximate solution as follows:

241 (2.7)
$$\boldsymbol{A}\boldsymbol{Z}_{k} = \boldsymbol{V}_{k+1}\boldsymbol{H}_{k}, \quad \boldsymbol{x}_{k} = \boldsymbol{Z}_{k}\boldsymbol{y}_{k}, \text{ where } \boldsymbol{y}_{k} = \arg\min_{\boldsymbol{y} \in \mathbb{R}^{k}} \|\boldsymbol{H}_{k}\boldsymbol{y} - \|\boldsymbol{b}\|_{2}\boldsymbol{e}_{1}\|_{2}$$

where $V_{k+1} = [v_1, \ldots, v_{k+1}] \in \mathbb{R}^{N \times (k+1)}$ has orthonormal columns, $H_k \in \mathbb{R}^{(k+1) \times k}$ is upper Hessenberg, and $Z_k = [P_1 v_1, \ldots, P_k v_k] \in \mathbb{R}^{N \times k}$ has columns that span the approximation subspace for the solution (P_i is an iteration-dependent preconditioner that is applied to v_i and, in the particular case of low-rank truncation, $P_i v_i = \tau_{\kappa_B}(v_i)$, is the truncation operator defined in (2.4), so that rank(vec⁻¹($Z_k e_i$)) = κ_B , i = 1, ..., k). The subscript B for the truncation rank κ_B suggests that the truncation is done on the original basis vectors v_i 's. The resulting algorithm is dubbed "LR-FGMRES", and it is summarized in Algorithm 2.1. Note that the approximate solution computed as in (2.7) is also truncated to guarantee rank κ (in general, we assume $\kappa_B \neq \kappa$). Also LR-FGMRES is started with $x_0 = 0$, to guarantee that the basis vectors for the solution (rather than a correction thereof) are low-rank.

Algorithm 2.1 LR-FGMRES

1: Inputs: $A, b, \tau_{\kappa_B}, \tau_{\kappa}$ 2: Take $v_1 = b/\|b\|_2$ 3: for i = 1, 2, ... until a stopping criterion is satisfied do 4: Compute $z_i = \tau_{\kappa_B}(v_i)$ and $w = Az_i$ 5: Compute $h_{ji} = w^T v_j$ for j = 1, ..., i and set $w = w - \sum_{j=1}^i h_{ji} v_j$ 6: Compute $h_{i+1,i} = \|w\|_2$, and if $h_{j+1,j} \neq 0$, take $v_{i+1} = w/h_{i+1,i}$ 7: end for 8: Compute $y_k = \arg\min_y \|H_k y - \|b\|_2 e_1\|_2^2$ and take $x_k = \tau_\kappa(Z_k y_k)$

A few remarks are in order. Differently from the kth iteration in the inner cycle of the RS-252253LR-GMRES method (2.5), the kth iteration of LR-FGMRES expands the approximation subspace by modifying (i.e., truncating) the previous orthonormal basis vector for the space $\mathcal{R}([b, AZ_k])$. 254Analogously to RS-LR-GMRES, the basis vectors for the approximate LR-FGMRES solution are 255all of rank κ_B , are not orthogonal, and do not span a Krylov subspace. Differently from RS-LR-256GMRES, the basis vector for the space $\mathcal{R}([b, AZ_k])$ are orthogonal. Also, the kth LR-FGMRES 257approximate solution is obtained by solving an order-k projected least squares problem that is 258formally analogous to the GMRES one (see (2.3) and (2.7)). 259

With LR-FGMRES in place, the extension to more general matrices $\mathbf{A} \in \mathbb{R}^{M \times N}$, with Mnot necessarily equal to N, can be naturally devised considering the flexible Golub-Kahan (FGK) process [4]. Taking $\mathbf{x}_0 = \mathbf{0}$ as initial guess, the *k*th FGK iteration updates partial factorizations of the form

264 (2.8)
$$AZ_k = U_{k+1}M_k$$
 and $A^T U_{k+1} = V_{k+1}T_{k+1}$,

where the columns of $U_{k+1} \in \mathbb{R}^{M \times (k+1)}$, $V_{k+1} \in \mathbb{R}^{N \times (k+1)}$ are orthonormal, $M_k \in \mathbb{R}^{(k+1) \times k}$ is upper Hessenberg, $T_{k+1} \in \mathbb{R}^{(k+1) \times (k+1)}$ is upper triangular, and $Z_k = [P_1v_1, \dots, P_kv_k] \in \mathbb{R}^{N \times k}$ 265266has columns that span the approximation subspace for the solution (P_i is an iteration-dependent 267preconditioner that is applied to v_i and, in the particular case of low-rank truncation, $P_i v_i$ 268 $\tau_{\kappa_B}(\boldsymbol{v}_i)$, as defined in (2.4), so that rank $(\operatorname{vec}^{-1}(\boldsymbol{Z}_k \boldsymbol{e}_i)) = \kappa_B, i = 1, \ldots, k)$. The flexible LSQR 269 method (FLSQR) uses the FGK process (2.8) to generate iterates of the form $\boldsymbol{x}_k = Z_k \boldsymbol{y}_k$, where 270the vector \boldsymbol{y}_k is computed as $\boldsymbol{y}_k = \arg\min_{\boldsymbol{y}} \left\| \boldsymbol{M}_k \boldsymbol{y} - \| \boldsymbol{b} \|_2 \boldsymbol{e}_1 \right\|_2^2$. When rank-truncation of the basis 271vectors takes place at each iteration, and the final approximate solution is rank-truncated as well, 272the resulting algorithm is dubbed "LR-FLSQR", and it is summarized in Algorithm 2.2. Note that, 273similarly to RS-LR-GMRES, both LR-FGMRES and LR-FLSQR are quite heuristic. Although the 274low-rank projection idea can be formulated in the flexible framework, we lack a formal formulation of 275

Algorithm 2.2 LR-FLSQR

1: Inputs: $\boldsymbol{A}, \boldsymbol{b}, \tau_{\kappa_B}, \tau_{\kappa}$ 2: Take $\boldsymbol{u}_1 = \boldsymbol{b}/\|\boldsymbol{b}\|_2$ 3: for $i = 1, 2, \ldots$, until a stopping criterion is satisfied do 4: Compute $\boldsymbol{w} = \boldsymbol{A}^T \boldsymbol{u}_i, t_{ji} = \boldsymbol{w}^T \boldsymbol{v}_j$ for $j = 1, \ldots, i-1$ 5: Set $\boldsymbol{w} = \boldsymbol{w} - \sum_{j=1}^{i-1} t_{ji} \boldsymbol{v}_j$, compute $t_{ii} = \|\boldsymbol{w}\|$ and take $\boldsymbol{v}_i = \boldsymbol{w}/t_{ii}$ 6: Compute $\boldsymbol{z}_i = \tau_{\kappa_B}(\boldsymbol{v}_i)$ and $\boldsymbol{w} = \boldsymbol{A}\boldsymbol{z}_i$ 7: Compute $m_{ji} = \boldsymbol{w}^T \boldsymbol{u}_j$ for $j = 1, \ldots, i$ and set $\boldsymbol{w} = \boldsymbol{w} - \sum_{j=1}^{i} m_{ji} \boldsymbol{u}_j$ 8: Compute $m_{i+1,i} = \|\boldsymbol{w}\|$ and take $\boldsymbol{u}_{i+1} = \boldsymbol{w}/m_{i+1,i}$ 9: end for 10: Compute $\boldsymbol{y}_k = \arg\min_{\boldsymbol{y}} \|\boldsymbol{M}_k \boldsymbol{y} - \|\boldsymbol{b}\|_2 \boldsymbol{e}_1\|_2^2$ and take $\boldsymbol{x}_k = \tau_{\kappa}(\boldsymbol{Z}_k \boldsymbol{y}_k)$

the problem that is being solved, and also a justification of why they work. Strategies for selecting κ_B and κ are not so clear either. To stabilize the behavior of LR-FGMRES as the iterations proceed, one may consider imposing additional Tikhonov regularization on the projected leastsquares problem in (2.7), in a hybrid fashion; the same holds for LR-FLSQR (see Sections 3.3 and 5 for more details).

3. Proposed Method. In this section, we first derive the IRN method for the solution of the 281 NNRp problem (1.3). The starting point for our derivations is the approximation of the nondifferen-282tiable nuclear norm regularizer by a smooth Schatten function (similarly to what is proposed in [25] 283for the affine rank minimization problem). The optimality conditions associated to the smoothed 284 285problem give rise to a nonlinear system of equations in X, which is handled by a fixed-point iteration scheme. We show that each iteration amounts to the solution of a Tikhonov-regularized 286 problem involving an iteratively reweighted 2-norm regularization term, which can be efficiently 287solved employing "preconditioned" Krylov methods. Flexible Krylov methods are introduced to 288 approximate the solution of the IRN problem within only one adaptively defined approximation 289subspace for the solution, bypassing the inner-outer iteration scheme required by standard Krylov 290methods. 291

3.1. Derivation. Define the smooth Schatten-*p* function as

293
$$\mathcal{S}_p^{\gamma}(\boldsymbol{X}) = \operatorname{Tr}((\boldsymbol{X}^T\boldsymbol{X} + \gamma \boldsymbol{I})^{p/2}), \text{ with } \gamma > 0$$

Note that $S_p^{\gamma}(\mathbf{X})$ is differentiable for p > 0 and convex for $p \ge 1$. In particular, for p = 1 and $\gamma = 0$ (i.e., no smoothing),

296
$$S_1^0(X) = \text{Tr}((X^T X)^{1/2}) = ||X||_*.$$

297 We start by considering the following smooth approximation to (1.3):

298 (3.1)
$$\min_{\boldsymbol{X} \in \mathbb{R}^{n \times n}} \| \mathcal{A}(\boldsymbol{X}) - \boldsymbol{B} \|_F^2 + \lambda \mathcal{S}_p^{\gamma}(\boldsymbol{X}) \,.$$

The following derivations are valid for p > 0 (and we keep them generic, being aware that p = 1approximates (1.2)). The optimality conditions associated to (3.1) read

301
$$0 = \nabla_{\boldsymbol{X}} \left(\| \mathcal{A}(\boldsymbol{X}) - \boldsymbol{B} \|_{F}^{2} + \lambda \mathcal{S}_{p}^{\gamma}(\boldsymbol{X}) \right)$$

302 (3.2)
$$= 2\mathcal{A}^*(\mathcal{A}(\mathbf{X}) - \mathbf{B}) + \lambda p(\mathbf{X}\mathbf{X}^T + \gamma \mathbf{I})^{p/2-1}\mathbf{X},$$

