GHRS: Graph-based Hybrid Recommendation System with Application to Movie Recommendation

Zahra Zamanzadeh Darban\textsuperscript{a,}\textsuperscript{*}, Mohammad Hadi Valipour\textsuperscript{b}

\textsuperscript{a}Faculty of Information Technology, Monash University, Melbourne, Australia
\textsuperscript{b}Department of Engineering and Product, Ostadkar Company, Tehran, Iran

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ABSTRACT

Research about recommender systems emerges over the last decade and comprises valuable services to increase different companies' revenue. While most existing recommender systems rely either on a content-based approach or a collaborative approach, there are hybrid approaches that can improve recommendation accuracy using a combination of both approaches. Even though many algorithms are proposed using such methods, it is still necessary for further improvement. This paper proposes a recommender system method using a graph-based model associated with the similarity of users' ratings in combination with users' demographic and location information. By utilizing the advantages of Autoencoder feature extraction, we extract new features based on all combined attributes. Using the new set of features for clustering users, our proposed approach (GHRS) outperformed many existing recommendation algorithms on recommendation accuracy. Also, the method achieved significant results in the cold-start problem. All experiments have been performed on the MovieLens dataset due to the existence of users' side information.

1. Introduction

Recommendation Systems (RS) are a type of choice advisor to overcome the explosive growth of information on the web. These systems facilitate users with personalized items (products or services), which are more likely to be interested in. RS have been employed to a wide variety of fields: movies (Wei et al., 2016; Moreno et al., 2016), music (Mao et al., 2016; Horsburgh et al., 2015), news (Shi et al., 2016; Wang and Shang, 2015), books, e-commerce, tourism, etc. An efficient RS may dramatically increase the number of sales of customers to boost business (Jannach et al., 2010; Ricci et al., 2015). In common, recommendations are generated based on user preferences, item features, user-item interactions, and some other information such as temporal and spatial data.

RS methods are mainly categorized into Collaborative Filtering (CF), Content-Based Filtering (CBF), and hybrid recommender system based on the input data (Adomavicius and Tuzhilin, 2005). CF models (Salah et al., 2016; Polatidis and Georgiadis, 2016; Koren and Bell, 2015) aim to exploit information about the rating history of users for items to provide a personalized recommendation. In this case, if someone rated a few items, CF relies on estimating the ratings he would have given to thousands of other items by using all the other users' ratings. On the other side, CBF uses the user-item side information to estimate a new rating. For instance, user information can be age, gender, or occupation. Item information can be the movie genre(s), director(s), or the tags. CF is more applied than CBF because it only aims at the users' ratings, while CBF requires advanced processing on items to perform well (Lops et al., 2011).

Although the CF model is preferred, it has some limitations. One of CF’s limitations is known as the cold-start problem: how to recommend an item when any rating does not exist for either the user or the item? One idea to overcome this issue is to build a hybrid model by combining CF and CBF, where side information can be utilized in the training process to compensate the lack of ratings through it. Some successful approaches extend the Probabilistic Matrix Factorization (Adams and Murray, 2010; Salakhutdinov and Mnih, 2008) to integrate side information. However, some algorithms outperform them in the general case.

There are tremendous achievements of deep learning (DL) in many applied domains in the past few decades, such as computer vision (Ding and Tao, 2015; Tian et al., 2016; Byeon et al., 2016; Huang and Sun, 2016) and speech tasks (Graves et al., 2013; Xue et al., 2016). Deep learning models have already been studied in a wider range of applications due to its capability in solving many complex tasks. Recently, DL has been inspiring the recommendation frameworks and brought us many performance improvements to the recommender. Deep learning can capture the non-linear user-item relationships and catches the complicated relationships within the data itself from different data sources such as visual, textual, and contextual.

