Applying Principal Component Analysis for Noise Reduction in Image Classification

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1 Introduction

As computer vision becomes more ubiquitous, expectations for speed and robustness continue to grow. This report focuses on the application of Principal component analysis (PCA) as a preprocessor to improve the robustness of image classifiers with respect to noisy data. Specifically, we consider convolutional neural networks (CNNs) trained on the MNIST dataset of handwritten characters. Methods for calculating principal components using singular value decomposition (SVD), eigenvector decomposition, and partial least squares minimization are introduced.

Each method is implemented in Python and compared on the basis of *Normalized Run-time*, *Loss*, *Error Rate*, and *Mean Squared Error (MSE)*. Methods for calculating principal components are also compared on the basis of asymptotic operation count with respect to data set size, image size, and number of principal components used. Finally, the advantages and disadvantages of each method applied to image noise reduction are discussed and conclusions are presented.

2 Overview of Principal Components

Principal component analysis is a mathematical transformation $X = TP^T$ that transforms a data set $X \in \mathbb{R}^{nxp}$ consisting of n samples and p variables with unknown correlation to a set of uncorrelated orthogonal loading vectors P that map X to a matrix T of principal component scores. The transformation operates such that the first vector of P corresponds to the direction of the original data set with the largest variance [?]. This property of principal components is useful for noise reduction in a data set as it allows for the correlated data to be separated from the uncorrelated data. X can be approximated as \hat{X} by using the first k principle components such that $\hat{X} = T_k P_k^T$ with $X = \hat{X} + E$ and E an error term representing the remainder of X in the directions with the lowest correlation.

In **Figure 1** on the following page, we present an example of PCA applied to the MNIST dataset. Each of the individual images is the result of projecting one noised digit onto the first n principal components, which are extracted across the entirety of the noised dataset.

Effect of Varying $n_{components}$ on PCA Output

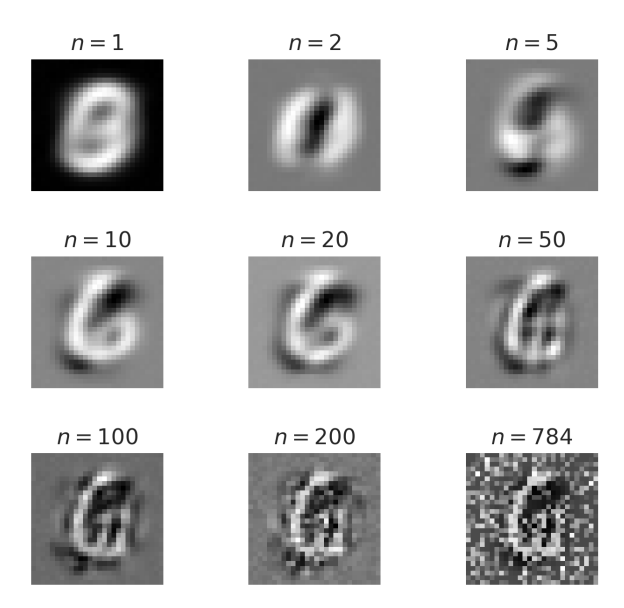


Figure 1: Result of projecting a noised image representing a 6 onto various numbers n of principal components

3 Algorithms

3.1 NIPALS

The nonlinear iterative partial least squares (NIPALS) algorithm is an iterative algorithm that has the advantage of calculating principal components consecutively and with arbitrary accuracy. These attributes are desirable when the number of required principal components is uncertain or variable and the requirements on accuracy are somewhat relaxed. As such, it is an algorithm well-suited for pre-processing noised images quickly.

The NIPALS algorithm consists of an outer loop that calculates the i^{th} next consecutive principal component. The outer loop begins with a guess of the i^{th} score vector. The initial value for t_i may be set to the the i^{th} column of the data set X_i where i is the column index of the principal component being calculated [?]. Next the inner product of t_i is taken with respect to itself as λ_i which represents the square of the magnitude of t_i . An inner loop next iteratively calculates the i^{th} loading vector p_i , score vector t_i and eigenvector λ_i . Within the inner loop p_i is calculated by projecting the transpose of the data set (X^T) onto the scores t_i . Vector p_i is then normalized to prevent overflow. The scores vector t_i is updated to be the projection of X onto p. A new λ_{new} is calculated as the inner product of t with itself. If $|\lambda - \lambda_{new}| < tol$ the iterative inner loop breaks whereas otherwise λ is set to λ_{new} . After a number of iterations t_i converges to the component score vector, p_i converges to the component loading vector, and λ_i converges to the i^{th} largest eigenvalue of X. After the inner loop converges, the outer product of t_i and p_i is subtracted from X and the outer loop is repeated. The code snippet in **Code Listing 1** below demonstrates an implementation of the NIPALS algorithm in Python with the Numba [?] package.

