Dictionary learning - from local towards global and adaptive

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Abstract

This paper studies the convergence behaviour of dictionary learning via the Iterative Thresholding and K-residual Means (ITKrM) algorithm. On one hand it is shown that there exist stable fixed points that do not correspond to the generating dictionary, which can be characterised as very coherent. On the other hand it is proved that ITKrM is a contraction under much relaxed conditions than previously necessary. Based on the characterisation of the stable fixed points, replacing coherent atoms with carefully designed replacement candidates is proposed. In experiments on synthetic data this outperforms random or no replacement and always leads to full dictionary recovery. Finally the question how to learn dictionaries without knowledge of the correct dictionary size and sparsity level is addressed. Decoupling the replacement strategy of coherent or unused atoms into pruning and adding, and slowly carefully increasing the sparsity level, leads to an adaptive version of ITKrM. In several experiments this adaptive dictionary learning algorithm is shown to recover a generating dictionary from randomly initialised dictionaries of various sizes on synthetic data and to learn meaningful dictionaries on image data.

Keywords: dictionary learning, sparse coding, sparse component analysis, Iterative Thresholding and K-residual Means (ITKrM), replacement, adaptive dictionary learning, parameter estimation.

1. Introduction

The goal of dictionary learning is to decompose a data matrix $Y = (y_1, \ldots, y_N)$, where $y_n \in \mathbb{R}^d$, into a dictionary matrix $\Phi = (\varphi_1, \ldots, \varphi_K)$, where each column also referred to as atom is normalised, $\|\varphi_k\|_2 = 1$ and a sparse coefficient matrix $X = (x_1, \ldots, x_N)$,

$$Y \approx \Phi X \quad \text{and} \quad X \text{ sparse.}$$

(1)

The compact data representation provided by a dictionary can be used for data restoration, such as denoising or reconstruction from incomplete information, \[12, 26, 28\] and data analysis, such as blind source separation, \[15, 25, 21, 22\]. Due to these applications dictionary learning is of interest to both the signal processing community, where it is also known as sparse coding, and the independent component analysis (ICA) and the blind source separation (BSS) community, where it is also known as sparse component analysis. It also means that there are not only many algorithms to choose from, \[15, 3, 14, 22, 25, 27, 38, 28, 33, 30\], but also that theoretical results have started to accumulate, \[17, 39, 1, 11, 34, 35, 19, 6, 5, 37, 40, 41, 9\]. As our reference list grows more
incomplete every day, we point to the surveys [32, 36] as starting points for digging into algorithms and theory, respectively.

One way to concretise the abstract formulation of the dictionary learning problem in (1) is to formulate it as optimisation programme. For instance, choosing a sparsity level $S$ and a dictionary size $K$, we define $\mathcal{X}_S$ to be the set of all columnwise $S$-sparse coefficient matrices, $\mathcal{D}_K$ to be the set of all dictionaries with $K$ atoms and for some $p \geq 1$ try to find

$$\arg\min_{\Psi \in \mathcal{D}_K, X \in \mathcal{X}_S} \sum_n \|y_n - \Psi x_n\|_2^p. \quad (2)$$

Unfortunately this problem is highly non-convex and as such difficult to solve even in the simplest and most commonly used case $p = 2$. However, randomly initialised alternating projection algorithms, which alternate between (trying to) find the best dictionary $\Psi$, based on coefficients $X$, and the best coefficients $X$, based on a dictionary $\Psi$, such as K-SVD (K Singular Value Decompositions) for $p = 2$, [3], and ITKrM (Iterative Thresholding and K residual Means) related to $p = 1$, [37], tend to be very successful on synthetic data - usually recovering 90 to 100% of all atoms - and to provide useful dictionaries on image data.

Apart from needing both the sparsity level and the dictionary size as input, the main drawback of these algorithms is that - assuming that the data $Y$ is synthesized from a generating dictionary $\Phi$ and randomly drawn $S$- sparse coefficients $X$ - they have almost no (K-SVD) or very weak (ITKrM) theoretical dictionary recovery guarantees. This is in sharp contrast to more involved algorithms, which - given the correct $S,K$ - have global recovery guarantees but due to their computational complexity can at best be used in small toy examples. [4, 2, 6].

One interesting exception is an algorithm developed by Sun, Qu and Wright, [40, 41]. The algorithm is gradient descent based with a Newton trust region method to escape saddle points and proven to recover the generating dictionary if it is a basis. This result together with several results in machine learning which prove that non-convex problems can be well behaved, meaning all local minima are global minima, gives rise to hope that a similar result can be proven for learning overcomplete dictionaries.

**Contribution:** In this paper we will first destroy all hope of proving global convergence of alternating projection algorithms by sketching the existence of stable fixed points, that are not equivalent to the generating dictionary. Conversely we will prove that ITKrM is a contraction towards the generating dictionary under much relaxed conditions compared to those from [37].

However, based on a characterisation of the fixed points, showing them to be very coherent, and an analysis of the residuals at these fixed points, we consider a replacement procedure for coherent atoms and develop a strategy for finding good replacement candidates. With the help of these replacement candidates we will then tackle one of the most challenging problems in dictionary learning - the automatic choice of the sparsity level $S$ and the dictionary size $K$. This will lead to a version of ITKrM that adapts both the sparsity level and the dictionary size in each iteration. To the best of our knowledge, this is the first dictionary learning algorithm that can recover a generating dictionary without prescribing its size in the presence of noise and outliers.

**Organisation:** In the next section we will summarise our notational conventions and introduce the sparse signal model on which all our theoretical results are based. In Section 3
we will familiarise the reader with the ITKrM algorithm and analyse situations where the
algorithm does resp. does not converge to a generating dictionary. In particular, we first
conduct a small experiment to identify dictionaries that are fixed points of ITKrM but not
equivalent to the generating dictionary through reordering and sign flips. We also provide
a proof sketch showing that these dictionaries are stable fixed points. Inspired by the
properties of the spurious fixed points we then prove that ITKrM is a contraction towards
the generating dictionary on an area much bigger than indicated by the convergence radius
proved in [37].
The insights from Section 3 will be used in Section 4 to develop a non-random replacement
strategy. Using the information at a spurious fixed point dictionary, we will construct
replacement candidates that allow ITKrM to escape towards the generating dictionary, and
test the resulting algorithm on synthetic data.
In Section 5 we will then address the big problem how to learn dictionaries without being
given the generating sparsity level and dictionary size. This will be done by slowly increasing
the sparsity level and by decoupling the replacement strategy into separate pruning of
the dictionary and adding of promising replacement candidates. Numerical experiments
will show that the resulting algorithm can indeed recover the generating dictionary from
initialisations with various sizes on synthetic data and learn meaningful dictionaries on
image data.
In the last section we will sketch how the concepts leading to adaptive ITKrM can be
extended to other algorithms such as (approximative) K-SVD. Finally, based on a discussion
of our results, we will map out future directions of research.

2. Notations and Sparse Signal Model

Before we hit the strings, we will fine tune our notation and introduce some definitions;
usually subscripted letters will denote vectors with the exception of \( \varepsilon, \alpha, \omega \), where they are
numbers, for instance \( x_n \in \mathbb{R}^K \) vs. \( \varepsilon_k \in \mathbb{R} \), however, it should always be clear from the
context what we are dealing with.

For a matrix \( M \) we denote its (conjugate) transpose by \( M^\star \) and its Moore-Penrose pseudo
inverse by \( M^\dagger \). We denote its operator norm by \( \| M \|_2 = \max_{\| x \|_2 = 1} \| M x \|_2 \) and its Frobenius
norm by \( \| M \|_F = \text{tr}(M^\star M)^{1/2} \), remember that we have \( \| M \|_2 \leq \| M \|_F \).

We consider a dictionary \( \Phi \) a collection of \( K \) unit norm vectors \( \phi_k \in \mathbb{R}^d \), \( \| \phi_k \|_2 = 1 \). By
abuse of notation we will also refer to the \( d \times K \) matrix collecting the atoms as its columns
as the dictionary, that is, \( \Phi = (\phi_1, \ldots, \phi_K) \). The maximal absolute inner product between
two different atoms is called the coherence \( \mu(\Phi) \) of a dictionary, \( \mu(\Phi) = \max_{k \neq j} | \langle \phi_k, \phi_j \rangle | \).

By \( \Phi_I \) we denote the restriction of the dictionary to the atoms indexed by \( I \), that is,
\( \Phi_I = (\phi_{i_1}, \ldots, \phi_{i_S}) \), \( i_j \in I \), and by \( P(\Phi_I) \) the orthogonal projection onto the span of the atoms
indexed by \( I \), that is, \( P(\Phi_I) = \Phi_I \Phi_I^\dagger \). Note that in case the atoms indexed by \( I \) are
linearly independent we have \( \Phi_I^\dagger = (\Phi_I^\dagger \Phi_I)^{-1} \Phi_I^\dagger \). We also define \( Q(\Phi_I) \) to be the orthogonal
projection onto the orthogonal complement of the span of \( \Phi_I \), that is, \( Q(\Phi_I) = I_d - P(\Phi_I) \),
where \( I_d \) is the identity operator (matrix) in \( \mathbb{R}^d \).

(Ab)using the language of compressed sensing we define \( \delta_I(\Phi) \) as the smallest number such
that all eigenvalues of \( \Phi_I^\dagger \Phi_I \) are included in \([1 - \delta_I(\Phi), 1 + \delta_I(\Phi)]\) and the isometry constant
\( \delta_S(\Phi) \) of the dictionary as \( \delta_S(\Phi) := \max_{|I| \leq S} \delta_I(\Phi) \). When clear from the context we
will usually omit the reference to the dictionary. For more details on isometry constants see for instance \[8\].

For a (sparse) signal \(y = \sum_k \phi_k x_k\) we will refer to the indices of the \(S\) coefficients with largest absolute magnitude as the \(S\)-support of \(y\). Again, we will omit the reference to the sparsity level \(S\) if clear from the context.

To keep the sub(sub)scripts under control we denote the **indicator function of a set** \(V\) by \(\chi(V, \cdot)\), that is \(\chi(V, v)\) is one if \(v \in V\) and zero else. The set of the first \(S\) integers we abbreviate by \(S = \{1, \ldots, S\}\).

We define the **distance** of a dictionary \(\Psi\) to a dictionary \(\Phi\) as

\[
d(\Phi, \Psi) := \max_k \min_\ell \|\phi_k \pm \psi_\ell\|_2 = \max_k \sqrt{2 - 2|\langle \phi_k, \psi_\ell \rangle|},
\]

Note that this distance is not a metric since it is not symmetric. For example, if \(\Phi\) is the canonical basis and \(\Psi\) is defined by \(\psi_i = \phi_i\) for \(i \geq 3\), \(\psi_1 = (e_1 + e_2)/\sqrt{2}\), and \(\psi_2 = \sum_i \phi_i/\sqrt{d}\) then we have \(d(\Phi, \Psi) = 1/\sqrt{2}\) while \(d(\Psi, \Phi) = \sqrt{2} - 2/\sqrt{d}\). The advantage is that this distance is well defined also for dictionaries of different sizes. A **symmetric distance** between two dictionaries \(\Phi, \Psi\) of the same size could be defined as the maximal distance between two corresponding atoms, that is,

\[
d_s(\Phi, \Psi) := \min_{p \in \mathcal{P}} \max_k \|\phi_k \pm \psi_{p(k)}\|_2,
\]

where \(\mathcal{P}\) is the set of permutations of \(\{1, \ldots, K\}\). The distances are equivalent whenever there exists a permutation \(p\) such that after rearrangement, the cross-Gram matrix \(\Phi^* \Psi\) is diagonally dominant, that is, \(\min_k |\langle \phi_k, \psi_k \rangle| > \max_{k \neq j} |\langle \phi_k, \psi_j \rangle|\). Since the main assumption for our results will be such a diagonal dominance we will state them in terms of the easier to calculate asymmetric distance and assume that \(\Psi\) is already signed and rearranged in a way that \(d(\Phi, \Psi) = \max_k \|\phi_k - \psi_k\|_2\). We then use the abbreviations \(\alpha_{\min} = \min_k |\langle \phi_k, \psi_k \rangle|\) and \(\alpha_{\max} = \max_k |\langle \phi_k, \psi_k \rangle|\). The maximal absolute inner product between two non-corresponding atoms will be called the **cross-coherence** \(\mu(\Phi, \Psi)\) of the two dictionaries, \(\mu(\Phi, \Psi) = \max_{k \neq j} |\langle \phi_k, \psi_j \rangle|\).

We will also use the following decomposition of a dictionary \(\Psi\) into a given dictionary \(\Phi\) and a perturbation dictionary \(Z\). If \(d(\Psi, \Phi) = \varepsilon\) we set \(\|\psi_k - \phi_k\|_2 = \varepsilon_k\), where by definition \(\max_k \varepsilon_k = \varepsilon\). We can then find unit vectors \(z_k\) with \(\langle \phi_k, z_k \rangle = 0\) such that

\[
\psi_k = \alpha_k \phi_k + \omega_k z_k, \quad \text{for} \quad \alpha_k := 1 - \varepsilon_k^2/2 \quad \text{and} \quad \omega_k := (\varepsilon_k^2 - \varepsilon_k^4/4)^{1/2}.
\]

Note that if the cross-Gram matrix \(\Phi^* \Psi\) is diagonally dominant we have \(\alpha_{\min} = \min_k \alpha_k\), \(\alpha_{\max} = \max_k \alpha_k\) and \(d(\Psi, \Phi) = \sqrt{2 - 2\alpha_{\min}}\).

### 2.1 Sparse signal model

As basis for our results we use the following signal model, already used in \[34, 35, 37\]. Given a \(d \times K\) dictionary \(\Phi\), we assume that the signals are generated as,

\[
y = \frac{\Phi x + r}{\sqrt{1 + \|r\|_2^2}},
\]
where \( x \in \mathbb{R}^K \) is a sparse coefficient sequence and \( r \in \mathbb{R}^d \) is some noise. We assume that \( r \) is a centered subgaussian vector with parameter \( \rho \), that is, \( \mathbb{E}(r) = 0 \) and for all vectors \( v \) the marginals \( \langle v, r \rangle \) are subgaussian with parameter \( \rho \), meaning they satisfy \( \mathbb{E}(e^{\langle v, r \rangle}) \leq e^{\rho^2 v^2/2} \) for all \( t > 0 \).

To model the coefficient sequences \( x \) we first assume that there is a measure \( \nu_c \) on a subset \( \mathcal{C} \) of the positive, non increasing sequences with unit norm, meaning for \( c \in \mathcal{C} \) we have \( c(1) \geq c(2) \ldots \geq c(K) \geq 0 \) and \( \|c\|_2 = 1 \). A coefficient sequence \( x \) is created by drawing a sequence \( c \) according to \( \nu_c \), and both a permutation \( p \) and a sign sequence \( \sigma \) uniformly at random and setting \( x = x_{c,p,\sigma} \) where \( x_{c,p,\sigma}(k) = \sigma(k)c(p(k)) \). The signal model then takes the form

\[
y = \frac{\Phi x_{c,p,\sigma} + r}{\sqrt{1 + \|r\|_2^2}}.
\]

Using this model it is quite simple to incorporate sparsity via the measure \( \nu_c \). To model approximately \( S \)-sparse signals we require that the \( S \) largest absolute coefficients, meaning those inside the support \( I = p^{-1}(S) \), are well balanced and much larger than the remaining ones outside the support. Further, we need that the expected energy of the coefficients outside the support is relatively small and that the sparse coefficients are well separated from the noise. Concretely we require that almost \( \nu_c \)-surely we have

\[
\frac{c(1)}{c(S)} \leq \gamma_{\text{dyn}}, \quad \frac{c(S+1)}{c(S)} \leq \gamma_{\text{gap}}, \quad \frac{\|c(S^\sigma)\|_2}{c(1)} \leq \gamma_{\text{app}} \quad \text{and} \quad \frac{\rho}{c(S)} \leq \gamma_{\rho}.
\]

We will refer to the worst case ratio between coefficients inside the support, \( \gamma_{\text{dyn}} \), as dynamic (sparse) range and to the worst case ratio between coefficients outside the support to those inside the support, \( \gamma_{\text{gap}} \), as the (sparse) gap. Since for a noise free signal the expected squared sparse approximation error is

\[
\mathbb{E}(\| \sum_{k \notin I} \sigma_k c(p(k)) \phi_k \|_2^2) = \| c(S^\sigma) \|_2^2,
\]

we will call \( \gamma_{\text{app}} \) the relative (sparse) approximation error. Finally, \( \gamma_{\rho} \) is called the noise to (sparse) coefficient ratio.

Apart from these worst case bounds we will also use three other signal statistics,

\[
\gamma_{1,S} := \mathbb{E}_c (\|c(S)\|_1), \quad \gamma_{2,S} := \mathbb{E}_c (\|c(S)\|_2^2), \quad C_r := \mathbb{E}_r \left( \frac{1}{\sqrt{1 + \|r\|_2^2}} \right).
\]

The constant \( \gamma_{1,S} \) helps to characterise the average size of a sparse coefficient, \( \gamma_{1,S} = \mathbb{E}(|x_i| : i \in I) \cdot S \leq \sqrt{S} \), while \( \gamma_{2,S} \) characterises the average sparse approximation quality, \( \gamma_{2,S} = \mathbb{E}(\|\Phi_I x\|_2^2) \leq 1 \). The noise constant can be bounded by

\[
C_r \geq \frac{1 - e^{-d}}{\sqrt{1 + 5d\rho^2}},
\]

and for large \( \rho \) approaches signal to noise ratio, \( C_r^2 \approx \frac{1}{d\rho} \approx \frac{\mathbb{E}(\|\Phi_I x\|_2^2)}{\mathbb{E}(\|r\|_2^2)} \), see [35] for details.

To get a better feeling for all the involved constants, we will calculate them for the case
**Algorithm 3.1: ITKrM (one iteration)**

**Input:** $\Psi, Y, S$ ;  
\text{ // dictionary, signals, sparsity}

Initialise: $\bar{\Psi} = 0$ ;

foreach $n$ do

$I^*_{hn} = \arg\max_{I \mid \|I\|_0 \leq S} \|\Psi^*_I y_n\|_1$ ;  
\text{ // thresholding}

$a_n = y_n - P(\Psi^*_I)n$ ;  
\text{ // residual}

foreach $k \in I^*_{hn}$ do

$\bar{\psi}_k \leftarrow \bar{\psi}_k + [a_n + P(\psi_k)y_n] \cdot \text{sign}(\langle \psi_k, y_n \rangle)$ ;  
\text{ // atom update}

end

$\Psi \leftarrow (\bar{\psi}_1/\|\bar{\psi}_1\|_2, \ldots, \bar{\psi}_K/\|\bar{\psi}_K\|_2)$ ;  
\text{ // atom normalisation}

**Output:** $\Psi$

of perfectly sparse signals where $c(i) = 1/\sqrt{S}$ for $i \leq S$ and $c(i) = 0$ else. We then have $\gamma_{dyn} = 1$, $\gamma_{gap} = 0$ and $\gamma_{app} = 0$ as well as $\gamma_{1,S} = \sqrt{S}$ and $\gamma_{2,S} = 1$. In the case of noiseless signals we have $C_r = 1$ and $\gamma_\rho = 0$. In the case of Gaussian noise the noise to coefficient ratio is related to the signal to noise ratio via $\text{SNR} = S/(\gamma^2_{1}\rho d)$.