303 where we have used that

$$\nabla_{\boldsymbol{X}} \operatorname{Tr}((\boldsymbol{X}^T \boldsymbol{X} + \gamma \boldsymbol{I})^{p/2}) = p \boldsymbol{X} (\boldsymbol{X}^T \boldsymbol{X} + \gamma \boldsymbol{I})^{p/2-1} = p (\boldsymbol{X} \boldsymbol{X}^T + \gamma \boldsymbol{I})^{p/2-1} \boldsymbol{X}.$$

Equivalently, the nonlinear system of equations (3.2) with respect to X can be expressed as

306
$$\boldsymbol{X} = \left(\mathcal{A}^*\mathcal{A} + \hat{\lambda}(\boldsymbol{X}\boldsymbol{X}^T + \gamma\boldsymbol{I})^{p/2-1}\right)^{-1}\mathcal{A}^*\boldsymbol{B}$$

304

307
$$= \left(\mathcal{A}^*\mathcal{A} + \hat{\lambda}((\boldsymbol{X}\boldsymbol{X}^T + \gamma\boldsymbol{I})^{p/4-1/2})^T(\boldsymbol{X}\boldsymbol{X}^T + \gamma\boldsymbol{I})^{p/4-1/2}\right)^{-1}\mathcal{A}^*\boldsymbol{B}, \quad \text{with } \hat{\lambda} = \lambda p/2,$$

308 which is naturally associated to the following fixed-point iteration scheme

309 (3.3)
$$X_{k+1} = \left(\mathcal{A}^* \mathcal{A} + \hat{\lambda} ((X_k X_k^T + \gamma I)^{p/4 - 1/2})^T (X_k X_k^T + \gamma I)^{p/4 - 1/2} \right)^{-1} \mathcal{A}^* B ,$$

310 which leads to the solution of (3.1). Equivalently,

311
$$\boldsymbol{X}_{k+1} = \arg\min_{\boldsymbol{X}} \left\| \begin{bmatrix} \boldsymbol{\mathcal{A}} \\ \sqrt{\hat{\lambda}} (\boldsymbol{X}_k \boldsymbol{X}_k^T + \gamma \boldsymbol{I})^{p/4-1/2} \end{bmatrix} \boldsymbol{X} - \begin{bmatrix} \boldsymbol{B} \\ \boldsymbol{0} \end{bmatrix} \right\|_{F}^{2}$$

i.e., (3.3) are the normal equations associated to the penalized least squares problem written above or, equivalently,

314 (3.4)
$$\boldsymbol{X}_{k+1} = \arg\min_{\boldsymbol{X}} \left\| \boldsymbol{\mathcal{A}} \boldsymbol{X} - \boldsymbol{B} \right\|_{F}^{2} + \hat{\lambda} \left\| (\boldsymbol{X}_{k} \boldsymbol{X}_{k}^{T} + \gamma \boldsymbol{I})^{p/4 - 1/2} \boldsymbol{X} \right\|_{F}^{2}.$$

315 We now reformulate problem (3.4) in vectorial form.

Let $U_{X_k} \Sigma_{X_k} V_{X_k}^T = X_k$ be the SVD of X_k ; thanks to the invariance of the Frobenius norm under orthogonal transformations, the regularization term in the above problem can be rewritten as

319
$$\left\| (\boldsymbol{X}_{k}\boldsymbol{X}_{k}^{T} + \gamma \boldsymbol{I})^{p/4-1/2}\boldsymbol{X} \right\|_{F}^{2} = \left\| \boldsymbol{U}_{\boldsymbol{X}_{k}} (\boldsymbol{\Sigma}_{\boldsymbol{X}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2} \boldsymbol{U}_{\boldsymbol{X}_{k}}^{T} \boldsymbol{X} \right\|_{F}^{2} = \left\| (\boldsymbol{\Sigma}_{\boldsymbol{X}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2} \boldsymbol{U}_{\boldsymbol{X}_{k}}^{T} \boldsymbol{X} \boldsymbol{V}_{\boldsymbol{X}_{k}} \right\|_{F}^{2}$$

320 Using well-known Kronecker product properties

321
$$\left\| (\boldsymbol{\Sigma}_{\boldsymbol{X}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2} \boldsymbol{U}_{\boldsymbol{X}_{k}}^{T} \boldsymbol{X} \boldsymbol{V}_{\boldsymbol{X}_{k}} \right\|_{F}^{2} = \left\| \operatorname{vec} \left((\boldsymbol{\Sigma}_{\boldsymbol{X}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2} \boldsymbol{U}_{\boldsymbol{X}_{k}}^{T} \boldsymbol{X} \boldsymbol{V}_{\boldsymbol{X}_{k}} \right) \right\|_{2}^{2}$$

$$322 = \left\| \left(\boldsymbol{V}_{\boldsymbol{Y}}^{T} \otimes \left((\boldsymbol{\Sigma}_{\boldsymbol{Y}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2} \boldsymbol{U}_{\boldsymbol{Y}_{k}}^{T} \right) \right) \boldsymbol{x} \right\|^{2} = \left\| \left(\boldsymbol{I} \otimes (\boldsymbol{\Sigma}_{\boldsymbol{Y}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2} \right) \left(\boldsymbol{V}_{\boldsymbol{Y}}^{T} \otimes \boldsymbol{U}_{\boldsymbol{Y}_{k}}^{T} \right) \boldsymbol{x} \right\|^{2}.$$

322
$$= \left\| \left(\boldsymbol{V}_{\boldsymbol{X}_{k}}^{T} \otimes \left((\boldsymbol{\Sigma}_{\boldsymbol{X}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2} \boldsymbol{U}_{\boldsymbol{X}_{k}}^{T} \right) \right) \boldsymbol{x} \right\|_{2}^{2} = \left\| \left(\boldsymbol{I} \otimes (\boldsymbol{\Sigma}_{\boldsymbol{X}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2} \right) \left(\boldsymbol{V}_{\boldsymbol{X}_{k}}^{T} \otimes \boldsymbol{U}_{\boldsymbol{X}_{k}}^{T} \right) \boldsymbol{x} \right\|_{2}^{2}$$

323 Problem (3.4) is therefore equivalent to

324 (3.5)
$$\boldsymbol{x}_{k+1} = \arg\min_{\boldsymbol{x}} \|\boldsymbol{A}\boldsymbol{x} - \boldsymbol{b}\|_{2}^{2} + \widehat{\lambda}\| \underbrace{\left(\boldsymbol{I} \otimes (\boldsymbol{\Sigma}_{\boldsymbol{X}_{k}}^{2} + \gamma \boldsymbol{I})^{p/4-1/2}\right)}_{=:(\boldsymbol{W}_{p}^{\gamma})_{k}} \underbrace{\left(\boldsymbol{V}_{\boldsymbol{X}_{k}}^{T} \otimes \boldsymbol{U}_{\boldsymbol{X}_{k}}^{T}\right)}_{=:(\boldsymbol{W}_{p}^{\gamma})_{k}} \boldsymbol{x}\|_{2}^{2}.$$

In the above formulation, $(\boldsymbol{W}_p^{\gamma})_k$ is a diagonal weighting matrix and \boldsymbol{S}_k is an orthogonal matrix; both $(\boldsymbol{W}_p^{\gamma})_k$ and \boldsymbol{S}_k depend on the current approximation \boldsymbol{x}_k of the solution \boldsymbol{x} . Intuitively, the

matrix S_k maps x into the "singular value domain" of X_k (and acts as an iteration-dependent sparsity transform), and the matrix $(W_p^{\gamma})_k$ assigns suitable weights that allow to approximate a p-norm of the singular values. Therefore, the penalization term in (3.5) can be interpreted as a reweighted vectorial 2-norm, with respect to a transformation of the solution x. For this reason, the proposed approach is dubbed "IRN-NNRp" and is summarized in Algorithm 3.1.

Algorithm 3.1 IRN-NNRp

1: Inputs: $\boldsymbol{A}, \boldsymbol{b}, (\boldsymbol{W}_p^{\gamma})_0 = \boldsymbol{I}, \boldsymbol{S}_0 = \boldsymbol{I}$ 2: for $k = 0, 1, \ldots$ until a stopping criterion is satisfied do 3: Solve problem (3.5) 4: "Decrease" γ 5: Update $(\boldsymbol{W}_p^{\gamma})_{k+1}$ and \boldsymbol{S}_{k+1} 6: end for

The next subsection derives new strategies for the efficient solution of the sequence of subproblems (3.5) appearing in Algorithm 3.1.

334 3.2. Solution of problem (3.5) via Krylov methods. First rewrite problem (3.5) using **335** an appropriate change of variable as

336 (3.6)
$$\widehat{\boldsymbol{x}}_{k+1} = \arg\min_{\widehat{\boldsymbol{x}}} \|\boldsymbol{A}\boldsymbol{S}_k^T(\boldsymbol{W}_p^{\gamma})_k^{-1}\widehat{\boldsymbol{x}} - \boldsymbol{b}\|_2^2 + \widehat{\lambda}\|\widehat{\boldsymbol{x}}\|_2^2, \text{ with } \widehat{\boldsymbol{x}} = (\boldsymbol{W}_p^{\gamma})_k \boldsymbol{S}_k \boldsymbol{x}.$$

337 Note that

338 (3.7)
$$\boldsymbol{S}_{k}^{T} = \boldsymbol{S}_{k}^{-1} = \boldsymbol{V}_{\boldsymbol{X}_{k}} \otimes \boldsymbol{U}_{\boldsymbol{X}_{k}} \quad \text{and} \quad (\boldsymbol{W}_{p}^{\gamma})_{k}^{-1} = \boldsymbol{I} \otimes (\boldsymbol{\Sigma}_{\boldsymbol{X}_{k}}^{2} + \gamma \boldsymbol{I})^{1/2 - p/4},$$

so that the above transformations (inversion of an orthogonal and a diagonal matrix) are numeri-339 340 cally affordable by exploiting properties of Kronecker products. The Tikhonov-regularized problem (3.6) in standard form is equivalent to the Tikhonov-regularized problem (3.5) in general form. 341 Many Krylov subspace methods based on the Golub-Kahan Bidiagonalization (GKB) or Arnoldi 342 algorithms can be employed to approximate the solution of (3.6). Moreover, if the regularization 343 parameter $\hat{\lambda}$ is not known a priori, many efficient strategies to set its value adaptively within the 344 sequence of projected problems can be used (i.e., in the framework of hybrid methods; see [18, 8]). The matrices S_k and $(W_p^{\gamma})_k^{-1}$ can be formally thought of as preconditioners for the original problem 346 (1.1), whose purpose is to enforce additional regularization into the solution subspace, rather than 347 348 speeding-up the convergence of linear solvers applied to (1.1).

