
DUAL-LOCO: Distributing Statistical Estimation Using Random Projections

Christina Heinze
Seminar for Statistics, ETH Zürich

Brian McWilliams
Disney Research

Nicolai Meinshausen
Seminar for Statistics, ETH Zürich

Abstract

We present DUAL-LOCO, a communication-efficient algorithm for distributed statistical estimation. DUAL-LOCO assumes that the data is distributed across workers according to the features rather than the samples. It requires only a single round of communication where low-dimensional random projections are used to approximate the dependencies between features available to different workers. We show that DUAL-LOCO has bounded approximation error which only depends weakly on the number of workers. We compare DUAL-LOCO against a state-of-the-art distributed optimization method on a variety of real world datasets and show that it obtains better speedups while retaining good accuracy. In particular, DUAL-LOCO allows for fast cross validation as only part of the algorithm depends on the regularization parameter.

1 Introduction

Many statistical estimation tasks amount to solving an optimization problem of the form

$$\min_{\beta \in \mathbb{R}^p} J(\beta) := \sum_{i=1}^n f_i(\beta^\top \mathbf{x}_i) + \frac{\lambda}{2} \|\beta\|_2^2 \quad (1)$$

where $\lambda > 0$ is the regularization parameter. The loss functions $f_i(\beta^\top \mathbf{x}_i)$ depend on labels $y_i \in \mathbb{R}$ and linearly on the coefficients, β through a vector of covariates, $\mathbf{x}_i \in \mathbb{R}^p$. Furthermore, we assume all f_i to be convex and smooth with Lipschitz continuous gradients. Concretely, when $f_i(\beta^\top \mathbf{x}_i) = (y_i - \beta^\top \mathbf{x}_i)^2$,

Eq. (1) corresponds to ridge regression; for logistic regression $f_i(\beta^\top \mathbf{x}_i) = \log(1 + \exp(-y_i \beta^\top \mathbf{x}_i))$.

For large-scale problems, it is no longer practical to solve even relatively simple estimation tasks such as (1) on a single machine. To deal with this, approaches to distributed data analysis have been proposed that take advantage of many cores or computing nodes on a cluster. A common idea which links many of these methods is stochastic optimization. Typically, each of the workers only sees a small portion of the data points and performs incremental updates to a global parameter vector. It is typically assumed that the number of data points, n , is very large compared with the number of features, p , or that the data is extremely sparse. In such settings – which are common, but not ubiquitous in large datasets – distributed stochastic optimization algorithms perform well but may converge slowly otherwise.

A fundamentally different approach to distributing learning is for each worker to only have access to a portion of the available features. Distributing according to the features could be a preferable alternative for several reasons. Firstly, for **high-dimensional data**, where p is large relative to n , better scaling can be achieved. This setting is challenging, however, since most loss functions are not separable across coordinates. High-dimensional data is commonly encountered in the fields of bioinformatics, climate science and computer vision. Furthermore, for a variety of prediction tasks it is often beneficial to map input vectors into a higher dimensional feature space, e.g. using deep representation learning or considering higher-order interactions. Secondly, **privacy**. Individual blocks of features could correspond to sensitive information (such as medical records) which should be included in the predictive model but is not allowed to be communicated in an un-disguised form.

Our contribution. In this work we introduce DUAL-LOCO to solve problems of the form (1) in the distributed setting when each worker only has access to a subset of the *features*. DUAL-LOCO is an exten-

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sion of the LOCO algorithm [1] which was recently proposed for solving distributed ridge regression in this setting. We propose an alternative formulation where each worker instead locally solves a *dual* optimization problem. DUAL-LOCO has a number of practical and theoretical improvements over the original algorithm:

- DUAL-LOCO is applicable to a wider variety of smooth, convex ℓ_2 penalized loss minimization problems encompassing many widely used regression and classification loss functions, including ridge regression, logistic regression and others.
- In §4 we provide a more intuitive and tighter theoretical result which crucially does not depend on specific details of the ridge regression model and has weaker dependence on the number of workers, K .
- We also show that *adding* (rather than concatenating) random features allows for an efficient implementation yet retains good approximation guarantees.

In §5 we report experimental results with high-dimensional real world datasets corresponding to two different problem domains: climate science and computer vision. We compare DUAL-LOCO with CoCoA⁺, a recently proposed state-of-the-art algorithm for distributed dual coordinate ascent [2]. Our experiments show that DUAL-LOCO demonstrates better scaling with K than CoCoA⁺ while retaining a good approximation of the optimal solution. We provide an implementation of DUAL-LOCO in Apache Spark¹. The portability of this framework ensures that DUAL-LOCO is able to be run in a variety of distributed computing environments.

2 Related work

2.1 Distributed Estimation

Recently, several asynchronous stochastic gradient descent (SGD) methods [3, 4] have been proposed for solving problems of the form (1) in a parallel fashion in a multi-core, shared-memory environment and have been extended to the distributed setting. For such methods, large speedups are possible with asynchronous updates when the data is sparse. However, in some problem domains the data collected is dense with many correlated features. Furthermore, the $p \gg n$ setting can result in slow convergence. In the distributed setting, such methods can be impractical since the cost of communicating updates can dominate other computational considerations.

