

Improved Neighborhood-Based Algorithms for Large-Scale Recommender Systems

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ABSTRACT

Neighborhood-based algorithms are frequently used modules of recommender systems. Usually, the choice of the similarity measure used for evaluation of neighborhood relationships is crucial for the success of such approaches. In this article we propose a way to calculate similarities by formulating a regression problem which enables us to extract the similarities from the data in a problem-specific way. Another popular approach for recommender systems is regularized matrix factorization (RMF). We present an algorithm – neighborhood-aware matrix factorization – which efficiently includes neighborhood information in a RMF model. This leads to increased prediction accuracy. The proposed methods are tested on the Netflix dataset.

Categories and Subject Descriptors

H.2.8 [Database Applications]: [Data mining, Recommender Systems, Collaborative Filtering, Netflix Competition]

General Terms

Latent factor model, Similarity matrix, Ensemble performance

Keywords

recommender systems, matrix factorization, KNN, Netflix, collaborative filtering

1. INTRODUCTION

Due to the increasing popularity of e-commerce, there is growing demand of algorithms that predict the interest of customers (called *users* in the following) in some product (called *item* in the following). Such interest is commonly

quantified by a non negative number r which we call a *rating* in this article. Algorithms which predict a rating for each user-item pair are called recommender systems [1]. The predictions of recommender systems are in general based on a database which contains information about users and items. The *Collaborative Filtering* (CF) approach to recommender systems relies only on information about the behavior of users in the past. The pure CF approach is appealing because past user behavior can easily be recorded in web-based commercial applications and no additional information about items or users has to be gathered. CF algorithms for recommender systems are therefore easily portable.

More abstractly, the goal of CF is missing value estimation. Consider a system consisting of m users and n items. We define the set of users as the set of integers $\{1, \dots, m\}$ and the set of items as the set of integers $\{1, \dots, n\}$. The $m \times n$ rating matrix $\mathbf{R} = [r_{ui}]_{1 \leq u \leq m; 1 \leq i \leq n}$ stores the ratings of users for items where r_{ui} is the rating of user u for item i . The input to a CF algorithm is a set $\mathcal{L} = \{(u_1, i_1), \dots, (u_L, i_L)\}$ of L user-item tuples referred to as *votes* and the corresponding ratings in the rating matrix. We assume that ratings in the rating matrix are non-zero if they are in the training set and zero if they are not in the training set, i.e., we assume $r_{ui} \neq 0$ if $(u, i) \in \mathcal{L}$ and $r_{ui} = 0$ otherwise. Such ratings not in the training set are called missing values. The goal of the system is to predict the missing values of R .

In fall 2006, the movie rental company Netflix started a competition, *the Netflix Prize*. The goal of the competition is to design a recommender system which improves on the Netflix recommender system *Cinematch* by 10% with regard to the root mean squared error (RMSE) on a published database. This database contains training data in the form of about 100 million ratings from about 480,000 users on 17,770 movies. Each rating in this database is an integer between 1 and 5. A probe set is provided which can be used to test algorithms. Furthermore, Netflix published a qualifying set which consists of user-item pairs but no ratings (the items correspond to movies in this database). The ranking of a submitted solution is based on this data set. The Netflix dataset captures the difficulties of large recommender systems. First, the dataset is huge and therefore the runtime and memory usage of potential algorithms become important factors. Second, the ranking matrix is very sparse with about 99 percent of its entries being missing such that

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many users have voted for just a few movies. The algorithms presented in this article were tested on the Netflix dataset. However, their design is not specific to this dataset, thus the algorithms can be applied to other CF problems as well.

An obvious way to generate predictions of ratings is to calculate similarities between users and deduce a rating of some user u for an item i from ratings of that item by users with high similarity to user u . Similarly, a rating for some item i by user u can be predicted from ratings by that user for similar items. Such approaches have successfully been applied to the Netflix dataset. We deal with pure neighborhood-based approaches in Section 2. It turned out that they benefit from simple preprocessing where the ratings are first cleaned from so called *global effects* [2]. Global effects are linear relationships between the ratings and some simple variables like the time of the rating (which is provided in the Netflix dataset). We introduce in Section 2.1 four global effects which have not been considered before.