### 3. Global behaviour patterns of ITKrM

The iterative thresholding and K residual means algorithm (ITKrM) was introduced in [37] as modification of its much simpler predecessor ITKsM, which uses signal means instead of residual means. As can be seen from the summary in Algorithm 3.1 the signals can be processed sequentially, thus making the algorithm suitable for an online version and parallelisation. The determining factors for the computational complexity are the matrix vector products $\Psi^*_I y_n$ between the current estimate of the dictionary $\Psi$ and the signals, $O(dKN)$, and the projections $P(\Psi^*_I y_n)$. If computed with maximal numerical stability these would have an overall cost $O(S^2dN)$, corresponding to the QR decompositions of $\Psi^*_I y_n$. However, since usually the achievable precision in the learning is limited by the number of available training signals rather than the numerical precision, it is computationally more efficient to precompute the Gram matrix $\Psi^*\Psi$ and calculate the projections less stably via the eigenvalue decompositions of $\Psi^*_I \Psi^*_I$, corresponding to an overall cost $O(S^3N)$.

Another good property of the ITKrM algorithm is that it is proven to converge locally to a generating dictionary. Concretely this means that if the data is homogeneously $S$-sparse in a dictionary $\Phi$, where $S \lesssim \mu^{-2}$, and we initialise with a dictionary $\Psi$ within radius $O(1/\sqrt{S})$, $d(\Psi, \Phi) \lesssim 1/\sqrt{S}$, then ITKrM using $N = O(K \log K)$ samples in each iteration will converge to the generating dictionary, [37]. In simulations on synthetic data ITKrM shows even better convergence behaviour. Concretely, if the atoms of the generating dictionary are perturbed with vectors $z_k$ chosen uniformly at random from the sphere, $\psi_k = \alpha_k \phi_k + \omega_k z_k$, ITKrM converges also for ratios $\alpha_k : \omega_k = 1 : 4$. For completely random initialisations, $\psi_k = z_k$, it finds between 90% and 100% of the atoms - depending on the noise and sparsity level.

Last but not least, ITKrM is not just a pretty toy with theoretical guarantees but on image data produces dictionaries of the same quality as K-SVD in a fraction of the time, [29].
Considering the good practical performance of ITKrM, it is especially frustrating that we only get a convergence radius of size $O(1/\sqrt{S})$, while for its simpler cousin ITKsM, which when initialised randomly performs much worse both on synthetic and image data, we can prove a convergence radius of size $O(1/\sqrt{\log K})$. To get a better understanding why ITKrM has a better practical performance, we will first have a closer look at what happens when things go wrong and things go right. Based on the intuition gained there we will then prove that ITKrM behaves well under much more relaxed conditions.

3.1 When things go wrong

We start with a closer inspection of what happens when ITKrM does not find all atoms using a random initialisation. From [37] we know that this is most likely to happen when the signals are very sparse ($S$ small) and the noise level is small. For better visualisation we only run a small experiment in $\mathbb{R}^{32}$, where we try to recover a very incoherent dictionary from $2$-sparse vectors $\mathbf{1}$. The dictionary, containing 48 atoms, consists of the Dirac basis and the first half of the vectors from the Hadamard basis, and as such has coherence $\mu = 1/\sqrt{32} \approx 0.18$.

The signals follow the model in (24), where the coefficient sequences $c$ are constructed by choosing $b \in [0.9, 1]$ uniformly at random and setting $c_1 = 1/\sqrt{1 + b^2}$; $c_2 = bc_1$ and $c_j = 0$ for $j \geq 3$. The noise is chosen to be Gaussian with variance $\rho^2 = 1/(16d)$, corresponding to SNR = 16. Running ITKrM with 20000 new signals per iteration for 25 iterations and 10 different random initialisations we recover 4 times 46 atoms and 6 times 44 atoms. An immediate observation is that we always miss an even number of atoms. Taking a look at the recovered dictionaries - examples for recovery of 44 and 46 atoms are shown in Figure 3.1 - we see that this is due to their special structure; in case of $2n$ missing atoms, we always observe $n$ atoms that are recovered twice and $n$ atoms that are 1:1 linear combinations of 2 missing atoms, respectively. To get a better understanding why these configurations are stable, we will sketch that for noiseless signals that are perfectly sparse in a dictionary $\Phi$,

1. All experiments and resulting figures can be reproduced using the matlab toolbox available at https://www.uibk.ac.at/mathematik/personal/schnass/code/adl.zip

![Figure 1: Cross-Gram matrices $\Psi^*\Phi$ for recovered dictionaries with 2 (left) and 4 (right) missing atoms.](image)
meaning $y = \Phi_1 \sigma(I)$, one iteration of ITkrM will stay near $\Psi$ where $\psi_1 = \psi_2 = \phi_1, \psi_i = \phi_i$ for $i > 3$ and $\psi_3 = (\phi_2 + h \cdot \phi_3)/\sqrt{2 + 2h\theta}$ with $h = 1$ if $\theta = \langle \phi_2, \phi_3 \rangle \geq 0$ and $h = -1$ else.

**Proof sketch:** We start by analysing which support thresholding will recover depending on whether the generating support contains the (indices of the) double or the missing atoms and how the residual, $a := y - P(\Psi_I)y$, will look like. Without loss of generality we assume that $\langle \phi_2, \phi_3 \rangle \geq 0$ and therefore $h = 1$. To better deal with the recovered support sets we use the following notation for an index set $I$ where the index $i \in I$ has been replaced by an index $j \notin I$, that is, $I_{i \leftarrow j} := I \setminus i \cup j$. Note that we have $1/\sqrt{2} \leq |\langle \phi_i, \psi_3 \rangle| \leq \sqrt{(1 + \mu)/2} \leq (1 + \mu)/\sqrt{2}$, for $i = 2, 3$ and $|\langle \phi_i, \psi_3 \rangle| \leq \sqrt{2} \mu \leq 2 \mu$ else, as well as $|\langle \phi_i, \psi_j \rangle| \leq \mu$ for $i \neq j$ and $j \neq 3$.

1. $2, 3 \notin I$: In the most common case we have $I^t = I$ since $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \leq S\mu$ if $i \notin I$ and $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \geq 1 - S\mu$ if $i \in I$. In consequence we have $a = y - P(\Phi_I)y = 0$.

2. $1 \in I, 2, 3 \notin I$: We have $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \leq S\mu$ for all $i \notin I \cup 2$ and $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \geq 1 - S\mu$ for all $i \in I \cup 2$. Therefore, the recovered support depends on the order of the inner products $|\langle \Phi_1 \sigma(I), \phi_i \rangle|$. Let’s assume that each inner product is equally likely to be the smallest. Then with probability $1/S$ we take both copies of $\phi_i$ but miss the generating atom $\psi_3$, meaning $I^t = I_{i \leftarrow 2}$ for $i \neq 1$, leading to $a = y - P(\Phi_{I_{1 \leftarrow 2}})y \approx \phi_1 \sigma(i)$. Further with probability $1/(2\mu)$ we take one copy of $\phi_3$ and all the other generating atoms, $I^t = I$ or $I^t = I_{1 \leftarrow 3}$, leading to $a = y - P(\Phi_I)y = 0$.

3. $3 \in I, 1, 2 \notin I$: As in the first case we have $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \leq S\mu$ for all $i \notin I$. Further, we have $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \geq 1 - S\mu$ for all $i \in I \setminus 3$ and $|\langle \Phi_1 \sigma(I), \psi_3 \rangle| \geq 1/\sqrt{2} - S\mu$, leading to $I^t = I$. For the residual we have $a = y - P(\Phi_I)y \approx (\phi_3 - P(\psi_3)2\phi_3)\sigma(3) \approx (\phi_3 - \phi_2)\sigma(3)/2$. The case $2 \in I, 1, 3 \notin I$ is analogue, meaning we get $I^t = I_{2 \leftarrow 3}$ and $a \approx (\phi_2 - \phi_3)\sigma(2)/2$.

4. $1, 3 \in I, 2 \notin I$: We have $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \leq S\mu$ for all $i \notin I \cup 2$, as well as $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \geq 1 - S\mu$ for all $i \in I_{3 \leftarrow 2}$. For $i = 3$ we have $|\langle \Phi_1 \sigma(I), \psi_3 \rangle| \leq 1/\sqrt{2} + S\mu$, meaning we will recover $I^t = I_{3 \leftarrow 2}$ and the residual will have the shape $a = y - P(\Phi_{I_{3 \leftarrow 2}})y \approx \phi_3 \sigma(3)$. In the analogue case $2, 1 \in I, 3 \notin I$ we will recover $I^t = I$ with residual $a \approx \phi_2 \sigma(2)$.

5. $2, 3 \in I, 1 \notin I$: We have $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \leq S\mu$ for all $i \notin I$ as well as for $i = 2$ and $|\langle \Phi_1 \sigma(I), \psi_i \rangle| \geq 1 - S\mu$ for $i \in I \setminus \{2, 3\}$. For $i = 3$ we have to distinguish whether the signal coefficients $\sigma(2), \sigma(3)$ have the same sign or not. If they have the same sign, we have $|\langle \Phi_1 \sigma(I), \psi_3 \rangle| \geq \sqrt{2} - S\mu$, thus $3 \in I^t$ and $a = y - P(\psi_I^t)y = (\|\phi_3 - P(\psi_3)\|\phi_2 + \phi_3)\sigma(2) = 0$. To have correct size the recovered support will also contain an index $j \in I^c \cup 2$, $I^t = I$ or $I^t = I_{2 \leftarrow j}$. However, since the residual is zero, this will hardly affect the update of the respective atom $\psi_j$. If conversely $\sigma(2), \sigma(3)$ have different signs, then the contribution of $\phi_2, \phi_3$ to the signal is orthogonal to $\psi_3$ and we have $|\langle \Phi_1 \sigma(I), \psi_3 \rangle| \geq (S - 2)\mu$. It is thus highly improbably that the thresholded

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2. To simplify the proof sketch, we assume that the non-zero coefficients $|x(k)|$ are equal to 1 in magnitude rather than $1/\sqrt{S}$, so that the signals have average energy $S$ instead of 1.
support will contain $ψ_3$. Instead it will contain two atoms $j,k \in I^c \cup 2$, likely those most correlated with the residual $a = y - P(Ψ_{I^c})y \approx y - P(Φ_{I(2,3)})y \approx σ(2)(φ_2 - φ_3)$.

1, 2, 3 $∈ I$: This quite rare case works analogue to the one before, taking into account that $ψ_2 = φ_1$ is sure to be selected in the thresholding.

Based on the knowledge of the outcome of thresholding and the shape of the residuals we can now estimate the updated atoms $ψ_k$, where

$$
\bar{ψ}_k = \frac{1}{N} \sum_{n,k \in I_n^t} \left[ y_n - P(Ψ_{I^c})y_n + P(ψ_k)y_n \right] \cdot \text{sign}(⟨ψ_k, y_n⟩).
$$

(11)

We have to distinguish between three types of atoms, the correct singles for $k > 3$, where $ψ_k = φ_k$, the double atoms $k = 1, 2$ where $ψ_1 = ψ_2 = φ_1$, and the combined atom $k = 3$, where $ψ_3 \propto φ_2 + φ_3$.

$k > 3$: We start with the correct singles, where $ψ_k = φ_k$. To calculate the sum we first have a look at the cases where $k$ was correctly recovered, meaning $k ∈ I$ and $k ∈ I^t$. From the analysis above we know that this occurs whenever 1, 2, 3 $∉ I$, so with probability $\frac{S(K-4)}{K} \left( \frac{K}{S-1} \right)^{-1} \approx \frac{S}{K} (1 - 3 \frac{S}{K})$. We then have $a_n = 0$ and since the recovery is independent of the sign patterns the sum over these signals should be close to its expectation, $\frac{NS^2}{K}(1 - 3 \frac{S}{K})φ_k$.

Given that $k ∈ I$ the next case which guarantees recovery of $k$ is that either 2, 3 $∈ I$ but not 1, which has probability $\approx 2 \frac{S^2}{K^2}$. Then the residual has the form $a_n = σ_n(j)(φ_2 - φ_3)$ for $j = 2, 3$. Since the product of the signs $σ(j)σ_n(k)$ for $j = 2, 3$ is equally likely to be positive or negative, the contribution of $(φ_2 - φ_3)$ will average out and we again get that the sum over these signals is close to its expectation $2 \frac{NS^2}{K^2} φ_k$.

The only situation where we might not recover $k$ is when 1 $∈ I$, that is the second case described above, occurring with probability $\approx \frac{S^2}{K^2}$. Under the assumption that all atoms in the support are equally likely to have smallest inner product, this happens with probability $1/S$. Most often we recover $k$ but fail to recover some other $j \in I$, so the residual has the form $a_n = σ_n(j)φ_j$. In the worst case the signs of $φ_3$ and the missing atom are very correlated so that they never cancel. Then we get that the sum over these signals is of the form

$$
\frac{NS^2}{K^2} \left( \left( 1 - \frac{1}{S} \right) φ_k + \frac{S - 2}{S(K - 2)} \sum_{j \neq 1,k} m_j^k φ_j \right) \text{ with } \sum_j |m_j^k| = K - 2.
$$

(12)

Note that if we are less pessimistic about the correlation of the signs, for instance because we have decaying coefficients, the contributions of the missing atoms will often cancel, meaning $\sum_j |m_j^k| = M_k < K - 2$, and the sum will be closer to $\frac{NS^2}{K^2} \psi_k$.

Finally, we have to check how often $k$ will be falsely recovered, meaning $k \notin I$ but $k ∈ I^t$. From the analysis of thresholding we see that this is possible whenever both 2, 3 $∈ I$ and 1 $∉ I$ (probability $\approx \frac{NS^2}{K^2}$). With probability 1/2 we have $σ(2)σ(3) > 0$, so that the residual is zero and adding $|⟨φ_k, y_n⟩| φ_k$ will not affect the sum negatively. However, also with probability 1/2, the residual has the shape $a_n = σ_n(2)(φ_2 - φ_3)$.
and at least for the atom $\psi_k$, which is most correlated with $(\phi_2 - \phi_3)$ and therefore picked with the highest probability $\bar{m}_k \approx 1$, we have to add to the sum a term 
$\approx \frac{NS^2}{2K^2} \bar{m}_k (\phi_2 - \phi_3)$. 
Puting it all together we see that 

$$
\bar{\psi}_k \approx \frac{S}{K} \left(1 - \frac{1}{K}\right) \phi_k + \frac{S^2}{K^3} \sum_{j \neq 1,k} m_j^k \phi_j + \frac{S^2}{2K^2} \bar{m}_k (\phi_2 - \phi_3). 
$$

This means that after normalisation we have $\|\phi_k - \bar{\psi}_k\|_2 \leq \frac{S}{K}$ and that for $k > 3$ the updated atoms will stay (critically) close to the generating atoms.

$k = 1, 2$: Next we have a look how the double atom $\psi_1 = \phi_1$ evolves; the case of $\psi_2$ is exactly the same. We again estimate the sum starting with the cases where $1 \in I$ and $1 \in I'$. For $1 \in I$ we most commonly have that $2, 3 \notin I$, probability $\approx \frac{S}{K}(1 - 2\frac{S}{K})$. Because of our assumption that all inner products $\langle \langle \phi_i, y_n \rangle \rangle_i \in I$ are equally likely to be the smallest, we recover $\psi_1$ uniquely with probability $1/(2S)$ and have $a_n = 0$. With probability $1 - 1/S$ we recover both copies of $\phi_1$ and the residual has the shape $a_n = \sigma_n(j) \phi_j$, where $j$ is the atom with smallest inner product. As before the question is how correlated the signs $\sigma_n(j)$ of the missing atoms are with the signs $\sigma_n(1)$ of $\phi_1$. If they are very correlated and never cancel, the sum over these signals has the form

$$
\frac{NS}{K} \left(1 - 2\frac{S}{K}\right) \left(1 - \frac{1}{2S}\right) \phi_1 + \frac{S - 1}{S(K - 3)} \sum_{j \neq 1,2,3} m_j^1 \phi_j,
$$

for $\sum_j |m_j^1| = K - 3$. We will additionally assume that they are quite uncorrelated, meaning that $\sum_j |m_j^1| = M_1 \ll K - 3$. For balanced coefficients this is likely to happen whenever $\phi_1$ is orthogonal (or comparatively incoherent) to all other atoms in $\Phi$. In this case the sign $\sigma_n(1)$ does not influence the order of the inner products $\langle \langle \phi_i, y_n \rangle \rangle_i$ for $i \neq 1$ (as much as the signs of the more coherent atoms) and we get the same residual $a_n = \sigma_n(j) \phi_j$ for both choices $\sigma_n(1) = \pm 1$ (most of the time).

The second case where $1 \in I$ that we have to take into account is when $2 \in I$ or $3 \in I$, happening with probability $\approx 2\frac{S^2}{K^2}$. In this case we always recover both copies of $\phi_1$ and the residuals have the shape $a_n \approx \sigma_n(j) \phi_j$ for $j = 2$ resp. $j = 3$. Since $\sigma_n(1)\sigma_n(j)$ is equally likely to be positive or negative the contribution of the residuals will average out and the sum over these signals will be $\approx \frac{2NS^2}{K^2} \phi_1$.

Lastly, we have to estimate the effects of wrongly recovering $\psi_1$, that is $1 \notin I$ but $1 \in I'$. As for $\psi_k$ with $k > 3$ this can only happen when both $2, 3 \in I$, and so the worst contribution to the sum is the term $\approx \frac{NS^2}{2K^2} (\phi_2 - \phi_3)$. Combining the three estimates and normalising, we get that $\|\phi_1 - \bar{\psi}_1\|_2 \leq \frac{M_1 + 8S}{K}$, so both updated copies of $\phi_1$ stay near $\phi_1$.

$k = 3$: Finally we will verify that $\psi_3 \propto \phi_2 + \phi_3$, the atom doing the jobs of two generating atoms, is stable. The analysis of thresholding tells us that $\psi_3$ is never falsely recovered, meaning $2, 3 \notin I$ but $3 \in I'$, and that whenever it is correctly recovered the residual is approximately a linear combination of $\phi_2$ and $\phi_3$. In particular if $2 \in I$ but $3, 1 \notin I$
we have \( a_n \approx (\phi_2 - \phi_3)\sigma_n(2)/2 \) so that \( a_n\sigma_n(2) + |\langle \psi_3, y_n \rangle| \psi_3 \approx \phi_2 \), and vice versa for \( 3 \in I \) but \( 2, 1 \notin I \). On the other hand if both \( 2, 3 \in I \) then either \( \psi_3 \) is not recovered or \( a_n \approx 0 \) and \( |\langle \psi_3, y_n \rangle| \approx \sqrt{2} \). Adding the terms scaled with their respective probabilities we get

\[
\overline{\psi}_3 \approx \frac{S}{K} \left( 1 - \frac{2S}{K} \right) \phi_2 + \frac{S}{K} \left( 1 - \frac{2S}{K} \right) \phi_3 + \frac{S^2}{2K^2} \sqrt{2} \psi_3 \propto \psi_3,
\]

showing that the updated atom is again a one to one combination of the missing atoms.