In recent years, the DL-based recommendation models achieve state-of-the-art recommendation tasks, and many companies apply deep learning for enhanced quality of their recommendation (Covington et al., 2016; Okura et al., 2017). For example, Salakhutdinov tackled the Netflix challenge using Restricted Boltzmann Machines (RBM-CF) (Salakhutdinov et al., 2007; Georgiev and Nakov, 2013). AutoRec is an Autoencoder for collaborative filtering (Sedhain et al., 2015), which uses Autoencoder to predict missing ratings. Autoencoders are stacked denoising Autoencoders with sparse...
inputs for collaborative filtering (Strub and Mary, 2015). Covington et al. (2016) proposed a DNN-based recommendation algorithm for video recommendation on YouTube, Cheng et al. (2016) presented an application recommender system for Google Play, and Okura et al. (2017) presented an RNN-based recommender system for Yahoo News. All of these models have shown significant improvement over traditional models. However, the existing deep learning models have not regarded the side information about the users or items, which is highly correlated to the users’ rating. Indeed, combining deep learning and side information may help us to discover a surpass solution for the considered challenges.

In this paper, we introduce a hybrid approach using Autoencoder, which tackles both challenges: learning a non-linear representation of users-items and dominating the cold start problem by integrating side information. Compared to previous models in that direction (Sedhain et al., 2015; Strub and Mary, 2015; Wu et al., 2016), our framework integrates the users’ preferences, similarities, and side information in a unique matrix. This conjunction leads to improved results in CF.

The outline of the paper is organized as follows. First, Section 2 discusses related works in both Autoencoder-based and hybrid recommendation models. Then, our proposed model is described in Section 3. Finally, experimental results are given and discussed in Section 4 and followed by a conclusion section.

2. Related Works

This section introduces the categories of DL-based recommendation models and then focuses on advanced research to identify the most outstanding and promising progress in recent years.

2.1. Deep Learning based Recommendation Models

Deep learning is a research field of machine learning. It learns multiple levels of representations and abstractions from data and it can solve both supervised and unsupervised learning tasks. We can categorize the existing recommendation models based on the types of employed deep learning approaches into the following two classes (Zhang et al., 2019)

- The recommendation with Neural Building Blocks; In this category, the deep learning technique determines the recommendation model’s applicability. For example, MLP can simply model the non-linear interactions between users and items; CNNs can extract local and global representations from heterogeneous data sources like text and image; recommender system can model the temporal dynamics and sequential evolution of content information using RNNs.
- The recommendation with Deep Hybrid Models; Some DL-based recommendation models utilize more than one deep learning technique. Deep neural networks’ flexibility makes it possible to combine several neural building blocks to complement one another and form a more powerful hybrid model. There are many possible combinations of these deep learning techniques, but not all have been exploited.

Additionally, we review and summarize some publications which utilize Autoencoder, and they will be discussed in the following sub-sections.

2.2. Autoencoder based Recommendation Models

Autoencoder is an unsupervised model attempting to reconstruct its input data in the output layer. In general, the bottleneck layer (the middle-most layer) is used as a salient feature representation of the input data (Zhang et al., 2019). The schematic of basic Autoencoder is illustrated in Figure 1, which output $X'$ should become closer to the input $X$ and the bottleneck layer is shown by $z$. The main variants of Autoencoders can be considered as denoising Autoencoder, marginalized denoising Autoencoder, sparse Autoencoder, contractive Autoencoder and variational Autoencoder (Goodfellow et al., 2016).

There are two general ways to apply Autoencoder to a recommender system (Zhang et al., 2019):

1. Using Autoencoder to learn lower-dimensional feature representations at the bottleneck layer; or
2. Filling the blanks of the interaction matrix directly in the reconstruction layer.

Almost all the Autoencoder variants such as denoising Autoencoder, variational Autoencoder, contractive Autoencoder, and marginalized Autoencoder can be applied to the recommendation task. In this paper, we employed the first technique to extract new low-dimension features. Figure 1 illustrates the structure of different recommendation models based on Autoencoder (Zhang et al., 2019).
Figure 2: Illustration of: (a) Item based AutoRec; (b) Collaborative denoising Autoencoder; (c) Deep collaborative filtering framework (Zhang et al., 2019)

2.2.1. Autoencoder based Collaborative Filtering Models

One of the successful applications is to consider collaborative filtering from the Autoencoder perspective. AutoRec (Sedhain et al., 2015) took user partial vectors $\rho(u)$ or item partial vectors $\rho(i)$ as input and attempted to reconstruct them in the output layer. Indeed, it has two variants: item-based AutoRec (I-AutoRec) and user-based AutoRec (U-AutoRec), corresponding to the two types of inputs.