¹<http://spark.apache.org/>

Jaggi et al. proposed a communication-efficient distributed dual coordinate ascent algorithm (CoCoA resp. CoCoA⁺) [5, 2]. Each worker makes multiple updates to its local dual variables before communicating the corresponding primal update. This allows for trading off communication and convergence speed. Notably they show that convergence is actually independent of the number of workers, thus CoCoA⁺ exhibits *strong scaling* with K .

Other recent work considers solving statistical estimation tasks using a single round of communication [6, 7]. However, all of these methods consider only distributing over the rows of the data where an i.i.d. assumption on the observations holds.

On the other hand, few approaches have considered distributing across the columns (features) of the data. This is a more challenging task for both estimation and optimization since the columns are typically assumed to have arbitrary dependencies and most commonly used loss functions are not separable over the features. Recently, LOCO was proposed to solve ridge regression when the data is distributed across the features [1]. LOCO requires a single round to communicate small matrices of randomly projected features which approximate the dependencies in the rest of the dataset (cf. Figure 1). Each worker then optimizes its own subproblem independently and finally sends its portion of the solution vector back to the master where they are combined. LOCO makes no assumptions about the correlation structure between features. It is therefore able to perform well in challenging settings where the features are correlated between blocks and is particularly suited when $p \gg n$. Indeed, since the relative dimensionality of local problems decreases when splitting by columns, they are easier in a statistical sense. LOCO makes no assumptions about data sparsity so it is also able to obtain speedups when the data is dense.

One-shot communication schemes are beneficial as the cost of communication consists of a fixed cost and a cost that is proportional to the size of the message. Therefore, it is generally cheaper to communicate a few large objects than many small objects.

2.2 Random projections for estimation and optimization

Random projections are low-dimensional embeddings $\Pi : \mathbb{R}^T \rightarrow \mathbb{R}^{T_{subs}}$ which approximately preserve an entire subspace of vectors. They have been extensively used to construct efficient algorithms when the sample size is large in a variety of domains such as: nearest neighbours [8], matrix factorization [9], least squares [10, 11] and recently in the context of optimization [12].

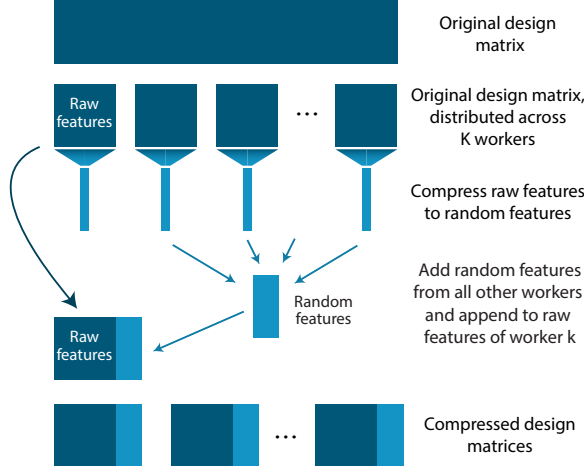


Figure 1: Schematic for the distributed approximation of a large data set with random projections, used by DUAL-LOCO.

We concentrate on the Subsampled Randomized Hadamard Transform (SRHT), a structured random projection [13]. The SRHT consists of a projection matrix, $\mathbf{\Pi} = \sqrt{\tau/\tau_{subs}} \mathbf{DHS}$ [9] with the definitions: (i) $\mathbf{S} \in \mathbb{R}^{\tau \times \tau_{subs}}$ is a subsampling matrix. (ii) $\mathbf{D} \in \mathbb{R}^{\tau \times \tau}$ is a diagonal matrix whose entries are drawn independently from $\{-1, 1\}$. (iii) $\mathbf{H} \in \mathbb{R}^{\tau \times \tau}$ is a normalized Walsh-Hadamard matrix. The key benefit of the SRHT is that due to its recursive definition the product between $\mathbf{\Pi}^\top$ and $\mathbf{u} \in \mathbb{R}^\tau$ can be computed in $O(\tau \log \tau)$ time while never constructing $\mathbf{\Pi}$ explicitly.

For moderately sized problems, random projections have been used to reduce the dimensionality of the data prior to performing regression [14, 15]. However after projection, the solution vector is in the compressed space and so interpretability of coefficients is lost. Furthermore, the projection of the low-dimensional solution back to the original high-dimensional space is in fact guaranteed to be a *bad* approximation of the optimum [16].

Dual Random Projections. Recently, [16, 17] studied the effect of random projections on the *dual* optimization problem. For the primal problem in Eq. (1), defining $\mathbf{K} = \mathbf{X}\mathbf{X}^\top$, we have the corresponding dual

$$\max_{\alpha \in \mathbb{R}^n} - \sum_{i=1}^n f_i^*(\alpha_i) - \frac{1}{2n\lambda} \alpha^\top \mathbf{K} \alpha \quad (2)$$

where f^* is the conjugate Fenchel dual of f and $\lambda > 0$. For example, for squared loss functions $f_i(u) = \frac{1}{2}(y_i - u)^2$, we have $f_i^*(\alpha) = \frac{1}{2}\alpha^2 + \alpha y_i$. For problems of this form, the dual variables can be directly mapped to the primal variables, such that for a vector α^* which attains the maximum of (2), the optimal

primal solution has the form $\beta^*(\alpha^*) = -\frac{1}{n\lambda} \mathbf{X}^\top \alpha^*$.

