Divide-and-Conquer Learning with Nyström: Optimal Rate and Algorithm

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Abstract
Kernel Regularized Least Squares (KRLS) is a fundamental learner in machine learning. However, due to the high time and space requirements, it has no capability to large scale scenarios. Therefore, we propose DC-NY, a novel algorithm that combines divide-and-conquer method, Nyström, conjugate gradient, and preconditioning to scale up KRLS, has the same accuracy of exact KRLS and the minimum time and space complexity compared to the state-of-the-art approximate KRLS estimates. We present a theoretical analysis of DC-NY, including a novel error decomposition with the optimal statistical accuracy guarantees. Extensive experimental results on several real-world large-scale datasets containing up to 1M data points show that DC-NY significantly outperforms the state-of-the-art approximate KRLS estimates.

Introduction
In nonparametric statistical learning, kernel methods (Liu et al. 2018; Li et al. 2018; Liu and Liao 2014; Ding et al. 2018; Liu et al. 2019) have made remarkable achievements for kernel regularized least squares (KRLS) (Liu et al. 2017; Taylor and Cristianini 2004; Yin et al. 2019; Liu and Liao 2015; Liu, Jiang, and Liao 2014) by projecting data into high-dimensional space. Unfortunately, due to high time and memory consumption, typically at least quadratic in the number of examples, KRLS is unfeasible to deal with large-scale learning despite excellent theoretical guarantee.

To address the problems, a variety of practical approximate approaches have been designed to avoid the costly expense of finding an exact minimizer: (1) Iterative optimization (Lo et al. 2008). It provides regularization against over-fitting and improves computational efficiency by limited and small iterations. The representative include gradient descent (Carratino, Rudi, and Rosasco 2018; Lin and Cevher 2018), preconditioned conjugate gradient (Fasshauer and Mccourt 2012; Yang, Pilanci, and Wainwright 2015; Gonen, Orabona, and Shalev-Shwartz 2016; Ma and Belkin 2017), and accelerated extensions (Raskutti, Wainwright, and Yu 2014; Cutajar et al. 2016; Bo et al. 2014); (2) Random projections (Williams and Seeger 2001; Smola 2000). It reduces dimensions of data to reduce the cost of matrix multiplication. Classical examples include Nyström (Rudi, Carratino, and Rosasco 2017; Tu et al. 2016) and random features (Rudi, Camoriano, and Rosasco 2016; Rahimi and Recht 2007); (3) Distributed learning (Guo, Lin, and Shi 2017). It computes KRLS in parallel by dividing into some subsets and then merge the result from each subset to get the final approximation. Recently, combinations of those accelerated algorithms have also captured a lot of attention, of which learning properties have been explored including the combination of divide-and-conquer and SGD (Lin and Cevher 2018) and the combination of divide-and-conquer and random features (Li, Liu, and Wang 2019). Even though the state-of-the-art KRLS estimates can preserve the same optimal statistical accuracy of exact KRLS, the computational requirements of them are still prohibitive faced with large-scale datasets, namely, there are no corresponding computational lower bounds.

In this paper, we investigate the algorithm of combining divide-and-conquer, Nyström, conjugate gradient and preconditioning to deal with extremely large-scale applications, which achieves the same accuracy of exact kernel regularized least squares with only a fraction of computations. Complexity analysis shows that the proposed algorithm solve KRLS with max(\(\frac{Nm}{p^2}, m^3\)) time and \(\frac{Nm}{p}\) space, where \(N\) is the number of data points, \(m\) is the sampling scale, and \(p\) is the number of partitions. Our theoretical analysis derives optimal statistical rates in a basic setting. Under benign conditions, the regularization parameter \(\lambda \approx 1/\sqrt{N}\), the proposed algorithm can reach the optimal convergence rate \(1/N\). Most importantly, extensive experimental results on large-scale datasets containing up to 1M data points show that the proposed algorithm can process millions of data points in just several seconds and has absolute advantage over the state-of-the-art approximate KRLS in terms of efficiency and accuracy. To the best of our knowledge, it is the first time utilizing these approximate methods and achieving fruitful results.

The reminder of the paper is organized as below: Section 2 introduces the related work. Section 3 states the background and the proposed approximate KRLS estimator. The corresponding theoretical assessments follows in section 4. Finally, we present the experiments and conclusions.
Table 1: Computational complexity of the classical approximation algorithms in KRLS estimates under the condition of the same statistical accuracy. The second, third, fourth, fifth and sixth columns correspond to the training time complexity, space complexity, the number of partitions $p$, sampling scale $m$, and the value of $r$ and $\gamma$, respectively. $N$ denotes the number of training data, where $N/p > m$.