There are essential points about AutoRec that worth noticing (Zhang et al., 2019). First, I-AutoRec performs better than U-AutoRec, which may be due to the higher variance of user partially observed vectors. Second, a different combination of activation functions will influence the performance significantly. Third, moderately increasing the hidden unit size will improve the result as expanding the hidden layer dimensionality gives AutoRec more capacity to model the input characteristics. Furthermore, adding more layers to formulate a deep network can lead to slight improvement.

CFN (Strub et al., 2016; Strub and Mary, 2015) is a continuation of AutoRec, and posses the following two improvements:

1. Deploying the denoising techniques makes CFN more robust.
2. Incorporating the side information such as user profiles and item descriptions mitigates the sparsity and cold start influence.

The CFN input is also partially observed vectors, so it also has two variants: I-CFN and U-CFN, taking $\rho(i)$ and $\rho(u)$ as input, respectively. Masking noise is imposed as a great regularizer to better deal with missing elements (with zero value). Further extension of CFN also incorporates side information. However, instead of just integrating side information in the first layer, CFN injects side information in every layer (Zhang et al., 2019).

Collaborative Denoising Autoencoder (CDAE). The three models reviewed earlier are mainly designed for rating prediction, while CDAE (Wu et al., 2016) is principally used for ranking prediction. The input of CDAE is user partially observed implicit feedbacks. If the user likes the movie, the entry value is one, otherwise zero. It can also be considered as a preference vector that reflects the user’s interest in items (Zhang et al., 2019). Figure 1b illustrates the structure of CDAE.

This model uses a unique weight matrix for each user and has a notable impact on model performance. Parameters of CDAE are also learned by minimizing the reconstruction error. CDAE initially updates its parameters using SGD over all feedbacks. However, it is impractical to consider all ratings in real-world applications. A negative sampling technique has been proposed to sample a small subset from the negative set (items with which the user has not interacted), which reduces the time complexity substantially without degrading the ranking quality (Zhang et al., 2019).

Multi-VAE and Multi-DAE (Liang et al., 2018) proposed a variant of variational Autoencoder for recommendation with implicit data, showing better performance than CDAE. These methods introduced a principled Bayesian inference approach for parameter estimation and showed agreeable results than generally used likelihood functions.

Based on a survey by Zhang et al. (2019), Autoencoder-based Collaborative Filtering (ACF) (Ouyang et al., 2014) is the first Autoencoder based collaborative recommendation model. Instead of using the original partial observed vectors, it decomposes them by integer ratings. Like AutoRec and CFN, ACF aims at reducing the mean squared error as the cost function. But, there are two demerits of ACF; it loses to deal with non-integer ratings, and the decomposition of partially observed vectors increases the sparseness of input data and drives to worse prediction accuracy.

2.2.2. Feature Representation Learning with Autoencoder

Autoencoder is a dominant feature representation learning approach, and it can be used in recommender systems to learn feature representations from users-items content features. In the following, we will summarize some of the related methods.
Collaborative Deep Learning (CDL) (Wang et al., 2015) is a hierarchical Bayesian model that integrates stacked denoising Autoencoder (SDAE) into probabilistic matrix factorization. The method proposed a general Bayesian deep learning framework (Wang and Yeung, 2016) to combine the deep learning and recommendation model. The framework consists of two tightly coupled parts: the perception component (deep neural network) and task-specific component. Mainly, CDL’s perception component is a probabilistic representation of ordinal SDAE, and PMF (Probability Mass Function) works as the task-specific component. This tight combination enables CDL to balance the impacts of side information and interaction records.

Collaborative Deep Ranking (CDR). CDR (Ying et al., 2016) is devised specifically in a pairwise framework for top-n recommendation. Some studies have demonstrated that the pairwise model is more suitable for ranking lists generation. Experimental results also show that CDR outperforms CDL in terms of ranking prediction (Zhang et al., 2019).