349 Methods based on the GKB algorithm. The mth step of the GKB algorithm applied to the 350 matrix $AS_k^T(W_p^{\gamma})_k^{-1}$ with starting vector **b** (i.e., taking $\boldsymbol{x}_0 = \boldsymbol{0}$) can be expressed by the following 351 partial matrix factorizations

352 (3.8)
$$(AS_k^T(W_p^{\gamma})_k^{-1})V_m = U_{m+1}\bar{B}_m \text{ and } ((W_p^{\gamma})_k^{-1}S_kA^T)U_{m+1} = V_{m+1}B_{m+1}^T$$

where $U_j \in \mathbb{R}^{M \times j}$ and $V_j \in \mathbb{R}^{N \times j}$ (with j = m, m + 1 and $U_j e_1 = b/||b||_2$) have orthonormal columns, and $B_{m+1} \in \mathbb{R}^{(m+1) \times (m+1)}$ is lower bidiagonal (with \overline{B}_m obtained by removing the last column of B_{m+1}). The orthonormal columns of V_m are such that

356
$$\mathcal{R}(\boldsymbol{V}_m) = \mathcal{K}_m\left(((\boldsymbol{W}_p^{\gamma})_k^{-1}\boldsymbol{S}_k\boldsymbol{A}^T)(\boldsymbol{A}\boldsymbol{S}_k^T(\boldsymbol{W}_p^{\gamma})_k^{-1}), ((\boldsymbol{W}_p^{\gamma})_k^{-1}\boldsymbol{S}_k\boldsymbol{A}^T)\boldsymbol{b}\right).$$

This manuscript is for review purposes only.

We find an approximate solution of (3.6) by imposing $\hat{\boldsymbol{x}} \in \mathcal{R}(\boldsymbol{V}_m)$, i.e., $\hat{\boldsymbol{x}}_m = \boldsymbol{V}_m \boldsymbol{y}_m$, where, by exploiting the first decomposition in (3.8) and the properties of the matrices appearing therein, $\boldsymbol{y}_m \in \mathbb{R}^m$ is such that

360 (3.9)
$$\boldsymbol{y}_m = \arg\min_{\boldsymbol{y} \in \mathbb{R}^m} \|\bar{\boldsymbol{B}}_m \boldsymbol{y} - \|\boldsymbol{b}\|_2 \boldsymbol{e}_1 \|_2^2 + \widehat{\lambda}_m \|\boldsymbol{y}\|_2^2.$$

We used the notation $\hat{\lambda}_m$ for the regularization parameter to highlight that its value can be adaptively set within the iterations. The approximate solution to problem (3.5) is such that

363 (3.10)
$$\boldsymbol{x} = \boldsymbol{S}_k^T (\boldsymbol{W}_p^{\gamma})_k^{-1} \hat{\boldsymbol{x}} \in \mathcal{K}_m \left((\boldsymbol{S}_k^T (\boldsymbol{W}_p^{\gamma})_k^{-2} \boldsymbol{S}_k) \boldsymbol{A}^T \boldsymbol{A}, (\boldsymbol{S}_k^T (\boldsymbol{W}_p^{\gamma})_k^{-2} \boldsymbol{S}_k) \boldsymbol{A}^T \boldsymbol{b} \right)$$

Looking at the above approximation subspace for the solution \boldsymbol{x} , it is evident that the "preconditioner" acts by first mapping into the "singular value domain" (by applying \boldsymbol{S}_k), enforcing sparsity in the singular values (by reweighting with $(\boldsymbol{W}_p^{\gamma})_k^{-2}$), and eventually transforming back into the "solution domain" (by applying \boldsymbol{S}_k^T).

Methods based on the Arnoldi algorithm. If \mathbf{A} is square, the *m*th step of the Arnoldi algorithm applied to the matrix $\mathbf{AS}_k^T(\mathbf{W}_p^{\gamma})_k^{-1}$ with starting vector \mathbf{b} (i.e., taking $\mathbf{x}_0 = \mathbf{0}$) can be expressed by the following partial matrix factorization

371 (3.11)
$$(AS_k^T(W_p^{\gamma})_k^{-1})V_m = V_{m+1}H_m$$

372 where $V_j \in \mathbb{R}^{N \times j}$ (with j = m, m + 1 and $V_j e_1 = b/\|b\|_2$) have orthonormal columns such that

373
$$\mathcal{R}(\boldsymbol{V}_m) = \mathcal{K}_m\left(\boldsymbol{A}\boldsymbol{S}_k^T(\boldsymbol{W}_p^{\gamma})_k^{-1}, \boldsymbol{b}\right),$$

and $\mathbf{H}_m \in \mathbb{R}^{(m+1) \times m}$ is upper Hessenberg. Similarly to the GKB case, we find an approximate solution of (3.6) by imposing $\hat{\mathbf{x}} \in \mathcal{R}(\mathbf{V}_m)$ and by solving a projected Tikhonov problem of order m. The approximate solution to problem (3.5) is such that

$$oldsymbol{x} = oldsymbol{S}_k^T (oldsymbol{W}_p^\gamma)_k^{-1} \widehat{oldsymbol{x}} \in oldsymbol{S}_k^T (oldsymbol{W}_p^\gamma)_k^{-1} \mathcal{K}_m \left(oldsymbol{A} oldsymbol{S}_k^T (oldsymbol{W}_p^\gamma)_k^{-1}, oldsymbol{b}
ight) \,,$$

374 where

375 $S_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\mathcal{K}_{m}\left(\boldsymbol{A}\boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1},\boldsymbol{b}\right) = \operatorname{span}\{\boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\boldsymbol{b},\ldots,\left(\boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\boldsymbol{A}\right)^{m-1}\boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\boldsymbol{b}\}$ 376 $= \mathcal{K}_{m}\left(\boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\boldsymbol{A},\boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\boldsymbol{b}\right).$

Contrarily to the GKB case, we immediately notice that, in this context, \boldsymbol{x} does not belong to a meaningful approximation subspace. Indeed, just by looking at the first vector: \boldsymbol{b} is in the image 378 space and $(\boldsymbol{W}_p^{\gamma})_k^{-1}$ is supposed to act on the singular value space of \boldsymbol{X}_k , so $(\boldsymbol{W}_p^{\gamma})_k^{-1}\boldsymbol{b}$ is hard to interpret; furthermore, \boldsymbol{S}_k^T is supposed to link the singular value space of \boldsymbol{X}_k to the image space, so $\boldsymbol{S}_k^T(\boldsymbol{W}_p^{\gamma})_k^{-1}\boldsymbol{b}$ is also hard for us to interpret. Although the generated solution subspace is not 379 380 381 meaningful for our applications, it may still have the potential to be a good subspace in other 382 contexts. Similarly to what is proposed in [1, 4], where the Arnoldi algorithm is applied to a 383 regularized problem that enforces sparsity in the wavelet domain, we propose to fix this issue by 384 incorporating S_k also as an orthogonal left "preconditioner" for the original system (1.1) so that, by 385 exploiting the invariance of the vectorial 2-norm under orthogonal transformations, problem (3.6) 386 can be equivalently reformulated as 387

388 (3.12)
$$\widehat{\boldsymbol{x}}_{k+1} = \arg\min_{\widehat{\boldsymbol{x}}} \|\boldsymbol{S}_k(\boldsymbol{A}\boldsymbol{S}_k^T(\boldsymbol{W}_p^{\gamma})_k^{-1}\widehat{\boldsymbol{x}} - \boldsymbol{b})\|_2^2 + \widehat{\lambda}\|\widehat{\boldsymbol{x}}\|_2^2, \text{ with } \widehat{\boldsymbol{x}} = (\boldsymbol{W}_p^{\gamma})_k \boldsymbol{S}_k \boldsymbol{x}.$$

The (right and left) preconditioned Arnoldi algorithm applied to problem (3.12) can now be expressed by the following partial matrix factorization

391 (3.13)
$$(S_k A S_k^T (W_p^{\gamma})_k^{-1}) V_m = V_{m+1} H_m.$$

We find an approximate solution of (3.12) by imposing $\hat{\boldsymbol{x}} \in \mathcal{R}(\boldsymbol{V}_m) = \mathcal{K}_m(\boldsymbol{S}_k \boldsymbol{A} \boldsymbol{S}_k^T \boldsymbol{W}^{-1}, \boldsymbol{S}_k \boldsymbol{b})$, i.e., $\hat{\boldsymbol{x}}_m = \boldsymbol{V}_m \boldsymbol{y}_m$, where, by exploiting (3.13) and the properties of the matrices appearing therein, $\boldsymbol{y}_m \in \mathbb{R}^m$ is such that

395 (3.14)
$$\boldsymbol{y}_m = \arg\min_{\boldsymbol{y}\in\mathbb{R}^m} \|\boldsymbol{H}_m\boldsymbol{y} - \|\boldsymbol{b}\|_2 \boldsymbol{e}_1\|_2^2 + \widehat{\lambda}_m \|\boldsymbol{y}\|_2^2$$

396 Hence

408

397 (3.15)
$$\boldsymbol{x} \in \boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1} \mathcal{K}_{m}(\boldsymbol{S}_{k}\boldsymbol{A}\boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}, \boldsymbol{S}_{k}\boldsymbol{b}) = \mathcal{K}_{m}(\boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\boldsymbol{S}_{k}\boldsymbol{A}, \boldsymbol{S}_{k}^{T}(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\boldsymbol{S}_{k}\boldsymbol{b}) ,$$

which is suitable for approximating the solution. The new methods based on the GKB algorithm (for generic matrices) and Arnoldi algorithm (only if $\mathbf{A} \in \mathbb{R}^{N \times N}$) are dubbed "IRN-LSQR-NNRp" and "IRN-GMRES-NNRp", respectively, and are summarized in Algorithm 3.2.

Algorithm 3.2 IRN-LSQR-NNRp and IRN-GMRES-NNRp 1: Inputs: $\boldsymbol{A}, \boldsymbol{b}, (\boldsymbol{W}_{p}^{\gamma})_{0} = \boldsymbol{I}, \boldsymbol{S}_{0} = \boldsymbol{I}$ 2: for $k = 0, 1, \ldots$ until a stopping criterion is satisfied do for $m = 1, 2, \ldots$ until a stopping criterion is satisfied do 3: Update the factorizations (3.8) and (3.13), respectively 4: Solve the projected problem (3.9) and (3.14), respectively, tuning $\widehat{\lambda}_m$ if necessary 5:6: end for "Decrease" γ 7: Update the new $(\boldsymbol{W}_{p}^{\gamma})_{k+1}$ and \boldsymbol{S}_{k+1} 8: 9: end for

3.3. Solution through flexible Krylov subspaces. Problem (1.3) reformulated as (3.6) allows us to naturally apply the flexible Golub-Kahan (FGK) and flexible Arnoldi algorithms. Indeed, instead of updating the "preconditioners" S_k and $(W_p^{\gamma})_k$ at the *k*th outer iteration of the nested iteration schemes of Algorithm 3.2, we propose to consider new "preconditioners" as soon as a new approximation of the solution is available, i.e., at each iteration of a Krylov subspace solver. Therefore, at the (i + 1)th iteration of the new solvers, the "preconditioners" $(W_p^{\gamma})_i$ and S_i are computed as in (3.7), but using the SVD of the *i*th approximate solution

$$\boldsymbol{X}_i = \operatorname{vec}^{-1}(\boldsymbol{x}_i) = \boldsymbol{U}_{\boldsymbol{X}_i} \boldsymbol{\Sigma}_{\boldsymbol{X}_i} \boldsymbol{V}_{\boldsymbol{X}_i}^T, \quad \text{for } i = 1, \dots, k-1,$$

with $(W_p^{\gamma})_0 = I$ and $S_0 = I$. In order to incorporate iteration-dependent preconditioning, the flexible versions of the Golub-Kahan and Arnoldi factorizations have to be used.