After sketching that there exist bad dictionaries from which ITKrM, or for that matter any alternating optimisation algorithm, is not likely to escape in reasonably many iterations, the next interesting question is how we end up near these bad dictionaries in the first place. The intuition from the sketch is the following; if we have two estimated atoms pointing to the same generating atom but nowhere else, meaning they are relatively close to an \( \phi_j \) but very incoherent to all the other atoms \( \phi_i \) for \( i \neq j \), they will quickly get contracted to this generating atom. At the same time we have two possible scenarios; a) one estimated atom is pointing to two generating atoms and drawn equally to both of them or b) once all estimated atoms have been drawn to a generating atom, the missing generating atom \( \phi_k \) will start to attract the estimator of the generating atom to which it is most coherent, that is \( \psi_{\overline{k}} \) where \( \overline{k} = \arg \max_{i \neq j,k} |\langle \phi_k, \phi_i \rangle| \), and they will start to efficiently share the same estimator \( \psi_{\overline{k}} \approx \phi_k + \phi_{\overline{k}} \).

Observe also that the two estimated atoms pointing to the same generating atom can be very incoherent even if they are both already quite close to \( \phi_j \). For instance, if \( \psi_{\overline{j}} \approx \alpha_j \phi_j \pm \omega_j z_j \) where \( z_j \) is a balanced sum of the other atoms \( z_j \approx \sum_{i \neq j} \phi_i \sigma(i) \), we have \( |\langle \psi_{\overline{i}}, \psi_{\overline{j}} \rangle| = \alpha_j^2 - \omega_j^2 \), meaning approximate orthogonality at \( \alpha_j = 1/\sqrt{2} \). Using these ideas we can construct well-conditioned and incoherent initial dictionaries \( \Psi \), with arbitrary distances \( d(\Psi, \Phi) \gtrsim 1/\sqrt{2} \) to the generating dictionary, so that things will go maximally wrong, meaning we end up with a lot of double and 1:1 atoms. Figure shows an example of a bad initialisation with coherence \( \mu = 0.52 \), leading to 16 missing atoms. The accompanying
Matlab toolbox provides more examples of these bad initializations to observe convergence, play around with and inspire more evil constructions. Here we will follow a less destructive route and ask ourselves, whether excluding the bad situation of two estimated atoms pointing to the same generating atom is sufficient to guarantee good behavior of ITKrM.

3.2 When things go right

Going back to theory we can prove a refined theorem. To keep the flow of the paper we will state it in an informal version and refer the reader to the appendix for the exact statement and its proof.

**Theorem 1** Assume that the sparsity level of the training signals scales as $S \lesssim \mu(\Phi)^{-2} / \log K$ and that the number of training signals scales as $N \approx SK \log K$. Further, assume that the coherence and operator norm of the current dictionary estimate $\Psi$ satisfy,

$$\mu(\Psi) \lesssim \frac{1}{\log K} \quad \text{and} \quad \|\Psi\|_{2,2}^2 \lesssim \frac{K}{S \log K}. \quad (16)$$

If the distance of $\Psi$ to the generating dictionary $\Phi$ satisfies either

a) $\frac{1}{\sqrt{S}} \lesssim d(\Psi, \Phi) \lesssim \frac{1}{\sqrt{\log K}}$ or

b) $d(\Psi, \Phi) \gtrsim \frac{1}{\sqrt{\log K}}$ but the cross-Gram matrix $\Phi^* \Psi$ is diagonally dominant in the sense that

$$\min_k |\langle \phi_k, \psi_k \rangle| \gtrsim (\log K)^{3/2} \cdot \max \left\{ \mu(\Phi, \Psi), \mu(\Phi), \|\Phi\|_{2,2} \sqrt{S/(K \log K)} \right\}, \quad (17)$$

then one iteration of ITKrM will reduce the distance by at least a factor $\kappa < 1$, meaning

$$d(\Psi, \Phi) < \kappa \cdot d(\Psi, \Phi).$$

The first part of the theorem simply says that, excluding dictionaries $\Psi$ that are coherent or have large operator norm, ITKrM is a contraction on a ball of radius $1/\sqrt{\log K}$ around the generating dictionary $\Phi$. To better understand the second part of the theorem, we have a closer look at the conditions on the cross-Gram matrix $\Phi^* \Psi$ in (17). The fact that the diagonal entries have to be larger than $(\log K)^{3/2} \mu(\Phi)$ and $\|\Phi\|_{2,2} \log K \sqrt{S/K}$ puts a constraint on the admissible distance $d(\Phi, \Psi)$ via the relation $d(\Phi, \Psi)^2 = 2 - 2 \min_k |\langle \phi_k, \psi_k \rangle|$. For an incoherent dictionary with $\mu(\Phi) \approx 1/\sqrt{d}$ and $\|\Phi\|_{2,2} \approx K/d$ and a moderate sparsity level $S \geq \log K$ this means that

$$d(\Phi, \Psi) \lesssim \left( 2 - 2 \sqrt{\frac{S(\log K)^2}{d}} \right)^{1/2}. \quad (18)$$
Considering that the maximal distance between two dictionaries is $\sqrt{2}$, this is a large improvement over the admissible distance $1/\sqrt{\log K}$ in a). However, the additional price to pay is that also the intrinsic condition on the cross-Gram matrix needs to be satisfied,

$$\min_k |\langle \phi_k, \psi_k \rangle| \gtrsim (\log K)^{3/2} \cdot \max_{j \neq k} |\langle \phi_k, \psi_j \rangle|.$$  

This condition captures our intuition that no two estimated atoms should point to the same generating atom and provides a bound for sufficient separation.

One thing that has to be noted about the result above is that it does not guarantee convergence of ITKrM since it is only valid for one iteration. To prove convergence of ITKrM, we need to additionally prove that $\bar{\Psi}$ inherits from $\Psi$ the properties that are required for being a contraction, which is part of our future goals. Still, the result goes a long way towards explaining the good convergence behaviour of ITKrM.

For example, it allows us to briefly sketch why the algorithm always converges in experiments where the initial dictionary is a large but random perturbation of a well-behaved generating dictionary $\Phi$ with coherence $\mu(\Phi) \approx 1/\sqrt{d}$ and operator norm $\|\Phi\|_2^2 \approx K/d$. If $\psi_k = \alpha_k \phi_k + \omega_k z_k$, where the perturbation vectors $z_k$ are drawn uniformly at random from the unit sphere orthogonal to $\phi_k$, then with high probability for all $j \neq k$ we have

$$|\langle \phi_k, z_j \rangle| \lesssim \sqrt{\log K/d} \quad \text{and} \quad |\langle z_k, z_j \rangle| \lesssim \sqrt{\log K/d} \quad \text{for all possible } \alpha_k$$

and consequently for all possible $\alpha_k$

$$\mu(\Psi) \lesssim \sqrt{4 \log K/d} \quad \text{and} \quad \mu(\Phi, \Psi) \lesssim \sqrt{2 \log K/d}. \quad (21)$$

Also with high probability the operator norm of the matrix $Z = (z_1, \ldots, z_K)$ is bounded by $\|Z\|_{2,2} \lesssim \sqrt{\log K}$, [12], so that for $\Psi$ we get $\|\Psi\|_{2,2} \lesssim \sqrt{K/d} + \sqrt{\log K}$, again independent of $\alpha_k$. Comparing these estimates with the requirements of the theorem we see that for moderate sparsity levels, $S \geq \log K$, we get a contraction whenever

$$\alpha_{\min} \gtrsim \sqrt{S(\log K)^3/d} \quad \iff \quad d(\Phi, \Psi) \lesssim \left(2 - 2\sqrt{S(\log K)^3/d}\right)^{1/2}. \quad (22)$$

Summarising the two last subsections we see that ITKrM has convergence problems if the current dictionary estimate is too coherent, has large operator norm or if two atoms are close to one generating atom. Both coherence and operator norm of the estimate could be calculated after each iteration to check whether ITKrM is going in a good direction. Unfortunately, the diagonal dominance of the cross-Gram matrix, which prevents two estimated atoms to be close to the same generating atom, cannot be verified. However, the most likely outcome of this situation is that both these estimated atoms converge to the same generating atom, meaning that eventually the estimated dictionary will be coherent. This suggests that in order to improve the global convergence behaviour of ITKrM, we should control the coherence of the estimated dictionaries. One strategy to incorporate incoherence into ITKrM could be adding a penalty for coherent dictionaries. The main disadvantages of this strategy, apart from the fact that ITKrM is not straightforwardly associated to an optimisation programme, are the computational cost and the fact that
penalties tend to complicate the high-dimensional landscape of basins of attractions which slows up convergence. Therefore, we will use a different strategy which allows us to keep the high percentage of correctly recovered atoms and even use the information they provide for identifying the missing ones: replacement.

4. Replacement

Replacement of coherent atoms with new, randomly drawn atoms is a simple clean-up step that most dictionary learning algorithms based on alternating minimisation, e.g. K-SVD [2], employ additionally in each iteration. While randomly drawing a replacement is cost-efficient and democratic, the drawback is that the new atom converges only very slowly to the missing generating atom. Thinking back to our example of a dictionary with one double atom, $\psi_1 = \psi_2 = \phi_1$ and one 1:1 atom, $\psi_3 \propto \phi_2 + \phi_3$, we immediately see the reason why. A randomly drawn replacement $\psi_{\text{new}}$ for $\psi_2$ will be quite incoherent to all generating atoms, meaning $|\langle \phi_k, \psi_{\text{new}} \rangle| \lesssim \sqrt{2\log(2K)/d}$ for all $k$. This means that the only time it has a chance to be picked is when the signal contains the rare constellation $\phi_2 - \phi_3$. Unfortunately, with high probability $\psi_{\text{new}}$ is also incoherent to this linear combination and so might actually never be picked. Looking on the bright side, we see that once it is picked, the updated atom $\tilde{\psi}_2$ will be very close to $\phi_2 - \phi_3$ since we have $y - P(\Psi I) y \approx (\phi_2 - \phi_3)$ and $|\langle y, \psi_{\text{new}} \rangle| \lesssim \sqrt{2\log(2K)/d}$. Thus in the next iteration the updated atom $\tilde{\psi}_2 \approx (\phi_2 - \phi_3)/\sqrt{2 - 2\theta}$ will be serious competition for $\tilde{\psi}_3 \approx (\phi_2 + \phi_3)/\sqrt{2 + 2\theta}$ in the thresholding of all signals containing either $\phi_2$ or $\phi_3$. This iteration will then create a first imbalance of the ratio between $\phi_2$ and $\phi_3$ within one or both of the estimated atoms, making one the more likely choice for $\phi_2$ and the other the more likely choice for $\phi_3$ in the subsequent iteration. There the imbalance will be further increased until a few iterations later we finally have $\psi_2 \approx \phi_2$ and $\psi_3 \approx \phi_3$ or the other way around.

The only problem is that in practical situations, where not all other atoms have already converged or the signals contain noise, the initial convergence of the randomly drawn atom to $\psi_2 - \psi_3$ might be very slow. This is because the atom is not updated at all or updated from correct but rarely occurring and noisy signals ($2, 3 \in I$) and equally many or more false positives ($2, 3 \notin I$).

Thus the natural next question is whether we can do better than a random replacement. To find a smarter strategy we again look back to our analysis of thresholding. Going through the various cases we see that whenever the residual $a = y - P(\Psi I) y$ is not zero, it is most likely to be proportional to $\phi_2 - \phi_3$. To be more precise, we have $a \approx \pm (\phi_2 - \phi_3)/2$ with probability about $\frac{2S}{K}$ compared to $a \approx \phi_k$ for $k > 3$ with probability about $\frac{S}{K^2}$. Extending our thought experiment to situations with several doubles and 1:1 combinations $\phi_{i_1} + h_i \phi_{i_2}$, we see that the residuals tend to be 1-sparse in the 1:1 complements, $\phi_{i_1} - h_i \phi_{i_2}$.

This suggests that we can directly recover the complements $\phi_{i_1} - h_i \phi_{i_2}$, by running ITKrM, which for $S = 1$ reduces to ITKsM, on the residuals. Concretely, we choose the number $L \ll K$ of candidate atoms, meaning the maximal number of atoms we can replace after each iteration, initialise a $d \times L$ dictionary $\Gamma = (\gamma_1 \ldots \gamma_L)$ of candidates and in each iteration of ITKrM add the following clean-up steps. For all signals we find $i_n = \arg \max_{\ell} |\langle \gamma_\ell, a_n \rangle|$, where $a_n = y_n - P(\Psi I_n) y_n$ and update the candidate atoms as $\bar{\gamma}_\ell = \sum n; i_n = \ell a_n \cdot \text{sign}(\langle \gamma_\ell, a_n \rangle)$ with subsequent normalisation.
The advantage of this strategy over even simpler approaches such as choosing the largest residual or principal components of the worst approximated residuals, [33, 30], is that it is still sequential and memoryless, robust to outliers in the training signals, allows to replace several atoms at once and can already work in early stages of the algorithm. As we have argued in the last section, the 1:1 atoms may be a secondary effect to the double atoms and are created because the missing generating atom \( \phi_k \) will start to attract the estimator of the generating atom to which it is most coherent, that is \( \bar{\psi}_k \), resulting in \( \psi_k \sim \phi_k + \phi_\bar{k} \). Repeating the thresholding analysis above for a recovered dictionary of the form \( \Psi = (\phi_1, \phi_1, \phi_3, \ldots, \phi_K) \), shows that in this case the residuals actually encode the missing atom \( \phi_2 \). This means that in early stages, where we have some very coherent atoms on one hand and consequently must have some very badly approximated atoms on the other hand, the residuals are 1-sparse in these badly approximated atoms and our strategy will recover them directly.

Now that we have laid out the basic strategy, it remains to deal with all the details. For instance, if we have used all replacement candidates after one iteration, after the next iteration the replacement candidates might not be mature yet, meaning they might not have converged yet. To solve this problem, observe that the number of replacement candidates will be much smaller than the dictionary size, \( L \ll K \). Therefore, we need less training signals per iteration to learn the candidates or equivalently we can update \( \Gamma \) more frequently, meaning we renormalise after each batch of \( N_\Gamma < N \) signals and set \( \Gamma = \bar{\Gamma} \). Like this, every augmented iteration of ITKrM will produce \( L \) replacement candidates.

The next questions concern the actual replacement procedure. Assume we have fixed a threshold \( \mu_{\text{max}} \) for the maximal coherence. If our estimate \( \Psi \) contains two atoms whose mutual coherence is above the threshold, \( |\langle \psi_k, \psi_{k'} \rangle| > \mu_{\text{max}} \), which atom should we replace? One strategy that has been employed for instance in the context of analysis operator learning, [11], is to average the two atoms, that is to set \( \psi_{k'}^{\text{new}} = \psi_k + \text{sign}(\langle \psi_k, \psi_{k'} \rangle)\psi_{k'} \). The reasoning is that if both atoms are good approximations to the generating atom \( \phi_k \) then their average will be an even better approximation. However, if one atom \( \psi_k \) is already a very good approximation to the generating atom \( \psi_k \approx \psi_k \) while \( \psi_{k'} \approx \mu_{\text{max}} \phi_k + \sqrt{1 - \mu_{\text{max}}^2}z_k \), then the averaged atom will be a worse approximation than \( \psi_k \) and it would be preferable to simply keep \( \psi_k \). To determine which of two coherent atoms is the better approximation, we note that the better approximation to \( \phi_k \) should be more likely to be selected during thresholding. This means that we can simply count how often each atom is contained in the thresholded supports \( I^n_t \),

\[
v(k) = \sharp \{n : k \in I^n_t\}
\]

and in the case of two coherent atoms keep the more frequently used one. Based on the value function \( v \) we can also employ a weighted merging strategy and set \( \psi_k^{\text{new}} = v(k)\psi_k + \text{sign}(\langle \psi_k, \psi_{k'} \rangle)v(k')\psi_{k'} \). If both atoms are equally good approximations, then their value functions should be similar and the balanced combination will be a better approximation. If one atom is a much better approximation it will be used much more often and the merged atom will correspond to this better atom.

Having chosen how to merge two coherent atoms, we next need to decide which of our \( L \) replacement candidates we are going to use. To keep the dictionary incoherent, we first discard all candidates \( \gamma_\ell \), whose maximal coherence with the remaining dictionary atoms is larger than our threshold, that is, \( \max_k |\langle \gamma_\ell, \phi_k \rangle| \geq \mu_{\text{max}} \). Note that in a perfectly \( S \)-sparse setting this is not very likely since the residuals we are summing up contain only noise and
therefore add up to noise but might be a problem if we underestimate the sparsity level in the learning. If we use $\hat{S} < S$, the residuals are still $S - \hat{S}$ sparse in the dictionary so some of our replacement candidates might be near copies of already recovered atoms in the dictionary.

To decide which remaining candidate is likely to be the most valuable, we use a counter similar to the one for the dictionary atoms. However, we have to be more careful here since every residual is added to one candidate. If the residual contains only noise, which happens in most cases, and the candidates are reasonably incoherent to each other, then each candidate is equally likely to have its counter increased. This means that the candidate atom that actually encodes the missing atom (or 1:1 complement) will only be slightly more often used than the other candidates. So to better distinguish between good and bad candidates, we additionally employ a threshold $\tau$ and set $v_\Gamma(\ell) = \sharp\{n : \ell = i_n, |\langle \gamma_\ell, a_n \rangle| \geq \tau \|a_n\| \}$. To determine the size of the threshold, observe that for a residual consisting only of Gaussian noise, $a = r$, we have for any $\gamma_\ell$ the bound

$$P(|\langle \gamma_\ell, r \rangle| \geq \tau \|r\|_2) \leq 2 \exp\left(-\frac{d \tau^2}{2}\right).$$

(23)

which for $\tau = \sqrt{2 \log(2K)/d}$ becomes $1/K$. This means that the contribution to $v_\Gamma(\ell)$ from all the pure noise residuals is at best $N/K$. On the other hand, with probability $S/K$, the residual will encode the missing atom or 1:1 complement $a \approx (\phi_i - \phi_j) \cdot |x_i|/2$. For reasonable sparsity levels, $S \lesssim \frac{d}{\log(2K)}$, and signal to noise ratios, the candidate $\gamma_\ell$ closest to the missing atom will be picked and should have inner product of the size $|\langle \gamma_\ell, a \rangle| \approx |x_i|/2 \approx \frac{1}{\sqrt{S}} \geq \tau \|a\|_2$. This means that for a good candidate the value function will be closer to $NS/K$. The threshold should also help in the earlier mentioned case of underestimating the sparsity level. There one could imagine the candidates to be poolings of already recovered atoms, that is, $\gamma_\ell \approx \sum_{j \in J_\ell} \pm \phi_j / \sqrt{|J_\ell|}$, which are sufficiently incoherent to the dictionary atoms to pass the coherence test. If the residuals are homogenously $S - \hat{S}$ sparse in the original dictionary, the candidate atom $\gamma_\ell$ will be picked if $\phi_j$ approximates the residual best for $a_j \in J_\ell$. If additionally the sets $J_\ell$ are disjoint, atoms corresponding to a bigger atom pool are (up to a degree) more likely to be choosen than those corresponding to a smaller pool. The threshold helps favour candidates associated to small pools, which have bigger inner products, since $|\langle \gamma_\ell, a_n \rangle| \approx 1/\sqrt{S|J_\ell|}$. This is desirable since the candidate closest to the missing atom will correspond to a smaller pool. After all, a candidate containing in its pool the missing atom (1:1 complement) will be soon distorted towards this atom since the sparse residual coefficient of the missing atom will be on average larger than those of the other atoms, thus reducing the effective size of the pool.