The computational complexity of the NIPALS algorithm is dominated by the matrix vector product operations used to project X^T onto t and X onto P with $\mathcal{O}(\mathbf{mn})$ complexity for a data set with \mathbf{m} samples and \mathbf{n} variables. Both of these operations take place within the inner loop of k iterations to convergence and an outer loop of \mathbf{c} principal components. The entire algorithm is then $\mathcal{O}(k\mathbf{c} \times \mathbf{mn})$ where \mathbf{m} is images, \mathbf{n} is pixels per image, \mathbf{c} is number of principal components and k is number of inner loop iterations necessary to reach desired accuracy.

Code Listing 1: NIPALS Algorithm

```
# Nonlinear Iterative Partial Least Squares
@jit(nopython=True) # Numba
def NIPALS(X, scores, loadings, lambdas, n_components,
            threshold=1e-6, max_iter=200):
    # Iterate over each calculated component
    for i in range(n_components):
        t = X[:, i]
        lambda_ = np.dot(t, t)
        # Iterate until convergence
        for j in range(max_iter):
            # Compute loadings
            p = np.dot(X.T, t)
            p /= lambda_
            loadings[:, i] = p
            p /= np.linalg.norm(p)
            \# Project X onto p to find score vector t
            t = np.dot(X, p)
            t /= np.dot(p, p)
            scores[:, i] = t
            lambda_n = np.dot(t, t)
            # Add score vector to matrix of score vectors
            lambdas[i] = lambda_n
            # Check for convergence
            diff = np.abs(lambda_ - lambda_n)
            if diff < threshold:</pre>
                break
            lambda_ = lambda_n
        # Update X
        X -= np.outer(t, p)
```

3.2 SVD

Singular Value Decomposition is a commonly implemented method for calculating principal components [?] [?]. The SVD transformation satisfies the equality $X = U\Sigma V^*$ for $X \in \mathbb{C}^{m,n}$, where $U \in \mathbb{C}^{m,n}$ is a unitary matrix representing the left singular vectors of X, $V \in \mathbb{C}^{n,n}$ is a unitary matrix representing the right singular vectors of X, and $\Sigma \in \mathbb{C}^{m,n}$ is a diagonal matrix of the singular values of X [?]. The principal component score vectors t_i of a mean-centered dataset X can be calculated as $U\Sigma$ and the loading vectors are VX [?].

SVD decomposition was chosen as a benchmark algorithm to highlight the computational complexity implications of calculating only a few principal components versus calculating all components at once via a method such as SVD. A common algorithm for calculating singular value decomposition of a matrix is the gebrd algorithm in LAPACK. The gebrd algorithm calculates all of the singular values and singular vectors simultaneously with $\mathcal{O}\left(\frac{4}{3}\mathbf{n}^2(3\mathbf{m}-\mathbf{n})\right)$ complexity [?]. Thus the NIPALS algorithm is computationally less complex than calculating SVD when $\frac{4}{3}\mathbf{n}(3\mathbf{m}-\mathbf{n})>(k\mathbf{c}\times\mathbf{m})$. An implementation of principal component analysis by SVD using Python and the *Numba* package [?] is given below in **Code Listing 2**.

Code Listing 2: Calculating Principal Components with SVD

```
@jit(nopython=True) # Numba
def SVD(X, scores, loadings, lambdas, n_components):

    u, s, vt = np.linalg.svd(X, full_matrices=False)
    m = X.shape[0]

    loadings[:] = (vt.T)[:, 0:n_components]
    scores[:] = np.dot(u, np.diag(s))[:,0:n_components]
    lambdas[:] = np.dot(s, s)
    lambdas /= (m - 1)
```

3.3 Eigenvector Decomposition (Simultaneous Iteration)

Eigenvector decomposition provides yet another method for calculating principal components. An iterative method for simultaneously calculating the the largest \mathbf{c} eigenvalues λ_c and eigenvectors Λ_c was selected as a benchmark algorithm to provide a comparative basis for iteratively calculating principal components sequentially as in NIPALS versus calculating them simultaneously.

Calculating principle components with eigenvector decomposition starts with calculation of the covariance $C \in \mathbf{R}^{n,n}$ of the zero-mean data set $X^T \in \mathbf{R}^{n,m}$ represented such that the rows \mathbf{n} are variables (pixels) and the columns \mathbf{m} are samples (images).

$$C = \frac{1}{n-1} X^T X$$

Calculating the covariance matrix is of $\mathcal{O}(\mathbf{mn^2})$. Note that for datasets with large images it may be unfeasible to calculate the pixel covariance and store it in memory. MNIST images with 28x28 pixels represented as int16 values on one color channel can be represented in an uncompressed form as a 9.8MB file whereas a VGA image with 640x480 pixels requires 188GB of in-memory storage.