<table>
<thead>
<tr>
<th>Algorithms</th>
<th>Time</th>
<th>Space</th>
<th>$p$</th>
<th>$m$</th>
<th>$r, \gamma$</th>
</tr>
</thead>
<tbody>
<tr>
<td>KRLS (Caponnetto and Vito 2007)</td>
<td>$N^3$</td>
<td>$N^2$</td>
<td>/</td>
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<td>/</td>
</tr>
<tr>
<td>Iterative (Avron, Clarkson, and Woodruff 2017)</td>
<td>$N^2$</td>
<td>/</td>
<td>/</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>Random Features (Rudi, Camoriano, and Rosasco 2016)</td>
<td>$Nm^2$</td>
<td>$Nm$</td>
<td>/</td>
<td>$N^{(2r-1)\gamma+1}$</td>
<td>[1/2,1], [0,1]</td>
</tr>
<tr>
<td>Nyström (Rudi, Camoriano, and Rosasco 2015)</td>
<td>$Nm^2$</td>
<td>$Nm$</td>
<td>/</td>
<td>$N^{1/(r+\gamma)}$</td>
<td>[1/2,1], (0,1)</td>
</tr>
<tr>
<td>Nyström-iterative (Rudi, Carratino, and Rosasco 2017)</td>
<td>$Nm + m^3$</td>
<td>$Nm$</td>
<td>/</td>
<td>$N^{(2r-1)\gamma+1}$</td>
<td>[1/2,1], (0,1)</td>
</tr>
<tr>
<td>DC (Guo, Lin, and Shi 2017)</td>
<td>$N^3$</td>
<td>$N^{2\gamma+1}$</td>
<td>/</td>
<td>/</td>
<td>/</td>
</tr>
<tr>
<td>DC-Random Features (Li, Liu, and Wang 2019)</td>
<td>$\frac{Nm^2}{p}$</td>
<td>$\frac{Nm}{p}$</td>
<td>$\frac{Nm}{p}$</td>
<td>$\frac{N^{2\gamma+1}}{p(2r-1)\gamma+1}$</td>
<td>[1/2,1], (0,1)</td>
</tr>
<tr>
<td>DC-NY (This paper)</td>
<td>$\max(\frac{Nm^2}{p}, m^3)$</td>
<td>$\frac{Nm}{p}$</td>
<td>$\frac{Nm}{p}$</td>
<td>$\frac{N^{2\gamma+1}}{p(2r-1)\gamma+1}$</td>
<td>(1/2,1), (0,1)</td>
</tr>
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Related Work

To overcome the computational and memory bottleneck of KRLS, practical algorithms are developed, including Nyström approach (Rudi, Carratino, and Rosasco 2017; Camoriano et al. 2016) and divide-and-conquer (Zhang, Duchi, and Wainwright 2013; Li, Liu, and Wang 2019) of which statistical properties are well studied. Nyström (Rudi, Camoriano, and Rosasco 2015; Camoriano et al. 2016) tactfully constructs some small-scale matrices, by sampling the dataset, to approximate the raw kernel matrix so that the time and space complexity can make a sudden drop. The typical Nyström method FALKON, proposed in (Rudi, Carratino, and Rosasco 2015), combines Nyström and preconditioned conjugate gradient (PCG) to obtain the optimal statistical accuracy. Divide-and-conquer methods divide the dataset into several small blocks so as to reduce the number of data in one processor. The representational method (Guo, Lin, and Shi 2017) utilizes divide-and-conquer to obtain a substantial reduction in computation time.

Further, we combine divide-and-conquer and Nyström to approximate KRLS, and accelerate the solution with PCG so as to obtain high computation gains and sound statistical guarantees. It is a non-trivial extension of these approximate approaches with technical challenges in algorithm design and theoretical analysis. Based on a novel partitioning, clever scaling, and the standard integral operator framework, the error is bounded tightly to obtain the optimal statistical performance. Table 1 shows the main information of the state-of-the-art KRLS estimators with the optimal statistical accuracy. Evidently, a substantial step in provably reducing the time complexity is taken by us when we have the same statistical accuracy. In detail, compared to the state-of-the-art approximate KRLS estimates, we reduce the time complexity at least by a factor of $\min(m, \frac{N}{p})$, where the number of data points in each processor $\frac{N}{p}$ is bigger than the sampling scale $m$. Considering the concrete values of $m$ and $p$ this leads to a computational cost for optimal generalization, we reduce the space complexity by a factor of $N^{(2r-1)\gamma+1}$ compared to the state-of-the-art KRLS estimate.