One problem of neighborhood-based approaches is the choice of the metric, or in other words, the measure of similarity between users or items.¹ A common way to measure the similarity between two users is the Pearson correlation between ratings of items which both users voted for. However, if two users have just a few common ratings (which is often the case in the Netflix dataset), the Pearson correlation is a bad estimate for their similarity. In Section 2.2 we propose a method where the similarities between items themselves are learned by gradient descent. Thus, the choice of a similarity measure is passed from the designer to the algorithm. One drawback of this method is the huge number of parameters which are fitted (the whole $n \times n$ matrix of item similarities has to be estimated) and thus its tendency to overfit the training data. This problem is tackled by a factorized version described in Section 2.3. This algorithm does not compute the whole similarity matrix but a low rank approximation in a linear latent factor model. This reduces the number of parameters significantly. A further advantage of factorized similarities is its memory efficiency. While the whole similarity matrix between users cannot be precomputed because of memory restrictions of computers to date, the online computation of correlations can be very time demanding. This makes naïve neighborhood-based approaches infeasible for large sets of elements like the set of users in the Netflix database. The factorized similarity model overcomes this problem. First, for a reasonable number of factors per user, the factor matrix can easily be held in memory, and second, for any two users the similarity is computed by just the inner product of two vectors. This model can also easily be extended to include information about unknown ratings (i.e., user-item pairs for which one knows that this user rated that item but the actual rating is unknown). The inclusion of unknown ratings in the prediction was first discussed in [7].

A regularized matrix factorization (RMF) produces a rank K approximation of the $m \times n$ rating matrix by factorizing it into a $m \times K$ matrix of user features and a $K \times n$ matrix of item features. RMF models are easy to implement and achieve good performance. Consequently, they are of

¹Obviously, one can also define a similarity measure for user-item pairs, an approach which is not discussed in this article.

Nb.	Side	Effect	RMSE
10	both	Previous effects	0.9659
11	item	average movie year	0.9635
12	user	movie production year	0.9623
13	user	STD of movie ratings	0.9611
14	item	STD of user ratings	0.9604

Table 1: Preprocessing for neighborhood models. The table shows the RMSE on the probe set of the Netflix dataset when accounting for 10 to 14 global effects (i.e., the row with Nb. i shows the error when one accounts for global effects 1 to i). The second column (“Side”) specifies whether the effect defined in the third column (“Effect”) needs the estimation of parameters for the users or for the items.

ten used in CF algorithms. Overspecialization on the data points can be avoided by careful use of regularization techniques. We show in Section 3 that a hybrid approach which is partly neighborhood-based and RMF-based is a very effective CF method. This method performs slightly better than well-tuned RMF methods or restricted Boltzmann machines (RBMs). Additionally, the model can be trained very quickly.

2. NEIGHBORHOOD-BASED MODELS

Neighborhood-based models for recommender systems commonly compute the similarity between users or items and use these similarities to predict unknown ratings. It is difficult to pre-calculate correlations between users for the Netflix database because the user correlation matrix can not be held in main memory of current computers. This makes the evaluation of naïve user-based neighborhood approaches very slow and therefore unpractical. We will discuss an efficient algorithm which is based on user similarities in Section 2.4. Before that, we discuss our algorithms for the case of item-based similarities and note that the principles apply to user-based similarities in the same way. The similarity c_{ij} between two items i and j is often estimated by the Pearson correlation between common ratings of that items, i.e., the correlation between the list of ratings of users $U(i, j) = \{u | (u, i) \in \mathcal{L} \text{ and } (u, j) \in \mathcal{L}\}$ which voted for both items i and j . The full neighborhood information is stored in the symmetric similarity matrix $\mathbf{C} = [c_{ij}]_{1 \leq i, j \leq n}$. Other similarity measures like the mean squared error (MSE) or a distance derived from the movie titles can also be used.

Algorithms in the flavor of the k -nearest neighbor (kNN) algorithm produce ratings based on the ratings of the k most similar items, i.e., the predicted rating \hat{r}_{ui} of an user u for item i is computed as

$$\hat{r}_{ui} = \frac{\sum_{j \in N_k(u, i)} c_{ij} r_{uj}}{\sum_{j \in N_k(u, i)} c_{ij}}, \quad (1)$$

where $N_k(u, i)$ denotes the set of the k items most similar to item i that were rated by user u .