Eigenvector decomposition is a transformation of the form $C = Q\lambda Q'$ where Q are the eigenvectors of C and λ are the eigenvalues. Simultaneous iteration solves for the k largest eigenvalue eigenvector pairs simultaneously by repeatedly multiplying the covariance matrix $C \in \mathbb{R}^{n,n}$ by a matrix $Q^{n,c}$ which is initialized with random values. Matrix Q converges to the c largest eigenvectors of the covariance matrix C [?]. In order to prevent the largest eigenvector from dominating all columns in Q, Matrix A is reorthogonalized every iteration using the modified gram-schmidt procedure adapted from [?]. Q_0 is then set to Q and the value of Q is tested for convergence. Convergence conditions are considered satisfied when the orthogonal projection of Q onto Q_0 is sufficiently small as discussed in [?].

$$||(I - Q_k Q_k^T)Q_{k-1}|| < tol$$

The principal component score matrix T and loading vector P can be constructed from the covariance C, Eigenvectors Q and Eigenvalues R as follows.

$$T = QC$$

$$P = V$$

The outer loop of the simultaneous power iteration algorithm is dominated in computational complexity by the multiplication of X with Q which is $\mathcal{O}(\mathbf{cn}^2)$ and the inner loop in which modified gram schmidt is used to orthogonalize A is of $\mathcal{O}(2\mathbf{nc}^2)$. The total complexity is then approximately $\mathcal{O}((2\mathbf{nc}^2 + \mathbf{n}^2\mathbf{c})k)$ and is quadratic with the number of pixels chosen for simultaneous computation. In the limit where \mathbf{c} approaches \mathbf{n} , we obtain the time complexity $\lim_{c\to n} \mathcal{O}((2\mathbf{nc}^2 + \mathbf{n}^2\mathbf{c})k) = 3\mathbf{n}^3k$. Hence, the algorithm becomes cubic in \mathbf{n} and as such it is clear that this method should be used only for low values of \mathbf{c} . Code Listing 3 below shows a Numba implementation of simultaneous iteration with orthogonalization at each step by MGS.

Code Listing 3: Calculating Principal Components with Eigenvalue Decomposition via Simultaneous Power Iteration

```
@jit(nopython=True) # Numba
def SI(X, Q, Q_o, R, I, threshold=1e-6, max_iter=200):
    for k in range(max_iter):
        A = operator.matmul(X, Q)
        Q_0[:] = Q
        _MGS(A, Q, R)
        delta = operator.matmul(I - np.dot(Q, Q.T), Q_o)
        if np.linalg.norm(delta) <= threshold:</pre>
            break
@jit(nopython=True) # Numba
def MGS(A, Q, R):
    (m, n) = A.shape
    for k in range(n):
        R[k, k] = np.sqrt(np.dot(A[:, k], A[:, k]))
        Q[:, k] = A[:, k] / R[k, k]
        for j in range (k + 1, n):
            R[k, j] = np.dot(Q[:, k].T, A[:, j])
            A[:, j] -= R[k, j] * Q[:, k]
```

3.4 GPU Paralellization

Consumer grade graphics processing units have been released with the capability to perform parallel computation of floating point arithmetic at high speed. In this study GPU hardware was used for the neural network classifier and so it was a natural extension to implement the image preconditioner in GPU-executable code as well. It was hypothesized that at least some reduction in computation time could be achieved by taking advantage of parallelization of matrix operations.

PyTorch was used as a high-end language to execute the NIPALS algorithm on a consumer-grade gaming GPU. Conversion from numpy to PyTorch proved to be a trivial development exercise and the resulting algorithm ran significantly faster on the GPU when compared to execution on a CPU. Figure 2 shows a comparison of the CPU and GPU implemented time to process a range of dataset sample sizes \mathbf{n} . The number of components calculated, $n_{components}$, is set to 5 and full images were used $(n_{pixels} = 784)$.

The data shows that runtime on the GPU is a factor of 4-5 times less than that of the CPU. Note that the GPU advantage is dependent on the number of images and for smaller datasets the CPU implementation is faster. This is likely due to the overhead associated with memory bandwidth and transfer between RAM on the CPU and GPU. The graph is also representative of the notion that runtime is not necessarily a good indicator of computational complexity.