Deep Collaborative Filtering Framework is a general framework for unifying deep learning approaches with a collaborative filtering model (Li et al., 2015). This framework makes it easy to utilize deep feature learning techniques to build hybrid collaborative models (Zhang et al., 2019). There is a marginalized denoising Autoencoder-based collaborative filtering model (mDA-CF) on top of this framework. In comparison to CDL, mDA-CF explores more computationally efficient variants of the Autoencoder. The method saves the computational costs of searching sufficient corrupted input by marginalizing the corrupted input, and it makes the mDA-CF more scalable than CDL. Plus, mDA-CF embeds content information of items and users, while CDL only regards item features’ effects.

AutoSVD++ (Zhang et al., 2017) uses contractive Autoencoder (Rifai et al., 2011) to learn item feature representations, then integrates them into the classic recommendation model, SVD++. The model posses the following advantages (Zhang et al., 2019)

1. Compared to other Autoencoder variants, contractive Autoencoder captures the infinitesimal input variations.
2. It models the implicit feedback to enhance the accuracy further.
3. An efficient training algorithm is designed to reduce training time.

HRCD (Wei et al., 2017) is a hybrid collaborative model based on Autoencoder and timeSVD++. It is a time-aware model that uses SDAE to learn item representations from raw features and solve the cold item problem (Zhang et al., 2019).

3. Graph-based Hybrid Recommendation System

In this section, we focus on our proposed method which can be categorized as a hybrid recommendation system. First we define the basic notations used throughout the paper. Next, we describe the proposed model in an architectural view and algorithmic steps. Then, graph-based features will be declared separately. Finally, we will explain about the clustering method and how we find the optimum number of clusters.

We first define the basic notations used throughout this paper. Given the set of n users, $U = \{u_1, \ldots, u_n\}$, and the set of m item, $I = \{i_1, \ldots, i_m\}$, all user-item pairs can be denoted by an n-by-m matrix $R = U \times I$, where the entry $r_{ui}$ indicates the assigned value of implicit feedback of user $u$ to item $i$. If $r_{ui}$ has been observed (or known), it is represented by a specified rating associated in a specific range and interval; otherwise, a global default rating is zero. We used this matrix to find similarity between users’ preferences. After generating the similarity graph which represents users as nodes and the relations as edges, we extract the features from this graph, $F_g = \{f_1, \ldots, f_g\}$, and preserve them in the n-by-g matrix. We collect some users’ features from the dataset, which are called side information, $F_u = \{f_1, \ldots, f_g\}$, some items’ side information $F_I = \{f_1, \ldots, f_I\}$ and obtain the combined feature matrix which is n-by-g+r+s. Without loss of generality, we categorized all the features (both graph features and side information) as binary which enlarged final feature vector for each user.

3.1. Architecture

The overall structure of our aggregated recommender system (GHRS) is presented in Figure 3. The Graph-based Hybrid Recommender System comprises the following seven steps:

1. In the first step, we build a graph with the number of users’ as nodes. Two users will be connected based on their similarities. The edge connects a pair of users who have more than a percent of items with similar ratings.
2. In the second step, a set of information will be extracted from the similarity graph for each user. For instance, we compute PageRank of the nodes, degree centrality, closeness centrality, the shortest-path betweenness centrality, load centrality, and the average degree of each node’s neighborhood in the graph. As a result, this matrix relies on the different data processing magnitude using a preference-based collaborative approach.
3. In the third step, we combine side information such as gender and age with graph-based features to retrieve the most relevant movies for users. Therefore, we have one combined matrix from different types of features, which is then used as the Autoencoder stage input.
4. In the fourth step, we apply the Autoencoder to extract new features and reduce the dimension. It includes selecting a proper optimizer, using a proper loss function and neural network architecture, and preventing the overfitting issue.
5. In the fifth step, we utilize the new features encoded by Autoencoder for user clustering, using the K-means algorithm to create a small number of peer groups. It
includes finding an appropriate number of clusters for each dataset.

6. In the sixth step, we assign new users to clusters based on encoded features and compute the new item rating based on similarity with other items.

7. In the seventh step, we compute the estimated rates of all items for each user according to its cluster’s average rating.

Algorithm 1 declared the total workflow in details.

3.2. Graph-based Features

This section reviews the intuition of some graph features that represent similarities and their general computational process.