2.1 Preprocessing

In [2], so called “global effects” of the data were discussed and the removal of such effects from the data was proposed as an effective preprocessing step for the Netflix dataset. Such

simple preprocessing turns out very useful if applied prior to kNN methods. As defined by [2], a global effect is a linear relationship between the ratings and some simple property of the votes. For each of the effects, the goal is to estimate one parameter per user or per item (see [2] for details). We found four effects not described before which lower the RMSE on the probe set to 0.9604, see Table 1. The effects considered are the effect of the average production year of the movies the given user voted for ("average movie year"), the production year of the given movie ("movie production year"), the standard deviation of the ratings for the given movie ("STD of movie ratings") and the standard deviation of the ratings of the given user ("STD of user ratings").

The algorithms described below are tested for different pre-processings in order to facilitate comparison with other algorithms.

2.2 Regression on similarity

One problem of approaches based on the Pearson correlation between item ratings is that for many item pairs, there are only a few users which rated both items. For any two items i, j , we define the *support* s_{ij} for these items as the number of users which rated both items. The reliability of the estimated correlation grows with increasing support s_{ij} . For the Netflix dataset most correlations between movies are around 0. In order to decrease the influence of estimated correlations with low support on the prediction, we found it useful to weight correlations according to their support such that the similarity c_{ij} between item i and j is given by $c_{ij} = \tilde{c}_{ij} \frac{s_{ij}}{s_{ij} + \alpha}$ where \tilde{c}_{ij} is the Pearson correlation between common ratings of the two items and α is a constant in the range of 100 to 1000 (this procedure was introduced in [3]).

In any case, the choice of the similarity measure is critical for the success of neighborhood-based algorithms. In this section we describe an alternative approach where the matrix of similarities \mathbf{C} between items is learned by the algorithm itself. The matrix can be initialized with small random values around 0 or with Pearson correlations². A prediction of the rating of user u for item i is calculated similar to equ. (1), but over the set $N(u, i) = \{j \neq i | (u, j) \in \mathcal{L}\}$ of all items different from i which user u voted for in the training set

$$\hat{r}_{ui} = \frac{\sum_{j \in N(u, i)} c_{ij} r_{uj}}{\sum_{j \in N(u, i)} |c_{ij}|}. \quad (2)$$

Since similarities can become negative, we normalize by the sum of absolute similarities.

The objective function to minimize is given by the MSE with an additional regularization term

$$E(\mathbf{C}, \mathcal{L}) = \frac{1}{2} \sum_{(u, i) \in \mathcal{L}} (\hat{r}_{ui} - r_{ui})^2 + \gamma \sum_{j < k} c_{jk}^2,$$

where γ is a regularization constant. The model is trained by stochastic gradient descent on the objective function. For each training example we update only those similarities relevant for the example, i.e., for example (u, i) we update c_{ij}

²We used a uniform distribution in $[-0.1, 0.1]$ or Pearson correlations, with similar results.

Preprocessing	probe RMSE	qual. RMSE
Raw data	0.9574	0.9487
1GE	0.9458	0.9384
2GE	0.9459	0.9384
6GE	0.9372	0.9281
10GE	0.9349	0.9256
14GE	0.9331	0.9239

Table 2: The RMSE of the similarity regression model on the probe set (middle column) and the qualifying set (right column) of the Netflix dataset for pre-processings that accounted for 0 to 14 global effects (GE).

if $j \in N(u, i)$. The update of similarity c_{ij} is then given by

$$c_{ij}^{new} = c_{ij}^{old} - \eta \cdot \text{sign} \left((\hat{r}_{ui} - r_{ui}) \frac{\partial \hat{r}_{ui}}{\partial c_{ij}} \right) - \eta \gamma c_{ij}^{old}. \quad (3)$$

We opted to use the sign of the error gradient in (3) because this update turned out to be much more stable than the gradient itself. This choice was inspired by the sign-sign LMS rule (see, e.g., [5]). The number of trainable parameters in this model for the Netflix dataset is around 157 million which is very large. Hence training of the model is prone to early overfitting. Typical values of γ and η are 0.01. Results on the Netflix data are shown in Table 2. At a single epoch, approximately $L \cdot s_M$ similarities are updated, where L is the size of the training set \mathcal{L} and s_M is the average number of votes per user (s_M is around 200 for the Netflix dataset). The training time for one epoch is in the range of one hour on a standard PC and usually a single epoch suffices.