Clearly, a similar dual problem to (2) can be defined in the projected space. Defining $\tilde{\mathbf{K}} = (\mathbf{X}\mathbf{\Pi})(\mathbf{X}\mathbf{\Pi})^\top$ we have

$$\max_{\alpha \in \mathbb{R}^n} - \sum_{i=1}^n f_i^*(\alpha_i) - \frac{1}{2n\lambda} \alpha^\top \tilde{\mathbf{K}} \alpha. \quad (3)$$

Importantly, the vector of dual variables does not change dimension depending on whether the original problem (2) or the projected problem (3) is being solved. Under mild assumptions on the loss function, by mapping the solution to this new problem, $\tilde{\alpha}$, back to the original space one obtains a vector $\tilde{\beta}(\tilde{\alpha}) = -\frac{1}{n\lambda} \mathbf{X}^\top \tilde{\alpha}$, which is a *good* approximation to β^* , the solution to the original problem (1) [16, 17].

3 The DUAL-LOCO algorithm

Algorithm 1 DUAL-LOCO

Input: Data: \mathbf{X}, Y , no. workers: K

Parameters: τ_{subs}, λ

- 1: Partition $\{p\}$ into K subsets of equal size τ and distribute feature vectors in \mathbf{X} accordingly over K workers.
- 2: **for each** worker $k \in \{1, \dots, K\}$ **in parallel do**
- 3: Compute and send random features $\mathbf{X}_k \mathbf{\Pi}_k$.
- 4: Receive random features and construct $\tilde{\mathbf{X}}_k$.
- 5: $\tilde{\alpha}_k \leftarrow \text{LocalDualSolver}(\tilde{\mathbf{X}}_k, Y, \lambda)$
- 6: $\hat{\beta}_k = -\frac{1}{n\lambda} \mathbf{X}_k^\top \tilde{\alpha}_k$
- 7: Send $\hat{\beta}_k$ to driver.
- 8: **end for**

Output: Solution vector: $\hat{\beta} = [\hat{\beta}_1, \dots, \hat{\beta}_K]$

In this section we detail the DUAL-LOCO algorithm. DUAL-LOCO differs from the original LOCO algorithm in two important ways. (i) The random features from each worker are summed, rather than concatenated, to obtain a τ_{subs} dimensional approximation allowing for an efficient implementation in a large-scale distributed environment. (ii) Each worker solves a local *dual* problem similar to (3). This allows us to extend the theoretical guarantees to a larger class of estimation problems beyond ridge regression (§4).

We consider the case where p features are distributed across K different workers in non-overlapping subsets $\mathcal{P}_1, \dots, \mathcal{P}_K$ of equal size², $\tau = p/K$.

Since most loss functions of interest are not separable across coordinates, a key challenge addressed by DUAL-LOCO is to define a local minimization problem for each worker to solve *independently* and *asynchronously* while still maintaining important depen-

²This is for simplicity of notation only, in general the partitions can be of different sizes.

dencies between features in different blocks and keeping communication overhead low. Algorithm 1 details DUAL-LOCO in full.

We can rewrite (1) making explicit the contribution from block k . Letting $\mathbf{X}_k \in \mathbb{R}^{n \times \tau}$ be the sub-matrix whose columns correspond to the coordinates in \mathcal{P}_k (the “raw” features of block k) and $\mathbf{X}_{(-k)} \in \mathbb{R}^{n \times (p-\tau)}$ be the remaining columns of \mathbf{X} , we have

$$J(\beta) = \sum_{i=1}^n f_i \left(\mathbf{x}_{i,k}^\top \beta_{\text{raw}} + \mathbf{x}_{i,(-k)}^\top \beta_{(-k)} \right) + \lambda (\|\beta_{\text{raw}}\|_2^2 + \|\beta_{(-k)}\|_2^2). \quad (4)$$

Where $\mathbf{x}_{i,k}$ and $\mathbf{x}_{i,(-k)}$ are the rows of \mathbf{X}_k and $\mathbf{X}_{(-k)}$ respectively. We replace $\mathbf{X}_{(-k)}$ in each block with a low-dimensional randomized approximation which preserves its contribution to the loss function. This procedure is described in Figure 1.

In **Step 4**, these matrices of random features are communicated and worker k constructs the matrix

$$\bar{\mathbf{X}}_k \in \mathbb{R}^{n \times (\tau + \tau_{\text{subs}})} = \left[\mathbf{X}_k, \sum_{k' \neq k} \mathbf{X}_{k'} \mathbf{\Pi}_k \right], \quad (5)$$

which is the concatenation of worker k ’s raw features and the *sum* of the random features from all other workers. $\mathbf{\Pi}$ is the SRHT matrix introduced in §2.2.