- **Page Rank**: Page Rank is an algorithm that measures the transitive influence or connectivity of nodes. It was initially designed as an algorithm to rank web pages (Xing and Ghorbani, 2004). We can compute the Page Rank by either iteratively distributing one node’s rank (based on the degree) over the neighbors or randomly traversing the graph and counting the frequency of hitting each node during these paths. In this paper, we used the first method.

- **Degree Centrality**: Degree centrality measures the number of incoming and outgoing relationships from a node. The Degree Centrality algorithm can be used to find the popularity of individual nodes (Freeman, 1979). The degree centrality values are normalized by dividing by the maximum possible degree in a simple graph n-1, where n is the number of nodes.

- **Closeness Centrality**: Closeness centrality is a way
Algorithm 1 Proposed method detailed workflow

Input: $U, I, R, F_u, F_i$

Output: Estimated rates for user-item

1: Set alpha = percentage of items with similar ratings between two users
2: Compute the aggregated similarity between users based on $\alpha$ (the percentage of items which two users rated them similarly)
3: Construct the Similarity Graph and consider the users as nodes
4: $F_u$: Extracted graph-based features for users (nodes)
5: $F_i$: Preprocessed and categorized users’ side information (demographic informations)
6: Combine $F_u$ and $F_i$ in a single feature vector $F_U$. Apply the Autoencoder on $F_U$ and train the model with the best settings
7: Encode the $F_i$ using the Autoencoder an extract the low dimensional feature vector $F_e$
8: Find the optimum clusters for clustering users with $F_e$
9: Perform user clustering using extracted features vector $F_e$ and find clusters $C$
10: Generate the user-cluster matrix $UC$
11: Estimate clusters’ ratings for items matrix $CI$
12: if there are users rated the item $i$ before in the cluster $c$ then
13: $CI_{ci} = \text{average (users’ rates of the item } i \text{ in the cluster } c)$
14: else if there are similar items to the item $i$, rated by users in the cluster $c$ then
15: $CI_{ci} = \text{average (users’ rates of similar items in the cluster } c)$
16: else
17: $CI_{ci} = \text{average (all users’ rates in the cluster } c)$
18: end if
19: Estimate users’ ratings’ matrix $R’ = UC \times CI$
20: Compute the recommendation list for target user $u$

3.3. User Clustering

As we mentioned before in section 3.1, each user belongs to a specific cluster and the cluster rate for an item will be considered as the estimated rating for the user-item pair. In the proposed method we use K-Mean algorithm to cluster the users based on extracted features by Autoencoder. One important issue in using such algorithms is to find the proper number of clusters regarding performance factors. We use two methods to choose the number of clusters; Elbow method and Average Silhouette algorithm.

In this section we will explain the summary of K-Mean algorithms and how we tackle and solve the number of clusters issue with both mentioned methods.
3.3.1. K-Means Algorithm
The K-means algorithm is a simple iterative clustering algorithm. Using the distance as the metric and given the K classes in the data set, calculate the distance mean, giving the initial centroid, with each class described by the centroid (Yuan and Yang, 2019; Awad and Khanna, 2015). For a given data set X with n data samples and the number of category K, the Euclidean distance is the measure of the similarity index, and the clustering method aims to minimize the sum of the squares of the various types. It means that it minimizes (Wang et al., 2012)

\[ d = \sum_{k=1}^{K} \sum_{i=1}^{n} \| x_i - u_k \|^2 \]  

(5)

where \( k \) represents \( K \) cluster centers, \( u_k \) represents the \( k^{th} \) center, and \( x_i \) represents the \( i^{th} \) point in the data set.

3.3.2. Elbow Method
The basic idea behind cluster partitioning methods, such as k-means clustering, is to define clusters such that the total intra-cluster variation (known as a total within-cluster variation or total within-cluster sum of squares) is minimized. It measures the compactness of the clusters, and it should be as small as possible (Kaufman and Rousseeuw, 2009). The elbow method is based on plotting the explained variation as a function of the number of clusters, and picking the elbow of the output curve as the proper number of clusters. Adding another cluster after the elbow point doesn’t give much better modeling of the data and may causes over-fitting.

3.3.3. Average Silhouette
Briefly, the average silhouette approach measures the quality of a clustering. It means that it determines how well each object occupies within its cluster. A high average silhouette width intimates a valuable clustering.