Before implementing our new replacement strategy let us address another less frequently activated safeguard included in most dictionary learning algorithms: the handling of dictionary atoms that are never selected and therefore have a zero update. As for coherent atoms the standard procedure is replacement of such an atom with a random redraw, which however comes with the problems discussed above. Fortunately our replacement candidates again provide an efficient alternative. If an atom has never been updated, or more generally, if the norm of the new estimator is too small, we simply do not update this atom but set the associated value function to zero. After replacing all coherent atoms we then proceed...
to replace these unused atoms.
The combination of all the above considerations leads to the augmented ITKrM algorithm, which is summarised in Algorithm [C.1] while the actual procedure for replacing coherent atoms is described in Algorithm [C.2] both to be found in Appendix [C] With these details fixed, the next step is to see how much the invested effort will improve dictionary recovery.

4.1 Numerical Simulations

In this subsection we will verify that replacing coherent atoms improves dictionary recovery and test whether our strategy improves over random replacement. Our main setup is the following:

**Generating dictionary:** As generating dictionary \( \Phi \) we use a dictionary of size \( K = 192 \) in \( \mathbb{R}^d \) with \( d = 128 \), where the atoms are drawn i.i.d. from the unit sphere.

**(Sparse) training signals:** We generate \( S \)-sparse training signals according to our signal model in (24) as

\[
y = \frac{\Phi x_{c,p,\sigma} + r}{\sqrt{1 + \|r\|_2^2}}. \tag{24}
\]

For every signal a new sequence \( c \) is generated by drawing a decay factor \( q \) uniformly at random in \([0.9, 1]\) and setting \( c_i = c_q q^{i-1} \) for \( i \leq S \) and 0 else, where \( c_q := \frac{1-q}{1-q^S} \) so that \( \|c\|_2 = 1 \). The noise is centered Gaussian noise with variance \( \rho^2 = (16d)^{-1} \), leading to a signal to noise ratio of \( \text{SNR} = 16 \). We will consider two types of training signals. The first type consists of 6-sparse signals with 5% outliers, that is, we randomly select 5% of the sparse signals and replace them with pure Gaussian noise of variance \( 1/d^2 \). The second type consists of 25% 4-sparse signals, 50% 6-sparse signals and 25% 8-sparse signals, where again 5% are replaced with pure Gaussian noise. In each iteration of ITKrM we use a fresh batch of \( N = 120000 \) training signals. Unless specified otherwise, the sparsity level given to the algorithm is \( S_e = 6 \).

**Replacement candidates:** During every iteration of ITKrM we learn \( L = \lceil \log d \rceil = 5 \) replacement candidates using \( m = \lceil \log d \rceil = 5 \) iterations each with \( N_\Gamma = \lfloor N/m \rfloor \) signals.

**Initialisations:** The dictionary \( \Psi \) containing \( K \) atoms as well as the replacement candidates are initialised by drawing vectors i.i.d. from the unit sphere. In case of random replacement we use the initialisations of the replacement candidates. All our results are averaged over 20 different initialisations.

**Replacement thresholds:** We will compare the dictionary recovery for various coherence thresholds \( \mu_{\text{max}} \in \{0.5, 0.7, 0.9\} \), and all three combination strategies, adding, deleting and merging. We also employ an additional safeguard and replace atoms, which have not been used at all or which have energy smaller than 0.001 before normalisation, if after replacement of coherent atoms we have candidate atoms left.

**Recovery threshold:** We use the convention that a dictionary atom \( \phi_k \) is recovered if

\[
\max_j |\langle \phi_k, \psi_j \rangle| \geq 0.99.
\]

The results of our first experiment[3] which explores the efficiency of replacement using our candidate strategy in comparison to random or no replacement on 6-sparse signals as

3. As already mentioned all experiments can be reproduced using the matlab toolbox available at [https://www.uibk.ac.at/mathematik/personal/schnass/code/adl.zip](https://www.uibk.ac.at/mathematik/personal/schnass/code/adl.zip)
described above, are depicted in Figure 3. We can see that for all three considered coherence thresholds \( \mu_{\text{max}} \in \{0.5, 0.7, 0.9\} \), our replacement strategy improves over random or no replacement. So while after 100 iterations ITKrM without replacement misses about 1\% of the atoms and with random replacement about 0.1\%, it always finds the full dictionary after at worst 55 iterations using the candidate atoms. Contrary to random replacement the candidate based strategy also does not seem sensitive to the combination method. Another observation is that candidate replacement leads to faster recovery the lower the coherence threshold is, while the average performance for random replacement is slightly better for the higher thresholds. This is connected to the average number of replaced atoms in each run, which is around 16 for \( \mu_{\text{max}} = 0.5 \), around 3.8 for \( \mu_{\text{max}} = 0.7 \) and around 0.8 for \( \mu_{\text{max}} = 0.9 \), since for the candidate replacement there is no risk of replacing a coherent atom that might still change course and converge to a missing generating atom with something useless. For completeness sake, we also mention that in none of the trials replacement of unused atoms is ever activated.

In our second experiment we explore the performance of candidate replacement for the more interesting (realistic) type of signals with varying sparsity levels. Since the signals can be considered 4, 6 or 8 sparse we compare the performance of ITKrM using all three possibilities, \( S_e \in \{4, 6, 8\} \) and a fixed replacement threshold \( \mu_{\text{max}} = 0.7 \). The results are shown in Figure 4. As before, candidate replacement outperforms random or no replacement and leads to 100\% recovery in all cases. Comparing the speed of convergence we see that it is higher the lower the sparsity level is, so for \( S_e = 4 \) we get 100\% recovery after about 30 iterations, for \( S_e = 6 \) after about 65 iterations and for \( S_e = 8 \) after 75 iterations. This would suggest that for the best performance we should always pick a lower than average sparsity level. However, the speed of convergence for \( S_e = 4 \) comes at the price of precision, as can be seen in the small table below, which lists both the average distance \( d(\Psi, \Phi) \) of the recovered dictionaries from the generating dictionary after 100 iterations as well as the mean atom distances \( d_1(\Psi, \Phi) := \frac{1}{K} \sum_k \| \phi_k - \psi_k \|_2 \). For both distances there is an increase

| \( S_e \) | 4 | 6 | 8 |
|---------|---|---|---|
| \( d(\Psi, \Phi) \) | 0.0392 | 0.0312 | 0.0304 |
| \( d_1(\Psi, \Phi) \) | 0.0322 | 0.0256 | 0.0250 |
in precision, going from $S_e = 4$ to $S_e = 6$, but hardly any improvement by going from $S_e = 6$ to $S_e = 8$. This suggests to choose the average or a slightly higher than average sparsity level. Alternatively, to get the best of both worlds, one should start with a smaller sparsity level and then slowly increase to the average sparsity level. Unfortunately this approach relies on the knowledge of the average sparsity level, which in practice is unknown. Considering that also the size of the dictionary is unknown this can be considered a minor problem. After all, if we underestimate the dictionary size, this will limit the final precision more severely. Assume for instance that we set $K - 1$ instead of $K$. In this case recovering a dictionary $\Psi$ with $K - 2$ generating atoms plus one 1:1 combination of two generating atoms leading to $d(\Psi, \Phi) \gtrsim \frac{1}{2}$ is actually the best we can hope for. Therefore, in the next section we will use our candidate atoms to make the big step towards adaptive selection of both sparsity level and dictionary size.

5. Adaptive dictionary learning

We first investigate how to adaptively choose the sparsity level for a dictionary of fixed size.

5.1 Adapting the sparsity level

In the numerical simulations of the last section we have seen that the sparsity level $S$ given as parameter to the ITKrM algorithm influences both the convergence speed and the final precision of the learned dictionary.

When underestimating the sparsity level, meaning providing $S_e < S$ instead of $S$, the algorithm tends to recover the generating dictionary in less iterations than with the true sparsity level. Note also that the computational complexity of an iteration increases with $S_e$, so a smaller sparsity level leads to faster convergence not only in terms of iterations but also reduces the computation time per iteration. The advantage of overestimating the sparsity level, $S_e > S$ on the other hand, is the potentially higher precision, so the final error between the recovered and the generating dictionary (atoms), can be smaller than for the true level $S$. Intuitively this is due to the fact that for $S_e > S$, thresholding with the generating dictionary is more likely to recover the correct support, in the sense that $I \subset I^t$.

![Figure 4: Recovery rates of ITKrM without replacement, random and candidate replacement for various input sparsity levels $S_e$ and atom combination strategies, with coherence threshold $\mu_{\text{max}} = 0.7$.](image-url)
For a clean signal, \( y = \Phi I x_I \) this means that the residual is zero, so that the estimate of every atom in \( I^t \), even if not in \( I \), is simply reinforced by itself \( \langle \phi_i, y \rangle \phi_i \). However, in a noisy situation \( y = \Phi I x_I + r \), where the residual has the shape \( a = Q r \) the estimate of the additional atom \( i \in I^t / I \) is not only reinforced but also disturbed by adding noise in form of the residual once more than necessary. Depending on the size of the noise and the inner product this might not always be beneficial to the final estimate. Indeed, we have seen that for large \( S \), where the smallest coefficients in the support are already quite small, overestimating the support does not improve the final precision.

To further see that both under- and overestimating the sparsity level comes with risks, assume that we allow \( S + 1 \) instead of the true sparsity level \( S \) for perfectly sparse, clean signals. Then any dictionary, derived from the generating dictionary by replacing a pair of atoms \( (\phi_i, \phi_j) \) by \( (\hat{\phi}_i, \hat{\phi}_j) = A(\phi_i, \phi_j) \) for an invertible (well conditioned) matrix \( A \), will provide perfectly \( S + 1 \)-sparse representations to the signals and be a fixed point of ITKrM. Providing \( S - 1 \) instead of \( S \) can have even more dire consequences since we can replace any generating atom with a random vector and again have a fixed point of ITKrM. If the original dictionary is an orthonormal basis and the sparse coefficients have equal size in absolute value any such disturbed estimator even gives the same approximation quality. However, in more realistic scenarios, where we have coherence, noise or imbalanced coefficients and therefore the missing atom has the same probability as the others to be among the \( S - 1 \) atoms most contributing to a signal, the generating dictionary should still provide the smallest average approximation error. Indeed, whenever we have coherence, noise or imbalanced coefficients the signals can be interpreted as being \( 1 \)-sparse (with enormous error and minuscule gap \( c(1)/c(2) \)) in the generating dictionary, so learning with \( S_c = 1 \) should lead to a reasonable first estimate of most atoms. Of course if the signals are not actually \( 1 \)-sparse this estimate will be somewhere between rough, for small \( S \), and unrecognisable, for larger \( S \), and the question is how to decide whether we should increase \( S_c \). If we already had the generating dictionary, the simplest way would be to look at the residuals and see how much we can decrease their energy by adding another atom to the support. A lower bound for the decrease of a residual \( a \) can be simply estimated by calculating \( \max_k (|\langle \phi_k, a \rangle|^2) \).

If we have the correct sparsity level and thresholding recovers the correct support \( I^t = I \), the residual consists only of noise, \( a = Q(\Phi_I) (\Phi_I x_I + r) = Q(\Phi_I) r \approx r \). For a Gaussian noise vector \( r \) and a given threshold \( \theta \cdot \|r\|_2 \), we now estimate how many of the remaining \( K - S \) atoms can be expected to have inner products larger than \( \theta \cdot \|r\|_2 \) as

\[
E \left( n \{ k : |\langle r, \phi_k \rangle|^2 > \theta^2 \cdot \|r\|_2^2 \} \right) = \sum_k \mathbb{P} \left( |\langle r, \phi_k \rangle|^2 > \theta^2 \cdot \|r\|_2^2 \right) < 2(K - S) e^{- \frac{d \theta^2}{2}}. \tag{25}
\]

In particular setting \( \theta = \theta_K = \frac{1}{\sqrt{2 \log(4K)/d}} \) the expectation above is smaller than \( \frac{1}{2} \).

This means that if we take the empirical estimator of the expectation above, using the approximation \( r_n \approx a_n \), we should get

\[
\frac{1}{N} \sum_n n \{ k : |\langle a_n, \phi_k \rangle|^2 > \theta^2 \cdot \|a_n\|_2^2 \} \approx \frac{1}{2}, \tag{26}
\]

which rounds to zero indicating that we have the correct sparsity level.

Conversely, if we underestimate the correct sparsity level, \( S_c = S - m \) for \( m > 0 \), then
thresholding can necessarily only recover part of the correct support, \( I^t \subset I \). Denote the set of missing atoms by \( I^m = I/I^t \). The residual has the shape

\[
a = Q(\Phi_{I^t})(\Phi_{I^t}x_I + r) = Q(\Phi_{I^t})(\Phi_{I^m}x_{I^m} + r) \approx \Phi_{I^m}x_{I^m} + r
\]

For all missing atoms \( i \in I^m \) the squared inner products are approximately

\[
|\langle a, \phi_i \rangle|^2 \approx (x_i + \langle r, \phi_i \rangle)^2.
\]

Assuming well-balanced coefficients, where \( |x_i| \approx 1/\sqrt{S} \) and therefore \( \|\Phi_{I^m}x_{I^m}\|^2 \approx m/S \), a sparsity level \( S \lesssim \frac{d}{2\log(4K)} \), and reasonable noise levels, this means that with probability at least \( \frac{1}{2} \) we have for all \( i \in I^m \)

\[
|\langle a, \phi_i \rangle|^2 \gtrsim |x_i|^2 \gtrsim \frac{1}{2m}(\|\Phi_{I^m}x_{I^m}\|^2 + \|r\|^2) \gtrsim \theta_K^2 \|a\|^2_2,
\]

and in consequence

\[
\frac{1}{N} \sum_n \mathbb{1}\{k : |\langle a_n, \phi_k \rangle|^2 > \theta_K^2 \|a_n\|^2_2\} \gtrsim \frac{m}{2}.
\]

This rounds to at least 1, indicating that we should increase the sparsity level.

Based on the two estimates above and starting with sparsity level \( S_e = 1 \) we should now be able to arrive at the correct sparsity level \( S \). Unfortunately, the indicated update rule for the sparsity level is too simplistic in practice as it relies on thresholding always finding the correct support given the correct sparsity level. Assume that \( S_e = S \) but thresholding fails to recover for instance one atom, \( I^t = I_{i+1} \). Then we still have \( a = Q(\Phi_{I^t})(x_i \phi_i + r) \approx x_i \phi_i + r \) and \( |\langle \phi_i, a \rangle|^2 \gtrsim \theta_K^2 \|a\|^2_2 \). If thresholding constantly misses one atom in the support, for instance because the current dictionary estimate is quite coherent, \( \mu \gg 1/\sqrt{d} \), or not yet very accurate, this will lead to an increase \( S_e = S + 1 \). However, as we have discussed above, while increasing the sparsity level increases the chances for full recovery by thresholding, it also increases the atom estimation error which decreases the chances for full recovery. Depending on which effect dominates, this could lead to a vicious circle of increasing the sparsity level, which decreases the accuracy leading to more failure of thresholding and increasing the sparsity level. In order to avoid this risk we should take into account that thresholding might fail to recover the full support and be able to identify such failure. Further, we should be prepared to also decrease the sparsity level.

The key to these three goals is to also look at the coefficients of the signal approximation. Assume that we are given the correct sparsity level \( S_e = S \) but recovered \( I^t = I_{i+1} \). The corresponding coefficients \( \tilde{x}_{I^t} \) have the shape

\[
\tilde{x}_{I^t} = \Phi_{I^t}^\dagger(\Phi_{I^t}x_I + r) = \Phi_{I^t}^\dagger(\Phi_{I_{i+1}}x_{I_{i+1}} + \phi_i x_i + r)
\]

\[
= (x_{I_{i+1}}, 0) + (\Phi_{I^t}^\dagger \Phi_{I^t})^{-1} \Phi_{I^t}^\dagger (\phi_i x_i + r),
\]

meaning \( |\tilde{x}_{I^t}(j)|^2 \lesssim \mu^2 (1+S\mu)^2 |x_i|^2 + |\langle \phi_j, r \rangle|^2 \) or even \( |\tilde{x}_{I^t}(j)|^2 \lesssim \mu^2 |x_i|^2 + |\langle \phi_j, r \rangle|^2 \). Since the residual is again approximately \( a \approx \phi_i x_i + r \), this means that for incoherent dictionaries
the coefficient of the wrongly chosen atom is likely to be below the threshold \(\theta_K^2 \|a\|_2\), while the one of the missing atom will be above the threshold, so we are likely to keep the sparsity level the same.