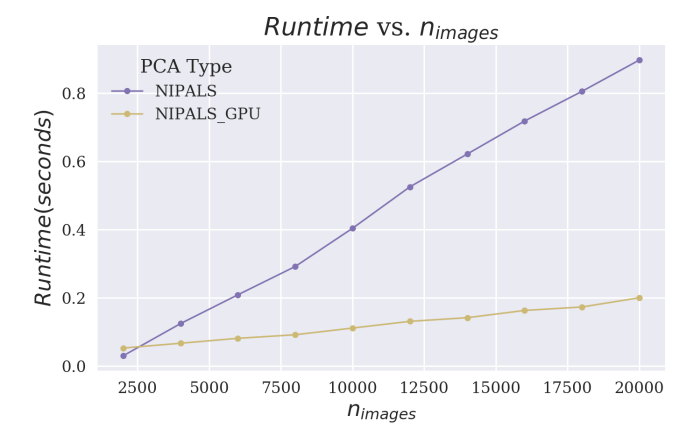


Figure 2: Runtimes of the CPU- and GPU-implemented versions of the NIPALS algorithms as observed on a GTX 1080

4 Methodology

In order to assess the relative performance of each of the algorithms, we measure four separate metrics: *Normalized Run-time*, *Loss*, *Error Rate*, and *Mean Squared Error* (*MSE*). **Figure 3** below provides an overview of how this is achieved.

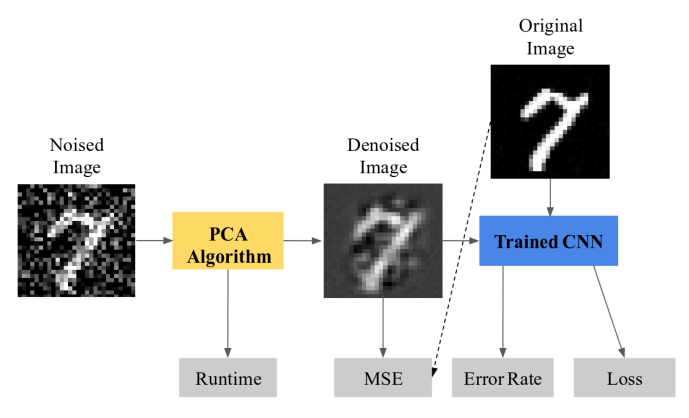


Figure 3: A flow chart of the bench-marking process

Direct comparison of runtime is not an adequate indicator of computational complexity as there are factors including hardware and code optimization that may far outweigh the computational advantages of one algorithm over another. In this study, three algorithms were implemented using Python and the built-in Python library NumPy. Because NumPy's built-in functions use a variety of different back-ends with varying efficiency, each of the original NumPy functions were wrapped inside Numba which compiles the code by LLVM and runs as machine code. In order to standardize results across hardware, only the CPU implementations were compared and all studies were performed on the same CPU. to unify the back-ends used, making it so that all three algorithms are

Comparing algorithms by counting floating point operations is unpractical and as an alternative, results in time are compared on the basis of *Normalized Run-time* as a proxy for computational cost. For each algorithm analyzed, all run-times recorded as a function of the variation of some parameter are normalized by the run-time recorded for a base-case that is a common point for all

parameter variation sets. With this methodology, the effect of non-controlled variables is significantly reduced and can compare their relative computational performance of the algorithm across a number of controlled variables. This method of evaluation was used to measure the computational complexities of the three algorithms with respect to dataset size, number of image samples, n_{images} , number of pixels per image, n_{pixels} , and number of principal components extracted, $n_{components}$. The combination of a standardized back end, standardized hardware, and run-time normalization are expected to be sufficient conditions for using time-based data as a first-order approximation for FLOP count in identifying trends in computational complexity and comparing complexity between algorithms.

Output quality is a second important factor when selecting a method for calculating principal components. In the context of the original problem, higher quality outputs should result in an improvement in the classifier's discriminatory ability. Since quality depends on $n_{components}$, each PCA algorithm is tested on a set of 20 unique $n_{components}$ ranging in size from 10 to 680. In addition, each of the 20 tests on quality versus $n_{components}$ is performed on 8 different noised datasets with mean noise mask magnitude ranging from $\mu_{noise} = 10$ to 80, resulting in a total of 160 different de-noised outputs sets for each algorithm. Each of the output data sets is processed through a set of five CNN classifiers, each trained using PyTorch on the same un-noised training data. The average Loss and average $Error\ Rate$ are observed and used to quantify quality.

The trained classifiers are unique and introduce additional variance into the results. As such, an objective evaluation of image quality is desirable in addition to performance on a neural network. An objective quality evaluation was achieved by comparing each denoised image D_i corresponding to a noised image N_i (where i is its index in the set), to the image it was derived from, O_i on the basis of MSE. These values are then averaged across the entire set to obtain the final value of MSE, as shown below:

$$MSE = \frac{1}{n_{images}} \frac{1}{n_{pixels}} \sum_{i=1}^{n_{images}} \sum_{j=1}^{n_{pixels}} (D_i[j] - O_i[j])^2$$

As with Loss and $Error\ Rate$ we calculate MSE for each of the 8 magnitude of noise and and several different values for $n_{components}$.

