Applications of the Compact Matrix Decomposition to the MNIST Dataset

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Introduction

The MNIST dataset is a popular dataset for research in machine learning. Datasets like MNIST are used to check the validity of techniques and models before applying them to real-world prediction situations. The dataset consists of 70,000 images, each a digitization of a handwritten digit (LeCun, Cortes, and Burges 1998). The standard for working with this data is to use 60,000 of the image for training, reserving the other 10,000 for testing and validating the predictive models. Each image in MNIST is 28 pixels by 28 pixels, yielding 784 distinct values. These values range from 0 to 255 in base 10, each representing the amount of black in that pixel. Since the images are grayscale, we can represent them with only one matrix rather than more complicated color images, which can either be represented with three distinct matrices (one for each of the color channels) or by a single matrix with all values in base 16. This reduces the amount of computation needed to work with the dataset, with the discarded color information being classified as out of scope for the project. Predictive results on the data set by various groups across the world can be found on the internet, sorted by type of classification algorithm used to make predictions. Simple models such as linear classifiers have error rates as high as 12.0%, while the most complicated models listed are able to drive the error rate as low as .23%. Some of the classification models use pre-processing to make the data more suitible for use with the chosen method. Especially popular are normalization techniques such as deskewing, Since the dataset has 784 predictors, feature selection methods such as least absolute shrinkage and selection operators (LASSO) are frequently used to eliminate variables that do not contribute to the final sublution. Examples of such variables in the case of the MNIST data would be the value in the very first pixel - it is totally white the vast majority of the time. Such a value does not improve our ability to predict what class our data belongs to, so we can safely ignore it. While some models are adept at handling large sparse data like MNIST, decompositions rising from linear algebra can perform similar functions while taking much less time to compute, a major victory that allows for computation on larger datasets. One such method, first published by

Compact Matrix Decomposition

In an effort to improve computation speed, specifically around the training of new models, optimization techniques are applied. These techniques range in effectiveness and complexity, and are a very active area of mathematic research. One recently developed technique is the Compact Matrix Decomposition (CMD). The CMD was developed at Carnegie Mellon University, and first published in the paper "Less is More: Compact Matrix Decomposition for Large Sparse Graphs". This paper has been cited 94 times since its publication in 2007 according to Google Scholar, increasing interest in the topic. The paper claims that the CMD requires less than 1/10 of the space of the more traditional Singular Value Decomposition (SVD) and is 10x more computationally efficient, while maintaining the same level of reconstruction accuracy. Here I seek to apply the CMD to the MNIST dataset. This is a prudent operation since the matrices of images, while not as large as the data used in the original paper (Sun et al. 2007)(200+ GB of network traffic data), are large enough to benefit from this precomputation.

Motivation

Initial computation on the MNIST data backs up the use of CMD. Attempting to execute ridge regression in R on the dataset without any form of precomputation does not complete, the code simply crashes after an hour or so of executing on my computer. Using the singular value decomposition in this situation is not effective, since the data is too sparse.

Derivation

The derivation of CMD stems from a basic central question: how can one approximate the matrix $A \in \mathbb{R}^{m \times n}$ as the product of smaller matrices $C \in \mathbb{R}^{m \times c}$, $U \in \mathbb{R}^{c \times r}$ and $R \in \mathbb{R}^{r \times n}$ such that we minimize $|A - CUR|^2$. This computation effectively seeks to reduce the rank of A while maintaining enough similarity between the result and the original. Symbolically we represent the approximation of A as \tilde{A} , we then claim that $\tilde{A} = CUR$, with dimensions defined above. To begin computation, we construct a subspace of the input matrix A, then reduce the rank of that subspace. Since the subspace is spanned by the columns of the input matrix, we choose to sample from the input matrix. We choose to use a biased sampling method, favoring columns that have a higher squared Frobenius norm (Drineas, Kannan, and Mahoney 2006), which is defined by $|A| = \sum_{i,j} A(i,j)^2$. This selection method favors columns that have higher entries, and by allowing repeted selection of the same column we arrive at a set of column vectors whose distribution is modeled by the norms of the columns in A. We'll call this matrix C_d . We know and accept that the sampling method will result in repeated columns, which we remove without reducing accuracy. We choose to remove the duplicates so as to reduce storage space required to store the data and thus the computation time on the data.

We know that the columns that are duplicated contain more information than the non-repeated columns, as they were selected at a higher rate. This, of course, rests on the assumption that the higher values (when normed) are more important, which is appropriate since the goal is to apply this method to sparse matrices, which we expect to have many values at or near 0. To address the issues created if we simply remove the duplicate columns, we find the unique columns, and scale them based on the number of times they occur in the sampled set. For a scaling factor, we use the square root of the number of occurances. This process results in a much narrower matrix, which is preferable for computation. We call this matrix C_s . Before further computation, we want to ensure that we have not lost information in our process thus far. As we intend on preforming a singular value decomposition on C_s , we want to ensure that the singular values of C_s and C_d are the same. We first define another matrix $D = [e_1, \ldots, e_1, \ldots, e_{c'}, \ldots, e_{c'}]$. Each e_i is a column vector that is all 0s, except for the *i*-th element, which is 1. In our defintion of D, we also note that there is one e_i for each C_i in C_d , ensuring that C_d and D have the same number of columns.