⁴¹¹ Namely, at the *i*th iteration, the new instance of the FGK algorithm updates partial factoriza-⁴¹² tions of the form (2.8), i.e., $AZ_i = U_{i+1}M_i$ and $A^TU_{i+1} = V_{i+1}T_{i+1}$, where

413
$$\boldsymbol{Z}_{i} = \left[\boldsymbol{S}_{0}^{T}(\boldsymbol{W}_{p}^{\gamma})_{0}^{-2}\boldsymbol{S}_{0}\boldsymbol{v}_{1}, \dots, \boldsymbol{S}_{i-1}^{T}(\boldsymbol{W}_{p}^{\gamma})_{i-1}^{-2}\boldsymbol{S}_{i-1}\boldsymbol{v}_{i}\right], \quad \boldsymbol{v}_{1} = \boldsymbol{A}^{T}\boldsymbol{b}/\|\boldsymbol{A}^{T}\boldsymbol{b}\|_{2}.$$

LOW-RANK KRYLOV METHODS

414 Taking $x_0 = 0$, the *i*th approximate solution is such that $x_i = Z_i y_i$, where

415 (3.16)
$$\boldsymbol{y}_i = \arg\min_{\boldsymbol{y}\in\mathbb{R}^i} \|\boldsymbol{M}_i\boldsymbol{y} - \|\boldsymbol{b}\|_2 \boldsymbol{e}_1\|_2^2 + \widehat{\lambda}_i \|\boldsymbol{y}\|_2^2$$

416 Note that the subspace for the solution $\mathcal{R}(\mathbf{Z}_i)$ can be regarded as a generalization of the subspace

(3.10) computed when considering preconditioned GKB within the IRN-LSQR-NNRp method. The new method is dubbed "FLSQR-NNRp", and is summarized in Algorithm 3.3.

419 For $\mathbf{A} \in \mathbb{R}^{N \times N}$ and $\mathbf{x}_0 = \mathbf{0}$, at the *i*th iteration, the new instance of the flexible Arnoldi 420 algorithm updates a partial factorization of the form (2.7), with k = i, and generates

421
$$\boldsymbol{Z}_{i} = [\boldsymbol{S}_{0}^{T}(\boldsymbol{W}_{p}^{\gamma})_{0}^{-1}\boldsymbol{S}_{0}\boldsymbol{v}_{1}, \dots, \boldsymbol{S}_{i-1}^{T}(\boldsymbol{W}_{p}^{\gamma})_{i-1}^{-1}\boldsymbol{S}_{i-1}\boldsymbol{v}_{i}], \quad \boldsymbol{v}_{1} = \boldsymbol{b}/\|\boldsymbol{b}\|_{2},$$

where both right and left preconditioners are used analogously to IRN-GMRES-NNR*p*. The *i*th approximate solution is such that $x_i = Z_i y_i$, where

424 (3.17)
$$\boldsymbol{y}_i = \arg\min_{\boldsymbol{y}\in\mathbb{R}^i} \|\boldsymbol{H}_i\boldsymbol{y} - \|\boldsymbol{b}\|_2 \boldsymbol{e}_1\|_2^2 + \widehat{\lambda}_i \|\boldsymbol{y}\|_2^2$$

425 Note that the subspace for the solution $\mathcal{R}(\mathbf{Z}_i)$ can be regarded as a generalization of the subspace

(3.15) computed when considering the preconditioned Arnoldi algorithm within the IRN-GMRES-

NNRp method. The new method is dubbed "FGMRES-NNRp", and is summarized in Algorithm 3.3.

Algorithm 3.3 FLSQR-NNRp and FGMRES-NNRp

1: Inputs: $\boldsymbol{A}, \, \boldsymbol{b}, \, (\boldsymbol{W}_p^{\gamma})_0 = \boldsymbol{I}, \, \boldsymbol{S}_0 = \boldsymbol{I}$

2: for i = 1, 2, ... until a stopping criterion is satisfied do

- 3: Update a factorization of the form (2.8) and (2.7), respectively, to expand the space $\mathcal{R}(\mathbf{Z}_i)$
- 4: Solve the projected problem (3.16) and (3.17), respectively, tuning $\hat{\lambda}_i$ if necessary
- 5: "Decrease" γ
- 6: Update the new $(\boldsymbol{W}_{p}^{\gamma})_{i}$ and \boldsymbol{S}_{i} , using the SVD $\boldsymbol{X}_{i} = \operatorname{vec}^{-1}(\boldsymbol{x}_{i}) = \boldsymbol{U}_{\boldsymbol{X}_{i}}\boldsymbol{\Sigma}_{\boldsymbol{X}_{i}}\boldsymbol{V}_{\boldsymbol{X}_{i}}^{T}$.
- 7: end for

Note that, although the approach of Algorithm 3.3 is quite heuristic, it avoids nested iteration cycles and computes only one approximation subspace for the solution of (1.3), where low-rank penalization is adaptively incorporated. Because of this, in many situations, Algorithm 3.3 computes solutions of quality comparable to the ones computed by Algorithm 3.2, with a significant reduction in the number of iterations. We should also mention that, in the framework of affine rank minimization problems, [25] outlines an algorithm that avoids inner projected gradient iterations for the solution of each quadratic subproblem in the sequence generated within the IRN strategy.

Finally, we underline that, within the framework of flexible Krylov subspaces, the approximation 436 437 subspaces $\mathcal{R}(\mathbf{Z}_i)$ for the *i*th approximate solution can be further modified, with some insight into the desired properties of the solution. Indeed, since the ith basis vector for the solution is of the form 438 $\boldsymbol{z}_i = \boldsymbol{S}_{i-1}^T (\boldsymbol{W}_p^{\gamma})_{i-1}^{-2} \boldsymbol{S}_{i-1} \boldsymbol{v}_i$ for FLSQR-NNRp, and $\boldsymbol{z}_i = \boldsymbol{S}_{i-1}^T (\boldsymbol{W}_p^{\gamma})_{i-1}^{-1} \boldsymbol{S}_{i-1} \boldsymbol{v}_i$ for FGMRES-NNRp, 439 one can consider alternative "preconditioners" S_{i-1} and $(W_p^{\gamma})_{i-1}$ that are still effective in delivering 440low-rank solutions. For instance, focusing on FGMRES, and given $v_i = V_i e_i$, where V_i is the matrix 441 appearing on the right-hand side of the factorization (2.8), and given the SVD of $vec^{-1}(v_i) =$ 442443 $U_{V_i} \Sigma_{V_i} V_{V_i}^T$, one can take

444 (3.18)
$$S_{i-1} = V_{V_i}^T \otimes U_{V_i}^T$$
 and $(W_p^{\gamma})_{i-1}^{-1} = I \otimes (\Sigma_{V_i})^{1-p/2}$,

and as a result, 445

14

446
447

$$\mathbf{S}_{i-1}\boldsymbol{v}_{i} = \operatorname{vec}(\boldsymbol{U}_{\boldsymbol{V}_{i}}^{T}\operatorname{vec}^{-1}(\boldsymbol{v}_{i})\boldsymbol{V}_{\boldsymbol{V}_{i}}) = \operatorname{vec}(\boldsymbol{\Sigma}_{\boldsymbol{V}_{i}}),$$
447

$$(\boldsymbol{W}_{r}^{\gamma})_{i=1}^{-1}\boldsymbol{S}_{i-1}\boldsymbol{v}_{i} = \operatorname{vec}((\boldsymbol{\Sigma}_{\boldsymbol{V}_{i}})^{1-p/2}\boldsymbol{\Sigma}_{\boldsymbol{V}_{i}}) = \operatorname{vec}((\boldsymbol{\Sigma}_{\boldsymbol{V}_{i}})^{2-p/2}),$$

447
$$(\boldsymbol{W}_{p}^{\gamma})_{i=1}^{-1}\boldsymbol{S}_{i-1}\boldsymbol{v}_{i} = \operatorname{vec}((\boldsymbol{\Sigma}_{\boldsymbol{V}_{i}})^{1-p/2}\boldsymbol{\Sigma}_{\boldsymbol{V}_{i}}) = \operatorname{vec}((\boldsymbol{\Sigma}_{i})^{1-p/2}\boldsymbol{\Sigma}_{i})$$

448
$$\boldsymbol{S}_{i-1}^{T}(\boldsymbol{W}_{p}^{\gamma})_{i-1}^{-1}\boldsymbol{S}_{i-1}\boldsymbol{v}_{i} = \operatorname{vec}(\boldsymbol{U}_{\boldsymbol{V}_{i}}((\boldsymbol{\Sigma}_{\boldsymbol{V}_{i}})^{2-p/2})\boldsymbol{V}_{\boldsymbol{V}_{i}}^{T}) = \boldsymbol{z}_{i},$$

so that the singular values of vec⁻¹(v_i) are rescaled: taking $0 , the power of <math>\Sigma_{V_i}$, 2 - p/2, 449 is always larger than 1, which means that large singular values get magnified and small singular 450values become even smaller. In this way, the gaps between singular values are emphasized and to 451some extent contribute to the low rank properties of the basis vectors. Similar derivations hold for 452FLSQR. Hence, methods analogous to LR-FLSQR and LR-FGMRES are obtained, and are dubbed 453FGMRES-NNRp(v) and FLSQR-NNRp(v), respectively. 454

4. Implementation details. All the methods considered in this paper are iterative, and 455therefore at least one suitable stopping criterion should be set for the iterations. When considering 456hybrid formulations (like the ones in Algorithms 3.2 and 3.3), one could simultaneously set a good 457value for the regularization parameter $\hat{\lambda}_i$ at the *j*th iteration, as well as properly stop the iterations. 458Strategies for achieving this are already available in the literature (see [6, 8]). 459

Assuming that a good estimate for the norm of the noise η affecting the right-hand-side of (1.1) 460is available, i.e., $\varepsilon \simeq \|\boldsymbol{\eta}\|_2$, one can consider the discrepancy principle and stop the iterative scheme 461 at the first iteration j such that 462

463 (4.1)
$$\|\boldsymbol{b} - \boldsymbol{A}\boldsymbol{x}_j\|_2 \leq \theta \varepsilon$$
, where $\theta > 1, \theta \simeq 1$ is a safety threshold.