5 Algorithm Benchmark Results

5.1 Normalized Runtime

The asymptotic time complexity for each of the results can be used as a check on the accuracy of the experimental methods described. The reader may refer to sections **3.1**, **3.2**, and **3.3** for the code and description of each algorithm.

5.1.1 Asymptotic Complexity

Computational complexity of the three PCA algorithms is primarily a function of three known variables: $\mathbf{m} == n_{images}$, $\mathbf{n} == n_{pixels}$, and $\mathbf{c} == n_{components}$ and one unknown variable $k = n_{iterations}$. The asymptotic behavior of computational complexity as a function of \mathbf{m} , \mathbf{n} cand k is derived below for NIPALS, SVD, and Simultaneous Iteration algorithms for calculating principal components.

NIPALS: The operation count of the inner loop is dominated by the calculation of the two dot products involving the data array X. With X of size (\mathbf{m}, \mathbf{n}) , both of these function calls are $\mathcal{O}(\mathbf{mn})$. With the addition of the outer loop and inner loop, the asymptotic computational complexity of NIPALS approaches $\mathcal{O}(k \times \mathbf{c} \times \mathbf{mn})$.

SVD: The method's cost is dominated by producing the SVD of the original data, which is size (\mathbf{m}, \mathbf{n}) . Hence this method is of complexity $\mathcal{O}(\mathbf{n}^2\mathbf{m})$. Because the SVD algorithm computes all components at once, the complexity should not have a dependence on \mathbf{c} .

Simultaneous Iteration: The majority of the computation for each iteration is done to multiply the matrices X and Q and to complete a Modified Gram-Schmidt (MGS) orthogonalization. Given that the input matrix X has dimensions (\mathbf{n}, \mathbf{n}) and that Q has dimensions (\mathbf{n}, \mathbf{c}) , the time complexity of the multiplication is $\mathcal{O}(\mathbf{n}^2\mathbf{c})$. On the other hand, the complexity of each inner loop within the MGS iterations is $\mathcal{O}(\mathbf{n})$. Since both the inner and outer loops of MGS are called \mathbf{c} times, each MGS call has time complexity $\mathcal{O}(\mathbf{n}\mathbf{c}^2)$. Note that since the input to the algorithm has no relation to the number of images inputted, the runtime has no dependence on \mathbf{m} (The calculation of the covariance matrix, which does depend on \mathbf{m} is located outside of the implementation tested). Finally, as with NIPALS above, Simultaneous Iteration contains a loop with a check for convergence that repeats $k \le \max_i ter$ times until a convergence condition is satisfied. Simultaneous Iteration thus has a complexity $\mathcal{O}(k \times (\mathbf{n}\mathbf{c}^2 + \mathbf{n}^2\mathbf{c}))$. However, note that for the test cases below, the runtime

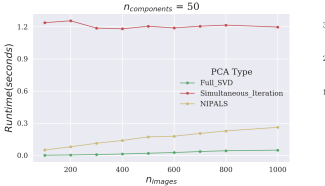
is be dominated by the former term.

The asymptotic complexity results are used as a check on the validity of the normalized time results shown in the next section. Variables **m**, **n**, and **c** are varied and measure the resulting *Normalized Runtime* of each of the algorithms as they are applied to a set of 70000 images noised with a mask of magnitude 30.

5.1.2 Runtime and $m == n_{images}$

The first variable we examine is $\mathbf{m} == n_{images}$. In this first test, full images are used ($n_{pixels} = 784$), and $n_{components}$ is set as 50. n_{images} is varied from 100 to 1000 and 15 trials are run for each of the algorithms. The median value of the resulting runtimes is then taken as the representative value. Also, the images used in each trial are randomly selected from the dataset to ensure that representative values of k are obtained for both **NIPALS** and **Simultaneous Iteration**.

The results of this test are shown in a raw and normalized form below in **Figures 4** and **5**, respectively. Note that in the normalized graph, all three methods are chosen to have runtime 1 at $n_{images} = 100$.



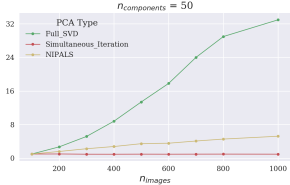


Figure 4: Unnormalized runtime behavior of PCA algorithms when n_{images} is varied.

Figure 5: Normalized runtime behavior of PCA algorithms when n_{images} is varied.