From the above definitions, we know that $C_d = CD^T$ where C is the matrix of unique columns of C_d . Then we have

$$C_d C_s^T = C D^T (C D^T)^T = C D^T D C^T = C \Lambda C^T = C \Lambda^{\frac{1}{2}} \Lambda^{\frac{1}{2}} C^T = C \Lambda^{\frac{1}{2}} (C \Lambda^{\frac{1}{2}})^T = C_s C_s^T C_s^T$$

where Λ is a $c' \times c'$ matrix that has the number of occurances of each column $e_i \in D$ on the diagonal and 0s elsewhere. From here, we can diagonalize either $C_d C_d^T$ or $C_s C_s^T$ to find the singular values of C_d and C_s .

The next steps in the process involve projecting the original matrix X onto the space spanned by C_s . We then will remove duplicate rows much like we did earlier with duplicate columns, reducing the computational cost. We begin by constructing an orthonormal basis for C using the SVD, so $C = U_C \Sigma_C V_C^T$. From there we could project the original matrix into the large and dense orthonormal basis U_C , but we choose to instead compute a low-rank approximation for A, stemming from the observation that $U_C = CV_C \Sigma_c^{-1}$ where $C \in \mathbb{R}^{m \times c}$ is large but sparse, $V_C \in \mathbb{R}^{c \times k}$ is dense but small, and $\Sigma \in \mathbb{R}^{k \times k}$ is small and diagonal. This approximation \tilde{A} is then defined as

$$\tilde{A} = U_c U_c^T A = C V_C \Sigma_C^{-1} (C V_C \Sigma_C^{-1})^T A = C (V_C \Sigma_C^{-2} V_C^T C^T) A = C T A$$

where $T = (V_C \Sigma_C^{-2} V_C^T C^T) \in \mathbb{R}^{c \times m}$ we know that C is sparse. and T is dense and large. We can continue optimiszing the calculation by reducing the overhead between T and A.

The process of reducing this overhead is defined in the ApprMultiplication process, which takes two matrices A and B, assuming that AB is defined, and then sampling the columns of A and the rows of B. We scale the rows and columns for multiplication, but still have the issue of duplicates. The CMD process removes duplicate rows by sampling and scaling rows from A and extracting the corresponding columns from C^T .

After removing the duplicate rows, we need to verify that our matrix multiplication is still correct. To do this we let I and J be the set of sampled rows, I with duplicates and J without duplicates. We define $J = [1, \ldots, 1, \ldots, r', \ldots, r']$ where there are d'_i instances of each element up to $d'_{r'}$. Additionally, we define $I = [1, \ldots, r']$. Letting A and B be matrices with dimensions $m_a \times n_a$ and $m_b \times n_b$ respectively. Allowing $i \leq \min(n_a, m_b)$ we have

$$A(:, J)B(J, :) = A(:, I)\Lambda'B(I, :)$$

where Λ' is a diagonal matrix with $1, \ldots, d_r'$. We can prove that the matrix multiplication is correct by stating that

$$A(:,J)B(J,:) = \sum_{k \in J} A(i,k)B(k,j) = \sum_{k \in J} d_{i_k} A(i,k)B(k,j) = A(:,I)\Lambda'B(I,:).$$

This verifies the multiplication.

Having proved that the multiplication is correct, we can now simply calculate the final portion of the decomposition, U, to complete the process. In this case

$$U = V_C \Sigma_C^{-2} V_C^T C_s.$$

Programming

To apply this to the MNIST dataset, we program a CMD implementation in R.

```
library(dplyr)
```

First we import libraries.

```
initial_subspace_construction <- function(mat, c) {</pre>
    columnDistribution <- vector(length = ncol(mat))</pre>
    matrixSum <- 0
    # sum the entrire matrix, we only do this once for efficiency
    for (i in 1:ncol(mat)) {
        columnSum <- 0
        for (j in 1:nrow(mat)) {
             element <- mat[j, i] ^ 2</pre>
             columnSum <- columnSum + element</pre>
        }
        matrixSum <- matrixSum + columnSum
    }
    # calculate the distribution for each column
    for (i in 1:ncol(mat)) {
        columnSum <- 0
        for (j in 1:nrow(mat)) {
             element <- mat[j,i] ^ 2</pre>
             columnSum <- columnSum + element</pre>
        }
        columnDistribution[i] <- columnSum / matrixSum</pre>
    }
    # the matrix that we'll return, of dimension m x c
    returner <- matrix(nrow = nrow(mat), ncol = c)
```

```
# perform biased sampling
for (i in 1:c) {
    j <- sample(1:ncol(mat), size = 1, replace = TRUE, prob = columnDistribution)
    returner[,i] <- mat[,j] / sqrt(c * columnDistribution[j])
}
return(returner)
}</pre>
```