Statistical and Computational Trade-offs in KRLS

Let $\rho$ be a probability measure on $\mathbf{X} \times \mathbb{R}$, which is fixed but unknown and where, $\mathbf{X}$ and $\mathbb{R}$ are the input and output spaces. Data $(x_i, y_i)_{i=1}^N$ are sampled identically and independently from $\mathbf{X} \times \mathbb{R}$ with respect to $\rho$. In the supervised learning, the problem of estimating a function from random noisy data can be formalized as minimizing the expected risk

$$\inf_{f \in H} \mathcal{E}(f), \quad \mathcal{E}(f) = \int (f(x) - y)^2 d\rho(x, y), \tag{1}$$

where $H$ is a space of candidate solutions. An ideal empirical solution $\hat{f}$ should correspond to small excess risk

$$\mathcal{R}(\hat{f}) = \mathcal{E}(\hat{f}) - \inf_{f \in H} \mathcal{E}(f). \tag{2}$$

Kernel Regularized Least Squares (KRLS)

Kernel Regularized Least Squares (KRLS) introduces the kernel trick which is based on choosing a separable reproducing kernel Hilbert space (RKHS) $\mathcal{H}$ with inner product $<\cdot, \cdot>_{\mathcal{H}}$. The reproducing kernel $K : \mathbf{X} \times \mathbf{X} \to \mathbb{R}$ is a positive definite kernel, measurable and uniformly bounded. We denote with $K_{ij}$ the function $K(x_i, \cdot)$ and have $(K_{ij})_{ij} = K(x_i, x_j)$ for all $x_1, \ldots, x_N \in \mathbf{X}$. The KRLS method for solving the problem in Eq.(1) can be expressed as

$$\hat{f}_{N,\lambda} = \arg\min_{f \in \mathcal{H}} \frac{1}{N} \sum_{i=1}^N (f(x_i) - y_i)^2 + \lambda \|f\|_{\mathcal{H}}^2, \lambda > 0. \tag{3}$$

It is obvious that a solution $\hat{f}_{N,\lambda}$ exists and is unique. According to the representer theorem (Schölkopf et al. 2002), Eq. (3) can be transferred to

$$\hat{f}_{N,\lambda}(x) = \sum_{i=1}^N \alpha_i K(x_i, x) \tag{4}$$

with

$$\alpha = (K_N + \lambda N)^{-1} y,$$
where \( x_1, \ldots, x_N \) are the data points, \( y = (y_1, \ldots, y_N) \) are the corresponding labels, and \( K_N \) is the kernel matrix.

Solving Eq.(4) is challenging as \( N \) increases: the time cost is \( O(N^3) \) for the linear system and the space cost is \( O(N^2) \) for storing the kernel matrix. To accelerate KRLS with accuracy guarantees, we proposed the following methods.

**Divide-and-Conquer KRLS with Nyström**

In this paper, we exploit divide-and-conquer learning, along with Nyström, preconditioning and conjugate gradient, to process the large-scale problem, which has the minimum time and space complexity compared to the state-of-the-art KRLS estimates, and has the same accuracy of exact kernel regularized least squares. Meanwhile, we provide a solid theoretical proof to guarantee the optimal convergence rate.

**KRLS with Nyström** We consider Nyström subsampling to reduce computational requirements, which uses a smaller matrix obtained from random sampling to approximate the empirical kernel matrix. Thus, a smaller hypothesis space \( \mathcal{H}_m \) is introduced

\[
\mathcal{H}_m = \{ f | f = \sum_{i=1}^{m} \alpha_i K(\tilde{x}_i, \cdot), \alpha \in \mathbb{R}^m \},
\]

where sampling scale \( m \leq N, \{ \tilde{x}_1, \ldots, \tilde{x}_m \} \) are Nyström centers sampled uniformly at random without replacement from the training set. The corresponding minimizer over the space \( \mathcal{H}_m \) is in the form:

\[
f^m_\lambda(x) = \sum_{i=1}^{m} \tilde{\alpha}_i K(\tilde{x}_i, x)
\]

with

\[
\tilde{\alpha} = \left( K_{Nm}^T K_{Nm} + \lambda N K_{mm} \right) \left( K_{Nm}^T y \right) z
\]

where \( \mathbf{H} \) is the Moore-Penrose pseudoinverse of kernel matrix \( K_{Nm} \).

**KRLS with Nyström and PCG** Gradient methods (Rudi, Carratino, and Rosasco 2017; Kumar, Mohri, and Talwalkar 2012; Dieuleveut and Bach 2016) are general strategies for the unconstrained optimization problem, which are simplicity, low iteration consumption. The conjugate gradient method (Saad 1996) is one of the most popular ones, which does not need to specify the step-size and whose speed of convergence can benefit from preconditioning. We can use it to compute the coefficients \( \tilde{\alpha} \) in Eq.(5).

The idea behind preconditioning is to use a suitable matrix \( \mathbf{P} \) to define an equivalent linear system with better condition number. Therefore, we consider random projections to approximately compute a preconditioner:

\[
\mathbf{P} = \frac{1}{\sqrt{N}} \mathbf{T}^{-1} \mathbf{A}^{-1},
\]

where \( \mathbf{T} = \text{chol}(K_{mm}) \) and \( \mathbf{A} = \text{chol}(\frac{1}{m} \mathbf{T}^T + \lambda I) \).