Applying the discrepancy principle to LR-FGMRES (Algorithm 2.1) and LR-FLSQR (Algorithm 464(2.2) is particularly convenient, as the norm of the residual on the left-hand side of (4.1) can be 465monitored using projected quantities, i.e., 466

467
$$\|\|\boldsymbol{b}\|_2 \boldsymbol{e}_1 - \boldsymbol{H}_j \boldsymbol{y}_j\|_2 \quad \text{for LR-FGMRES and} \quad \|\|\boldsymbol{b}\|_2 \boldsymbol{e}_1 - \boldsymbol{M}_j \boldsymbol{y}_j\|_2 \quad \text{for LR-FLSQR},$$

where decompositions (2.7) and (2.8), respectively, and the properties of the matrices appearing 468 therein, have been exploited. When running hybrid methods (see Algorithms 3.2 and 3.3), we 469employ the so-called "secant method", which updates the regularization parameter for the projected 470 471 problem in such a way that stopping by the discrepancy principle is ensured. We highlight again that the quantities needed to implement the "secant method" (namely, the norm of the residual and 472the discrepancy associated to (3.6) at each iteration) can be conveniently monitored using projected 473 quantities: this is obvious for IRN-LSQR-NNRp and FLSQR-NNRp, as only right-"preconditioning" 474is employed; it is less obvious for IRN-GMRES-NNRp and FGMRES-NNRp, but since the left-475"preconditioner" is orthogonal, one can still write 476

477
$$\|\boldsymbol{b} - \boldsymbol{A}\boldsymbol{x}_j\|_2 = \|\boldsymbol{S}_k \boldsymbol{b} - \boldsymbol{S}_k \boldsymbol{A} \boldsymbol{S}_k^T (\boldsymbol{W}_p^{\gamma})_k^{-1} \hat{\boldsymbol{x}}_j\|_2 = \|\|\boldsymbol{b}\|_2 \boldsymbol{e}_1 - \boldsymbol{H}_j \boldsymbol{y}_j\|_2$$

Note that all the methods in Algorithm 3.2 and 3.3 can also run with $\hat{\lambda} = 0$, and still achieve 478 low-rank approximate solutions: this is because the approximation subspace for the solution in-479corporates regularizing "preconditioning" (see [12, 14] for details on this approach in the case of 480 smoothing "preconditioning" with finite-difference approximations of derivatives operators). Fi-481 nally, when dealing with the inner-outer iteration scheme of Algorithm 3.2, in addition to a pa-482rameter choice strategy and stopping criterion for the hybrid projected problems (3.9) and (3.14), 483

one should also consider a stopping criterion for the outer iterations. We propose to do this by monitoring the norm of the difference of the singular values (normalized by the largest singular value so that $\sigma_1(\Sigma_{\mathbf{X}_{k+1}}) = \sigma_1(\Sigma_{\mathbf{X}_k}) = 1$) of two approximations of the solution of (1.3) obtained at two consecutive outer iterations of Algorithm 3.2, i.e., we stop as soon as

488 (4.2)
$$\|\operatorname{diag}(\boldsymbol{\Sigma}_{\boldsymbol{X}_{k+1}}) - \operatorname{diag}(\boldsymbol{\Sigma}_{\boldsymbol{X}_k})\|_2 < \tau_{\sigma}, \quad k = 1, 2, \dots,$$

where $\operatorname{vec}^{-1}(\boldsymbol{x}_i) = \boldsymbol{X}_i = \boldsymbol{U}_{\boldsymbol{X}_i} \boldsymbol{\Sigma}_{\boldsymbol{X}_i} \boldsymbol{V}_{\boldsymbol{X}_i}^T$ (i = k, k + 1), and $\tau_{\sigma} > 0$ is a user-specified threshold. If no significant changes happen in the rank and singular values of two consecutive approximations of the solution, then (4.2) is satisfied.

We conclude this section with a few remarks about the computational cost of the proposed 492methods. Note that, if $\mathbf{A} \in \mathbb{R}^{N \times N}$, IRN-GMRES-NNRp is intrinsically cheaper than IRN-LSQR-493 NNRp (since, at each iteration, the former requires only one matrix-vector product with A, while 494the latter requires one matrix-vector product with A and one with A^{T}). However, methods based 495on the Arnoldi algorithm are typically less successful than methods based on the GKB algorithm 496for regularization; see [15]. Other key operations for implementing our proposed methods are the 497computation of the SVDs of relevant quantities, and/or the application of the "preconditioners" in 498(3.18). Namely, each iteration of LR-FGMRES, LR-FLSQR, FLSQR-NNRp, and FGMRES-NNRp 499requires the computation of the SVD of an $n \times n$ matrix, which amounts to $\mathcal{O}(n^3) = \mathcal{O}(N^{3/2})$ 500 floating point operations. When considering IRN-LSQR-NNRp and IRN-GMRES-NNRp, only the 501SVD of the approximate solution should be computed once at each outer iteration. However, 502each inner iteration of IRN-LSQR-NNRp and IRN-GMRES-NNRp, as well as each iteration of 503FLSQR-NNRp and FGMRES-NNRp, requires the computation of matrix-vector products of the 504form $S_k^T (W_p^{\gamma})_k^{-1} v_i$: this can be achieved within a two-step process, where first the rescaling $\tilde{v}_i =$ 505 $(\boldsymbol{W}_{p}^{\gamma})_{k}^{-1}\boldsymbol{v}_{i}$ is applied with $\mathcal{O}(N) = \mathcal{O}(n^{2})$ floating-point operations, and then $\boldsymbol{S}_{k}^{T}\tilde{\boldsymbol{v}}_{i} = (\boldsymbol{V}_{\boldsymbol{X}_{k}}\otimes \boldsymbol{U}_{\boldsymbol{X}_{k}})\tilde{\boldsymbol{v}}_{i}$ is computed. While a straightforward implementation of the latter would require $\mathcal{O}(N^{2}) = \mathcal{O}(n^{4})$ floating-point operations, exploiting Kronecker product properties can bring down the cost of 506507508 this operation to $\mathcal{O}(n^3) = \mathcal{O}(N^{3/2})$, by computing $S_k^T \widetilde{v}_i = \text{vec}(U_{X_k}^T \text{vec}^{-1}(v_i)V_{X_k})$. We emphasize 509 that the incorporation of the flexible "preconditioners" does not increase the order of computational 510complexity and is very practical, since operations are done on matrices of size $n \times n$ (n is the 511dimension of the image). In particular, the full SVD's of $n \times n$ matrices can be computed easily 512with MATLAB's built-in svd function (this is what we used in our numerical experiments); one 513can also use Lanczos bidiagonalization [20] or randomized SVD [11] to compute the approximate 514leading singular values and vectors.

5. Experimental Results. In this section, we present results of numerical experiments on 5. several image processing problems to demonstrate the performance of the new IRN-GMRES-NNRp, 5.18 IRN-LSQR-NNRp, FGMRES-NNRp, and FLSQR-NNRp methods. Variants of FGMRES-NNRp5.19 and FLSQR-NNRp (marked with "(v)") are also tested. To shorten the acronyms, we omit p when 5.20 p = 1, which means IRN-GMRES-NNR denotes IRN-GMRES-NNRp when p = 1, etc. Examples 5.21 are generated using *IR Tools* [6].

In general, we compare the performances of the proposed methods to standard Krylov subspace methods GMRES and LSQR, also used in a hybrid fashion. We also test against the low-rank projection methods described in Section 2 and the singular value thresholding (SVT) algorithm [3], which was originally proposed for low-rank matrix completion problems, and can be extended to problems with linear constraints of the form

527 (5.1)
$$\min_{\boldsymbol{x}} \tau \| \operatorname{vec}^{-1}(\boldsymbol{x}) \|_{\ast} + \frac{1}{2} \| \operatorname{vec}^{-1}(\boldsymbol{x}) \|_{F}^{2}$$
 subject to $A\boldsymbol{x} = \boldsymbol{b}$, where $\operatorname{vec}^{-1}(\boldsymbol{x}) = \boldsymbol{X}$.

528 The kth iteration of the SVT algorithm for (5.1) reads

529 (5.2)
$$\begin{cases} \boldsymbol{X}_k = \mathcal{D}_{\tau}(\boldsymbol{A}^T \boldsymbol{y}_{k-1}) \\ \boldsymbol{y}_k = \boldsymbol{y}_{k-1} + \delta_k (\boldsymbol{b} - A \boldsymbol{x}_k) \end{cases}$$

530 where δ_k is a step size and \mathcal{D}_{τ} is the singular value shrinkage operator, defined as

31
$$\mathcal{D}_{\tau}(\boldsymbol{X}) = \boldsymbol{U}_{\boldsymbol{X}} \mathcal{D}_{\tau}(\boldsymbol{\Sigma}_{\boldsymbol{X}}) \boldsymbol{V}_{\boldsymbol{X}}^{T}, \quad \mathcal{D}_{\tau}(\boldsymbol{\Sigma}_{\boldsymbol{X}}) = \max\{\boldsymbol{\Sigma}_{\boldsymbol{X}} - \tau \boldsymbol{I}, \boldsymbol{0}\},\$$

532 where $X = U_X \Sigma_X V_X^T$ is the SVD of X, **0** is a matrix of zeros, and the maximum is taken 533 component-wise. Although (5.1) is not the same problem as (1.2), they are similar in that both 534 penalize the nuclear norm of $\operatorname{vec}^{-1}(x)$ and they respect the constraint Ax = b.

The Schatten-*p* function is introduced in Section 3.1 as a smooth approximation for $\|\cdot\|_{*,p}$. The smooth approximation allows for further derivations including computation of optimality conditions, where the "smoothing coefficient" γ is crucial. However, γ is not so crucial numerically, and we can set it to 0 without affecting the results (compared to using a very small γ). However, to be consistent with Algorithms 3.2 and 3.3, in our experiments, we have set the initial value of γ to 10^{-10} , and every time we need to decrease γ , we divide the current γ value by 2.

Regarding the comparisons with the low-rank projection methods presented in Section 2, there are no universal and theoretically informed ways of choosing the truncation ranks for the solutions and for the basis vectors of the solution subspace. Hence, for all test problems, we experiment on a reasonable number of trials, each with different truncation rank choices, and select the best performing rank out of all ranks tested. For simplicity, we consider the same truncation rank for basis vectors and solutions ($\tau_{\kappa_B} = \tau_{\kappa}$). We follow the same process to choose the number of restarts and the number of iterations for each restart for RS-LR-GMRES, as well as the shrinkage threshold τ in SVT; strategies to select the step size for SVT are described in [3].