Similarly if we overestimate the sparsity level \(S_e = S + 1\) and recover an extra atom \(I^i = I_{e-j}\), we have \(a = Q(\Phi_{I^j})r \approx r\) while the coefficient of the extra atom will be of size \(|\tilde{x}_{I^j}(j)|^2 \approx |\langle \phi_j, r \rangle|^2 < \theta^2 \|a\|_2^2\). All in all our estimates suggest that we get a more stable estimate of the sparsity level by averaging the number of coefficients \(\tilde{x}_{I^j} = \Phi_{I^j}^\dagger y\) and residual inner products \(|\langle \phi_i, r \rangle|_{i \notin I^j}\) that have squared value larger than \(\theta_K^2 \) times the residual energy. However, the last detail we need to include in our considerations is the reason for thresholding failing to recover the full support given the correct sparsity level in first place. Assume for instance, that the signal does not contain noise, \(y = \Phi_{I^j}x_I\) but that the sparse coefficients vary quite a lot in size. We know (from Appendix B or [7]) that in case of i.i.d. random coefficient signs, \(\mathbb{P}(\text{sign}(x_i) = 1) = 1/2\), the inner products of the atoms inside resp. outside the support concentrate around,

\[
\begin{align*}
    i \in I & \quad |\langle \phi_i, \Phi_{I^j}x_I \rangle| \approx |x_i| \pm \left(\sum_{k \neq i} x_k^2 |\langle \phi_i, \phi_k \rangle|^2\right)^{1/2} \approx |x_i| \pm \mu \|y\|_2 \\
    i \notin I & \quad |\langle \phi_i, \Phi_{I^j}x_I \rangle| \approx \left(\sum_{k} x_k^2 |\langle \phi_i, \phi_k \rangle|^2\right)^{1/2} \approx \mu \|y\|_2.
\end{align*}
\]

This means that we will only recover the atoms corresponding to the \(S_r\)-largest coefficients for \(S_r < S\), that is, \(I_r = \{i \in I : |x_i| \geq \mu \|y\|_2\}\). The good news is that these will capture most of the signal energy, \(\|P(\Phi_{I_r})y\|_2^2 \approx \|\Phi_{I_r}x_{I^r}\|_2^2 \approx \|y\|_2^2\), meaning that in some sense the signal is only \(S_r\) sparse. It also means that for \(\mu^2 \approx 1/d\) we can estimate the recoverable sparsity level of a given signal as the number of squared coefficients/residual inner products that are larger than

\[
\frac{1}{d} \|P(\Phi_{I^j})y\|_2^2 + \frac{2 \log(4K)}{d} \|Q(\Phi_{I^j})y\|_2^2.
\] (29)

If \(S_n\) is the estimated recoverable sparsity level of signal \(y_n\), a good estimate of the overall sparsity level \(S\) will be the rounded average sparsity level \(\overline{S} = \lfloor \frac{1}{n} \sum_n S_n \rfloor\). The corresponding update rule then is to increase \(S_e\) by one if \(\overline{S} > S_e\), keep it the same if \(\overline{S} = S_e\) and decrease it by one if \(\overline{S} < S_e\), formally

\[
S_{e_{\text{new}}} = S_e + \text{sign}(\overline{S} - S_e).
\] (30)

To avoid getting lost between numerical and explorative sections we will postpone an algorithmic summary and testing of our adaptive sparsity selection to Subsection 5.3 and next address the big question how to adaptively select the dictionary size.

### 5.2 Adapting the dictionary size

The common denominator of all popular dictionary learning algorithms, from MOD to K-SVD, is that before actually running them one has to choose a dictionary size. This choice might be motivated by a budget, such as being able to store \(K\) atoms and \(S\) values per signal, or application specific, that is, the expected number of sources in sparse source separation. In applications such as image restoration \(K\) (like \(S\)) is either chosen ad hoc or experimentally with an eye towards computational complexity, and one will usually find
$d \leq K \leq 4d$, and $S = \sqrt{d}$. If algorithms include some sort of adaptivity of the dictionary size, this is usually in the form of not updating unused atoms, a rare occurrence in noisy situations, and deleting them at the end. Also this strategy can only help if $K$ was chosen too large but not if it was chosen too small.

Underestimating the size of a dictionary obviously prevents recovery of the generating dictionary. For instance, if we provide $K - 1$ instead of $K$ the best we can hope for is a dictionary containing $K - 2$ generating atoms and a $1 : 1$ combination of the two missing atoms. The good news is that if we are using a replacement strategy one of the candidates will encode the $1 : 1$ complement, as in the situation discussed in the last section where we are given the correct dictionary size but had a double atom.

Overestimating the dictionary size does not prevent recovering the dictionary per se, but can decrease recovery precision, meaning that a bigger dictionary might not actually provide smaller approximation error. To get an intuition what happens in this case assume that we are given a budget of $K + 1$ instead of $K$ atoms and the true sparsity level $S$. The most useful way to spend the extra budget is to add a $1 : 1$ combination of two atoms, which frequently occur together, meaning $\phi_0 \propto \phi_i + h\phi_j$ for $h = \text{sign}(\langle \phi_i, \phi_j \rangle)$. The advantage of the augmented dictionary $\Psi = (\phi_0, \Phi)$ is that some signals are now $S - 1$ sparse. The disadvantage is that $\Psi$ is less stable since the extra atom $\phi_0$ will prevent $\phi_i$ or $\phi_j$ to be selected by thresholding whenever they are contained in the support in a $1 : h$ ratio. This disturbs the averaging process and reduces the final accuracy of both $\phi_i$ and $\phi_j$.

The good news is that the extra atom $\phi_0$ is actually quite coherent with the dictionary $|\langle \phi_0, \phi_{i(j)} \rangle| \geq 1/\sqrt{2}$, so if we have activated a replacement threshold of $\mu_{\text{max}} \leq 1/\sqrt{2}$, atom $\phi_0$ will be soon replaced, necessarily with another useless atom.

This suggests as strategy for adaptively choosing the dictionary size to decouple our replacement scheme into pruning and adding, which allows to both increase and decrease the dictionary size. We will first have a closer look at pruning.

**Pruning atoms:**

From the replacement strategy we can derive two easy rules for pruning: 1) if two atoms are too coherent, delete the less often used one or merge them, 2) if an atom is not used, delete it. Unfortunately, the second rule is too naive for real world signals, containing among other imperfections noise, which means also purely random atoms are likely to be used at least once by mistake. To see how we need to refine the second rule assume again that our sparse signals are affected by Gaussian noise (of a known level), that is, $y = \Phi_I x_I + r$ with $E(|r|^2) = \rho^2$ and that our current dictionary estimate has the form $\Psi = (\phi_0, \Phi)$ where $\phi_0$ is some vector with admissible coherence to $\Phi$. Whenever $\phi_0$ is selected this means that thresholding has failed. From the last subsection we also know that we have a good chance of identifying the failure of thresholding by looking at the coefficients $\Phi^\dagger_I (\Phi_I x_I + r)$.

The squared coefficient corresponding to the incorrectly chosen atom $\phi_0$ is likely to be smaller than $\lesssim \frac{\|\Phi_I x_I\|^2_2}{d} + |\langle \phi_0, r \rangle|^2$ while the squared coefficient of a correctly chosen atom $i \in I \cap I'$ will be larger than $|x_i|^2 + |\langle \psi_i, r \rangle|^2 \gtrsim \|\Phi_I x_I\|^2_2 / S + |\langle \phi_i, r \rangle|^2$ at least half the time. The size of the inner product of any atom with Gaussian noise can be estimated as

$$\mathbb{P}(|\langle \phi_k, r \rangle| > \tau \|r\|_2) \leq 2 \exp \left(-\frac{d \tau^2}{2} \right).$$

(31)
Taking again $\|P(\Phi_H)y\|_2$ as estimate for $\|\Phi_f x_i\|_2$ and $\|a\|_2 = \|Q(\Phi_H)y\|_2$ as estimate for $\|r\|_2$ we can define the refined value function $\tilde{v}(k)$ as the number of times an atom $\phi_k$ has been selected and the corresponding coefficient has squared value larger than $\|P(\Phi_H)y\|_2^2/d + \tau^2\|a_n\|_2^2$. Based on the bound above we can then estimate that for $N$ noisy signals the value function of the unnecessary or random atom $\phi_0$ is bounded by $\tilde{v}(0) \lesssim 2N \exp\left(-\frac{d\tau^2}{2}\right) := M$, leading to a natural criterion for deleting unused atoms. Setting for instance $\tau = \theta_K = \sqrt{2\log(4K)/d}$ we get $M = N/(2d)$. Alternatively, we can say that in order to accurately estimate an atom we need $M$ reliable observations and accordingly set the threshold to $\tau = \sqrt{2\log(2N/M)/d}$.

The advantage of a relatively high threshold $\tau \approx \sqrt{2\log(4K)/d}$ is that in low noise scenarios, we can also find atoms that are rarely used. The disadvantage is that for high $\tau$ the quantities $\tilde{v}(\cdot)$ we have to estimate are relatively small and therefore susceptible to random fluctuations. In other words, the number of training signals $N$ needs to be large enough to have sufficient concentration such that for unnecessary atoms the value function $\tilde{v}(\cdot)$ is actually smaller than $M$. Another consideration is that at the beginning, when the dictionary estimate is not yet very accurate, also the approximate versions of frequently used atoms will not be over the threshold often enough. This risk is further increased if we also have to estimate the sparsity level. If $S_e$ is still small compared to the true level $S$ we will overestimate the noise, and even perfectly balanced coefficients $1/\sqrt{S}$ will not yet be over the threshold. Therefore, pruning of the dictionary should only start after an embargo period of several iterations to get a good estimate of the sparsity level and most dictionary atoms.

Also we have seen in the replacement section that after replacing a double atom with the 1:1 complement $\phi_i - \phi_j$ of a 1:1 atom $\phi_i + \phi_j$, it takes a few iterations for the pair $(\phi_i \pm \phi_j)$ to rotate into the correct configuration $(\phi_i, \phi_j)$, where they are recovered most of the time. In the case of decoupled pruning and adding, we run the risk of deleting a missing atom or a 1 : 1 complement one iteration after adding it simply because it has not been used often enough. Therefore, every freshly added atom should not be checked for its usefulness until after a similar embargo period of several iterations, which brings us right to the next question when to add an atom.

**Adding atoms:**

To see when we should add a candidate atom to the dictionary, we have a look back at the derivation of the replacement strategy. There we have seen that the residuals are likely to be either 1-sparse in the missing atoms (or 1:1 complements of the atoms doing the job of two generating atoms), meaning $a \approx |x_i|/2(\phi_i - \phi_j)$ or in a more realistic situation $a \approx |x_i|/2(\phi_i - \phi_j) + r$, or zero, which again in the case of noise means $a \approx r$. To identify a good candidate atom we observe again that if the residual consists only of (Gaussian) noise we have for any vector/atom $\gamma_k$

$$P(\langle \gamma_k, r \rangle > \tau_r \|r\|_2) \leq 2 \exp\left(-\frac{d\tau_r^2}{2}\right). \tag{32}$$

If on the other hand the residual consists of a missing complement, the corresponding candidate $\gamma_i \approx (\phi_i - \phi_j)/\sqrt{2}$ should have $|\langle a, \gamma_i \rangle| \approx |x_i|/\sqrt{2} \gtrsim \|a\|_2 r$. This means that we can use a similar strategy as for the dictionary atoms to distinguish between useful and
useless candidates. In the last candidate iteration using \( N \) residuals, we count for each candidate atom \( \gamma_k \) how often it is selected and satisfies \( |\langle \gamma_k, a \rangle| > \tau \| a \|_2 \). Following the dictionary update and pruning we then add all candidates to the dictionary whose value function is higher than \( M = 2N \exp\left(-\frac{d\tau^2}{2}\right) \) and which are incoherent enough to atoms already in the dictionary.

Now, having dealt with all aspects necessary for making ITKrM adaptive, it is time to test whether adaptive dictionary learning actually works.

5.3 Experiments on synthetic data

We first test our adaptive dictionary learning algorithm on synthetic data\(^4\). The basic setup is the same as in Subsection 4.1. However, one type of training signals will again consist of 4, 6 and 8-sparse signals in a 1:2:1 ratio with 5% outliers, while the second type will consist of 8, 10 and 12-sparse signals in a 1:2:1 ratio with 5% outliers. Additionally, we will consider the following settings.

The minimal number of reliable observations \( M \) for a dictionary atom is set to either \( d, \lfloor d \log d \rfloor \) or \( \lfloor 2d \log d \rfloor \) with corresponding coefficient thresholds \( \tau = \sqrt{2 \log(2N/M)}/d \). For the candidate atoms the minimal number of reliable observations in the 4th (and last) candidate iteration is always set to \( M = d \).

The sparsity level is adapted after every iteration starting with iteration \( m = |\log d| = 5 \). The initial sparsity level is 1.

Promising candidate atoms are added to the dictionary after every iteration, starting again in the \( m \)-th iteration. In the last 3\( m \) iterations no more candidate atoms are added to the dictionary.

Coherent dictionary atoms are merged after every iteration, using the threshold \( \mu_{\text{max}} = 0.7 \). As weights for the merging we use the value function of the atoms from the most recent iteration.

Unused dictionary atoms are pruned after every iteration starting with iteration 2\( m \). An atom is considered unused if in the last \( m \) iterations the number of reliable observations has always been smaller than \( M \). Candidate atoms, freshly added to the dictionary, can only be deleted because they are unused at least \( m \) iterations later. In each iteration at most \( \lfloor d/5 \rfloor \) unused atoms are deleted, with an additional safeguard for very undercomplete dictionaries (\( K_e < d/10 \)) that at most half of all atoms can be deleted.

The initial dictionary is chosen to be either of size \( K_e = d = 128 \), \( K_e = 4d = 512 \) or the correct size \( K_e = K \), with the atoms drawn i.i.d. from the unit sphere as before. Figure 5 shows the results averaged over 10 trials each using a different initial dictionary.

The first observation is that all our effort paid off and that adaptive dictionary learning works. For the smaller average sparsity level \( S = 6 \), adaptive ITKrM always recovers all atoms of the dictionary and only overshoots and recovers more atoms for \( M = d \). The main difference in recovery speed derives from the size of the initial dictionary, where a larger dictionary size leads to faster recovery.

For the more challenging signals with average sparsity level \( S = 10 \), the situation is more diverse. So while the initial dictionary size mainly influences recovery speed but less the

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\(^4\) Again we want to point all interested in reproducing the experiments to the matlab toolbox available at [https://www.uibk.ac.at/mathematik/personal/schnass/code/adl.zip](https://www.uibk.ac.at/mathematik/personal/schnass/code/adl.zip)
Figure 5: Average recovery rates (top row) and dictionary sizes (bottom row) for adaptive dictionary learning based on ITKrM on signals with sparsity $S = \{4, 6, 8\}$ (left column) resp. $S = \{8, 10, 12\}$ (right column) in a 1:2:1 ratio for various initial dictionary sizes $K_e$ and required number of observations per atom $M$.

The final number of recovered atoms, the cut off threshold $M$ for the minimal number of reliable observations is critical for full recovery. So for $M = d$ adaptive ITKrM never recovers the full dictionary. We can also see that not recovering the full dictionary is strongly correlated with overestimating the dictionary size. Indeed, the higher the overestimation factor for the dictionary size is, the lower is the amount of recovered atoms. For example, for $M = d$, $K_e = 512$ the dictionary size is overestimated by a factor 1.5 and only about half of the dictionary atoms are recovered. To see that the situation is not as bad as it seems we have a look at the average sorted atom recovery error. That is, we sort the recovery errors $(d(\phi_k, \Psi))_k$ after 100 iterations in ascending order and average over the number of trials. The resulting curves are depicted in Figure 6. As we can see, overestimating the dictionary size degrades the recovery in a gentle manner. For the unrecovered atoms in case $M = d$, $K_e = 512$, the largest inner product with an atom in the recovered dictionary is below the cut-off threshold of 0.99 which corresponds to an error of size $\approx 0.14$ but for almost all of them it is still above 0.98 which corresponds to an error of 0.2. Also the oscillating recovery behaviour for $M = [d \log(d)]$ before the final phase, where no more atoms are added, can be explained by the fact that the worst approximated atoms have average best inner product very close to 0.99. So, depending on the batch of training signals in each iteration, their
inner product is below or above 0.99 and accordingly they count as recovered or not. In
general, we can see that the more accurate the estimate of the dictionary size is, the better
is the recovery precision of the learned dictionary. This is only to be expected. After all,
whenever thresholding picks a superfluous atom instead of a correct atom, the number of
observations for the missing atom is reduced and moreover, the residual error added to the
correctly identified atoms is increased.
The relative stability of these spurious atoms can in turn be explained by the fact that
$S = 10$ is at the limit of admissible sparsity for a generating dictionary with $\mu(\Phi) = 0.32$,
especially for sparse coefficients with a dynamic range of $0.9^{S-1} \approx 2.58$. In particular,
thresholding is not powerful enough to recover the full support, so the residuals still contain
several generating atoms. This promotes candidate atoms that are a sparse pooling of all
dictionary atoms. These poolings again have a good chance to be selected in the thresholding
and to be above the reliability threshold, thus positively reinforcing the effect. A quick look
at the estimated sparsity level as well as the average number of coefficients above the
threshold, or in other words, the average number of (probably) correctly identified atoms
in the support, denoted by $S_t$, also supports this theory. So for average sparsity level $S = 6$
the estimated sparsity level is $S_e = 6 = \lceil 5.7 \rceil$ and the average number of correctly identified
atoms is $S_t \approx 5$, regardless of the setting. This is quite close to the average number of
correctly identifiable atoms given $S_e = 6$, which is $0.95 \times (0.25 \times 4 + 0.75 \times 6) = 5.225$.
For average generating sparsity level $S = 10$ the table below lists $S_e : S_t$ for all settings.
We can see that even for the settings where the full dictionary is recovered, the estimated

|      | $d$ | $d \log(d)$ | $2d \log(d)$ |
|------|-----|-------------|--------------|
| 128  | 8 : 5.5 | 9 : 7.2 | 9 : 7.2 |
| 192  | 8 : 5.4 | 9 : 7.0 | 9 : 7.2 |
| 512  | 7 : 4.8 | 9 : 6.3 | 9 : 7.2 |

sparsity level is below 10 and the number of correctly identified atoms lags even more
behind. For comparison for $S_e = 9$ the average number of correctly identifiable atoms is
We will postpone a more in-depth discussion of how to further stabilise and improve adaptive

precision of the recovered dictionary. Also in order to estimate \( S_e \), we already have a good guess which atoms of the threshold support were correct and which atoms outside should have been included. This suggests to remove any atom from the support for which there is a more promising atom outside the support, or in other words, to update the support by thresholding (\( \Psi_{\hat{I}}^\dagger y, \Psi_{\hat{I}}^\dagger(I_d - P(\Psi_t)y) \)). Iterating this procedure until the support is stable is known as Hard Thresholding Pursuit (HTP), [16]. Using 2 iterations of HTP would not overly increase the computational complexity of adaptive ITKrM but could help to weed out spurious atoms. Also by not keeping the \( S_e \) best atoms but only those above the threshold \( \tau \) one could deal with varying sparsity levels, which would increase the final precision of the recovered dictionary.

Such a strategy might also help in addressing the only case where we have found our adaptive dictionary learning algorithm to fail spectacularly. This is - at first glance surprisingly - the most simple case of exactly 1-sparse signals and an initial dictionary size smaller than the generating size. At second glance it is not that surprising anymore. In case of underestimating the dictionary size \( K - K_e = K_m > 0 \) the best possible dictionary consists of \( K - 2K_m \) generating atoms and \( K_m \) 1:1 combinations of 2 non-orthogonal atoms of the form \( \phi_{ij} = (\phi_i + h\phi_j)/\alpha_{ij} \), where \( h = \text{sign}(\phi_i, \phi_j) \) and \( \alpha_{ij} = \sqrt{2 + 2|\langle \phi_i, \phi_j \rangle|} \).

In such a situation the (non-zero) residuals are again 1-sparse in the 1:1 complements \( \tilde{\phi}_{ij} = (\phi_i - h\phi_j)/\tilde{\alpha}_{ij} \), where \( \tilde{\alpha}_{ij} = \sqrt{2 - 2|\langle \phi_i, \phi_j \rangle|} \), and so the replacement candidates will be the 1:1 complements. However, the problem is that \( \tilde{\phi}_{ij} \) is never picked by thresholding since for both \( y \approx \phi_i \) and \( y \approx \phi_j \) the inner product with \( \phi_{ij} \) is larger,

\[
|\langle \phi_{ij}, \phi_i \rangle| = \sqrt{\frac{1 + |\langle \phi_i, \phi_j \rangle|}{2}} > \sqrt{\frac{1 - |\langle \phi_i, \phi_j \rangle|}{2}} = |\langle \tilde{\phi}_{ij}, \phi_i \rangle|.
\]

Still the inner product of \( \tilde{\phi}_{ij} \) with the residual has magnitude \( \approx 1/2 > \tau \) and so would be included in the support in a second iteration of HTP, thus keeping the chance that the pair \( (\phi_{ij}, \tilde{\phi}_{ij}) \) rotates into the correct configuration \( (\phi_i, \phi_j) \) alive.