As we prove in Lemma 2, summing $\mathbb{R}^\tau \rightarrow \mathbb{R}^{\tau_{\text{subs}}}$ -dimensional random projections from $(K-1)$ blocks is equivalent to computing the $\mathbb{R}^{(p-\tau)} \rightarrow \mathbb{R}^{\tau_{\text{subs}}}$ -dimensional random projection in one go. The latter operation is impractical for very large p and not applicable when the features are distributed. Therefore, summing the random features from each worker allows the dimensionality reduction to be distributed across workers. Additionally, the summed random feature representation can be computed and combined very efficiently. We elaborate on this aspect in §5.

For a single worker the local, approximate primal problem is then

$$\min_{\bar{\beta} \in \mathbb{R}^{\tau + \tau_{\text{subs}}}} J_k(\bar{\beta}) := \sum_{i=1}^n f_i(\bar{\beta}^\top \bar{\mathbf{x}}_i) + \frac{\lambda}{2} \|\bar{\beta}\|_2^2 \quad (6)$$

where $\bar{\mathbf{x}}_i \in \mathbb{R}^{\tau + \tau_{\text{subs}}}$ is the i^{th} row of $\bar{\mathbf{X}}_k$. The corresponding dual problem for each worker in the DUAL-LOCO algorithm is

$$\max_{\alpha \in \mathbb{R}^n} - \sum_{i=1}^n f_i^*(\alpha_i) - \frac{1}{2n\lambda} \alpha^\top \tilde{\mathbf{K}}_k \alpha, \quad \tilde{\mathbf{K}}_k = \bar{\mathbf{X}}_k \bar{\mathbf{X}}_k^\top. \quad (7)$$

The following steps in Algorithm 1 detail respectively how the solution to (7) and the final DUAL-LOCO estimates are obtained.

Step 5. LocalDualSolver. The LocalDualSolver computes the solution for (7), the local dual problem. The solver can be chosen to best suit the problem at hand. This will depend on the absolute size of n and $\tau + \tau_{\text{subs}}$ as well as on their ratio. For example, we could use SDCA [18] or Algorithm 1 from [16].

Step 6. Obtaining the global primal solution. Each worker maps its local dual solution to the primal solution corresponding only to the coordinates in \mathcal{P}_k . In this way, each worker returns coefficients corresponding only to its own raw features. The final primal solution vector is obtained by concatenating the K local solutions. Unlike LOCO, we no longer require to discard the coefficients corresponding to the random features for each worker. Consequently, computing estimates is more efficient (especially when $p \gg n$).

4 DUAL-LOCO Approximation Error

In this section we bound the recovery error between the DUAL-LOCO solution and the solution to Eq. (1).

Theorem 1 (DUAL-LOCO error bound). *Consider a matrix $\mathbf{X} \in \mathbb{R}^{n \times p}$ with rank at most r . Assume that the loss $f(\cdot)$ is smooth with Lipschitz continuous gradients. For a subsampling dimension $\tau_{\text{subs}} \geq c_1 p K$ where $0 \leq c_1 \leq 1/K^2$, let β^* be the solution to (1) and $\hat{\beta}$ be the estimate returned by Algorithm 1. We have with probability at least $1 - K(\delta + \frac{p-\tau}{e^r})$*

$$\|\hat{\beta} - \beta^*\|_2 \leq \frac{\varepsilon}{1 - \varepsilon} \|\beta^*\|_2, \quad (8)$$

where $\varepsilon = \sqrt{\frac{c_0 \log(2r/\delta)r}{c_1 p}} < 1$.

Proof. By Lemma 4 and applying a union bound we can decompose the global optimization error in terms of the error due to each worker as $\|\beta^* - \hat{\beta}\|_2 = \sqrt{\sum_{k=1}^K \|\beta_k^* - \hat{\beta}_k\|_2^2} \leq \sqrt{K} \frac{\rho}{1-\rho} \|\beta^*\|_2$, which holds with probability $1 - K(\delta + \frac{p-\tau}{e^r})$. The final bound, (8) follows by setting $\rho = \sqrt{\frac{c_0 \log(2r/\delta)r}{\tau_{\text{subs}}}}$ and $\tau_{\text{subs}} \geq c_1 p K$ and noting that $\sqrt{K} \cdot \frac{\frac{\varepsilon}{\sqrt{K}}}{1 - \frac{\varepsilon}{\sqrt{K}}} \leq \frac{\varepsilon}{1 - \varepsilon}$. \square

Theorem 1 guarantees that the solution to DUAL-LOCO will be close to the optimal solution obtained by a single worker with access to all of the data. Our result relies on the data having rank $r \ll p$. In practice, this assumption is often fulfilled, in particular when the data is high dimensional. For a large enough projection dimension, the bound has only a weak dependence on K through the union bound used to determine the probability with which Eq. (8) holds. The error is then mainly determined by the ratio between the

rank and the random projection dimension. When the rank of \mathbf{X} increases for a fixed p , we need a larger projection dimension to accurately capture its spectrum. On the other hand, the failure probability increases with p and decreases with r . However, this countering effect is negligible as typically $\log(p - \tau) \ll r$.