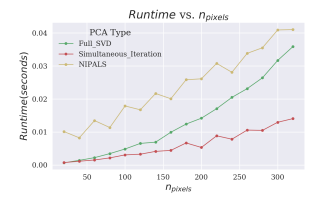
From our time complexity analysis, we expect to see a linearly increasing runtime for both **NIPALS** and **SVD**. And indeed, this trend is clearly visible on the normalized graph. Moreover, our expectation of constant runtime for the **Simultaneous Iteration** algorithm is also confirmed by the data.

These results suggest that for datasets in which the number of unique samples far exceeds the number of components calculated, the **Simultaneous Iteration** variant of PCA may be the most cost-efficient. However, we note that for the MNIST problem we explore, the number of computed components is usually quite large, and the number of images is relatively small, making **SVD** the most efficient.

5.1.3 Runtime and $n == n_{pixels}$

Next, we show how varying the value of $\mathbf{n} == n_{pixels}$ affects the time cost of the algorithms. In this test, we fix n_{images} to be 5000 and $n_{components}$ to be 5, and vary n_{pixels} from 20 to 320. We run 25 trials for each of the algorithms, selecting the median value as the representative value. As before, we randomly select the images (and also pixels) used in each individual trial to ensure that we obtain representative values of k.

The results of this test are shown in a raw and normalized form below in **Figures 6** and **7**, respectively. Note that in the normalized graph, all three methods are chosen to have runtime 1 at $n_{pixels} = 20$.



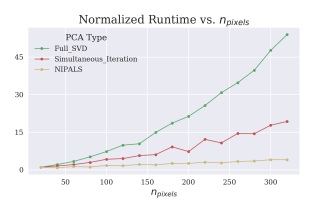


Figure 6: Unnormalized runtime behavior of PCA algorithms when n_{pixels} is varied.

Figure 7: Normalized runtime behavior of PCA algorithms when n_{pixels} is varied.

Using the complexity analysis above, we predict SVD to have quadratic complexity in n_{pixels} , which the data appears to confirm. Moreover, our expectations of linear runtimes for both **NIPALS** and **Simultaneous Iteration** are also supported by the graphs.

These data suggest that for datasets with large image sizes, unlike the MNIST dataset, SVD will no longer be the most cost-efficient algorithm. In particular, for datasets with many big images, it is likely that **Simultaneous Iteration** will be the most cost-efficient. On the other hand, if the images are sufficiently large and the number of components needed to be computed is relatively high, **NIPALS** may become the most efficient.

5.1.4 Runtime and $c == n_{components}$

Finally, we analyze $n_{components}$. Here, we fix n_{images} to be 1000 and again use full images (i.e. n_{pixels} =784). We run 15 trials, randomly selecting the images and taking the median runtime as the representative.

The results of this test are shown in a raw and normalized form below in **Figures 8** and **9**, respectively. Note that in the normalized graph, all three methods are chosen to have runtime 1 at $n_{components} = 10$.

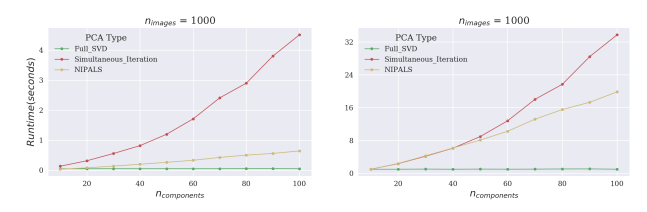


Figure 8: Unnormalized runtime behavior of PCA algorithms when n_{images} is varied.

Figure 9: Normalized runtime behavior of PCA algorithms when n_{images} is varied.

The data confirms the predicted linear relation between the cost and $n_{components}$ for **NIPALS**. The quadratic complexity of **Simultaneous Iteration** is also clearly visible in the normalized graph. Finally, the data confirms that the cost of **SVD** does not depend at all on the number components extracted.

These data reinforce the conclusions drawn in the previous two sections. In particular, it is

again clear that for problems in which large numbers of principal components must be computed, **SVD** will likely be the most efficient.

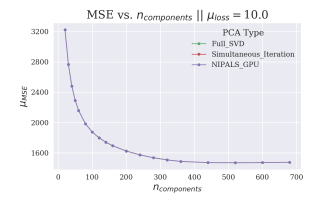
5.1.5 Discussion

In general, the above assessments confirm the derived theoretical complexities of three three PCA algorithms. The relationship between computational complexity and the size of the data sets as well as number of principal components is different for each algorithm. Each method has its advantages and disadvantages and all methods should be considered before implementing one to denoise image data.