We will postpone a more in-depth discussion of how to further stabilise and improve adaptive

\[
0.95 \times (0.25 \times 8 + 0.75 \times 9) = 8.3125.
\]
dictionary learning to the discussion in Section 6. Here we will first check whether adaptive dictionary learning is robust to reality by testing it on image data.

5.4 Experiments on image data

In this subsection we will learn dictionaries for the images Mandrill and Peppers. The training signals are created as follows. Given a $256 \times 256$ image, we contaminate it with Gaussian noise of variance $\rho^2 = \rho^2/255$ for $\rho^2 \in \{0, 5, 10, 15, 20\}$. From the noisy image we extract all $8 \times 8$ patches (sub-images), vectorise them and remove their mean. In other words, we assume that the constant atom, $\phi_0(k) = 1/8$, is always contained in the signal, remove its contribution and thus can only learn atoms that are orthogonal to it.

The set-up for adaptive dictionary learning is the same as for the synthetic data, taking into account that for the number of candidates $L$ and the memory $m$, we have $L = m = \lceil \log(d) \rceil = 4$, since the signals have dimension $d = 64$. Also based on the lesson learned on the more complicated data set with average sparsity level $S = 10$, we only consider as minimal number of observations $M = \lceil d \log(d) \rceil$ and $M = 2d \log(d)$. The initial dictionary size $K_e$ is either 8, 64 or 256 and in each iteration we use all available signals, $N = 62001$. All results are averaged over 10 trials, each using a different initial dictionary and - where applicable - a different noise-pattern. In the first experiment we compare the sizes of the

![Figure 7: Average dictionary sizes for adaptive dictionary learning based on ITKrM on all patches of Mandrill/Peppers for various initial dictionary sizes $K_e$ and required number of observations per atom $M$ (left). Average sparse approximation error of all patches of Mandrill/Peppers using OMP and the learned dictionaries with $K_e = 64$ and both choices of $M$ (right).](image)

dictionaries learned on both clean images with various parameter settings as well as their approximation powers. The approximation power of a dictionary augmented by flat atom $\phi_0$ for a given sparsity level $S$ is measured as $\|Y - \tilde{Y}\|_2^2/\|Y\|_2^2$, where $\tilde{Y} = (\tilde{y}_1, \ldots, \tilde{y}_n)$ and $\tilde{y}_n$ is the $S$-sparse approximation to $y_n$ calculated by Orthogonal Matching Pursuit, [31]. The results are shown in Figure 7. We can see that as for synthetic data the final size of the learned dictionary does not depend much on the initial dictionary size, but does depend on the minimal number of observations. So for $M = \lceil d \log(d) \rceil$ the average dictionary size is about 106 atoms for Mandrill and 55 atoms for Peppers, while for $M = 2d \log(d)$ we have about 54 atoms on Mandrill and 36 atoms on Peppers. The estimated sparsity level vs.
average number of correctly identified atoms for Mandrill is $S_e = \lfloor 2.1 \rfloor = 2$ vs. $S_t \approx 1.5$ and for Peppers $S_e = \lfloor 2.9 \rfloor = 3$ vs. $S_t \approx 2.25$. Comparing the approximation power, we see that for both images the smaller (undercomplete) dictionaries barely lag behind the larger dictionaries. The probably most interesting aspect is that despite being smaller, the Peppers-dictionaries lead to smaller error than the Mandrill-dictionaries. This confirms the intuition that the smooth image Peppers has a lot more sparse structure than the textured image Mandrill. To better understand why for both images the larger dictionaries do not improve the approximation much, we have a look at the number of reliable observations for each atom in the last trial of $K_e = 64$ in Figure 8. The corresponding dictionaries for Mandrill/Peppers can be found in Figures 9/10.

We can see that for both images the number of times each atom is observed strongly

![Figure 8: Final number of reliable observations of the atoms in the dictionaries learned on Mandrill/Peppers with initial dictionary size $K_e = 64$ in the last trial.](image)

varies. If we interpret the relative number of observations as probability of an atom to be used, the first ten atoms are more than 10 times more likely to be used/observed than the last ten atoms. This accounts for the fact that by increasing the threshold for the minimal number of observations, we reduce the dictionary size without much affecting the approximation power.

In our second experiment we learn adaptive dictionaries on Mandrill contaminated with Gaussian noise of variance $\tilde{\rho}^2 = \rho^2/255$ for $\sigma^2 \in \{5, 10, 15, 20\}$, corresponding to average peak signal to noise ratios $\{34.15, 28.13, 24.61, 22.11\}$. We again compare their sizes and their approximation power for the clean image patches. The results are shown in Figure 12 and two example dictionaries are shown in Figure 11.

We can see that the size of the dictionary decreases quite drastically with increasing noise, while the approximation power degrades only very gently. The sparsity level chosen by the algorithm is $S_e = \lfloor 1.80 \rfloor = 2$ for $\sigma^2 = 5$ and $S_e = \lfloor 1.11 \rfloor = \lfloor 0.89 \rfloor = 1$ for $\sigma^2 \in \{15, 20\}$. For $\sigma^2 = 10$ the average recoverable sparsity level is $\approx 1.5$ so that for all trials the estimated sparsity level $S_e$ alternates between 1 and 2 in consecutive iterations. The fact that with increasing noise both the dictionary size and the sparsity level decrease but not the approximation power indicates that less often used atoms also tend to capture less energy per observation. This suggests an alternative value function, where each reliable observation is additionally weighted, for instance, by the squared coefficient or inner product. The
Figure 9: Dictionaries learned on Mandrill with initial dictionary size $K_e = 64$ and required number of observations $M = \lceil d \log(d) \rceil$ (top) resp. $M = 2d \log(d)$ (bottom).

Figure 10: Dictionaries learned on Peppers with initial dictionary size $K_e = 64$ and required number of observations $M = \lceil d \log(d) \rceil$ (middle) resp. $M = 2d \log(d)$ (right).

Figure 11: Dictionaries learned on the Mandrill image contaminated with Gaussian noise of variance $\tilde{\rho}^2 = 10/255$ (left) and $\tilde{\rho}^2 = 20/255$ (right), initial dictionary size $K_e = 64$ and required number of observations $M = \lceil d \log(d) \rceil$. 
Figure 12: Average dictionary sizes for adaptive dictionary learning based on ITKrM on all patches of the Mandrill image contaminated with Gaussian noise of variance $\tilde{\rho}^2 = \rho^2 / 255$, initial dictionary size $K_e = 64$ and required number of observations $M = \lceil d \log(d) \rceil$ (left). Corresponding average sparse approximation error of all clean patches of Mandrill using OMP and the learned dictionaries. (right).

Now that we have seen that adaptive dictionary learning produces sensible and noise robust results not only on synthetic but also on image data, we will turn to a final discussion of our results.

6. Discussion

In this paper we have studied the global convergence behaviour of the ITKrM (Iterative Thresholding and K residual means) algorithm for dictionary learning. We have characterised stable fixed points of ITKrM that are not equivalent to a generating dictionary. Further, we have proved that ITKrM contracts a dictionary estimate $\Psi$ towards the generating dictionary $\Phi$ whenever the cross-Gram matrix $\Psi^\star \Phi$ is diagonally dominant and $\Psi$ is incoherent and well-conditioned.

Using our insights that the stable fixed points of ITKrM always contain several atoms twice, meaning they are coherent, and that the residuals contain information about the missing atoms, we have developed a heuristic for finding good candidates, which we can use to replace one of two coherent atoms in a dictionary estimate. Simulations on synthetic data showed that replacement using these candidates improved over random or no replacement, always leading to recovery of the full dictionary.

Armed with replacement candidates, we then addressed one of the most challenging problems in dictionary learning, how to automatically choose the sparsity level and dictionary size. We developed a strategy for adapting the sparsity level from the initial guess $S_e = 1$ and the dictionary size by decoupling replacement into pruning of coherent and unused atoms and adding of promising candidates. The resulting adaptive dictionary learning algorithm was
shown to perform very well in recovering a generating dictionary from random initialisations with various sizes on synthetic data with sparsity levels \( S \geq 2 \) and in learning meaningful dictionaries on image data.

One improvement suggested by the numerical experiments, in order to reduce the number of spurious atoms and to increase the precision for large generating sparsity levels \( S_t \), is to keep track of both the average sparsity level \( \bar{S} \) and the average number of correctly identified atoms \( S_t \). If \( \bar{S} \) has been stable for several iterations but \( S_t \) lags behind, this is an indication that thresholding fails consistently or that the sparsity level of the signals is quite variable. The solution to both problems is to use 2 iterations of an iterative type of thresholding, known as Hard Thresholding Pursuit (HTP), [16], and to include all reliable atoms - with coefficients or inner product above the threshold \( \tau \) - in the support.

If the signals are compressible, meaning the sparse coefficients are rapidly decaying, this approach can be taken one step further and more iterations can be used. This could lead to greedy dictionary learning, which might be the sensible approach to signal classes which are sparse in a dictionary where the atoms are used with different probabilities and their coefficients differ in average magnitude. This greedy approach should be able to avoid the concentration of many coherent atoms in high energy subspaces by allowing to first recover atoms that are often used and have high coefficients and then gradually less often used atoms with smaller coefficients. To better balance frequency of occurrence with magnitude of coefficients, we could also use a different value function for the dictionary atoms. The one proposed here is based on the assumption that all atoms are used equally often, which for image data was clearly not the case. Alternatively, the current function could be scaled, that is, every reliable occurrence of an atom is weighted with the squared coefficient, which is related to the signal energy lost without this atom. Preliminary experiments using this weighted value function and an accordingly scaled cut-off indicate that it indeed helps to remove spurious atoms on synthetic data and leads to smaller dictionaries with the same approximation power on image data.

In this context it will also be interesting to study the behaviour of ‘freeloader’ atoms that are sparse combinations of the (most often used) generating dictionary atoms and as such are likely to be picked by thresholding without contributing much to the approximation. This question leads us directly to ongoing and future research concerning theory.

Currently we are trying to further relax the contraction conditions by reducing all \( \log K \) to \( \log S \) factors, while at the same time studying the behaviour of ITrKrM given the correct dictionary size \( K \) but a smaller sparsity level \( S_e < S \). The next big goal will be to study the cross-Gram matrix of a (well-conditioned, incoherent) dictionary with a randomly drawn dictionary. We expect that for most but not all dictionary atoms there will be a one to one correspondence to a random atom. Estimating the percentage of these atoms and proving that for these pairs the random atom will converge to the corresponding generating atom, (starting from \( S_e = 1 \)), will anchor our replacement in theory. The final step would be to show that if \( K_e \) is large enough that most signals can be considered 1-sparse in a subdictionary of the generating dictionary of size \( K_e \), then convergence to the subdictionary will still hold, thus anchoring adaptive dictionary learning in theory. Once this analysis has been done under the assumption that all atoms are used with the same frequency and average magnitude, it will be interesting to study dictionaries where the atoms are used with different frequencies and average magnitudes. In particular, if two atoms are frequently
used, how can we avoid a 1:1 combination of these atoms.
Another set of questions, less relevant to the convergence radius but to the final precision,
is the average case performance of sparse approximation/compressed sensing algorithms,
such as Hard Thresholding Pursuit (HTP), [16], starting with 2 iterations or (Stagewise) Orthogonal Matching Pursuit OMP, [31, 13], for perturbed dictionaries. Stagewise OMP is particularly interesting since it can take several atoms at a time, thus being as fast as HTP, and because the support size can be chosen adaptively - according to the residual error or for dictionary learning the size of the next largest inner product - like OMP.

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Appendix A. Exact Statement and Proof of Theorem 1

We only state and prove the exact version of the second part, since the first part consists literally of considering just one iteration of ITKrM and replacing in Theorem 4.2 of [37] the assumption on the distance $d(\Psi, \Phi)$ with the assumptions on the coherence and the operator norm of $\Psi$, ie.

$$d(\Psi, \Phi) \leq \frac{1}{32\sqrt{S}} \quad \text{and} \quad \mu(\Psi) \leq \frac{1}{18 \log K} \quad \text{and} \quad \|\Psi\|_{2,2}^2 \leq \frac{K}{108e^2S\log K} - 1. \quad (34)$$

Similarly the amendment to the proof consists in using Lemma 5 instead of Lemma B.8 and potentially some tweaking of constants.

Proposition 2 (Theorem 1(b) exact) Assume that the signals $y_n$ follow model (24) for coefficients with gap $c(S+1)/c(S) \leq \gamma_{\text{gap}}$, dynamic sparse range $c(1)/c(S) \leq \gamma_{\text{dyn}}$, noise to coefficient ratio $\rho/c(S) \leq \gamma_{\rho}$ and relative approximation error $\|c(S')\|/c(1) \leq \gamma_{\text{app}} \leq \log K$. Further, assume that the coherence and operator norm of the current dictionary estimate $\Psi$ satisfy,

$$\mu(\Psi) \leq \frac{1}{18 \log K} \quad \text{and} \quad \|\Psi\|_{2,2}^2 \leq \frac{K}{108e^2S\log K} - 1. \quad (35)$$

If $d(\Psi, \Phi) \geq \frac{1}{32\sqrt{S}}$ but the cross Gram matrix $\Phi^*\Psi$ is diagonally dominant in the sense that

$$\min_k |\langle \psi_k, \phi_k \rangle| \geq \max_k \left\{ 24 \gamma_{\text{gap}} \cdot \max_k |\langle \psi_k, \phi_k \rangle| , \right.$$  

$$68 \gamma_{\rho} \cdot \sqrt{\log K} ,$$  

$$78 \gamma_{\text{dyn}} \cdot (\log K)^{3/2} \max \{ \mu(\Phi), \mu(\Phi, \Psi) \},$$  

$$87 \gamma_{\text{dyn}} \cdot \log K \sqrt{(S+1)(\|\Phi\|_{2,2}^2+1)/K} \right\} , \quad (36)$$
then one iteration of ITKrM using \( N \) training signals will reduce the distance by at least a factor \( \kappa \leq 0.96 \), meaning \( d(\bar{\Psi}, \Phi) \leq 0.96 \cdot d(\Psi, \Phi) \), except with probability

\[
3K \exp \left( \frac{-NC_r^2 \gamma_{1,S}^2 \cdot \varepsilon}{768K \max\{S, B+1\}^{3/2}} \right) + 2K \exp \left( \frac{-NC_r^2 \gamma_{1,S}^2 \cdot \varepsilon^2}{512K \max\{S, B+1\} (1 + d_r^2)} \right). \tag{37}
\]

**Proof** We follow the outline of the proof for Theorem 4.2 in [37]. However, to extend the convergence radius we need to introduce new ideas, first for bounding the difference between the oracle residuals based on \( \Psi \) and \( \Phi \), replacing Lemma B.8 of [37], and second for bounding the probability of thresholding with \( \Psi \) not recovering the generating support or preserving the generating sign, replacing Lemma B.3/4 of [37]. We denote the thresholding residual based on \( \Psi \) by

\[
R^l(\Psi, y_n, k) := [y_n - P(\Psi_{I_n})y_n + P(\psi_k)y_n] \cdot \text{sign}(\langle \psi_k, y_n \rangle) \cdot \chi(I_n^\Psi, k), \tag{38}
\]

and the oracle residual based on the generating support \( I_n = p_n^{-1}(S) \), the generating signs \( \sigma_n \) and \( \Psi \), by

\[
R^o(\Psi, y_n, k) := [y_n - P(\Psi_{I_n})y_n + P(\psi_k)y_n] \cdot \sigma_n(k) \cdot \chi(I_n, k). \tag{39}
\]

Abbreviating \( s_k = \frac{1}{N} \sum_n \langle y_n, \phi_k \rangle \cdot \sigma_n(k) \cdot \chi(I_n, k) \) and \( \mathcal{F} = \{ y : R^l(\Psi, y, k) \neq R^o(\Psi, y, k) \} \) and setting \( B := \|\Phi\|_2^2 \) as well as \( \varepsilon := d(\Psi, \Phi) \) for conciseness, we know from the proof of Theorem 4.2 in [37] that

\[
\|\bar{\psi}_k - s_k \phi_k\|_2 \leq \frac{2\sqrt{B+1}}{N} \cdot \mathbb{1}\{n : y_n \in \mathcal{F}\} + \frac{1}{N} \left\| \sum_n [R^o(\Psi, y_n, k) - R^o(\Phi, y_n, k)] \right\|_2 + \frac{1}{N} \left\| \sum_n [y_n - P(\Phi_{I_n})y_n] \cdot \sigma_n(k) \cdot \chi(I_n, k) \right\|_2. \tag{40}
\]

By Lemma B.6 from [37] we have

\[
P\left( \left\| \frac{1}{N} \sum_n \chi(I_n, k) \sigma_n(k) \langle y_n, \phi_k \rangle \right\| \leq (1 - t_0) \frac{C_r \gamma_{1,S}}{K} \right) \leq \exp \left( \frac{-NC_r^2 \gamma_{1,S}^2 \cdot t_0^2}{2K(1 + \frac{SB}{K} + S\rho^2 + t_0C_r \gamma_{1,S} \sqrt{B+1}/3)} \right). \tag{41}
\]

By Lemma 6 in Appendix B (substituting Lemma B.3/4 of [37]) we have

\[
P\left( \mathbb{1}\{n : y_n \in \mathcal{F}\} \geq N \left( \frac{4}{K^3} + \frac{216}{SK^3} + \frac{C_r \gamma_{1,S}}{K \sqrt{B+1}} \cdot t_1 \varepsilon \right) \right) \leq \exp \left( \frac{-NC_r^2 \gamma_{1,S}^2 \cdot t_1^2 \varepsilon^2}{(B+1)(4 + 216/S) + t_1 \varepsilon C_r \gamma_{1,S} K \sqrt{B+1}} \right). \tag{42}
\]

35
By Lemma 5 in Appendix B (substituting Lemma B.8 of [37]) we have that for \(0 \leq t_2 \leq 1/8\)

\[
\mathbb{P} \left( \frac{1}{N} \left\| \sum_n \left[ R^\alpha(\Psi, y_n, k) - R^\alpha(\Phi, y_n, k) \right] \right\|_2 \geq \frac{C_r \gamma_1 S}{K} (0.339 \varepsilon + t_2 \varepsilon) \right) \\
\leq \exp \left( -\frac{NC_r^2 \gamma_1^2 S \cdot t_2^2 \varepsilon}{12K \max \{S, B\}^{3/2}} + \frac{1}{4} \right),
\]

and by Lemma B.7 from [37] we have

\[
\mathbb{P} \left( \frac{1}{N} \sum_n [y_n - P(\Phi t_n) y_n] \cdot \sigma_n(k) \cdot \chi(I_n, k) \right) \geq \frac{C_r \gamma_1 S}{K} t_3 \varepsilon \\
\leq \exp \left( -\frac{NC_r^2 \gamma_1^2 S \cdot t_3 \varepsilon}{8K \max \{S, B+1\}} \min \left\{ \frac{t_3 \varepsilon}{(1 - \gamma_{2, S} + d \rho^2)^\cdot 1} \right\} + \frac{1}{4} \right). \quad (43)
\]