2.3 Regression on factorized similarity

Gradient descent on the elements of the symmetric item similarity matrix \mathbf{C} leads to early overfitting because of the huge number of trained parameters. In this section, we show that one can overcome this problem by learning a factorized version of \mathbf{C} . In other words, the algorithm learns two $K \times n$ matrices \mathbf{P} and \mathbf{Q} with $K \ll n$ and \mathbf{C} is computed as

$$\mathbf{C} = \mathbf{P}^T \mathbf{Q}. \quad (4)$$

Hence, we learn a rank- k approximation of \mathbf{C} which drastically reduces the number of parameters. Only the upper triangle of $\mathbf{P}^T \mathbf{Q}$ is used to calculate similarities, since similarities are assumed to be symmetric, i.e., $c_{ij} = c_{ji}$. The similarity c_{ij} between items i and j is then given by

$$c_{ij} = \begin{cases} \mathbf{p}_i^T \mathbf{q}_j, & \text{if } i < j \\ \mathbf{p}_j^T \mathbf{q}_i, & \text{if } i > j, \end{cases} \quad (5)$$

where \mathbf{p}_i and \mathbf{q}_i denote the i th column of \mathbf{P} and \mathbf{Q} respectively. Ratings are predicted as in the previous section by equ. (2).

The training schedule is similar to the direct similarity regression model with the difference that for every similarity c_{ij} (with $i < j$) we have to update the $2K$ parameters p_{1i}, \dots, p_{Ki} and q_{1j}, \dots, q_{Kj} . Hence the training is slowed down by a factor of K as compared to direct similarity regression. Results on the Netflix data are shown in Table 3. The results are marginally better compared to the non-factorized version.

Preprocessing	K	RMSE
10GE	80	0.9324
14GE	100	0.9313

Table 3: The RMSE of the factorized item similarity regression model on the probe set of the Netflix dataset for various preprocessings.

Preprocessing	K	RMSE
Raw	10	0.9951
2GE	10	0.9539
6GE	10	0.9469
14GE	20	0.9371

Table 4: The RMSE of the factorized user similarity regression model on the probe set of the Netflix dataset for various preprocessings.

2.4 Factorized user similarity matrix

An advantage of the factorized similarity model is that similarities between large sets of elements can be stored in memory. This enables us to store similarities between users in the factor matrices. In order to learn user similarities we perform gradient descent on the factorized user similarity matrix. All calculations and update rules are mirrored versions of those discussed above for the item similarity matrix. For the Netflix dataset the training time increases by a factor of 30 compared to the training time of the factorized item similarity model since there are approximately 30 times more users than items in the database.

The results of the model for a few different preprocessings of the data are shown in Table 4.³ Although the results are similar to those of the factorized item similarity model, the algorithm is still useful since the information extracted from user similarities is different from that when item similarities are used. This contributes to the performance if the models are finally combined for a single prediction.

2.5 Incorporating unknown ratings

The model can be extended to include unknown ratings. This helps in general on users with few ratings in the training set. Let \mathcal{L}' denote the set of votes for which the rating is unknown (for the Netflix dataset, our set \mathcal{L}' consisted of votes in the probe set as well as those in the qualifying set). Let $N'(u, i) = \{j \neq i | (u, j) \in \mathcal{L}'\}$ denote the set of items different from i user u has voted for with unknown rating. Then, the prediction \hat{r}_{ui} of a rating for user u on item i is given by

$$\hat{r}_{ui} = \frac{\sum_{j \in N(u, i)} c_{ij} r_{uj} + \sum_{j \in N'(u, i)} c_{ij} \tilde{r}_{uj}}{\sum_{j \in N(u, i)} |c_{ij}| + \sum_{j \in N'(u, i)} |c_{ij}|}, \quad (6)$$

where \tilde{r}_{uj} are estimates of the unknown ratings (they are parameters of the model which are trained, see below). Train-

³Because of the time demands of this algorithm, training was stopped after the presentation of only 30% of the training set.

ing of the similarities is done as in the basic model (see equ. (3)) with the difference that for training example (u, i) we update all c_{ij} for $j \in N(u, i) \cup N'(u, i)$. The unknown ratings \tilde{r}_{uj} are trained simultaneously with gradient descent. One can initialize each unknown rating with the mean rating of the corresponding item. Slightly better performance can be obtained if one initializes the unknown ratings with predictions of a neighborhood-based approach, see equ. (1) (the reported results were obtained in this manner). On the Netflix dataset, this model achieved a RMSE of 0.9278 on the probe set with a preprocessing that accounted for 14 global effects. This is an improvement of 0.053 over the model without unknown ratings (see Table 2).