The average silhouette method computes the average silhouette of observations for different values of \( k \). The optimal number of clusters \( k \) is the one that maximizes the average silhouette over a range of possible values for \( k \) (Kaufman and Rousseeuw, 2009).

4. Empirical Experiments and Performance Evaluation
In this section, the performance of the proposed model is evaluated, analyzed, and enumerated in separate parts. The dataset is processed and described in detail, followed by the requisite experimental setup. Due to the variation of steps and processes in the proposed method, we elaborate on the practical results in-depth. Finally, we compared the method with basics and modern methods, which we discussed most of them in related works.

4.1. Dataset
We have utilized two benchmark datasets (MovieLens 100K and MovieLens 1M) of the real-world in recommender systems to implement the model practically (Harper and Konstan, 2015). MovieLens 100K contains 100,000 ratings \( R \in \{1, 2, 3, 4, 5\} \), 1,682 movies (items) rated by 943 users. MovieLens 1M comprises of 1,000,209 ratings \( R \in \{1, 2, 3, 4, 5\} \) of approximately 3,900 movies made by 6,040 users. As discussed in Section 3, the proposed method uses users’ demographic data to solve the new users’ cold-start issue. Hence, due to the lack of users’ demographic data in larger datasets like MovieLens 10M, it would not be possible to evaluate the model more on larger datasets. We used the MovieLens 100K dataset for analyzing the proposed method’s steps. The final evaluations and comparisons have been done on the MovieLens 1M dataset. Table 1 shows the details of the mentioned datasets.

4.2. Features Statistics
As declared in Section 3, we use two types of features in the proposed method: side information (users’ demographic data) and features extracted from the similarity graph between users. We transformed the demographic data into a categorical format, concatenated both types of features, and made the raw feature set before dimension reduction with an Autoencoder. In this section, we discuss a little about the statistics of the raw features. We have declared before the only parameter we have used for generating the graph is \( \alpha \), the value of a threshold for connecting two users having at least several same movies in their ratings. This threshold is represented as a percentage of total movies in the dataset. Hence, we have an exploration of a very sparse graph to near a full-mesh graph. Figure 6 illustrates the similarity graph visualization for \( \alpha \in \{0.005, 0.01, 0.02, 0.03\} \) for 943 users in MovieLens 100K.

Figure 5 shows the normalized graph-based features’ distributions against each other for MovieLens 100K and MovieLens 1M with \( \alpha = 0.015 \). We can see correlations between these types of features in some cases.

As all the demographic features are transformed into a categorical format, the demographic features vector is one-hot encoded and has a specific sparsity level for each dataset. On the other hand, we declared that the graph-based features’ value is related to the similarity graph size, and the graph size is directly related to the factor \( \alpha \). In Figure 6, we can see that the feature set’s sparsity rises when the value of the \( \alpha \) increases.

4.3. Performance Metrics
We use 10-fold cross-validation on MovieLens 1M dataset and 5-fold cross-validation on MovieLens 100K dataset to partition the datasets into training and testing to measure the performance of the GHRS. The final prediction metrics are the average of the iterations of training and testing base on the number of folds in each dataset. The training set comprises the User-Item list with given ratings, user’s demographic information, and item’s side information. We consider the Root Mean Squared Error (RMSE) as the metric for evaluation. RMSE (Equation 6) is generally related to
Table 1
Details of the datasets used for evaluation

| Dataset       | Users | Items | Ratings     | Rating Scale | Density   | Source                        |
|---------------|-------|-------|-------------|--------------|-----------|-------------------------------|
| MovieLens 100K| 943   | 1,682 | 100,000     | [1-5]        | 6.304%   | (Harper and Konstan, 2015)    |
| MovieLens 1M  | 6,040 | 3,900 | 1,000,209   | [1-5]        | 1.431%   | (Harper and Konstan, 2015)    |

Figure 4: Visualization of Similarity Graph for $\alpha = \{0.005, 0.01, 0.02, 0.03\}$ for MovieLens 100K with 943 users.