Thus, with high probability we have \(s_k \geq (1 - t_0) \frac{C_r \gamma_1 S}{K}\) and

\[
\| \bar{\psi} - s_k \phi_k \|_2 \leq \frac{C_r \gamma_1 S}{K} \left( \frac{4 \sqrt{B + 1}}{K^2 C_r \gamma_1 S} \left( 1 + \frac{54}{S} \right) + t_1 + 0.339 + t_2 + t_3 \right) \varepsilon. \quad (44)
\]

Note that we only need to take into account distances \(\varepsilon > \frac{1}{32 \sqrt{S}}\), so we will use some crude bounds on \(C_r \gamma_1 S\) to show that the fraction with \(\varepsilon\) in the denominator above is small. The requirement that \(\|c(S')\|_2 / c(1) \leq \gamma_{\text{app}} \leq \log K\) ensures that \(\gamma_1 S \geq (1 + \log^2(K))^{-1/2}\) and we trivially have \(\gamma_1 S \geq Sc(S)\). Combining this with the bound on \(C_r\) in (10) we get

\[
\frac{1}{C_r \gamma_1 S} \leq \frac{\sqrt{1 + 5d \rho^2}}{(1 - e^{-d}) \gamma_1 S} \leq \frac{\sqrt{1 + \log^2(K)}}{(1 - e^{-d})} + \frac{\rho}{c(S) S (1 - e^{-d})}. \quad (45)
\]

The conditions in (36) imply that \(K \geq 87^2 \log^2(K)(S + 1)(B + 1)\), which in turn means that \(\log^2(K) > 16\), as well as \(\rho / c(S) \leq \gamma_{\rho} \leq 1/68\). Assuming additionally that \(K \geq \sqrt{d}\), meaning the dictionary is not too undercomplete, this leads to

\[
\frac{4 \sqrt{B + 1}}{K^2 C_r \gamma_1 S} \left( 1 + \frac{54}{S} \right) \leq \frac{4 \cdot 32 \cdot 55 \sqrt{(B + 1) S}}{K^2 C_r \gamma_1 S} \leq 0.001 \quad (46)
\]

Setting \(t_0 = t_1 = 1/20\) and \(t_2 = t_3 = 1/8\) we get

\[
\max_k \| \bar{\psi}_k - s_k \phi_k \|_2 \leq 0.64 \cdot \frac{C_r \gamma_1 S}{K} \varepsilon \quad \text{and} \quad \min_k s_k \geq 0.95 \cdot \frac{C_r \gamma_1 S}{K}, \quad (47)
\]

which by Lemma B.10 from [37] implies that

\[
d(\Psi, \Phi)^2 = \max_k \| \bar{\psi}_k - \phi_k \|_2 \leq 2 \left( 1 - \sqrt{1 - \frac{0.64^2 \varepsilon^2}{0.95^2}} \right) \leq 2 \cdot \frac{0.64^2 \varepsilon^2}{0.95^2} \leq 0.96^2 \varepsilon^2, \quad (48)
\]

36
except with probability

\[
K \exp \left( \frac{-NC_r^2 \gamma_{1,S}^2}{K(801 + 7C_r \gamma_{1,S} \sqrt{B+1})} \right) + \exp \left( \frac{-NC_r^2 \gamma_{1,S}^2 \cdot \varepsilon^2}{K(1 + 20rC_r \gamma_{1,S} \sqrt{B+1})} \right) + 2K \exp \left( \frac{-NC_r^2 \gamma_{1,S}^2 \cdot \varepsilon}{768K \max\{S,B\}^{3/2}} \right) + e^{-1/4}K \exp \left( \frac{-NC_r^2 \gamma_{1,S}^2 \cdot \varepsilon^2}{512K \max\{S,B+1\}(1 + d\rho^2)} \right).
\]

The final probability bound follows from the observations that \(C_r \gamma_{1,S} \leq \sqrt{S}, B+1 \geq 2\) and \(\varepsilon \leq \sqrt{2}\).

**Appendix B. Technical Lemmata**

To prove the two lemmata characterising the behaviour of thresholding and the difference between the oracle residuals based on the generating dictionary and a perturbation we need both the vector and the scalar version of Bernstein’s inequality.

**Theorem 3 (Vector Bernstein, \[23, 20, 24\])** Let \((v_n)_n \in \mathbb{R}^d\) be a finite sequence of independent random vectors. If \(\|v_n\|_2 \leq M\) almost surely, \(\|\mathbb{E}(v_n)\|_2 \leq m_1\) and \(\sum_n \mathbb{E}(\|v_n\|^2) \leq m_2\), then for all \(0 \leq t \leq m_2/(M+m_1)\), we have

\[
P\left( \left\| \sum_n v_n - \sum_n \mathbb{E}(v_n) \right\|_2 \geq t \right) \leq \exp \left( -\frac{t^2}{8m_2} + \frac{1}{4} \right),
\]

and, in general,

\[
P\left( \left\| \sum_n v_n - \sum_n \mathbb{E}(v_n) \right\|_2 \geq t \right) \leq \exp \left( -\frac{t}{8} \cdot \min \left\{ \frac{t}{m_2}, \frac{1}{M+m_1} \right\} + \frac{1}{4} \right).
\]

Note that the general statement is simply a consequence of the first part, since for \(t \geq m_2/(M+m_1)\) we can choose \(m_2 = t(M+m_1)\).

**Theorem 4 (Scalar Bernstein, \[7\])** Let \(v_n \in \mathbb{R}, n = 1\ldots N\), be a finite sequence of independent random variables with zero mean. If \(\mathbb{E}(v_n^2) \leq m\) and \(\mathbb{E}(|v_n|^k) \leq \frac{1}{2!}k!mM^{k-2}\) for all \(k > 2\), then for all \(t > 0\) we have

\[
P\left( \sum_n v_n \geq t \right) \leq \exp \left( -\frac{t^2}{2(Nm+Mt)} \right).
\]

We first prove that, assuming incoherence and good conditioning of the perturbed dictionary, the oracle residuals based on the perturbed dictionary \(\Psi\) and the generating dictionary \(\Phi\) are close to each other.
Lemma 5 Assume that the signals $y_n$ follow the random model in [24]. Further, assume that $S \leq \min \left\{ \frac{K}{98\|\Phi\|_{2,2}^2}, \frac{1}{98\rho^2} \right\}$ and that the current estimate of the dictionary $\Psi$ has distance $d(\Phi, \Psi) = \varepsilon \geq \frac{1}{32\sqrt{3}}$, but is incoherent and well-conditioned, meaning its coherence $\mu(\Psi)$ and its operator norm $\|\Psi\|_{2,2}$ satisfy
\[
\mu(\Psi) \leq \frac{1}{18\log K} \quad \text{and} \quad \|\Psi\|_{2,2}^2 \leq \frac{K}{108e^2S\log K} - 1.
\] (51)

Then for all $0 \leq t \leq 1/8$ we have
\[
\mathbb{P}\left(\frac{1}{N} \left\| \sum_n [R^o(\Psi, y_n, k) - R^o(\Phi, y_n, k)] \right\|_2 \geq \frac{C_{r\gamma_1}S K}{N} (0.339\varepsilon + t\varepsilon) \right) \leq \exp \left(-\frac{C_{r\gamma_1}^2S^2t^2\varepsilon}{12K \max\{S, \|\Phi\|_{2,2}^3\}} + \frac{1}{4} \right).
\]

Proof Throughout the proof we will use the abbreviations $B = \|\Phi\|_{2,2}$ and $\bar{B} = \|\Psi\|_{2,2}$. We apply Theorem 3 to $v_n = R^o(\Phi, y_n, k) - R^o(\Psi, y_n, k)$, dropping the index $n$ for conciseness. From Lemma B.8 in [37] we know that $v = T(I, k)y \cdot \sigma(k) \cdot \chi(I, k)$, where $T(I, k) := P(\Phi_I) - P(\Psi_I) - P(\phi_k) + P(\psi_k)$, and that
\[
\mathbb{E}(v) = \frac{C_{r\gamma_1}S}{K} \left(\frac{K-1}{S-1}\right) \sum_{|I|=S, k \in I} \left[ P(\psi_k) - P(\Psi_I) \right] \phi_k.
\] (52)

Using the orthogonal decomposition $\phi_k = [P(\psi_k) + Q(\psi_k)]\phi_k$, where $P(\psi_k)Q(\psi_k) = 0$, we get
\[
\mathbb{E}(v) = \frac{C_{r\gamma_1}S}{K} \left(\frac{K-1}{S-1}\right) \sum_{|I|=S, k \in I} -P(\Psi_I)Q(\psi_k)\phi_k.
\] (53)

Since the perturbed dictionary $\Psi$ is well-conditioned and incoherent, for most $I$ the sub-dictionary $\Psi_I$ will be a quasi isometry and $P(\Psi_I) \approx \Psi_I \Psi_I^*$. We therefore expand the expectation above, using the abbreviation $p_{K,S} = \left(\frac{S-1}{S-1}\right)^{-1}$, as
\[
\frac{K}{C_{r\gamma_1}S} \mathbb{E}(v) = p_{K,S} \left( \sum_{|I|=S, k \in I} \left[ \Psi_I \Psi_I^* - P(\Psi_I) \right] Q(\psi_k) \phi_k - \sum_{|I|=S, k \in I} \Psi_{I \setminus k} \Psi_{I \setminus k}^* Q(\psi_k) \phi_k \right)
= p_{K,S} \left( \sum_{|I|=S, k \in I} \left[ \Psi_I \Psi_I^* - P(\Psi_I) \right] Q(\psi_k) \phi_k - \frac{K-2}{S-2} \sum_{j \neq k} \psi_j \psi_j^* Q(\psi_k) \phi_k \right)
= p_{K,S} \left( \sum_{|I|=S, k \in I} \left[ \Psi_I \Psi_I^* - P(\Psi_I) \right] Q(\psi_k) \phi_k - \frac{S-1}{K-1} (\Psi \Psi^* - \psi_k \psi_k^*) Q(\psi_k) \phi_k \right)
= p_{K,S} \left( \sum_{|I|=S, k \in I} \left[ \Psi_I \Psi_I^* - P(\Psi_I) \right] Q(\psi_k) \phi_k \right)
+ p_{K,S} \sum_{|I|=S, k \in I, \delta(\Psi_I) \leq \delta_0} \left[ \Psi_I \Psi_I^* - P(\Psi_I) \right] Q(\psi_k) \phi_k
- \frac{S-1}{K-1} (\Psi \Psi^* - \psi_k \psi_k^*) Q(\psi_k) \phi_k.
\]
Combining the estimates for \(\|Q(\psi_k)\phi_k\|_2 = \omega_k \leq \varepsilon\), we can bound the norm of the expectation above as

\[
\|E(v)\|_2 \leq \frac{C_r \gamma_{1,S}}{K} \left[ \delta_0 + \mathbb{P}(\delta(\Psi_I) > \delta_0) |I| = S, k \in I) \cdot (B + 1) + \frac{(S-1)B}{K-1} \right] \varepsilon. \tag{54}
\]

To estimate the probability of a subdictionary being ill-conditioned we use Chretien and Darses’s results on the conditioning of random subdictionaries, which are slightly cleaner and thus easier to handle than the original results by Tropp, [43]. Theorem 3.1 of [10] reformulated for our purposes and applied to \(\Psi\) states that

\[
\mathbb{P}(\delta(\Psi_I) > \delta_0) |I| = S) \leq 216K \exp \left( - \min \left\{ \frac{\delta_0}{2\mu(\Psi)}, \frac{\delta_0^2 K}{4e^2SB} \right\} \right). \tag{55}
\]

Together with the union bound,

\[
\mathbb{P}(\delta(\Psi_I) > \delta_0) |I| = S, k \in I) \leq \frac{K}{S} \cdot \mathbb{P}(\delta(\Psi_I) > \delta_0) |I| = S), \tag{56}
\]

this leads to

\[
\|E(v)\|_2 \leq \frac{C_r \gamma_{1,S}}{K} \left[ \delta_0 + \frac{216K^2(B + 1)}{S} \exp \left( - \min \left\{ \frac{\delta_0}{2\mu(\Psi)}, \frac{\delta_0^2 K}{4e^2SB} \right\} \right) + \frac{SB}{K} \right] \varepsilon. \tag{57}
\]

Choosing \(\delta_0 = 1/3\), as long as \(B \leq \frac{K}{108e^2S \log K} - 1\) and \(\mu(\Psi) \leq \frac{1}{18 \log K}\) we have

\[
\|E(v)\|_2 \leq 0.339 \cdot \frac{C_r \gamma_{1,S}}{K} \cdot \varepsilon. \tag{58}
\]

The second quantity we need to bound is the expected energy of \(v = T(I, k)y \cdot \sigma(k) \cdot \chi(I, k)\). Combining Eqs. (115-118) from Lemma B.8 in [37] we get that

\[
\mathbb{E}(\|v\|_2^2) \leq \mathbb{E}_{p} \left( \chi(I, k) \left[ 4\gamma_{2,S}\varepsilon^2 + \left( \frac{B(1 - \gamma_{2,S})}{K-S} + \rho^2 \right) \|T(I, k)\|_{\mathcal{F}}^2 \right] \right). \tag{59}
\]

Since we are only interested in the regime \(\varepsilon > O(1/\sqrt{S})\) we will accept an additional factor \(S\) in the final sample complexity in return for a crude but painless estimate. Concretely, we use that \(T(I, k)\) is the difference of two orthogonal projections onto subspaces of dimension \(S-1\), namely \(P(\Psi_I) - P(\phi_k)\) and \(P(\Psi_I) - P(\psi_k)\). This leads to the bound \(\|T(I, k)\|_{\mathcal{F}}^2 \leq 2(S-1) \leq 2S\) and we get

\[
\mathbb{E}(\|v\|_2^2) \leq \frac{S}{K} \left( 4\gamma_{2,S}\varepsilon^2 + \frac{2BS}{K-S}(1 - \gamma_{2,S}) + 2S\rho^2 \right) \leq \frac{S}{K} (4\varepsilon^2 + 1/24), \tag{60}
\]

where for the second inequality we have used the assumption \(S \leq \min\{\frac{K}{9BS}, \frac{1}{9S\rho^2}\}\). Combining the estimates for \(\|E(v)\|_2\) and \(\mathbb{E}(\|v\|_2^2)\) with the norm bound \(\|v\|_2 \leq 2\sqrt{B+1},\)
we get that for $\varepsilon \geq \frac{1}{32\sqrt{S}}$ and $0 \leq t \leq 1/8$

$$\mathbb{P}\left(\frac{1}{N}\left\| \sum_n \left[ R^o(\Psi, y_n, k) - R^o(\Phi, y_n, k) \right] \right\|_2 \geq \frac{C_r\gamma_1 S}{K} \left( 0.339\varepsilon + t\varepsilon \right) \right) \leq \exp\left( -\frac{C_r\gamma_1 S t^2 \varepsilon}{8K} \min\left\{ \frac{1}{4e + (24\varepsilon)^{-1}}, \frac{1}{3t\sqrt{2}} \right\} + \frac{1}{4} \right).$$

Next we prove the lemma estimating how often thresholding will fail to recover the generating supports and signs.

**Lemma 6** Assume that the signals $y_n$ follow model (24) for coefficients with gap $c(S + 1)/c(S) \leq \gamma_{gap}$, dynamic sparse range $c(1)/c(S) \leq \gamma_{dyn}$, noise to coefficient ratio $\rho/c(S) \leq \gamma_{app}$ and relative approximation error $\|c(Sc)\|_2/c(1) \leq \gamma_{app} \leq \log K$. If the cross Gram matrix $\Phi^* \Psi$ is diagonally dominant in the sense that

$$\min_k \left| \langle \psi_k, \phi_k \rangle \right| \geq \max_k \left\{ 24 \gamma_{gap} \cdot \max_k \left| \langle \psi_k, \phi_k \rangle \right|, \right.$$

$$68 \gamma_{app} \cdot \sqrt{\log K},$$

$$78 \gamma_{dyn} \cdot (\log K)^{3/2} \max\{\mu(\Phi), \mu(\Phi, \Psi)\},$$

$$87 \gamma_{dyn} \cdot (S + 1)(\|\Phi\|_{2,2}^2 + 1)/K \right\}.$$ (61)

then

$$\mathbb{P}\left( \sharp\{n : y_n \in \mathcal{F}\} \geq N \left( \frac{4}{K^3} + \frac{216}{SK^3} + \frac{t}{K} \right) \right) \leq \exp\left( -\frac{t^2 KN}{4 + 216/S + tK^2} \right).$$ (62)

**Proof** We apply Theorem 4 to the sum of recentered indicator functions $v_n = 1_\mathcal{F} - \mathbb{P}(\mathcal{F})$ to get

$$\mathbb{P}\left( \sharp\{n : y_n \in \mathcal{F}\} \geq N\mathbb{P}(\mathcal{F}) + tN/K \right) \leq \exp\left( -\frac{t^2 N}{2K^2\mathbb{P}(\mathcal{F}) + tK} \right).$$ (63)

Next we need to estimate the probability $\mathbb{P}(\mathcal{F})$ of the oracle residual not being equal to the thresholding residual. This event is contained in the event of thresholding failing for a signal $y$ or of the generating sign being changed. Contrary to [37], we do not look at the
inner products of the signals with the generating atoms and the perturbations, which make up $\Psi$, separately but directly bound the inner products with $\psi_k$. In order for thresholding to succeed we need to have for $I = p^{-1}(S)$
\[
\min_{i \in I} |\langle \psi_i, \Phi x_{c,p,\sigma} + r \rangle| > \max_{i \notin I} |\langle \psi_i, \Phi x_{c,p,\sigma} + r \rangle|.
\]
(64)

Expanding the inner product and bounding it from below resp. above for an atom inside resp. outside the support,
\[
i \in I : \quad |\langle \psi_i, \Phi x_{c,p,\sigma} + r \rangle| \geq c(S)\alpha_{\min} - |\sum_{j \neq i} \sigma(j)c(p(j))\langle \psi_i, \phi_j \rangle| - |\langle \psi_i, r \rangle|,
\]
(65)
\[
i \notin I : \quad |\langle \psi_i, \Phi x_{c,p,\sigma} + r \rangle| \leq c(S+1)\alpha_{\max} + \left|\sum_{j \neq i} \sigma(j)c(p(j))\langle \psi_i, \phi_j \rangle\right| + |\langle \psi_i, r \rangle|,
\]
(66)

we see that the following conditions are sufficient for thresholding to succeed and for preservation of the generating signs. For all atoms $\psi_i$ we have
\[
\left|\sum_{j \neq i} \sigma(j)c(p(j))\langle \psi_i, \phi_j \rangle\right| < \theta_1 \cdot c(S)\alpha_{\min} \quad \text{and} \quad |\langle \psi_i, r \rangle| < \theta_2 \cdot c(S)\alpha_{\min},
\]
(67)

where $\theta_1, \theta_2$ satisfy
\[
2\theta_1 + 2\theta_2 + \frac{c(S+1)\alpha_{\max}}{c(S)} \frac{\alpha_{\min}}{\alpha_{\min}} \leq 1.
\]
(68)