5 Implementation and Experiments

In this section we report on the empirical performance of DUAL-LOCO in two sets of experiments. The first demonstrates the performance of DUAL-LOCO in a large, distributed classification task. The second is an application of ℓ_2 penalized regression to a problem in climate science where accurate recovery of the coefficient estimates is of primary interest.

Cross validation. In most practical cases, the regularization parameter λ is unknown and has to be determined via v -fold cross validation (CV). The chosen algorithm is usually run entirely once for each fold and each of l values of λ , leading to a runtime that is approximately $v \cdot l$ as large as the runtime of a single run³. In this context, DUAL-LOCO has the advantage that steps 3 and 4 in Algorithm 1 are independent of λ . Therefore, these steps only need to be performed *once per fold*. In step 5, we then estimate $\tilde{\alpha}_k$ for each value in the provided sequence for λ . Thus, the runtime of DUAL-LOCO will increase by much less than $v \cdot l$ compared to the runtime of a single run. The performance of each value for λ is then not only averaged over the random split of the training data set into v parts but also over the randomness introduced by the random projections which are computed and communicated once per fold. The procedure is provided in full detail in Algorithm 2 in Appendix C.

Implementation details. We implemented DUAL-LOCO using the Apache Spark framework⁴. Spark is increasingly gaining traction in the research community as well as in industry due to its easy-to-use high-level API and the benefits of in-memory processing. Spark is up to 100× faster than Hadoop MapReduce. Additionally, Spark can be used in many different large-scale computing environments and the various, easily-integrated libraries for a diverse set of tasks greatly facilitate the development of applications.

When communicating and summing the random features in Spark, DUAL-LOCO leverages the `treeReduce` scheme as illustrated in Figure 2(b). Summing has the advantage that increasing the number of workers simply introduces more layers in the tree struc-

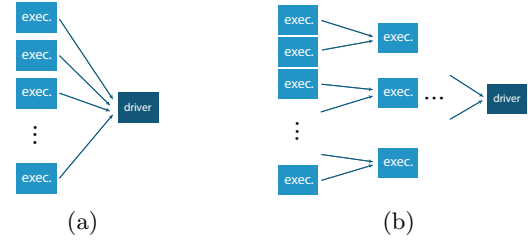


Figure 2: Schematic for the aggregation of the random features in Spark. (a) When concatenating the random features naively, every worker node (exec.) sends its random features to the driver from where they are broadcasted to all workers. (b) Using the `treeReduce` scheme we can reduce the load on the driver by summing the random features from each worker node as this operation is associative and commutative. Worker k is only required to subtract its own random features locally.

ture (Figure 2b) while the load on the driver remains constant and the aggregation operation also benefits from a parallel execution. Thus, when increasing K only relatively little additional communication cost is introduced which leads to speedups as demonstrated below.

In practice, we used the discrete cosine transform (DCT) provided in the FFT library `jTransforms`⁵⁶ and we ran DUAL-LOCO as well as `CoCoA`⁺ on a high-performance cluster⁷.

Competing methods. For the classification example, the loss function is the hinge loss. Although the problem is non-smooth, and therefore not covered by our theory, we still obtain good results suggesting that Theorem 1 can be generalized to non-smooth losses. Alternatively, for classification the smoothed hinge or logistic losses could be used. For the regression problem we use the squared error loss and modify `CoCoA`⁺ accordingly. As the `LocalDualSolver` we use SDCA [18].

We also compared DUAL-LOCO against the reference implementation of distributed loss minimization in the `MLlib` library in Spark using SGD and L-BFGS. However, after extensive cross-validation over regularization strength (and step size and mini-batch size in case of SGD), we observed that the variance was still very large and so we omit the `MLlib` implementations from the figures. A comparison between `CoCoA` and variants of SGD and mini-batch SDCA can be found in

⁵<https://sites.google.com/site/piotrwendykier/software/jtransforms>

⁶For the Hadamard transform, τ must be a power of two. For the DCT there is no restriction on τ and very similar theoretical guarantees hold.

⁷`CoCoA`⁺ is also implemented in Spark with code available from <https://github.com/gingsmith/cocoa>.

³“Approximately” since the cross validation procedure also requires time for testing. For a single run we only count the time it takes to estimate the parameters.

⁴Available from: <http://christinaheinze.github.io/loco-lib/>.

[5].

Kaggle Dogs vs Cats dataset. This is a binary classification task consisting of 25,000 images of dogs and cats⁸. We resize all images to 430×430 pixels and use OVERFEAT [19] – a pre-trained convolutional neural network – to extract $p = 200,704$ *fully dense* feature vectors from the 19th layer of the network for each image. We train on $n_{train} = 20,000$ images and test on the remaining $n_{test} = 5,000$. The size of the training data is 37GB with over 4 billion non-zero elements. All results we report in the following are averaged over five repetitions and by “runtime” we refer to wall clock time.