Therefore, the Nyström KRLS with conjugate gradient and preconditioning can be seen as solving the following system

\[
\mathbf{P}^T \mathbf{H} \mathbf{\tilde{\alpha}} = \mathbf{P}^T \mathbf{z},
\]

with

\[
f^m_\lambda(x) = \sum_{i=1}^{m} \tilde{\alpha}_i K(\tilde{x}_i, x),
\]

where \( \mathbf{\tilde{\alpha}} \) is solved via \( t \)-step conjugate gradient algorithm and \( t \in \mathbb{N} \).

**Divide-and-Conquer KRLS with Nyström and PCG** For further saving computational costs, we consider divide-and-conquer based on the solver in Eq.(7). Given the training dataset \( D \), \( \{ (x_i, y_i) \}_{i=1}^N \), the data in \( D \) can be randomly partitioned into \( p \) disjoint subsets \( \{ D_j \}_{j=1}^p \) with \( |D_1| = |D_2| = \ldots = |D_p| = n \). The \( j \)-th data subset is sent to the \( j \)-th local processor, where \( j \in \{1, p\} \), which is separately used to solve the solver in Eq.(7) to get \( \tilde{\alpha}_j \) and the estimator \( f^m_{D_j, \lambda} \).

Namely, in each local processor, we use the Nyström method in Eq.(7) to solve KRLS. Then, the final training estimator \( \hat{f}^m_{D, \lambda} \) is obtained by combining the individual estimators to the center, whose formula is as follows:

\[
\hat{f}^m_{D, \lambda} = \frac{1}{p} \sum_{j=1}^{p} f^m_{D_j, \lambda}.
\]

**Algorithm 1 Divide-and-Conquer KRLS with Nyström (DC-NY)**

**Input:** Training dataset \( \{ (x_i, y_i) \}_{i=1}^N \in \mathbb{R}^{N \times d} \), \( \{ y_i \}_{i=1}^N \in \mathbb{R}^N \), kernel parameter, regularization parameter \( \lambda \), sampling scale \( m \), number of iterations \( t \), number of partitions \( p \).

**Output:** \( \hat{f}^m_{D, \lambda}(x) \)

1. randomly partition the training dataset into \( p \) disjoint subsets \( \{ D_j \}_{j=1}^p \);
2. sent each data subset to each local processor;
3. // In parallel: take \( j \)-th local processor for example
4. get \( \tilde{\alpha}_j \) based on \( j \)-th training subset by solving the problem in Eq.(7);
5. get the estimator \( \hat{f}^m_{D_j, \lambda}(x) = \sum_{i=1}^{m} \hat{\alpha}_j i K(x_i, x) \) in \( j \)-th local processor;
6. // End parallelism
7. compute the final estimator by synthesizing each one:

\[
\hat{f}^m_{D, \lambda}(x) = \frac{1}{p} \sum_{j=1}^{p} \hat{f}^m_{D_j, \lambda}(x).
\]

In the prediction stage, based on the trained \( \tilde{\alpha}_j \), the new query point \( x_{test} \) is transmitted to each local processor to get an estimation \( \hat{f}^m_{D_j, \lambda}(x_{test}) \), then we get the finally predicted estimation \( \hat{f}^m_{D, \lambda}(x_{test}) = \frac{1}{p} \sum_{j=1}^{p} \hat{f}^m_{D_j, \lambda}(x_{test}) \) by synthesizing each \( \hat{f}^m_{D_j, \lambda}(x_{test}) \).

The detailed process of proposed algorithm (DC-NY) is summarized in Algorithm 1.
Theoretical Assessments

In this section, for exploring the generalization ability, we firstly introduce four standard assumptions, which are widely used in statistical learning of squared loss (Smale and Zhou 2007; Caponnetto and Vito 2007; Rudi, Carratino, and Rosasco 2017; Li, Liu, and Wang 2019). Under the basic assumptions, the theoretical bound of the proposed algorithm is provided, which is the same as that of the exact Kernel Regularized Least Squares (KRLS).

**Assumption 1** There exists an \( f_\mathcal{H} \in \mathcal{H} \) such that

\[
\mathcal{E}(f_\mathcal{H}) = \min_{f \in \mathcal{H}} \mathcal{E}(f).
\]  

(9)

The above assumption is standard in kernel-based nonparametric regression (Smale and Zhou 2007; Caponnetto and Vito 2007), which shows that the problem in Eq.(1) has at least a solution. We also need a basic assumption on data distribution to derive probabilistic results.