In particular, we would expect **NIPALS** to excel in problems where $n_{components}$ is low and the total data size of the problem is sufficiently large. However, if the number of unique data samples is sufficiently large, we would expect **Simultaneous Iteration** to be the most efficient. Finally, for problems like image de-noising, where the the value of $n_{components}$ is high and the dataset size relatively small, we should expect to see that **SVD** be the most efficient. Indeed, as observed in the following sections, de-noising the MNIST dataset often required up to 100 principal components to achieve the best noise reduction, And as shown by the unnormalized graphs above, the raw runtime for **SVD** was often only a fraction of that of the other two methods.

5.2 Mean Squared Error

The obtained values of MSE for both the lowest noise ($\mu_{noise} = 10$) and highest noise ($\mu_{noise} = 80$) datasets are shown below in **Figures 10** and **11**, respectively.



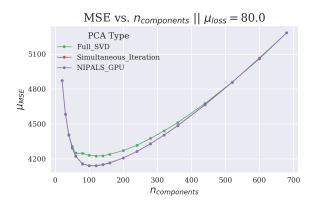


Figure 10: The relationship between MSE and $n_{components}$ for the dataset with $\mu_{noise} = 10$

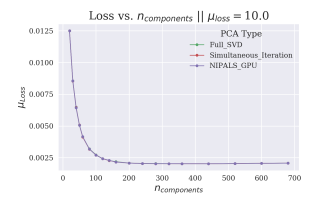
Figure 11: The relationship between MSE and $n_{components}$ for the dataset with $\mu_{noise} = 80$

The three different PCA algorithms appear to perform nearly identically on the least noisy dataset, but for the noisiest dataset, there is a non-trivial difference between the algorithms. In particular, it appears that the de-noised images from the SVD algorithm are consistently more distant from the ground truth than the de-noised images from the other two variants.

Additionally, for the least noisy dataset, as $n_{components}$ increases, the value of MSE generally decreases. However, for the dataset with $\mu_{noise} = 80$ and the other high noise datasets, there appears to be an optimal value for $n_{components}$. That is, there is a definite minimum value of $n_{components}$ that minimizes the value of MSE. Therefore, we can conclude that the choice of $n_{components}$ not only has a significant impact on the algorithms' run-times but also has a large influence on the quality of images they produce.

5.3 Loss

As in the previous section, the obtained values of Loss for both the lowest noise ($\mu_{noise} = 10$) and highest noise ($\mu_{noise} = 80$) datasets are shown below in **Figures 12** and **13**, respectively.



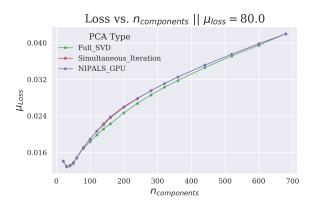


Figure 12: The relationship between Loss and $n_{components}$ for the dataset with $\mu_{noise} = 10$

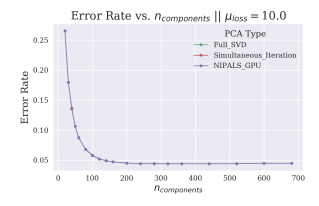
Figure 13: The relationship between Loss and $n_{components}$ for the dataset with $\mu_{noise} = 80$

Like before, we see that the different PCA variants perform nearly identically for the lowest noise dataset. However, for the highest noise dataset something is quite different. Specifically, whereas before the SVD algorithm performed worse than its peers on the MSE metric, on the Loss metric, it appears to consistently perform the best out of the three. The reason for both of these discrepancies is unclear, though it may result from the small non-zero value added to the mean of both the NIPALS and Simultaneous Iteration to prevent division by zero-norm vectors. In any case, we believe that the relative differences are not significant enough to have a noticeable impact on the algorithms' performance.

Additionally, a curious difference in the optimal value of $n_{components}$ between the Loss and MSE metrics appears. In the case of Loss, it appears that the optimal value is somewhat smaller and the local minimum somewhat steeper. This may result from the fact that classifiers are quite sensitive to the noise that a larger number of $n_{components}$ retains.

5.4 Error Rate

The data concerning the *Error Rate*, again for the least and most noisy datasets are shown below in **Figures 14** and **15**, respectively.



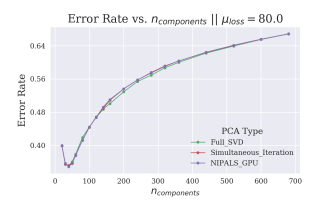


Figure 14: The relationship between Error Rate and $n_{components}$ for the dataset with $\mu_{noise} = 10$

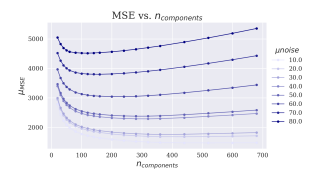
Figure 15: The relationship between Error Rate and $n_{components}$ for the dataset with $\mu_{noise} = 80$

These results roughly mirror those concerning Loss. This is hardly surprising as the two metrics are inextricably linked by the classifiers. The main point of interest here is the fact that the relative discrepancy that was seen in the Loss graphs is somewhat smaller in the Error Rate graphs. This probably results from the fact that small changes in loss do not necessarily result in the reclassification of a sample.