3. NEIGHBORHOOD-AWARE MATRIX FACTORIZATION

In this section we present an algorithm – neighborhood-aware matrix factorization (NAMF) – which efficiently incorporates a linear regularized matrix factorization (RMF) in a neighborhood-based model. More specifically, for a given vote (u, i) , the algorithm computes three predictions: a prediction \hat{r}_{ui}^{MF} which is based on a RMF, a prediction \hat{r}_{ui}^{user} is based on a user-neighborhood model, and a prediction \hat{r}_{ui}^{item} which is based on an item-neighborhood model. Both neighborhood-based models utilize predictions from the RMF model if needed. The final prediction of the algorithm is a combination of the three predictions.

3.1 Regularized matrix factorization model

A RMF computes a rank K approximation $\mathbf{R}' = \mathbf{A}\mathbf{B}^T$ of the rating matrix \mathbf{R} , where $\mathbf{A} \in \mathbb{R}^{m \times K}$ is the user factor matrix and $\mathbf{B} \in \mathbb{R}^{n \times K}$ is the item factor matrix. The entries of these matrices are determined such that $r_{ui} \approx r'_{ui}$ for all votes $(u, i) \in \mathcal{L}$. After the factor matrices \mathbf{A} and \mathbf{B} have been determined by the training algorithm, the prediction \hat{r}_{ui}^{MF} for a vote (u, i) is given by $\hat{r}_{ui}^{MF} = r'_{ui} = \sum_{k=1}^K a_{uk} b_{ik}$. Because the rating matrix is usually sparse, additional regularization is needed. Using a regularization as proposed in [8], [6] leads to the error function

$$E(\mathbf{A}, \mathbf{B}, \mathcal{L}) = \sum_{(u, i) \in \mathcal{L}} (r_{ui} - \hat{r}_{ui}^{MF})^2 + \frac{\lambda}{2} (\|\mathbf{A}\|_F^2 + \|\mathbf{B}\|_F^2), \quad (7)$$

where $\|\cdot\|_F$ denotes the Frobenius norm and λ is the regularization parameter. We use stochastic gradient descent to minimize this error function. The update equations for a training example (u, i) are therefore

$$a_{uk}^{new} = a_{uk}^{old} + \eta \cdot (e_{ui} b_{ik}^{old} - \lambda a_{uk}^{old}) \quad (8)$$

$$b_{ik}^{new} = b_{ik}^{old} + \eta \cdot (e_{ui} a_{uk}^{old} - \lambda b_{ik}^{old}), \quad (9)$$

for $k = 1, \dots, K$ and

$$e_{ui} = r_{ui} - \sum_{k=1}^K a_{uk}^{old} b_{ik}^{old}. \quad (10)$$

3.2 User-neighborhood model

The similarity of two users can be measured by the Pearson correlation $\hat{\rho}_{uv}^{user}$ between the list of ratings for items which were rated by both users. In order to decrease the influence of correlations with low support we shrink each correlation

according to their support s_{uv} [3]:

$$\rho_{uv}^{user} = \frac{s_{uv} \tilde{\rho}_{uv}^{user}}{s_{uv} + \alpha^{user}}, \quad (11)$$

where the parameter α^{user} is determined as discussed below (in order to facilitate readability we denote all variables and parameters of the user-neighborhood model by a ‘‘user’’ superscript and those of the item-neighborhood model by a ‘‘item’’ superscript). However, the use of the Pearson correlation may be problematic in some cases. The most severe problem occurs if the number of common ratings is small for most user pairs (i.e., the support for most user pairs is low). In this case, the Pearson correlation between these ratings is a very unreliable measure of similarity. For many datasets however there exist for most users other users such that the number of common rated items is large, and reliable correlations can be calculated for these pairs. We will make use of this observation below. Another problem of employing correlations between users is the size of the correlation matrix. For the Netflix dataset where the number of users is around 480,000, the whole matrix needs about one TByte of memory. We overcome this problem by storing for each user u only the correlations with the J users with highest correlation to u . A rating prediction \hat{r}_{ui}^{user} is then computed as the weighted sum over the ratings of these best correlating users where the rating r_{vi} is given by the predicted rating of the RMF model or a rating from a training example if it exists

$$\hat{r}_{ui}^{user} = \frac{\sum_{v \in U_J(u)} c_{uv}^{user} r_{vi}}{\sum_{v \in U_J(u)} c_{uv}^{user}}, \quad (12)$$