Example 1: Binary Star. We consider an image deblurring problem involving a binary star test 549image of size 256×256 : this test image has rank 2. The true image is displayed in the leftmost 550frame of Figure 2. A standard Gaussian blur is applied to the test image, and Gaussian white 551noise of level $\|\boldsymbol{\eta}\|_2 / \|\boldsymbol{b}^{\text{ex}}\|_2 = 10^{-3}$ is added. The blurred and noisy image is shown in Figure 2, 552second frame from the left. Due to the presence of noise, the blurred image has full rank. For this 553 example, the blurring operator A is square of size 65536 \times 65536, hence GMRES-related methods 554are used for comparison, namely: GMRES, IRN-GMRES-NNR, FGMRES-NNR, LR-FGMRES and 555RS-LR-GMRES (i.e., we only consider the case p = 1 here). SVT is also taken into consideration. 556The truncation rank for LR-FGMRES and RS-LR-GMRES is set to 30 for both basis vectors and 557approximate solutions (i.e., $\tau_{\kappa_B} = \tau_{\kappa} = 30$). RS-LR-GMRES is restarted every 40 iterations. The step size for SVT is set to be $\delta_k = \delta = 2$ and the singular value shrinkage threshold τ is 1. Note 558559that, although the true solution has only rank 2, setting truncation rank to 2 for low rank methods 560 produces solutions of worse quality (compared to setting the rank to 30). This might be because 561 of the inherent ill-posedness of the problem, which makes it harder to obtain solutions with desired 562563 properties (e.g., with rank 2): indeed, if we do truncate to rank 2, a lot of information about the solution might be lost. 564

Figure 1 displays the histories of relative errors $\|\boldsymbol{x}^{\text{ex}} - \boldsymbol{x}_m\|_2 / \|\boldsymbol{x}^{\text{ex}}\|_2$ for the first 200 iterations (i.e., m = 1, ..., 200) of these methods. For IRN-GMRES-NNR, 4 outer cycles were run, each with a maximum of 50 iterations: a new outer cycle is initiated as soon as the discrepancy principle is satisfied in the inner cycle. No additional regularization is used (i.e., $\hat{\lambda} = 0$ for all methods).

This manuscript is for review purposes only.



Fig. 1: Example 1. Relative errors vs. number of iterations for GMRES-based methods and SVT.

We can observe from Figure 1 that when the truncation ranks are chosen reasonably, LR-FGMRES and RS-LR-GMRES both produce a less pronounced semi-convergence behavior than GMRES, with LR-FGMRES attaining a smaller relative error than RS-LR-GMRES. FGMRES-NNR, on the other hand, shows slower semi-convergence than GMRES, but it also converges to a slightly better relative error. IRN-GMRES-NNR behaves especially well in this case, with significantly reduced relative errors even at the end of the second outer cycle. The "jumps" at the beginning of each outer IRN-GMRES-NNR iteration are due to the strategy used for restarts (the older basis vectors are cleared at each restart).

Figure 2 displays the exact and the corrupted images, as well as the best reconstructions computed by LR-FGMRES and IRN-GMRES-NNR: these are obtained at the 47th and the 189th (total) iteration of LR-FGMRES and IRN-GMRES-NNR, respectively. By looking at relative errors in Figure 1, we see that LR-FGMRES is the second best out of all methods, and yet the quality of the solution is inferior compared to IRN-GMRES-NNR. Compared to the LR-FGMRES solution, the IRN-GMRES-NNR one is a more truthful reconstruction of the exact image: it not only has less artifacts immediately around the stars, but also has less background noise, in the sense that the pixel intensities in the background are closer to the true ones (as it can be seen by looking





Fig. 2: *Example 1*. Exact and corrupted test images, together with the best reconstructions obtained by the LR-FGMRES and the IRN-GMRES-NNR methods.

S. GAZZOLA, C. MENG AND J. NAGY

pixels) of the computed images, as shown in Figure 3: here the best LR-FGMRES reconstruction, as well as the IRN-GMRES-NNR reconstructions at the end of the 2nd, 3rd and 4th inner cycles are displayed. It is clear that the IRN reconstructions are improving over each outer cycle, and that even the solution at the end of the 2nd cycle is significantly better than the LR-FGMRES solution, which means that not all four outer iterations need to be run to achieve solutions of superior qualities (even if more outer iterations allow further improvement in the solution). Figure



Fig. 3: *Example 1*. Zoom-ins of the LR-FGMRES best solution, and the IRN-GMRES-NNR solutions at the end of each inner cycle.

591

592 4 displays surfaces plots of the central part $(51 \times 51 \text{ pixels})$ of the test problem data, as well as the best reconstructed images (for RS-LR-GMRES and FGMRES-NNR these are obtained at the 165th and the 63th (total) iterations, respectively). It can be seen that for all the solutions shown here, the 594reconstructed central two stars approximately have the same intensity, although they are somewhat less intense than in the exact image. These surface plots also confirm our earlier observation that 596 IRN-GMRES-NNR does an exceptional job removing background noise. In addition, FGMRES-NNR also gives a good background reconstruction. Finally, Figure 5 displays the singular values 598 of the best solutions obtained adopting different GMRES-based solvers, as well as the evolution of 599 600 the singular values of the solution at the end of each inner IRN-GMRES-NNR cycle (matching the reconstructions displayed in Figure 3). The singular values are "normalized" (i.e., divided by the 601 largest one), and the graphs are cropped to focus on the relevant values. Looking at the displayed 602 values, we can conclude that the solutions computed by all the low-rank solvers have indeed some 603 low-rank properties, with very quickly-decaying large singular values followed by slowly-decaying 604 smaller singular values. Compared to GMRES, the new FGMRES-NNR and IRN-GMRES-NNR 605 methods give solutions that have a more pronounced low rank, as shown by the large gaps between 606 the smaller singular values of the solutions computed by these methods. Regarding IRN-GMRES-607 NNR, the evolution of the singular values stabilizes as we move toward later outer iterations, which 608 validates the stopping criterion proposed in Section 4. 609

Example 2: Limited angle parallel-ray tomography. We consider a computed tomography (CT) 610 test problem, modeling an undersampled X-ray scan with parallel beam geometry. This is a so 611 612 called "limited angle" CT reconstruction problem, where the viewing angles for the object span 613 less than 180 degrees. A smooth and rank-4 phantom is considered, as shown in the leftmost frame of Figure 7 (note that the yellow straight lines in the northwestern corner do not belong to the 614 phantom; they are shown for later purposes). Gaussian white noise of level 10^{-2} is added to the 615 data. The coefficient matrix A has size 32942×65536 . Because of this, among the new solvers, 616 only LR-FLSQR, FLSQR-NNRp, FLSQR-NNRp(v), and IRN-LSQR-NNRp will be tested, against 617



Fig. 4: *Example 1*. Zoomed-in surfaces of the exact solution and the available data, as well as the best reconstructions obtained by the new GMRES-based methods.



Fig. 5: *Example 1*. Left frame: normalized singular values of the best solutions computed by each GMRES-based method. Right frame: evolution of the singular values of the solutions computed by IRN-GMRES-NNR at each outer iteration. Singular values less than 10^{-3} are omitted.

- 618 their standard counterpart LSQR. Recall that FLSQR-NNRp(v) is the FLSQR-NNRp variant that
- 619 defines the preconditioners using the basis vectors of the solution subspace. The hybrid strategy 620 is not used here, meaning that we set $\hat{\lambda} = 0$ for all methods. For this test problem, we consider 621 both the values p = 1 and p = 0.75 (recall that, when p = 1, we omit p from the notation). The
- results obtained running the available low-rank solvers SVT and RS-LR-GMRES are shown, too. Note that RS-LR-GMRES only works for square matrices \boldsymbol{A} , hence this solver is tested on the
- normal equations $A^T A x = A^T b$, which is not the problem solved by the other methods (therefore

this comparison may not be completely fair). Parameters for SVT are chosen to be: step size $\delta_k = \delta = 8 \times 10^{-5}$ and threshold $\tau = 100$. RS-LR-GMRES is set to restart every 20 iterations. The truncation rank is 10 for both basis vectors and solutions, and for both the LR-FLSQR and the RS-LR-GMRES methods. The maximum number of iterations is 100 for all methods.

Figure 6 displays the history of the relative errors for LSQR, LR-LSQR, FLSQR-NNRp, FLSQR-NNRp(v), and IRN-LSQR-NNRp, for p = 1 and p = 0.75. Figure 7 displays the exact phantom together with the best reconstructions obtained by LSQR, FLSQR-NNRp(v), and IRN-LSQR-NNR. Figure 8 displays surface plots of the northwestern corner of the exact and reconstructed phantoms (64 × 64 pixels, as highlighted in the leftmost frame of Figure 7).

Looking at relative errors in Figure 6, it is obvious that the winners are the FLSQR-NNRp(v)634 methods, with both p = 1 and p = 0.75: they give the lowest relative errors, and the fastest semi-635 convergences. For this test problem, using a value of p < 1 lowers the relative error of FLSQR-636 NNRp(v); however, the same does not hold for IRN-LSQR-NNRp. Therefore we can conclude that 637 the the choice of p is problem and solver dependent, and using p < 1 does not necessarily improve 638 639 the quality of the solution. We regard p = 1 as a safe choice for this parameter. Although both the FLSQR-NNR $p(\mathbf{v})$ methods with p = 1 and p = 0.75 perform well, the latter is able to further 640 reduce the noise in the reconstructed solution, especially on the boundary. 641



Fig. 6: *Example 2.* Relative errors vs. number of iterations for different solvers. Upper frame: some of the new solvers are compared to the already available solvers. Lower frame: comparisons of different instances of the new solvers (here p = 0.75).



Fig. 7: Example 2. Exact phantom and best reconstructions obtained by different solvers.



Fig. 8: *Example 2*. Surface plots of the northwestern corner of the exact phantom (highlighted in Figure 7) and the best reconstructed phantoms computed by different solvers.

642 Looking at all the displayed results, the advantages of our new FLSQR-NNRp(v) and IRN-643 LSQR-NNR methods are evident. Namely, they produce smooth solutions that preserve the original 644 concave shape of the exact phantom, and they retain similar intensities of pixels at the same loca-645 tions of the exact phantom (although the LR-FLSQR solution is smooth within the boundary, it fails 646 to reconstruct intensity at the high point). Differences between FLSQR-NNRp(v) and IRN-LSQR-647 NNR reconstructions are clear, too: while both are smooth, the IRN-LSQR-NNR reconstruction 648 has a less concave shape compared to that of FLSQR-NNRp(v), but a smoother boundary.