**Assumption 2** Let \( z_x \) be the random variable \( z_x = y - f_\mathcal{H}(x) \), with \( x \in X \), and \( y \) distributed according to \( \rho(y|x) \). Then, there exists \( b, \sigma > 0 \) such that

\[
\mathbb{E}|z_x|^r \leq \frac{1}{\sigma^r} b^r e^{-2\sigma^2}
\]

for any \( r \geq 2 \), almost everywhere on \( X \).

This assumption is related to a noise assumption in the regression model, used to control random quantities, and holds the bounded \( y \). When \( |y| \leq b, \forall b > 1 \), the assumption is satisfied with \( \sigma = b \).

**Assumption 3** Let \( C \) be the covariance operator as

\[
C : \mathcal{H} \rightarrow \mathcal{H}, (f,g)_{\mathcal{H}} = \int_X f(x)g(x)d\rho_X(x), \forall f, g \in \mathcal{H}.
\]

For \( \lambda > 0 \), we define the random variable \( \mathcal{N}_x(\lambda) = (K_x, (C + \lambda I)^{-1}K_x)_{\mathcal{H}} \) with \( x \in X \) distributed according to \( \rho_X \) and let \( \mathcal{N}(\lambda) = \mathbb{E}\mathcal{N}_x(\lambda), \mathcal{N}_\infty(\lambda) = \sup_{x \in X} \mathcal{N}_x(\lambda) \).

The kernel \( K \) is measurable, \( C \) is bounded. Moreover, for all \( \lambda > 0 \) and a \( Q > 0 \),

\[
\mathcal{N}_\infty(\lambda) < \infty,
\]

(10)

\[
\mathcal{N}(\lambda) \leq Q\lambda^{-\gamma}, \quad 0 < \gamma \leq 1.
\]

(11)

This assumption controls the variance of the estimator (Rudi, Camoriano, and Rosasco 2015). Typically, this assumption ensures that the covariance operator is a well defined linear, continuous, self-adjoint, positive operator. If the kernel satisfied \( \sup_{x \in X} K(x, x) = \kappa^2 < \infty \), we have \( \mathcal{N}_\infty(\lambda) \leq \kappa^2/\lambda \) for all \( \lambda > 0 \), \( \gamma \) affects the size of RKHS \( \mathcal{H} \), namely it quantifies the capacity assumption. The more benign situation with smaller \( \mathcal{H} \) is obtained when \( \gamma \rightarrow 0 \). Note that, because the operator \( C \) is trace class, Eq.(11) always holds for \( \gamma = 1 \).

The bias/approximation error of KRLS can be controlled by the following assumption (Rudi, Camoriano, and Rosasco 2015).

**Assumption 4** There exists \( s > 0, 1 \leq R < \infty \), such that

\[
\|C^{-s} f_\mathcal{H}\|_\mathcal{H} < R.
\]

The degree, \( f_\mathcal{H} \), can be well approximated by functions in the RKHS \( \mathcal{H} \), can be quantified by this assumption. This assumption can be seen as regularity of \( f_\mathcal{H} \).

In the following, we quantify the quality of empirical solutions of Eq.(1) obtained by schemes of Eq.(8) in terms of the quantities in Assumptions 1, 2, 3 and 4.

**Theorem 1.** Under Assumptions 1,2,3 and 4, let \( \delta \in (0,1], r = 1/2 + \min(s,1/2), \gamma \in (0,1], \lambda = N^{-\frac{1}{r+1}}, N \geq n_0 \) with \( n_0 \in \mathbb{N} \), and \( f_{\mathcal{H},\lambda}^n \) be the estimator. When

\[
t \geq \mathcal{O}(\log(N)), \quad p \leq N^{\frac{1}{2(r+1)}}, \quad m \geq \mathcal{O}(N^{\frac{1}{r+1}}),
\]

with probability \( 1 - \delta \), we have

\[
\mathbb{E}[\|\hat{f}_{\mathcal{H},\lambda}^n\| - \mathcal{E}(f_\mathcal{H})] = \mathcal{O}(N^{\frac{1}{2(r+1)}}).
\]

(13)

**Remark 1.** Note that, \( \mathcal{O}(N^{-\frac{1}{r+1}}) \) is the optimal convergence rate (Caponnetto and Vito 2007). Under the same convergence rates, the convergence rate of the proposed algorithm (DC-NY) is the same as the bounds of the exact KRLS (Steinwart et al. 2009), DC (Guo, Lin, and Shi 2017), Nyström (Rudi, Camoriano, and Rosasco 2015) and Nyström-iterative (Rudi, Carratino, and Rosasco 2017) by \( p \leq N^{\frac{1}{2(r+1)}}, \) and \( m \geq \mathcal{O}(N^{\frac{1}{r+1}}) \). Although Random Features (Rudi, Camoriano, and Rosasco 2016) and DC-Random Features (Li, Liu, and Wang 2019) can also obtain the same optimal convergence rate as DC-NY, their corresponding \( m \geq \mathcal{O}(N^{\frac{1}{2(r+1)}}) \) is bigger than our \( m \geq \mathcal{O}(N^{\frac{1}{r+1}}) \). Namely, under the same \( m \), the accuracy of the proposed method is theoretically better than that of Random Features and DC-Random Features. In the best case \( r = 1 \) and \( \gamma = 0 \), \( \lambda \approx 1/\sqrt{N} \), DC-NY can reach the convergence rate \( 1/N \). Theoretical analysis demonstrate that our algorithm is sound and effective.