6 Application to Classifier

In the **analysis** of the MSE data, we noted that the ideal value of $n_{components}$ appears to vary with the magnitude of noise applied. We also noted that this value appears to change depending on the metric used.

We show this below for all 8 noise levels, using both *MSE* and *Loss*, in **Figures 16** and **17**, respectively. Note that in these figures, the metrics are averaged over both the 5 classifiers and the 3 different PCA methods.



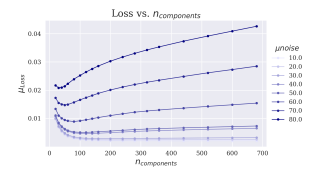


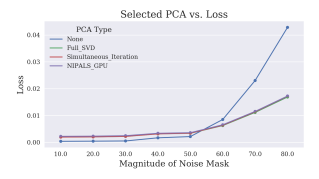
Figure 16: The relationship between MSE and $n_{components}$ for all 8 noised datasets

Figure 17: The relationship between Loss and $n_{components}$ for all 8 noised datasets

From these data, we can again determine that the optimal value of $n_{components}$ decreases as noise increases. And as before, we note that the curves for Loss (and also for $Error\ Rate$) reach their minimum earlier and are steeper than those relating to MSE.

Nevertheless, since the end goal of using these PCA algorithms is to improve the performance of the classifier, we utilize the optimal value of $n_{components}$ with respect to Loss in order to select the final PCA algorithm. In particular, for each noise value and each classifier, we select the value of $n_{components}$ that resulted in the lowest Loss value.

We give an example of how the selected PCA algorithms impact the classifiers' performances below in **Figures 18** and **19** below, which show the average *Loss* and *Error Rate* respectively of the classifiers when each of the selected PCA algorithms is applied. The graphs also show the average performance of the classifiers on the raw noised data for comparison.



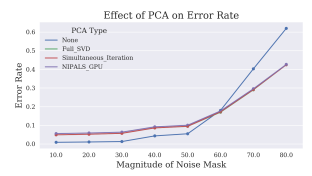


Figure 18: The average *Loss* of the classifiers when a PCA pre-processor is applied to the noised data

Figure 19: The average *Error Rate* of the classifiers when a PCA pre-processor is applied to the noised data

The use of any of the three PCA algorithms results in roughly the same result in the classifier. Specifically, for lower values of noise, it appears that the use of a PCA pre-processor does not help the classifier at all; in fact, both the average Loss and Error Rate of the classifiers increases when the noise is low. However, at sufficiently high values of noise, it appears that using a PCA pre-processor has a significant positive impact on classifier performance. Indeed for the most noised data available, the use of any of the PCA algorithms resulted in a > 20% increase in the accuracy of the classifier. And we surmise that if still more noise were to be added to the data, an even more significant difference could be observed. Hence, it appears that PCA can function as an image denoiser, particularly for data with extremely high levels of noise.

7 Conclusion

Our results demonstrate that principal component analysis (PCA) can be an effective tool at de-noising images. Indeed, our data suggests that applying a PCA pre-processor to a set of noised images can significantly improve data quality and aid in classification tasks, particularly when noise levels are significant.

Moreover, we find that while the output qualities of the three algorithms we analyzed, **NIPALS**, **SVD**, and **Simultaneous Iteration**, are quite similar, we find that their computational complexity differs greatly. For image de-noising, due to the large number of components being relatively close to the total number of pixels in an image, **SVD** appears to be by far the lowest in cost. Nevertheless, our results suggest that this will not always be the case, and in problems with different data requirements, one of the other variants may be the most efficient.

Finally, we raise several possible avenues of further research. One of the main characteristics of our approach to PCA is that it makes no assumptions about the input data. In particular, our PCA filter is not specific to a single classifier and relies only on the noised data. Tailoring PCA to the specific classification task, perhaps by including the unnoised training data may lead to larger improvements in classifier performance. Furthermore, the improvements in performance we found when converting NumPy code to Numba, and the speedups obtained by utilizing a GPU suggest that there are many options to further optimize implementations of the studied algorithms.

Determining the optimal number of principal components in real-time is another avenue for future research that would lead to further advantages as one could dynamically choose the ideal implementation method based on the data set size and the number of principal components needed.