Note that for perfectly $S$-sparse signals the constraint on the parameters $\theta_i$ reduces to $2\theta_1 + 2\theta_2 \leq 1$. In the case of not perfectly $S$-sparse signals, we see that in order to allow for relatively large $\theta_i$, we need the recovery ratio between best and worst approximated atom to be balanced. Via these sufficient conditions we can now upperbound the failure probability of thresholding by
\[
P(\exists i : \left|\sum_{j \neq i} \sigma(j)c(p(j))\langle \psi_i, \phi_j \rangle\right| \geq \theta_1 \cdot c(S)\alpha_{\min}) + P(\exists i : |\langle \psi_i, r \rangle| \geq \theta_2 \cdot c(S)\alpha_{\min}).
\]
(69)

Using a union bound and the sub-Gaussian property of the noise vector $r$, the second term in the sum above can be straightforwardly bounded as
\[
P(\exists i : |\langle \psi_i, r \rangle| \geq \theta_2 \cdot c(S)\alpha_{\min}) \leq 2K \exp\left(-\frac{\left(\theta_2 \cdot c(S)\alpha_{\min}\right)^2}{2\rho^2}\right).
\]
(70)

The first term is a bit more tricky and interesting to control. To see what has to be done, we first have a look at the probability that the condition is violated for one index $i$. Using Hoeffding’s inequality we get,
\[
P\left(\sum_{j \neq i} \sigma(j)c(p(j))\langle \psi_i, \phi_j \rangle \geq \theta_1 \cdot c(S)\alpha_{\min}\right) \leq 2 \exp\left(-\frac{\left(\theta_1 \cdot c(S)\alpha_{\min}\right)^2}{2\sum_{j \neq i} c(p(j))^2 |\langle \psi_i, \phi_j \rangle|^2}\right).
\]
(71)
The sum in the denominator of the right hand side is split into two parts and bounded as,
\[
\sum_{j \neq i} c(p(j))^2 | \langle \psi_i, \phi_j \rangle |^2 \leq c(1)^2 \sum_{j \in I \setminus \{i\}} | \langle \psi_i, \phi_j \rangle |^2 + \max_{j \neq i} | \langle \psi_i, \phi_j \rangle |^2 \sum_{j > S} c(j)^2 \quad (72)
\]
\[
\leq c(1)^2 \| \Phi^*_I \psi_i \|_2^2 + \mu(\Phi, \Psi)^2 \cdot \| c(S') \|_2^2. \quad (73)
\]
The residual energy \( \| c(S') \|_2^2 \) is again zero for perfectly \( S \)-sparse signals and can be assumed to be small otherwise, so the interesting question is how big \( \| \Phi^*_I \psi_i \|_2 \) is for most supports \( I \). Using the fact that for \( J \subset I \) we have \( \| \phi^*_J \Phi_{I \setminus J} \|_{2,2} \leq \delta(\Phi_I), \quad [18], \) we can turn this into a question about the conditioning of subdictionaries. Denote by \( \Phi^{(i)} \) the dictionary \( \Phi \) with the \( i \)-th atom \( \phi_i \) replaced by \( \psi_i \). We have
\[
P \left( \| \Phi^*_I \psi_i \|_2 \geq \delta_0 \left| I \right| = S \right) \leq P \left( \delta(\Phi^{(i)}_I) \geq \delta_0 \left| I \right| = S \right) = \frac{\# \{ I : \delta(\Phi^{(i)}_I) \geq \delta_0, \left| I \right| = S \}}{\# \{ I : \left| I \right| = S \}}
\]
\[
= \frac{\# \{ I : \delta(\Phi^{(i)}_I) \geq \delta_0, \left| I \right| = S, i \in I \}}{\# \{ I : \left| I \right| = S \}} + \frac{\# \{ I : \delta(\Phi^{(i)}_I) \geq \delta_0, \left| I \right| = S, i \notin I \}}{\# \{ I : \left| I \right| = S \}}
\]
\[
\leq \frac{\# \{ I : \delta(\Phi^{(i)}_I) \geq \delta_0, \left| I \right| = S \}}{\# \{ I : \left| I \right| = S \}} + \frac{\# \{ I : \delta(\Phi^{(i)}_I) \geq \delta_0, \left| I \right| = S + 1 \}}{\# \{ I : \left| I \right| = S \}}
\]
\[
\leq P \left( \delta(\Phi^{(i)}_I) \geq \delta_0 \left| I \right| = S \right) + \frac{K - S}{S + 1} \cdot P \left( \delta(\Phi^{(i)}_I) \geq \delta_0 \left| I \right| = S + 1 \right)
\]
\[
\leq \frac{K + 1}{S + 1} \cdot P \left( \delta(\Phi^{(i)}_I) \geq \delta_0 \left| I \right| = S + 1 \right). \quad (74)
\]
Note that we can bound the coherence of \( \Phi^{(i)} \) by the coherence of \( \Phi \) and the cross-coherence of \( \Psi \) and \( \Phi \), meaning \( \mu(\Phi^{(i)}) \leq \max \{ \mu(\Phi), \mu(\Phi, \Psi) \} =: \hat{\mu} \). Similarly, since we replace only one vector, we have \( \| \Phi^{(i)} \|_{2,2} \leq \| \Phi \|_{2,2} + 1 \). Using again Theorem 3.1 of [10] we get that
\[
P \left( \| \Phi^*_I \psi_i \|_2 \geq \delta_0 \left| I \right| = S \right) \leq \frac{216K^2}{S} \exp \left( - \min \left\{ \frac{\delta_0}{2\hat{\mu}}, \frac{\delta_0^2K}{4c^2(S + 1)(\| \Phi^* \|_{2,2} + 1)} \right\} \right). \quad (75)
\]
Next we use this estimate to bound the first term in (69) as
\[
P \left( \exists i : \left| \sum_{j \neq i} \sigma(j) c(p(j)) \langle \psi_i, \phi_j \rangle \right| \geq \theta_1 \cdot c(S) \alpha_{\min} \right)
\]
\[
\leq P \left( \exists i : \left| \sum_{j \neq i} \sigma(j) c(p(j)) \langle \psi_i, \phi_j \rangle \right| \geq \theta_1 \cdot c(S) \alpha_{\min} \left| \| \Phi^*_I \psi_i \|_2 \leq \delta_0, \forall i \right) \right)
\]
\[
\leq 2K \exp \left( - \frac{\left( \theta_1 \cdot c(S) \alpha_{\min} \right)^2}{2c(1)^2 \delta_0^2 + 2\hat{\mu}^2 \cdot \| c(S') \|_2^2} \right)
\]
\[
+ \frac{216K^3}{S} \exp \left( - \min \left\{ \frac{\delta_0}{2\hat{\mu}}, \frac{\delta_0^2K}{4c^2(S + 1)(\| \Phi^* \|_{2,2} + 1)} \right\} \right). \quad (76)
\]
In order to have the first term reasonably small, we need to choose \( \delta_0 = c(S)\alpha_{\min}\theta_1 \frac{n_1}{\sqrt{\log K}} \), for some \( n_1 > 1 \). To have also the second term small, we need to have \( \bar{\mu} = \frac{\delta_0}{n_2 \log K} \) for \( n_2 > 1 \). Concretely, we have

\[
\mathbb{P}\left( \exists i : \left| \sum_{j \neq i} \sigma(j)c(p(j))\langle \psi_i, \phi_j \rangle \right| \geq \theta_1 \cdot c(S)\alpha_{\min} \right) \leq 2K \cdot K^{2n^2/n_2} + \frac{216K^3}{S} \cdot K^{-n_2}, \tag{77}
\]

whenever \( \frac{\|c(S^c)\|_2}{c(1)} \leq \log K \), \( \alpha_{\min} \geq n_1 n_2 \frac{c(1)}{c(S)} (\log K)^{3/2} \cdot \bar{\mu} \) and

\[
\alpha_{\min} \geq \sqrt{2e^2n_1^2n_2^2\theta_1^2c(1)} \frac{c(1)}{c(S)} \log K \cdot \left( \frac{(S+1)(\|\Phi\|_2^2+1)}{K} \right)^{1/2}. \tag{78}
\]

To choose the constants \( n_i, \theta_i \), we recall that for the second term in (69) we have by (70)

\[
\mathbb{P}\left( \exists i : |\langle \psi_i, r \rangle| \geq \theta_2 \cdot c(S)\alpha_{\min} \right) \leq 2K \cdot K^{-n_3^2}, \tag{79}
\]

whenever

\[
\alpha_{\min} \geq \frac{\sqrt{2n_3}}{\theta_2} \frac{1}{c(S)} \sqrt{\log K \cdot \rho} \quad \text{and} \quad 2\theta_1 + 2\theta_2 + \frac{c(S+1)}{c(S)} \frac{\alpha_{\max}}{\alpha_{\min}} \leq 1. \tag{80}
\]

Setting \( \theta_1 = 7/16, \theta_2 = 1/24, n_1^2 = 145/18, n_2 = 12 \) and \( n_3 = 2 \), we get that \( \mathbb{P}(\mathcal{F}) < (4 + 216/S)K^{-3} \) as long as \( \frac{\|c(S^c)\|_2}{c(1)} \leq \log K \) and

\[
\alpha_{\min} \geq \max \left\{ 24\alpha_{\max} \frac{c(S+1)}{c(S)}, 68\sqrt{\log K \cdot \rho} \frac{1}{c(S)}, 78 \frac{c(1)}{c(S)} (\log K)^{3/2} \cdot \bar{\mu}, 87 \frac{c(1)}{c(S)} \log K \cdot \sqrt{(S+1)(\|\Phi\|_2^2+1) K} \right\} \tag{81}
\]

The statement follows substituting the bounds on the gap \( \frac{c(S+1)}{c(S)} \), the noise to coefficient ratio \( \frac{\rho}{c(S)} \), the dynamic sparse range \( \frac{c(1)}{c(S)} \), and the relative approximation error \( \frac{\|c(S^c)\|_2}{c(1)} \). \( \blacksquare \)
Appendix C. Pseudocode

Algorithm C.1: ITKrM augmented for replacement/adaptivity - one iteration

**Input:** $\Psi, Y, S, \Gamma, M$;  
  // dictionary, signals, sparsity, candidates,  
  // minimal observations (only for adaptive)

Set: $m = \lfloor \log d \rfloor$, $N_{\Gamma} = \lfloor N/m \rfloor$;  
Initialise: $\Psi = 0, \Gamma = 0, S = 0$;

foreach $n$ do
  // basic ITKrM steps
  $I_n = \arg \max_{I \subseteq S} ||\Psi_I y_n||_1$;  
  // thresholding
  $x_n = \Psi_{I_n}^\dagger y_n$;  
  // sparse coefficients
  $a_n = y_n - \Psi_{I_n} x_n$;  
  // residual
  $\tau = 0$;  
  // simple counter for replacement
  $\tau = (2\log\left(\frac{2N_{\Gamma}}{d}\right)||a_n||_2^2 + ||\Psi_{I_n} x_n||_2^2) / d$;  
  // advanced counter for adaptivity
  foreach $k \in I_n$ do
    $\tilde{\psi}_k \leftarrow \psi_k + [a_n + P(\psi_k)y_n] \cdot \text{sign}(\langle \psi_k, y_n \rangle)$;  
    // atom update
    if $|x_n(k)|^2 \geq \tau$ then
      $v(k) \leftarrow v(k) + 1$;  
      // atom value update
    end
  end
  // steps for replacement candidates
  $i_n = \arg \max_I |\langle \gamma_I, a_n \rangle|$;  
  // residual thresholding
  $\tilde{\gamma}_n \leftarrow \gamma_n + a_n \cdot \text{sign}(\langle \gamma_n, a_n \rangle)$;  
  // candidate update
  $\tau = 2\log(2K)/d$;  
  // simple counter for replacement
  $\tau = 2\log\left(\frac{2N_{\Gamma}}{d}\right)/d$;  
  // advanced counter for adaptivity
  if $|\langle \gamma_n, a_n \rangle|^2 \geq \tau \cdot ||a_n||_2^2$ then
    $v_I(i_n) \leftarrow v_I(i_n) + 1$;  
    // candidate value update
  end
  if $n \mod N_{\Gamma} == 0 \land n < mN_{\Gamma}$ then
    $\Gamma \leftarrow (\gamma_1/||\gamma_1||_2, \ldots, \gamma_L/||\gamma_L||_2)$;  
    // candidate normalisation
    $\Gamma = 0$;  
    // cand. iteration restart
    $v_I = 0$;  
    // skip for replacement
  end
  // steps for estimating sparsity level, skip for replacement
  $\theta = (2\log(4K)||a_n||_2^2 + ||\Psi_{I_n} x_n||_2^2) / d$;
  $\tilde{S} \leftarrow \tilde{S} + \{k : |x_n(k)|^2 \geq \theta\}$;  
  // correct atoms
  $\tilde{S} \leftarrow \tilde{S} + \{k : |\langle \psi_k, a_n \rangle|^2 \geq \theta\}$;  
  // missed atoms
end

$\Psi \leftarrow (\tilde{\psi}_1/||\tilde{\psi}_1||_2, \ldots, \tilde{\psi}_K/||\tilde{\psi}_K||_2)$;  
// atom normalisation
$S \leftarrow \lfloor S/N \rfloor$;  
// average sparsity level, skip for replacement

**Output:** $\Psi, v, \Gamma, v_I, \tilde{S}$;  
// estimated sparsity only for adaptivity
Algorithm C.2: Replacing coherent atoms (delete, merge, add)

Input: $\Psi, v, \Gamma, v_T, \mu_{\text{max}}$; // dict., score, cand., cand.score, threshold
Find: $(k, k') = \text{arg max}_{i < j} |\langle \psi_i, \psi_j \rangle|; \quad \text{// most coherent atom pair}
while $|\langle \psi_k, \psi_{k'} \rangle| > \mu_{\text{max}} \land \Gamma \neq []$ do
  $h = \text{sign}(\langle \psi_k, \psi_{k'} \rangle)$;
  $\psi_k = |\psi_k| \frac{v(k')}{v(k) + v(k')} \psi_{k'} + h |\psi_k| \frac{v(k)}{v(k) + v(k')} \psi_k$; // delete
  $\psi_k = v(k') \psi_{k'} + h v(k) \psi_k$; // or merge
  $\psi_k = \psi_{k'} + h \psi_k$; // or add
  $\Gamma = \{ \mu \in \Gamma : \mu > \mu_{\text{max}} \}$;
  $\Gamma_{\Lambda} = [\Lambda, \Lambda] \leftarrow []$; // discard coherent candidates
  if $\Gamma = []$ then
    $\psi_k \leftarrow \psi_k$; // replace with merged atom
    $v(k) \leftarrow v(k) + v(k')$; // update score of merged atom
    $\gamma_1 \leftarrow [\Lambda, \Lambda] \leftarrow []$; // replace with most useful candidate
    if $\mu_1 < \mu_{\text{max}}$ then
      $v(k') \leftarrow v_T(1)$; // update with candidate score
    else
      $v(k') \leftarrow 0$; // preferably replaced again
    end
    $\gamma_1 \leftarrow [], v_T(1) \leftarrow []$; // discard used candidate
  end
  Find: $(k, k') = \text{arg max}_{i < j} |\langle \psi_i, \psi_j \rangle|; \quad \text{// update most coherent atom pair}
end
Output: $\Psi, v, \Gamma, v_T$

Algorithm C.3: Pruning coherent atoms (merge)

Input: $\Psi, V = (v_1, \ldots, v_K), \mu_{\text{max}}$; // dictionary, last $m$ scores, threshold
$\Delta = []$; // initialise set of atoms to delete
$H = \Psi^* \Psi - I_K$; // hollow Gram matrix in absolute
Find: $(k, k') = \text{arg max}_{i < j} |H(i, j)|; \quad \text{// most coherent atom pair}
while $|H(k, k')| > \mu_{\text{max}}$ do
  $\psi_k \leftarrow v(k) \psi_k + \text{sign}(\langle \psi_k, \psi_{k'} \rangle) v_{k'} \psi_k$; // merge according to most recent score
  $v_k \leftarrow |v_k| |\psi_k|_2$;
  $v_{k'}(1) \leftarrow v_{k'}(1) + v_k(1)$; // update most recent score
  $\Delta \leftarrow \Delta \cup \{k'\}$;
  $H(k, \cdot) \leftarrow 0, H(k', \cdot) \leftarrow 0, H(\cdot, k) \leftarrow 0, H(\cdot, k') \leftarrow 0$; // update hollow Gram matrix
  Find: $(k, k') = \text{arg max}_{j > i} |H(i, j)|; \quad \text{// update most coherent atom pair}
end
$\Psi_{\Delta} \leftarrow [], V_{\Delta} \leftarrow []$; // delete atoms
Output: $\Psi, V$
**Algorithm C.4:** Pruning unused atoms

**Input:** $\Psi, V = (v_1, \ldots, v_K), M, \delta$;  
// dictionary, last $m$ scores, threshold,  
// maximally pruned atoms

**foreach** $k$ do

$\hat{v}(k) = \max_i v_k(i)$;  
// maximum of last $m$ scores

end

$\Delta = \{ k : \hat{v}(k) < M \}$;  
// atoms with max.scores below threshold

**if** $|\Delta| > \delta$ then

Sort: $\hat{v}(i_1) \leq \hat{v}(i_2) \leq \cdots \leq \hat{v}(i_K)$;  
// sort max.score

$\Delta = \{ i_1 \ldots i_\delta \}$;  
// $\delta$ atoms with smallest max.scores

end

$\Psi_\Delta \leftarrow \emptyset, V_\Delta \leftarrow \emptyset$;  
// delete atoms

**Output:** $\Psi, V$

**Algorithm C.5:** Adding atoms

**Input:** $\Psi, V, \Gamma, v_\Gamma, \mu_{\max}, M$

Sort: $v_\Gamma(i_1) \geq v_\Gamma(i_2) \geq \cdots \geq v_\Gamma(i_L)$;  
// sort according to score

$L = |\{ \ell : v_\Gamma(\ell) \geq d \}|$;  
// score above $d$

**for** $\ell = 1 \ldots L$ do

**if** $\max_k |\langle \psi_k, \gamma_{i_\ell} \rangle| \leq \mu_{\max}$ then

$\Psi \leftarrow (\Psi, \gamma_{i_\ell})$;  
// in order of score add if incoherent

$V \leftarrow (V, M \cdot 1)$;  
// set last $m$ scores of added atoms to $M$

end

end

**Output:** $\Psi, V$

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