where $U_J(u)$ denotes the set of J users with highest correlation to u . Each weighting coefficient c_{uv}^{user} is computed from the Pearson correlation ρ_{uv}^{user} by applying a squashing function

$$c_{uv}^{user} = (\sigma(s^{user} \rho_{uv}^{user} - b^{user}))^{\gamma^{user}}, \quad (13)$$

where the scaling factor s^{user} , the bias b^{user} , and the exponent γ^{user} are global parameters which were determined as described below. The sigmoidal squashing function $\sigma(\cdot)$ is given by

$$\sigma(x) = \frac{1}{(1 + \exp(-x))}. \quad (14)$$

3.3 Item-neighborhood model

For the item side the same principle can be applied. Correlations $\tilde{\rho}_{ij}^{item}$ between common rated items are shrunk

$$\rho_{ij}^{item} = \frac{s_{ij} \tilde{\rho}_{ij}^{item}}{s_{ij} + \alpha^{item}}. \quad (15)$$

For each item i only the correlations with the J items with highest correlation to i are stored. A rating prediction is then computed as the weighted sum over the ratings of these best correlating items $I_J(i)$. The weighting coefficients are given by

$$c_{ij}^{item} = \left(\sigma \left(s^{item} \rho_{ij}^{item} - b^{item} \right) \right)^{\gamma^{item}}, \quad (16)$$

where ρ_{ij}^{item} denotes the Pearson correlation between the ratings of users that rated both items i and j , and α^{item} , s^{item} , b^{item} , and γ^{item} are constants. The rating prediction

\hat{r}_{ui}^{item} for a vote (u, i) is given by

$$\hat{r}_{ui}^{item} = \frac{\sum_{j \in I_J(i)} c_{ij}^{item} r_{uj}}{\sum_{j \in I_J(i)} c_{ij}^{item}}. \quad (17)$$

3.4 Combining the information

The predictions from the RMF model, the user neighborhood model and the item neighborhood model are combined in a single rating. The obvious way to archive this is an optimal linear combination of the three predictions. Experiments have shown that the predictive accuracy of the models strongly depends on the support and the number of ratings from the training data (as opposed to those from the RMF model) used in the neighborhood models. So we use a weighted sum, based on this information to combine the predictions:

$$\hat{r}_{ui} = \frac{\tilde{S}(u, i)^\delta \cdot \hat{r}_{ui}^{RMF} + \hat{\beta} \hat{S}(u, i)^\delta \cdot \hat{r}_{ui}^{user} + \bar{\beta} \bar{S}(u, i)^\delta \cdot \hat{r}_{ui}^{item}}{\tilde{S}(u, i)^\delta + \hat{\beta} \hat{S}(u, i)^\delta + \bar{\beta} \bar{S}(u, i)^\delta} \quad (18)$$

$$\tilde{S}(u, i) = \min\{N_u, N_i\}. \quad (19)$$

In the equation above, $N_u = |\{i | (u, i) \in \mathcal{L}\}|$ denotes the number of votes of user u , and $N_i = |\{u | (u, i) \in \mathcal{L}\}|$ denotes the number of votings for item i . $\tilde{S}(u, i) = |\{v \in U_J(u) | (v, i) \in \mathcal{L}\}|$ and $\hat{S}(u, i) = |\{j \in I_J(i) | (u, j) \in \mathcal{L}\}|$ denote the number of votes from the training set used to calculate the corresponding ratings.

The training schedule can be summarized as follows. First, correlations $\tilde{\rho}_{uv}^{user}$ between users and correlations $\tilde{\rho}_{ij}^{item}$ between items are computed. The best correlating users/items are computed according to the shrunken correlations (we used $\alpha^{user} = 10$ for user correlations and for item correlations $\alpha^{item} = 30$) and the corresponding correlations (not shrunken) are stored. This step is the computationally most demanding one. Then the RMF is computed. Once this is done, the predictions of the neighborhood models can be computed very efficiently. Then, good values for the 13 constants α^{user} , α^{item} , $\tilde{\beta}$, $\hat{\beta}$, s^{user} , s^{item} , b^{user} , b^{item} , γ^{user} , γ^{item} , δ , $\bar{\delta}$, and $\hat{\delta}$ in the model are determined with a genetic algorithm. Because the evaluation of individuals is very fast, this optimization step can be done quite efficiently (on a standard PC this step needed 1-2 hours). Once the model is trained, predictions can be generated very quickly.