Figure 3 shows the median normalized training and test prediction MSE of DUAL-LOCO and CoCoA⁺ for different numbers of workers⁹. For DUAL-LOCO, we also vary the size of the random feature representation and choose $\tau_{subs} = \{0.005, 0.01, 0.02\} \times (p - \tau)$. The corresponding errors are labeled with DUAL-LOCO 0.5, DUAL-LOCO 1 and DUAL-LOCO 2. Note that combinations of K and τ_{subs} that would result in $\tau < \tau_{subs}$ cannot be used, e.g. $K = 192$ and $\tau_{subs} = 0.01 \times (p - \tau)$. We ran CoCoA⁺ until a duality gap of 10^{-2} was attained so that the number of iterations varies for different numbers of workers¹⁰. Notably, for $K = 48$ more iterations were needed than in the other cases which is reflected in the very low training error in this case. The fraction of local points to be processed per round was set to 10%. We determined the regularization parameter λ via 5-fold cross validation.

While the differences in training errors between DUAL-LOCO and CoCoA⁺ are notable, the differences between the test errors are minor as long as the random feature representation is large enough. Choosing τ_{subs} to be only 0.5% of $p - \tau$ seems to be slightly too small for this data set. When setting τ_{subs} to be 1% of $p - \tau$ the largest difference between the test errors of DUAL-LOCO and CoCoA⁺ is 0.9%. The averaged mean squared prediction errors and their standard deviations are collected in Table 1 in Appendix C.

Next, we would like to compare the wall clock time needed to find the regularization parameter λ via 5-fold cross validation. For CoCoA⁺, using the number of iterations needed to attain a duality gap of 10^{-2} would lead to runtimes of more than 24 hours for $K \in \{48, 96, 192\}$ when comparing $l = 20$ possible values for λ . One might argue that using a duality gap of 10^{-1} is sufficient for the cross validation

⁸<https://www.kaggle.com/c/dogs-vs-cats>

⁹In practice, this choice will depend on the available resources in addition to the size of the data set.

¹⁰For K ranging from 12 to 192, the number of iterations needed were 77, 207, 4338, 1966, resp. 3199.

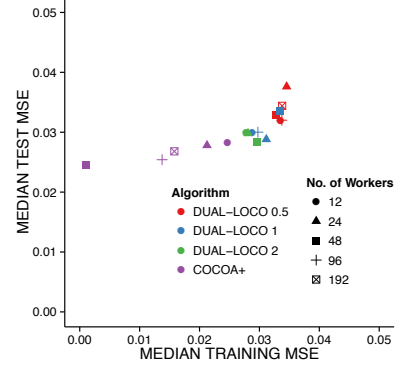


Figure 3: Dogs vs Cats data: Median normalized training and test prediction MSE based on 5 repetitions.

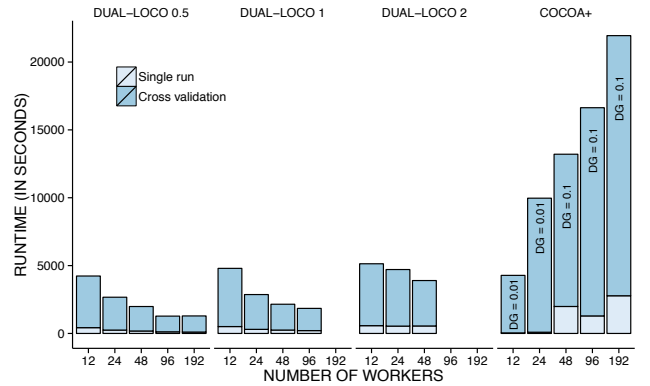


Figure 4: Total wall clock time including 5-fold CV over $l = 20$ values for λ . For CoCoA⁺, we use a duality gap (DG) of 10^{-1} for the CV runs when $K \geq 48$.

runs which would speed up the model selection procedure significantly as much fewer iterations would be required. Therefore, for $K \geq 48$ we use a duality gap of 10^{-1} during cross validation and a duality gap of 10^{-2} for learning the parameters, once λ has been determined. Figure 4 shows the runtimes when $l = 20$ possible values for λ are compared; Figure 6(a) compares the runtimes when cross validation is performed over $l = 50$ values. The absolute runtime of CoCoA⁺ for a single run is smaller for $K = 12$ and $K = 24$ and larger for $K \in \{48, 96, 192\}$, so using more workers increased the amount of wall clock time necessary for job completion. The total runtime, including cross validation and a single run to learn the parameters with the determined value for λ , is always smaller for DUAL-LOCO, except when $K = 12$ and $l = 20$.

Figures 5 and 6(b) show the relative speedup of DUAL-LOCO and CoCoA⁺ when increasing K . The speedup is computed by dividing the runtime for $K = 12$ by the runtime achieved for the corresponding $K = \{24, 48, 96, 192\}$. A speedup value smaller than 1 implies an *increase* in runtime. When considering a single run, we run CoCoA⁺ in two different settings: (i) We use the number of iterations that are needed to