4.4. Impact of similarity graph size

In this experiment, we check the impact of graph size on rating accuracy. As we use graph features for every node (users) in the similarity graph, it's important to produce a similarity graph in a state that represents the similarity between nodes as optimized as it can be. For this purpose, we experimented with searching in parameter space, which impacts the size and the shape of the similarity graph. Figure 7 shows the RMSE vs. $\alpha$ in both dataset we used for the evaluation. As it can be seen in the figure, there is no direct relation between the result of the method. But, the minimum value of RMSE achieved on a specific value of alpha in the middle of the experiment range.

The main reason for this result is that when the alpha’s value is very small, all users can be connected due to this value because we consider just a very little common items in their ratings to connect them to each other in the similarity graph. Hence, most of the users are similar to each other in this condition and the difference will be missed in some cases. On the other hand, when the alpha’s value raises the similarity graph become more sparse (As it is shown in Figure 6). So, we consider the most of users not related to each other when the $\alpha$ value increases to very large values. There is an optimum point for the size of the similarity graph near the $\alpha = 0.01$ for dataset MovieLens 100K and near the $\alpha = 0.005$ for MovieLens 1M. The result of this parameter tuning has been produced with the exact condition of the final evaluation with k-fold cross-validation (Figure 6).

4.5. Dimension Reduction

We declared that we use Autoencoder to simultaneously extract new features and reduce the raw feature set dimension before clustering. In this experiment, we examine the learning algorithms for Autoencoder and check each algorithm’s ability to minimize the loss function on our input raw feature set. In all experiments, we have used a 5-layer Autoencoder with the structure shown in Figure 8. The activation function of all layers is ReLU (Nair and Hinton, 2010).

Both input and output size (raw feature vector’s sizes)
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(a) MovieLens 100K

Figure 5: Graph-based features for (a) MovieLens 100K and (b) MovieLens 1M both for $\alpha = 0.015$. Abbreviation used in the figure: PR (page rank), CD (degree centrality), CC (closeness centrality), CB (betweenness centrality), AND (average neighbor degree), and LC (load centrality). Centrality measures like CD and PR have more correlation than others, and it may be caused by the correlation of eigenvector centrality and degree centrality in our user graph. It means that users with a high degree of connection are likely connected in many cases.

are 35 for MovieLens 100K and 36 for MovieLens 1M.

We experiment with a set of optimizers to train the Autoencoder. Optimizers include Adagrad (Duchi et al., 2011), Adadelta (Zeiler, 2012), RMSProp (Hinton et al., 2012), Adam (Kingma and Ba, 2017), AdaMax (Kingma and Ba, 2017), Nadam (Dozat, 2016) and SGD (Bottou and Bousquet, 2008) with loss function of Mean Squared Error.

Figure 9 shows the result of the experiment and compares the optimizers in minimizing the loss function on validation data over 100 epochs of training. We have randomly selected 10% of users in MovieLens 1M and 20% of users in MovieLens 100K as validation data and exclude them from the training set. We can see the best results for Adam, Adadelta, and RMSProp optimizers. As a discussion about the result, RMSProp can be considered as an extension of Adagrad that deals with its radically diminishing learning rates. It is identical to Adadelta, except that Adadelta uses the RMS of parameter updates in the nominator update rule. Adam, finally, adds bias-correction and momentum to RMSProp (Ruder, 2017).

RMSprop, Adadelta, and Adam are very similar algorithms that do well in similar conditions. Kingma and Ba (2017) showed that the bias-correction helps Adam slightly exceed RMSprop towards the end of optimization when the gradients become sparser. Hence, Adam seems to be the best option for the optimizer. Recently, many researchers use vanilla SGD without momentum and a simple learning rate annealing schedule (Ruder, 2017). Nevertheless, in our experiment, SGD approaches to achieves a minimum, but it

Figure 6: Sparsity of combined features’ dataset vs. $\alpha$ before dimension reduction

Figure 7: RMSE vs. $\alpha$. There is an optimum point for $\alpha$ near the alpha = 0.01 for dataset MovieLens 100K and near the alpha = 0.005 for MovieLens 1M.
Figure 8: Structure of Autoencoder used for dimension reduction. Both input and output size are 35 for MovieLens 100K and 36 for MovieLens 1M.

Figure 9: The value of loss function on validation data over 100 epochs of training with seven target optimizers in the experiment.

may take longer than other methods.