**Remark 2.** The proposed algorithm DC-NY is an extraordinary expansion of approximate KRLS. We succeed in conquering the bottleneck that optimal learning rate for the combination of distributed learning algorithm, Nyström algorithm, preconditioner and conjugate gradient. By introducing a novel technique of error decomposition and applying standard integral operator framework, the proposed algorithm achieves a tight bound under some basic assumptions. This is the first time that combining these approximate methods, achieving the optimal statistical accuracy and the minimum time and space complexity.

Complexity Analysis

The dataset is divided into \( p \) subsets so that the number of data in each processor is \( N/p \).

The matrices \( \mathbf{T} \) and \( \mathbf{A} \) in Eq.(6) are upper-triangular matrices, so the time complexity of \( \mathbf{A}^{-1}\mathbf{x} \) and \( \mathbf{A}^{-T}\mathbf{x} \) (\( \mathbf{x} \) is a vector) are all \( n^2 \). The matrix \( \mathbf{P} \) does not need to be represented explicitly. The time cost for the preconditioner, namely computing the matrices \( \mathbf{T} \) and \( \mathbf{A} \), is \( \frac{1}{3}n^3 \).
which includes two Cholesky decompositions and one product of two triangular matrices. The time cost in Nyström with conjugate gradient is $O(Nm t/p)$. As stated in Theorem 1, using the conjugate gradient solver typically requires $O(\log(N))$ iterations. Therefore, the time complexity in DC-NY is $O(\max(N/m, p/m))$. Compared to exact kernel regularized least squares whose time complexity is $O(N^3)$, we reduce the time by a factor of $\min(N^2p, N^3)$. Compared to the state-of-the-art approximate KRLS estimates whose time complexity is $O(Nm^2/p)$, we reduce the time by a factor of $\min(m, N/p m)$, where the number of data in each processor $N/p$ is bigger than the sampling scale $m$.

In space complexity, the decisive element is the scale of matrix $K_{nm}$ in Eq.(7). Therefore, the required space complexity of the proposed algorithm is only $O(Nm/p)$ which is the minimum. Compared to the exact kernel regularized least squares whose space complexity is $O(N^2)$, we reduce the space complexity by a factor of $N^2/m$. Considering the concrete values of $m$ and $p$, this leads to a computational cost for optimal generalization, we reduce the space complexity by a factor of $\min(m, N/p m)$ compared to the state-of-the-art KRLS estimates, where $\frac{2r(2r+1)}{2r+1} > 0$. Details are shown in Table 1.

To the best of our knowledge, DC-NY currently possesses the best time and space complexity to achieve the optimal statistical accuracy of KRLS.

**Error Decomposition**

In this section, a novel technique of error decomposition is introduced for DC-NY. The main task of proving the generalization performance is to bound excess risk $R(f) = \mathcal{E}(f) - \inf_{f \in H} \mathcal{E}(f)$.

In order to describe the decomposition of excess risk clearly, we provide some estimators in advance. $\hat{f}_{m,D,\lambda}$ is our estimator in Eq.(8), namely the estimator in Eq.(8) after $t$ iterations of the conjugate gradient algorithm. $\tilde{f}_{m,D,\lambda}$ is the corresponding estimator by exact Nyström in Eq.(5), namely the estimator in Eq.(8) after infinite iterations of the conjugate gradient algorithm. $f_{m,D,\lambda}$ focus on noise-free data on the $j$-th subset $D_j$, $f_{m,D,\lambda}$ is the estimator in Eq.(5) on the total dataset $D$. We obtain the error decomposition in Lemma 1.