Example 3: Inpainting. We consider two different inpainting test problems. Inpainting is the process of restoring images that have missing or deteriorated parts. These images are likely to

have quite a few lost pixels, either in the form of salt and pepper noise, or missing patches with 651 regular or irregular shapes. The two examples considered here are of different nature: the first one 652 has less structured and more randomly distributed missing patches, while the second one has more 653 structured and regularly shaped missing parts. The corrupted images (shown in top-middle frames 654of Figures 11 and 13) are constructed by first applying a blur operator, and then superimposing 655 the undersampling pattern to the ideally exact images (shown in the top-left frames of Figures 11 656 and 13). We follow this particular order of first blurring and then taking out pixels to simulate the 657 real process of photo-taking. For both these test problems, white noise of level 10^{-2} is added to 658 the data, and we consider purely iterative methods (i.e., $\hat{\lambda} = 0$). We always take p = 1, and we run 659 100 iterations of all the methods. 660

Firstly, we consider a test problem where 58.2% of the pixels are missing (following some 661 random and not very regular patterns). The exact image is commonly known as the house test 662 image, whose rank is 243 and has a total number of 65536 (256×256) pixels; the corrupted image 663 has the same size and number of pixels, but out of which only 27395 are non-zero. The singular 664 665 values of the exact image is shown in Figure 9(a). Correspondingly, the forward operator \boldsymbol{A} is of size 27395×65536 , so we have an underdetermined linear system: A is obtained by first applying 666 a shaking blur, and by then undersampling the blurred image. This can be easily coded within the 667 IR Tools framework. 668

Figure 10 displays the history of the relative errors for LSQR, LR-FLSQR (with truncation of the basis vectors for the solution, as well as the solution, to rank 20), FLSQR-NNR, FLSQR-NNR(v) and IRN-LSQR-NNR. Figure 11 displays the exact and corrupted images, together with the best reconstructions obtained by the methods listed above: these correspond to the 16th, 32nd, 67th, 30th and 62nd iterations of LSQR, LR-FLSQR, FLSQR-NNR, FLSQR-NNR(v) and IRN-LSQR-NNR, respectively (i.e., these are the iterations where the minimum relative error is attained over the total 100 iterations).



Fig. 9: *Example 3*. Singular values of exact test images house and peppers scaled by the largest singular values respectively.



Fig. 10: Example 3 (house). Relative errors vs. number of iterations for different solvers.



Fig. 11: *Example 3* (house). Exact and corrupted images; best reconstructions obtained by standard and new solvers.

Secondly, we consider a test problem similar to the previous one, i.e., we take an exact image 676 commonly known as the peppers test image, which has full rank (its singular values are shown 677 in Figure 9(b)), and we obtain the forward operator A by first applying a shaking blur, and 678 by then undersampling the blurred image. Here the exact image has a total number of 65536 679 (256×256) pixels, and only around 1.3% of pixels are missing and should be inpainted: differently 680 from the previous problem, the missing pixels follow particular patterns (e.g., circles, squares, and 681 rectangles), and this makes the inpainting task somewhat more challenging. Figure 12 displays the 682 history of the relative errors for LSQR, LR-FLSQR (with truncation of the basis vectors for the 683

684 solution, as well as the solution, to rank 50), FLSQR-NNR, FLSQR-NNR(v) and IRN-LSQR-NNR.

Figure 13 displays the exact and corrupted images, together with the best reconstructions obtained

by the methods listed above: these correspond to the 11th, 18th, 60th, 33rd and 34th iterations of

687 LSQR, LR-FLSQR, FLSQR-NNR, FLSQR-NNR(v) and IRN-LSQR-NNR, respectively (i.e., these

are the iterations where the minimum relative error is attained over the total 100 iterations).



Fig. 12: Example 3 (peppers). Relative errors vs. number of iterations for different solvers.



Fig. 13: *Example 3* (peppers). Exact and corrupted images; best reconstructions obtained by standard and new solvers.

It is evident that FLSQR-NNR(v) achieves reconstructions of superior quality, including clarity, brightness, and smoothness. Its ability to fill-in missing spots with pixels that are of similar intensity

to their surroundings is the best among all methods. The best reconstructions are computed by IRN-691 LSQR-NNR for the house test image, and by FLSQR-NNR for the pepper test image: in both cases, 692 these methods are also good at removing noise and restoring missing pixels. However, for both test 693 images, the reconstructions obtained by IRN-LSQR-NNR lack clarity compared to ones obtained 694 by both FLSQR-NNR and FLSQR-NNR(v) methods; compared to the reconstructions obtained by 695 LSQR and LR-FLSQR, they are anyway more desirable in terms of recovered brightness and fill-in of 696 the missing pixels. Moreover, we have seen in these two examples that our newly proposed methods 697 perform very well not only for low rank, but also for full or nearly full rank image reconstruction, 698 thanks to the regularizing properties of our newly derived "preconditioners" $(W_p^{\gamma})_k$ and S_k . Our 699 methods can also be extensively tested for higher noise levels (for example, 10^{-1}) and yield similar 700 results. However, for space considerations we are not able to show all of them here. 701

A study of regularization parameters. In the previous examples we have seen that the IRN-NNR 703 methods and the flexible Krylov NNR methods perform exceptionally well on image deblurring, tomography, and inpainting problems, producing superior reconstructions compared to existing 704 705 methods including SVT, RS-LR-GMRES and the low-rank flexible Krylov methods inspired by RS-LR-GMRES, even without the use of additional regularization. In this section, we explore 706 the effect of additional regularization (i.e., we set $\lambda \neq 0$) on the reconstructed images and the corresponding relative errors. In particular, additional regularization allows the new methods to be 708 used in a hybrid fashion. We are going to observe that there is only little to negligible room for the 709 methods to improve when they are used in a hybrid fashion (as their performance is already very 710 good with $\hat{\lambda} = 0$). 711

We consider three different ways of choosing the regularization parameter $\hat{\lambda}$. (i) We take the "secant method" mentioned in Section 4, which updates the regularization parameter at each iteration using the discrepancy. (ii) We select the optimal regularization parameter which minimizes the 2-norm of the difference between the exact solution and the regularized solution at each iteration. Namely, when using standard GMRES and LSQR, at the *m*th iteration we seek to minimize with respect to $\hat{\lambda}$

$$\|oldsymbol{x}^{ ext{ex}}-oldsymbol{x}_{m,\widehat{\lambda}}\|=\|oldsymbol{V}_m^Toldsymbol{x}^{ ext{ex}}-oldsymbol{V}_m^Toldsymbol{x}_{m,\widehat{\lambda}}\|=\|oldsymbol{V}_m^Toldsymbol{x}^{ ext{ex}}-oldsymbol{y}_{m,\widehat{\lambda}}\|;$$

when using the IRN methods we should incorporate the appropriate preconditioners $(W_p^{\gamma})_k$ and S_k and, for all the iterations in the inner iteration cycle corresponding to the kth outer iteration, we seek to minimize with respect to $\hat{\lambda}$

722
$$\|\hat{\boldsymbol{x}}^{\text{ex}} - \boldsymbol{V}_m \boldsymbol{y}_{m,\hat{\lambda}}\| = \|\boldsymbol{V}_m^T \hat{\boldsymbol{x}}^{\text{ex}} - \boldsymbol{V}_m^T \boldsymbol{V}_m \boldsymbol{y}_{m,\hat{\lambda}}\| = \|\boldsymbol{V}_m^T \hat{\boldsymbol{x}}^{\text{ex}} - \boldsymbol{y}_{m,\hat{\lambda}}\|, \text{ where } \hat{\boldsymbol{x}}^{\text{ex}} = (\boldsymbol{W}_p^{\gamma})_k \boldsymbol{S}_k \boldsymbol{x}^{\text{ex}}.$$

It is intrinsically difficult to implement this strategy for flexible Krylov subspace methods, because of 723 the complexity of changing preconditioners at each iteration. (iii) We perform a manual exhaustive 724 search. Namely, we first run the solvers multiple times using various regularization parameters λ , 725 starting with a larger range and narrowing down to a smaller range containing the best parameter; 726 we then record the minimum relative errors among all iterations for all values of $\hat{\lambda}$, and select the 727 corresponding λ . This approach is the most expensive one, and differs from the previous one in that 728 the (optimal) regularization parameter λ is fixed for all iterations. Of course, both the second and 729 third approaches require the knowledge of the exact solution and we test them only to investigate 730 the best possible performance of the hybrid approach.

Table 1 compares the performances (in terms of minimum relative error achieved by each method) of standard Krylov methods (GMRES and LSQR) and their IRN-NNR and flexible NNR

S. GAZZOLA, C. MENG AND J. NAGY

(F-NNR) counterparts, with and without using a hybrid approach. In this way we can understand how the use of additional regularization affects each solver differently. The three parameter
choice methods described above are dubbed "Secant (i)", "Optimal (ii)" and "Fixed (iii)", respectively. All the previous examples are considered here. GMRES and its counterparts IRN-GMRESNNR, FGMRES-NNR are used for Example 1, while LSQR and its counterparts IRN-LSQR-NNR,
FLSQR-NNR(v) are used for Examples 2 and 3.

| | | $\hat{\lambda} = 0$ | $\hat{\lambda} \neq 0$ |
|----------|--------------|---------------------|------------------------|---------------------|------------------------|---------------------|------------------------|---------------------|------------------------|
| | | Example 1 | | Example 2 | | Example 3 (house) | | Example 3 (peppers) | |
| Standard | Secant (i) | 0.2995 | 0.2528 | 0.1201 | 0.1389 | 0.2712 | 0.2715 | 0.1141 | 0.1138 |
| | Optimal (ii) | 0.2995 | 0.2268 | 0.1201 | 0.1201 | 0.2712 | 0.2710 | 0.1141 | 0.1138 |
| | Fixed (iii) | 0.2995 | 0.2268 | 0.1201 | 0.1183 | 0.2712 | 0.2710 | 0.1141 | 0.1138 |
| IRN-NNR | Secant (i) | 0.2081 | 0.2096 | 0.0685 | 0.0696 | 0.1249 | 0.1250 | 0.0964 | 0.0967 |
| | Optimal (ii) | 0.2081 | 0.2292 | 0.0685 | 0.0685 | 0.1249 | 0.1249 | 0.0964 | 0.0964 |
| | Fixed (iii) | 0.2081 | Х | 0.0685 | 0.0660 | 0.1249 | Х | 0.0964 | 0.0960 |
| F-NNR | Secant (i) | 0.2829 | 0.2658 | 0.0577 | 0.0684 | 0.1035 | 0.1046 | 0.0625 | 0.0618 |
| | Fixed (iii) | 0.2829 | 0.2640 | 0.0577 | 0.0568 | 0.1035 | Х | 0.0625 | 0.0618 |

Table 1: Minimum relative errors without $(\hat{\lambda} = 0)$ and with $(\hat{\lambda} \neq 0)$ a hybrid approach. The mark "X" means that the optimal regularization parameter found by the "Fixed (iii)" method is 10^{-16} , hence there is no need for additional regularization.