This function performs the initial subspace construction provided the initial matrix mat and a sample size c.

```
cmd_subspace_construction <- function(mat, c) {</pre>
    initSubspace <- initial_subspace_construction(mat, c)</pre>
    # round the initSubspace so that unique() will work
    initSubspace <- apply(initSubspace, c(1,2), round, digits = 5)</pre>
    uniqueCols <- t(unique(t(initSubspace)))</pre>
    numUniqueCols <- ncol(uniqueCols)</pre>
    returner <- matrix(nrow = nrow(initSubspace), ncol = numUniqueCols)</pre>
    for (i in 1:numUniqueCols) {
        numInstancesInInitSubspace <- 0</pre>
        for (j in 1:ncol(initSubspace)) {
             # count the instances of this column in the initial subspace
             if (identical(initSubspace[,j], uniqueCols[,i])) {
               numInstancesInInitSubspace <- numInstancesInInitSubspace + 1</pre>
             }
        returner[,i] <- sqrt(numInstancesInInitSubspace) * t(uniqueCols[,i])</pre>
    return(returner)
```

This function uses initial_subspace_construction() to construct the scaled version that we use, with the unique columns.

```
appr_multiplication <- function(matA, matB, sampleSize) {</pre>
    q <- vector(length = nrow(matB)) # the row distribution</pre>
    # sum the entrire matB matrix, we only do this once for efficiency
    matrixSum <- 0
    for (i in 1:ncol(matB)) {
        columnSum <- 0
        for (j in 1:nrow(matB)) {
             element <- matB[j, i] ^ 2</pre>
             columnSum <- columnSum + element</pre>
        }
        matrixSum <- matrixSum + columnSum</pre>
    # row distribution of matB
    for (x in 1:nrow(matB)) {
        rowSum <- 0
        for (i in 1:ncol(matB)) {
            rowSum <- rowSum + matB[x,i] ^ 2</pre>
        q[x] <- rowSum / matrixSum
```

```
R_d <- matrix(nrow = sampleSize, ncol = ncol(matB))</pre>
C_d <- matrix(nrow = nrow(matA), ncol = sampleSize)</pre>
for (i in 1:sampleSize) {
    j <- sample(1:nrow(matB), size = 1, replace = TRUE, prob = q)</pre>
    R_d[i,] <- matB[j,] / sqrt(sampleSize * q[j])</pre>
    C_d[,i] <- matA[,j] / sqrt(sampleSize * q[j])</pre>
uniqueColsC <- t(unique(t(C d)))</pre>
uniqueRowsR <- unique(R d)
R_s <- matrix(nrow = sampleSize, ncol = ncol(matB))</pre>
C_s <- matrix(nrow = nrow(matA), ncol = sampleSize)</pre>
for (i in 1:sampleSize) {
    u <- 0 # number of instances of this row in the full set of Rs
    for (j in 1:nrow(R_d)) {
        # count the instances of this column in the initial subspace
        if (identical(R_d[j,], uniqueRowsR[i,])) {
             u <- u + 1
    }
    R_s[i,] <- u * uniqueRowsR[i,]</pre>
    C_s[,i] <- uniqueColsC[,i]</pre>
}
return(list("C" = C_s, "R" = R_s))
```

This rather long function performs the ApprMultiplication process that we described above, creating two of the components that we need to use the decomposition in practice.

```
cmd_decomposition <- function(matA, c, r) {
    print("Constructing Subspace")
    cSubspace <- cmd_subspace_construction(matA, c)
    print("Calculating SVD")
    cSubspace.svd <- svd(cSubspace)
    cTranspose <- t(cSubspace)
    print("ApprMult")
    apprMult <- appr_multiplication(cTranspose, matA, r)
    c_s <- apprMult$C
    r_s <- apprMult$R
    print("Calculating U")
    u <- cSubspace.svd$v %*% qr.solve(diag(cSubspace.svd$d)) %*%
        qr.solve(diag(cSubspace.svd$d)) %*% t(cSubspace.svd$v) %*% c_s
    return(list("C" = cSubspace, "U" = u, "R" = apprMult$R))
}</pre>
```

This summary function makes calls to the above functions, and performs the calculation of U to complete the decomposition.

Application

Having derived and programmed the CMD algorithm, we move to applying it to the MNIST dataset. We begin by manipulating the data into a structure that makes sense in this situation:

References

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