**Lemma 1.** Let $\hat{f}_{m,D,\lambda}$, $\tilde{f}_{m,D,\lambda}$, $f_{m,D,\lambda}$, $f_{m,D,\lambda}$ and $f_{m,D,\lambda}$ be defined as above, we have

$$
\mathbb{E} \left[ \mathcal{E}(\hat{f}_{m,D,\lambda}) \right] - \inf_{f \in \mathcal{H}} \mathcal{E}(f) \\
\leq 2\mathbb{E} \left[ \left\| \hat{f}_{m,D,\lambda} - \tilde{f}_{m,D,\lambda} \right\|_\rho^2 + \frac{4}{p^2} \sum_{j=1}^p \mathbb{E} \left[ \left\| f_{m,D_j,\lambda} - f_{m,D_j,\lambda} \right\|_\rho^2 \right] \\
+ \left( \frac{8}{p^2} + \frac{4}{p} \right) \sum_{j=1}^p \mathbb{E} \left[ \left\| f_{m,D_j,\lambda} - f_{m,D_j,\lambda} \right\|_\rho^2 \right] \\
+ \left( \frac{8}{p^2} + \frac{4}{p} \right) \sum_{j=1}^p \mathbb{E} \left[ \left\| f_{m,D_j,\lambda} - f_{m,D_j,\lambda} \right\|_\rho^2 \right] 
$$

(14)

The error decomposition and the standard integral operator framework are the keys to guarantee the generalization performance of DC-NY algorithm. Our error bound is not a simple sum of block errors. Instead, it uses clever scaling and novel partitioning, then each error is transformed reasonably and bounded tightly based on the integral operator. The error bound obtained by the conventional summation of block errors is much larger than that obtained by our method.

**Experiment**

We empirically analyze the performance of the proposed algorithm compared with the classical algorithms, considering
a Gaussian kernel of width $\sigma$. The form of kernel function is as below $K(x_1, x_2) = e^{-\frac{1}{2\sigma^2} (x_1 - x_2)^2}$. Each experiment is measured on a server with 2.40GHz Intel(R) Xeon(R) E5-2630 v3 CPU and 32 GB of RAM in Matlab.

**Dataset Preparation**

The comparative experiments are based on four real-world datasets: SUSY, HIGGS, YearPredictionMSD and covtype, from website 1. Each feature has been normalized subtracting its mean and dividing for its variance. The details are shown in Table 2. We randomly sample $1 \times 10^6$ data points on SUSY and HIGGS, use the whole of YearPredictionMSD and covtype, and then randomly divide each experimental dataset into training set and prediction set, of which the training set accounts for 70%.

**Baselines and Parameters**

In order to avoid contingency, each experiment is repeated 10 times. For ensuring fairness, we use the same way to tune parameters $\sigma$ in $2^{-5:2^{-5:10}+0.5:10}$ and $\lambda$ in $2^{-21:+1:3}$, on each dataset and algorithm. Maybe the selected parameters are not optimal, but they are sufficient to achieve satisfactory results. The detail of parameters $\sigma$ and $\lambda$ in DC-NY are displayed in Table 2.

We will compare our method with 3 methods. (1) RF: it is the abbreviation of Random Features (Rudi, Camoriano, and Rosasco 2016), which is a classical approximate KRLS. We use the code from website 2. (2) NY: it represents Nyström-iterative (Rudi, Carratino, and Rosasco 2017), which combine Nyström with iterative methods. The code is from website 3. (3) DC-RF: it represents the algorithm (Li, Liu, and Wang 2019): combining Random Features with Divide-and-Conquer. The code is from website 4. (4) DC-NY: this is the proposed algorithm.

The error is measured with RMSE for regression problems, and with classification error for the classification problems, to be consistent with the literature.

**Parameters Evaluation**

As shown in (a) of Figure 1, when $t > 4$, the errors of DC-NY have converged on four datasets, which is consistent with our theoretical analysis $t > O(log(N))$.

In (b) and (c) of Figure 1, the test errors of DC-NY decline significantly when sampling scale $m$ is a small number, and when $m = \sqrt{N}$, DC-NY has converged on four datasets which is in line with the theoretical reasoning. The bigger the $m$ is, the longer the training time is. Therefore, in practice, we only need to take a small $m$ to obtain a satisfactory error, which will result in significant savings in resources.