3.5 Experimental results

The RMF model was trained on the residuals of the first global effect (movie effect) described in [2]. The use of this effect slightly improves RMF performance whereas the use of all global effects decreases the performance of the RMF model. All RMF models were trained with stochastic gradient descent using $\eta = 0.002$ and $\lambda = 0.02$. The weights were initialized to small values sampled from a normal distribution with zero mean and standard deviation 0.001. The neighborhood models were trained on preprocessed data that incorporated 10 global effects.

In comparison, a restricted Boltzmann machine on the Netflix probe data achieved a RMSE of 0.907 (see Fig. 4 in [7]). Another approach which combines a neighborhood model

Features K of the RMF	RMSE
10	0.9175
50	0.9069
100	0.9056
300	0.9046
600	0.9042

Table 5: RMSE of different neighborhood-aware matrix factorizations on the Netflix probe data. Pre-processing for the neighborhood model was done on 10 global effects, the neighborhood size was $J = 50$.

with a RMF was described in [2]. This algorithm obtained a RMSE of 0.9071 on the Netflix probe set.

4. ENSEMBLE PERFORMANCE

The final goal of each team that participates in the Netflix contest is the prediction of unknown ratings with optimal accuracy. In order to achieve maximal prediction accuracy, it is a common strategy to combine predictions of different algorithms into a final one. We did linear blending on the probe set, which was not used for training, similar to [4]. Whether an algorithm is particularly powerful on a given data point (u, i) depends strongly on the support of the vote, i.e., the number of votes of user u and the number of votes for item i . Consequently, a linear combination of predictions for data points with low support will be quite different from a linear combination for data points with high support. We therefore divided the probe set into *slots* based on the support of the data points. To obtain a single value from the user support and the item support we combined them by taking the minimum of both. This procedure is called “slot blending” [4]. The slot boundaries were chosen such that the number of ratings in the slots was approximately uniform.

For each slot, the final prediction is then computed as a linear combination of the predictions of the individual algorithms. Suppose one wants to combine the predictions of N algorithms. These predictions are first stored in a predictor matrix $\mathbf{P} \in \mathbb{R}^{l \times N}$ where l is the number of votes in the slot and p_{ij} is prediction of algorithm j for the i -th vote in the slot. The interpolation weights \mathbf{w} are computed with the pseudo-inverse of P as $\mathbf{w} = (\mathbf{P}^T \mathbf{P})^{-1} \mathbf{P}^T \mathbf{q}$ where \mathbf{q} is the column vector of probe ratings of the slot. The final prediction for a vote from the qualifying set which falls – according to its support – into this slot is then given by the linear combination of the individual predictions with the interpolation weights \mathbf{w} .

Using this method, we calculated the ensemble performance of the algorithms proposed in this article. The RMSE of the ensemble on the probe set was 0.8981 which results in a RMSE of 0.8919 on the qualifying set (for predictors which were re-trained after blending with the probe ratings included). This is an improvement of 6.25% over the Cinematch system. The proposed methods can well be combined with other powerful algorithms like different kinds of matrix factorizations and restricted Boltzmann machines to further improve prediction accuracy.

5. CONCLUSIONS

In this article, we proposed several neighborhood-based algorithms for large-scale recommender systems. An important property of these algorithms is that their memory usage scales linearly with the number of users or items as compared to a quadratic scaling of most other neighborhood-based approaches. This makes the algorithms scalable to large-scale problems. To date it seems that powerful solutions for collaborative filtering problems need to combine the predictions of a diverse set of single algorithms. This procedure is able to combine the specific advantages of single algorithms. The standard approach is linear blending, where the predictions are simply combined in a linear way after training. The neighborhood-aware matrix factorization algorithm tries to combine the advantages of two powerful methods – a RMF approach and neighborhood-based approach – in a more direct way: The predictions of one algorithm are used to estimate unknown variables in the other ones. One can therefore hope that the combination of them is more than the (weighted) sum of its parts. Such hybrid models are promising candidates for future research.

6. ACKNOWLEDGMENTS

We thank Netflix for providing this nice dataset. We also thank the participants of the previous KDD workshop in 2007 for providing their inspiring ideas in the field of recommender algorithms. Also thanks to the active Netflix community for discussing all kind of things regarding various implementation details on proposed algorithms.

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