Fig. 14: Reconstructions obtained by standard hybrid Krylov methods and by the new methods without using additional regularization. Left side: zoomed in surface plots of the reconstructions of *Example 1*; right side: reconstructions of *Example 3* (peppers).

It is easy to observe that the use of additional regularization is most effective for the standard 740 GMRES solver, where the minimum relative error is reduced significantly. However, for the other 741 solvers, the hybrid approach does not have a notable advantage over not using regularization. At 742 times the "Fixed (iii)" parameter choice strategy delivers a regularization parameter of the order of 10^{-16} , which is numerically equivalent to not having regularization. This indicates that our 744 new IRN-NNR and F-NNR methods are successful in computing good reconstructions and, even 745 without additional regularization, they perform much better than standard Krylov methods used 746 in a hybrid fashion (comparing IRN-GMRES-NNR to GMRES in Example 1, and FLSQR-NNR(v) 747748 to LSQR in the other examples). Figure 14 shows a couple of such comparisons.

6. Conclusions. This paper introduced new solvers, based on Krylov subspace methods, for the computation of approximate low-rank solutions to large-scale linear systems of equations. Our

main goal was to apply the new methods to regularize inverse problems arising in imaging applications. The starting point of our derivations was an IRN approach to the NNRp problem (1.3). In 752 this way, the original problem (1.3) is reduced to the solution of a sequence of quadratic problems, where an appropriate smoothed linear transformation is introduced to approximate the nondif-754ferentiable nuclear norm regularization term. Our new methods make smart use of Kronecker 756 product properties to reformulate each quadratic problem in the IRN sequence as a Tikhonovregularized problem in standard form. We use both Krylov methods with fixed "preconditioners" 757 within an inner-outer iteration scheme (namely, IRN-LSQR-NNRp and IRN-GMRES-NNRp), and 758 Krylov methods with flexible iteration-dependent "preconditioners" within a single iteration scheme 759(namely, FLSQR-NNRp, FGMRES-NNRp, LR-FGMRES, and LR-FLSQR). Some of these meth-760 ods (namely, IRN-LSQR-NNRp, IRN-GMRES-NNRp, FLSQR-NNRp, and FGMRES-NNRp) can 761 be used in a hybrid framework, so that the Tikhonov regularization parameter can be efficiently, 763 effectively, and adaptively chosen. These new solvers are shown to perform exceptionally well on the test problems described in Section 5, and they give reconstructions of significantly improved 764 765 quality over existing methods.

Future work includes the extension of the present methods to handle cases where the solution of (1.1) is low-rank but rectangular, i.e., $\operatorname{vec}^{-1}(x) = X \in \mathbb{R}^{m \times n}$ with $m \neq n$. Also, while a solid theoretical justification is provided for IRN-LSQR-NNR*p* and IRN-GMRES-NNR*p*, the same is not true for FGMRES-NNR*p* and FLSQR-NNR*p*: further analysis will be needed to deeply understand the regularization properties of these flexible solvers. Finally, the new IRN-LSQR-NNR*p* and IRN-GMRES-NNR*p* methods can be reformulated to work with well-posed problems and in the framework of matrix equations, possibly providing a valid and principled alternative to

⁷⁷³ the current popular methods based on low-rank-projected and restarted Krylov solvers.

774

REFERENCES

- [1] M. BELGE, M. E. KILMER, AND E. L. MILLER, Wavelet domain image restoration with adaptive edge-preserving regularization, IEEE Trans. Image Process., 9 (2000), pp. 597–608, https://doi.org/10.1109/83.841937.
- [2] J. D. BLANCHARD, J. TANNER, AND K. WEI, CGIHT: conjugate gradient iterative hard thresholding for compressed sensing and matrix completion, Information and Inference: A Journal of the IMA, 4 (2015), pp. 289– 327, https://doi.org/10.1093/imaiai/iav011.
- [3] J. CAI, E. CANDÈS, AND Z. SHEN, A singular value thresholding algorithm for matrix completion, SIAM Journal on Optimization, 20 (2010), pp. 1956–1982, https://doi.org/10.1137/080738970.
- [4] J. CHUNG AND S. GAZZOLA, Flexible Krylov Methods for ℓ_p Regularization, ArXiv e-prints, (2018), https: //arxiv.org/abs/1806.06502.
- [5] M. FORNASIER, H. RAUHUT, AND R. WARD, Low-rank matrix recovery via iteratively reweighted least squares minimization, SIAM Journal on Optimization, 21 (2011), pp. 1614–1640, https://doi.org/10.1137/ 100811404.
- [6] S. GAZZOLA, P. C. HANSEN, AND J. G. NAGY, IR Tools: A MATLAB Package of Iterative Regularization Methods and Large-Scale Test Problems, 2017, https://arxiv.org/abs/arXiv:1712.05602.
- [7] S. GAZZOLA AND J. G. NAGY, Generalized Arnoldi-Tikhonov Method for Sparse Reconstruction, SIAM J. Sci.
 Comput., 36 (2014).
- [8] S. GAZZOLA, P. NOVATI, AND M. R. RUSSO, On krylov projection methods and tikhonov regularization, Elec tronic Transactions on Numerical Analysis, 44 (2014), pp. 82–123.
- [9] S. GAZZOLA AND M. SABATÉ LANDMAN, Flexible gmres for total variation regularization, BIT Numerical Math ematics, (2019), https://doi.org/10.1007/s10543-019-00750-x.
- [10] D. GOLDFARB AND S. MA, Convergence of fixed-point continuation algorithms for matrix rank minimization, Foundations of Computational Mathematics, 11 (2011), pp. 183–210, https://doi.org/10.1007/ s10208-011-9084-6.
- [11] N. HALKO, P. G. MARTINSSON, AND J. A. TROPP, Finding structure with randomness: Probabilistic algorithms for constructing approximate matrix decompositions, SIAM Review, (2011).

S. GAZZOLA, C. MENG AND J. NAGY

- [12] M. HANKE AND P. C. HANSEN, Regularization methods for large-scale problems, Surv. Math. Ind., 3 (1993),
 pp. 253–315.
- [13] P. C. HANSEN, Discrete Inverse Problems: Insight and Algorithms, Society for Industrial and Applied Mathe matics, Philadelphia, PA, USA, 2010.
- [14] P. C. HANSEN AND T. K. JENSEN, Smoothing-norm preconditioning for regularizing minimum-residual methods,
 SIAM J. Matrix Anal. Appl., 29 (2006), pp. 1–14.
- [15] T. K. JENSEN AND P. C. HANSEN, Iterative regularization with minimum-residual methods, BIT, 47 (2007),
 pp. 103–120.
- [16] R. H. KESHAVAN, A. MONTANARI, AND S. OH, Matrix completion from noisy entries, J. Mach. Learn. Res., 11
 (2010), pp. 2057–2078, http://dl.acm.org/citation.cfm?id=1756006.1859920.
- [17] M. KILMER AND D. O'LEARY., Choosing regularization parameters in iterative methods for ill-posed problems,
 SIAM Journal on Matrix Analysis and Applications, 22 (2001), pp. 1204–1221, https://doi.org/10.1137/
 S0895479899345960.
- [18] M. E. KILMER AND D. O'LEARY, Choosing regularization parameters in iterative methods for ill-posed problems,
 SIAM J. Matrix Anal. Appl., 22 (2001), pp. 1204–1221.
- [19] D. KRESSNER AND C. TOBLER, Low-rank tensor Krylov suspace methods for parameterized linear systems,
 SIAM J. Matrix Anal. Appl., 32 (2011), pp. 1288–1316.
- [20] R. M. LARSEN, Lanczos bidiagonalization with partial reorthogonalization, Department of Computer Science,
 Aarhus University, Technical report, (1998).
- [21] B. J. LAST AND K. KUBIK, Compact gravity inversion, GEOPHYSICS, 48 (1983), pp. 713–721, https://doi.
 org/10.1190/1.1441501.
- [22] K. LEE AND H. C. ELMAN, A Preconditioned Low-Rank Projection Method with a Rank-Reduction Scheme for
 Stochastic Partial Differential Equations, SIAM J. Sci. Comput., 9 (2017).
- [23] C. LU, Z. LIN, AND S. YAN, Smoothed low rank and sparse matrix recovery by iteratively reweighted least squares minimization, IEEE Transactions on Image Processing, 24 (2015), pp. 646–654, https://doi.org/ 10.1109/TIP.2014.2380155.
- [24] S. MA, D. GOLDFARB, AND L. CHEN, Fixed point and bregman iterative methods for matrix rank minimization,
 Mathematical Programming, 128 (2011), pp. 321–353, https://doi.org/10.1007/s10107-009-0306-5.
- [25] K. MOHAN AND M. FAZEL, Iterative reweighted algorithms for matrix rank minimization, J. Mach. Learn. Res.,
 13 (2012), pp. 3441–3473, http://dl.acm.org/citation.cfm?id=2503308.2503351.
- [26] D. P. O'LEARY AND J. A. SIMMONS, A bidiagonalization-regularization procedure for large scale discretizations
 of ill-posed problems, SIAM J. Sci. Statist. Comput., 2 (1981), pp. 474–489.
- [27] B. RECHT, M. FAZEL, AND P. PARRILO, Guaranteed minimum-rank solutions of linear matrix equations via nuclear norm minimization, SIAM Review, 52 (2010), pp. 471–501, https://doi.org/10.1137/070697835.
- [28] R. A. RENAUT, S. VATANKHAH, AND V. E. ARDESTANI, Hybrid and iteratively reweighted regularization by
 unbiased predictive risk and weighted GCV for projected systems, SIAM J. Scientific Computing, 39 (2017),
 https://doi.org/10.1137/15M1037925.
- 837 [29] R. T. ROCKAFELLAR, Convex analysis, vol. 28, Princeton university press, 1970.
- [30] P. RODRIGUEZ AND B. WOHLBERG, An efficient algorithm for sparse representations with ℓ_p data fidelity term, (2008).
- [31] Y. SAAD, A flexible inner-outer preconditioned gmres algorithm, SIAM Journal on Scientific Computing, 14
 (1993), pp. 461–469, https://doi.org/10.1137/0914028.
- [32] V. SIMONCINI AND D. B. SZYLD, Recent computational developments in krylov subspace methods for linear
 systems, Numerical Linear Algebra with Applications, 14, pp. 1–59, https://doi.org/10.1002/nla.499.
- [33] M. STOLL AND T. BREITEN, A low-rank in time approach to PDE-constrained optimization, SIAM J. Sci.
 Comput., 27 (2015), pp. B1–B29.
- [34] S. VATANKHAH, R. RENAUT, AND S. LIU, Research note: A unifying framework for widely-used stabilization
 of potential field inverse problems, Geophysical Prospecting, (2019), https://doi.org/10.1111/1365-2478.
 12926.