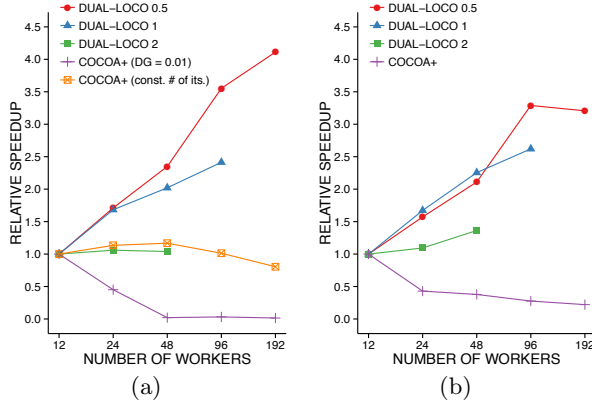


Figure 5: Relative speedup for (a) a single run and (b) 5-fold CV over $l = 20$ values for λ .

obtain a duality gap of 10^{-2} which varies for different number of workers¹⁰. Here, the speedup is smaller than 1 for all K . (ii) We fix the number of outer iterations to a constant number. As K increases, the number of inner iterations decreases, making it easier for CoCoA⁺ to achieve a speedup. We found that although CoCoA⁺ attains a speedup of 1.17 when increasing K from 12 to 48 (equivalent to a decrease in runtime of 14%), CoCoA⁺ suffers a 24% increase in runtime when increasing K from 12 to 192.

For DUAL-LOCO 0.5 and DUAL-LOCO 1 we observe significant speedups as K increases. As we split the design matrix by features the number of observations n remains constant for different number of workers. At the same time, the dimensionality of each worker’s local problem decreases with K . Together with the efficient aggregation of the random features, this leads to shorter runtimes. In case of DUAL-LOCO 2, the communication costs dominate the costs of computing the random projection and of the LocalDualSolver, resulting in much smaller speedups.

Although CoCoA⁺ was demonstrated to obtain speedups for low-dimensional data sets [2] it is plausible that the same performance cannot be expected on a very high-dimensional data set. This illustrates that in such a high-dimensional setting splitting the design matrix according to the columns instead of the rows is more suitable.

Climate data. This is a regression task where we demonstrate that the coefficients returned by DUAL-LOCO are interpretable. The data set contains the outcome of control simulations of the GISS global circulation model [20, 21] and is part of the CMIP5 climate modeling ensemble. We aim to forecast the monthly global average temperature Y in February using the air pressure measured in January. Results are very similar

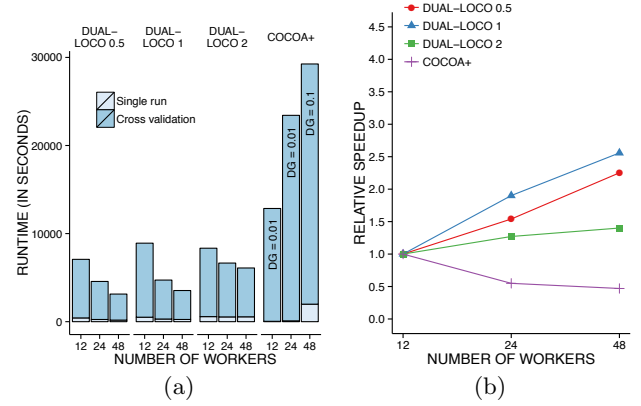


Figure 6: 5-fold CV over $l = 50$ values for λ : (a) Total wall clock time and (b) relative speedup.

for other months. The $p = 10,368$ features are pressure measurements taken at 10,368 geographic grid points in January. The time span of the climate simulation is 531 years and we use the results from two control simulations, yielding $n_{\text{train}} = 849$ and $n_{\text{test}} = 213$.

In Figure 7 we compare the coefficient estimates for four different methods. The problem is small enough to be solved on a single machine so that the full solution can be computed (using SDCA; cf. Figure 7(a)). This allows us to report the normalized parameter estimation mean squared error ($\text{MSE}_{\hat{\beta}}$) with respect to the full solution in addition to the normalized mean squared prediction error (MSE). The solution indicates that the pressure differential between Siberia (red area, top middle-left) and Europe and the North Atlantic (blue area, top left and top right) is a good predictor for the temperature anomaly. This pattern is concealed in Figure 7(b) which shows the result of up-projecting the coefficients estimated following a random projection of the columns. Using this scheme for prediction was introduced in [15]. Although the MSE is similar to the optimal solution, the recovered coefficients are not interpretable as suggested by [16]. Thus, this method should only be used if prediction is the sole interest. Figure 7(c) shows the estimates returned by DUAL-LOCO which is able to recover estimates which are close to the full solution. Finally, Figure 7(d) shows that CoCoA⁺ also attains accurate results.