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1https://www.csie.ntu.edu.tw/~cjlin/libsvmtools/datasets/
2https://github.com/superlj666
3https://github.com/LCSL/FALKON-paper
4https://github.com/superlj666
Table 3: Comparison of average training time (left) in seconds and average test error (right) in solving KRLS between RF, NY, DC-RF and DC-NY algorithms on YearPredictionMSD, covtype, SUSY and HIGGS datasets, with partitions $p = 2$, iteration times $t = 7$, sampling scale $m = 1500$ and 2000. We bold the numbers of the best algorithm. YPMSD is the abbreviation of YearPredictionMSD.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>RF(m = 1500)</th>
<th>NY(m = 1500)</th>
<th>DC-RF(m = 1500)</th>
<th>DC-NY(m = 1500)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>time</td>
<td>error</td>
<td>time</td>
<td>error</td>
</tr>
<tr>
<td>YPMSD</td>
<td>7.270 ± 0.061 ± 0.00008</td>
<td>2.90 ± 0.058 ± 0.00012</td>
<td>3.560 ± 0.062 ± 0.00007</td>
<td>1.60 ± 0.058 ± 0.00013</td>
</tr>
<tr>
<td>covtype</td>
<td>17.90 ± 0.233 ± 0.00079</td>
<td>5.02 ± 0.185 ± 0.00050</td>
<td>9.060 ± 0.234 ± 0.00013</td>
<td>2.56 ± 0.185 ± 0.00027</td>
</tr>
<tr>
<td>SUSY</td>
<td>19.24 ± 0.221 ± 0.00031</td>
<td>8.32 ± 0.197 ± 0.00068</td>
<td>9.020 ± 0.223 ± 0.00027</td>
<td>4.09 ± 0.197 ± 0.00064</td>
</tr>
<tr>
<td>HIGGS</td>
<td>20.87 ± 0.346 ± 0.00094</td>
<td>9.02 ± 0.323 ± 0.00048</td>
<td>10.84 ± 0.347 ± 0.00070</td>
<td>4.34 ± 0.323 ± 0.00057</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Dataset</th>
<th>RF(m = 2000)</th>
<th>NY(m = 2000)</th>
<th>DC-RF(m = 2000)</th>
<th>DC-NY(m = 2000)</th>
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<tr>
<td></td>
<td>time</td>
<td>error</td>
<td>time</td>
<td>error</td>
</tr>
<tr>
<td>YPMSD</td>
<td>10.68 ± 0.060 ± 0.00007</td>
<td>3.49 ± 0.058 ± 0.00016</td>
<td>5.590 ± 0.061 ± 0.00007</td>
<td>2.09 ± 0.058 ± 0.00003</td>
</tr>
<tr>
<td>covtype</td>
<td>25.79 ± 0.233 ± 0.00041</td>
<td>5.62 ± 0.185 ± 0.00067</td>
<td>12.88 ± 0.234 ± 0.00067</td>
<td>2.63 ± 0.185 ± 0.00047</td>
</tr>
<tr>
<td>SUSY</td>
<td>27.06 ± 0.220 ± 0.00035</td>
<td>12.9 ± 0.195 ± 0.00065</td>
<td>13.67 ± 0.222 ± 0.00061</td>
<td>6.40 ± 0.195 ± 0.00059</td>
</tr>
<tr>
<td>HIGGS</td>
<td>22.58 ± 0.343 ± 0.00118</td>
<td>9.14 ± 0.321 ± 0.00050</td>
<td>11.68 ± 0.345 ± 0.00108</td>
<td>5.19 ± 0.322 ± 0.00053</td>
</tr>
</tbody>
</table>

Comparison with Baselines

Figure 2 shows how the number of partition affects the error of the algorithms on test sets. The horizontal coordinate represents the number of partitions $p$, and the vertical coordinate the average test errors of different algorithms. DC-NY keeps the optimal accuracy level which is consistent with the theoretical analysis. With the increase of the number of partition $p$, the errors increase in each algorithms and our algorithm provides competitive accuracy. Taking the same $m$ and $p$, RF and DC-RF have bigger error than DC-NY, which is in line with the theoretical analysis.

Figure 3 shows the training time of the algorithms on train sets with respect to the number of partition $p$. The vertical coordinate is the training time (logarithmizing it) of different algorithms in seconds. With the increase of $p$, the training time decreases in divide-and-conquer algorithms (DC-RF and DC-NY). Our algorithm has a significant advantage over other algorithms in the training time. On covtype, the time cost of DC-RF with $p = 10$ is higher than that of DC-NY with $p = 2$, that is to say, our algorithm requires less expensive hardware devices, under the same scenario and time cost. The bigger the number of data in a subprocessor, the more obvious the time advantage of the proposed algorithm is. Apparently, combining Figure 2 with Figure 3, we get that DC-NY can use fewer hardware devices (processors) to reach a smaller error under the same time cost, which is consistent with the statistical analysis.

Table 3 shows the specific numerical information of experimental results when $m = 1500$ and $m = 2000$. Apparently, DC-NY always keeps the least time consumption than other algorithms on four datasets. Even on SUSY and HIGGS dataset of millions of points, the training time of DC-NY is always a few seconds. In test error, DC-NY keeps the optimal value or just has a little gap with the optimal, which validate the effectiveness of the proposed algorithm.

The empirical performance verifies our theoretical results that the proposed algorithm has a prominent advantage in speed while achieving satisfactory accuracy.

Conclusions

We focus on the trade-off between statistical performance and computational requirements to propose a novel approximate KRLS estimator DC-NY, which achieves the same accuracy of exact KRLS and has the minimum time complexity and space complexity, compared to the state-of-the-art approximate KRLS estimates. The empirical performance verifies the theoretical analysis. In the future, we try to explore other distributed methods to solve KRLS.

Acknowledgments

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