We selected Adam as the optimizer in Autoencoder to encode the raw feature set. The output of the encoding process shows a diverse distribution with a low correlation between the encoded features. Figure 11 shows the encoded features for both MovieLens 100K and MovieLens 1M, which will be used for clustering the users.

We use elastic net regularization (linear combination of $L_1$ and $L_2$ penalties) (Zou and Hastie, 2005) for Autoencoder to avoid the overfitting on the training data and improve the model’s performance.

4.6. Number of Clusters

This section examines the mentioned method in section 3.3 to find the correct number of clusters for both datasets MovieLens 1M and MovieLens 100K. As listed before, we applied two methods for this reason; the Elbow method and the Average Silhouette method. The input of both methods is the encoded feature sets from the previous state. For both methods, we consider the range of $K$ in $[1–30]$. Figure 11 and Figure 12 show the algorithms’ iteration for the Elbow method and Average Silhouette method, respectively. The best value of $K$ has been founded, as shown in Table 2.

4.7. Results and Comparison with Other Methods

Performance of the proposed model has been evaluated on the datasets mentioned in section 4.1. Table 3 shows the result of the proposed model based on the best setting derived from the experiments conducted to find the best values for parameters $\alpha$ (section 4.4) and $K$ (section 4.6), and best optimizer for Autoencoder (section 4.5).

In another experiment, we are going to assess the GHRS method in tackling the cold-start problem. For this reason, we had to produce a synthetic dataset from the original dataset like MovieLens 100k or 1M. In the synthetic dataset, regard-
Table 2
Value of K suggested by Elbow method and Average Silhouette method

| Dataset       | Elbow method | Average Silhouette method |
|---------------|--------------|---------------------------|
| MovieLens 100K| K = 8, Distortion Score = 37.11 | K = 7, Average Silhouette = 18.93 |
| MovieLens 1M  | K = 9, Distortion Score = 264.61 | K = 7, Average Silhouette = 17.53 |

Figure 12: Finding the elbow point as the optimum number of clusters of users. (a) K-Mean Algorithm. (b) MiniBatchK-Mean Algorithm

Table 3
Performance metrics value for the proposed method on target dataset

| Dataset | RMSE | Precision | Recall |
|---------|------|-----------|--------|
| 100K    | 0.887, $S^2=1.595 \times 10^{-4}$ | 0.771 | 0.799 |
| 1M      | 0.833, $S^2=2.815 \times 10^{-4}$ | 0.792 | 0.838 |

Figure 13: GHRG method RMSE result versus the percentage of users which have been randomly removed from the user-item rating matrix.

5. Conclusion and Future Works

We have proposed a method for the recommendation in user-item systems in this paper. The method can be used for every user-item system that provides side information for both users and items. The proposed method’s main idea is finding the relation between users based on their similarities as nodes in a similarity graph and combining them with the users’ side information to solve the cold-start issue. Plus, we applied Autoencoder to extract new low dimensional features with low correlation and more information. This made the final clustering step more accurate and highly performed in time consumption. Final experiments and comparison with other methods showed the competitive results for the selected datasets and improved the best result on MovieLens 1M dataset.

There are several lines of research arising from this work that should be pursued. Future research might apply for the work on the item properties like user side information to detect similarity between items precisely. Admittedly, it will be like considering similarities between two users who similarly rate the same items, and their rates and properties in the similarity graph are close to each other. Indeed, in this case, items will be considered similar if identical or similar users (based on similarity definition between users in this research) rate them with the same patterns. Thus, we will also have the approach using graph features and deep-learning for users, for items.

On the other hand, it is great to devote future research to developing and extracting more other features from the similarity graph, which we did not mention in the current study. Besides, the structure of the Autoencoder might be an important area for future research. The different structures should be examined regarding the Autoencoder structure affecting feature extraction, training duration, and the model’s final performance. In this article, we used a predefined structure...
for Autoencoder using the heuristic method and manual tuning. Also, there are many methods to cluster the users in this method. They should be investigated and measured to find the optimal one for this type of feature space and distribution. This assumptions might be addressed in future studies.

As discussed in section 4.1, few datasets have side information for users and items (e.g., demographic data for users). It will be desirable to assess the proposed method with other future datasets that include this information.

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