Considering a longer time period or adding additional model variables such as temperature, precipitation or salinity rapidly increases the dimensionality of the problem while the number of observations remains constant. Each additional variable adds 10,368 dimensions per month of simulation. Estimating very high-dimensional linear models is a significant challenge in climate science and one where distributing the problem across features instead of observations is

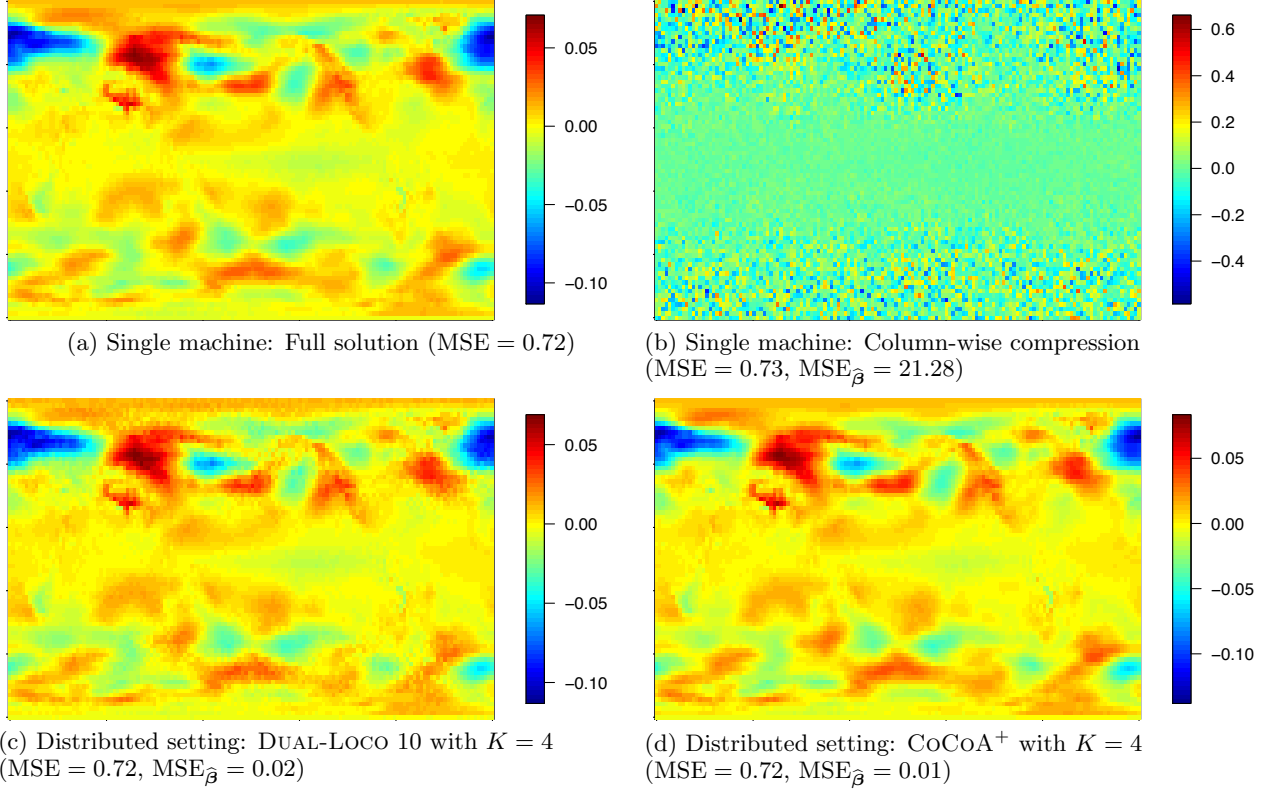


Figure 7: Climate data: The regression coefficients are shown as maps with the prime median (passing through London) corresponding to the left and right edge of the plot. The Pacific Ocean lies in the center of each map.

advantageous. The computational savings are much larger when distributing across features as $p \gg n$ and thus reducing p is associated with larger gains than when distributing across observations.

6 Conclusions and further work

We have presented DUAL-LOCO which considers the challenging and rarely studied problem of statistical estimation when data is distributed across features rather than samples. DUAL-LOCO generalizes LOCO to a wider variety of loss functions for regression and classification. We show that the estimated coefficients are close to the optimal coefficients that could be learned by a single worker with access to the entire dataset. The resulting bound is more intuitive and tighter than previous bounds, notably with a very weak dependence on the number of workers. We have demonstrated that DUAL-LOCO is able to recover accurate solutions for large-scale estimation tasks whilst also achieving better scaling than a state-of-the-art competitor, CoCoA⁺, as K increases. Additionally, we have shown that DUAL-LOCO allows for fast model selection using cross-validation.

The dual formulation is convenient for ℓ_2 penalized problems but other penalties are not as straightforward. Similarly, the theory only holds for smooth loss functions. However, as demonstrated empirically

DUAL-LOCO also performs well with a non-smooth loss function.

As n grows very large, the random feature matrices may become too large to communicate efficiently even when the projection dimension is very small. For these situations, there are a few simple extensions we aim to explore in future work. One possibility is to first perform row-wise random projections (c.f. [22]) to further reduce the communication requirement. Another option is to distribute \mathbf{X} according to rows and columns.

Contrary to stochastic optimization methods, the communication of DUAL-LOCO is limited to a single round. For fixed n , p and τ_{subs} , the amount of communication is deterministic and can be fixed ahead of time. This can be beneficial in settings where there are additional constraints on communication (for example when different blocks of features are distributed *a priori* across different physical locations).

Clearly with additional communication, the theoretical and practical performance of DUAL-LOCO could be improved. For example, [16] suggest an iterative dual random projection scheme which can reduce the error in Lemma 4 exponentially. A related question for future research involves quantifying the amount of communication performed by DUAL-LOCO in terms of known minimax lower bounds [23].

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