

Predicting the Spatio-Temporal Evolution of Chronic Diseases in Population with Human Mobility Data

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Abstract

Chronic diseases like cancer and diabetes are major threats to human life. Understanding the distribution and progression of chronic diseases of a population is important in assisting the allocation of medical resources as well as the design of policies in preemptive healthcare. Traditional methods to obtain large scale indicators on population health, e.g., surveys and statistical analysis, can be costly and time-consuming and often lead to a coarse spatio-temporal picture. In this paper, we leverage a dataset describing the human mobility patterns of citizens in a large metropolitan area. By viewing local human lifestyles we predict the evolution rate of several chronic diseases at the level of a city neighborhood. We apply the combination of a collaborative topic modeling (CTM) and a Gaussian mixture method (GMM) to tackle the data sparsity challenge and achieve robust predictions on health conditions simultaneously. Our method enables the analysis and prediction of disease rate evolution at fine spatio-temporal scales and demonstrates the potential of incorporating datasets from mobile web sources to improve population health monitoring. Evaluations using real-world check-in and chronic disease morbidity datasets in the city of London show that the proposed CTM+GMM model outperforms various baseline methods.

1 Introduction

Recent studies show that many chronic and malignant diseases, e.g., heart disease, diabetes, and cancer, are extremely prevalent in our society [Siegel *et al.*, 2015]. To estimate disease penetration in a population and to be then able to implement appropriate health-care and preventative measures, healthcare administrators often perform statistical analysis from the records of hospital visits or conduct survey among a sample of residents. The analysis and survey usually incur high labor costs and are time-consuming. In addition, they often lead to coarse estimates both spatially and temporally.

*This work was done while Yingzi Wang was a visitor at the University of Cambridge

Although chronic diseases are, to some extent, related to patients' genetics, recent studies have shown that 70% to 90% of chronic diseases can be attributed to other factors [Rapaport and Smith, 2010]. For example, there is a causal relationship between chronic diseases and our daily habits, such as diet [McCullough *et al.*, 2002] and alcohol consumption [Martínez, 2005], which reflect different aspects of humans' lifestyles. *Lifestyle* depicts typical routine lives of people. Large-scale human mobility data collected through location-based social network services can act as a proxy for human lifestyle [Yuan *et al.*, 2013]. For instance, regular visits to college libraries, gyms, and lecture theaters, may correspond to the lifestyle of a college student, while a record of constant visits to meeting rooms and restaurants may indicate the lifestyle of a white-collar employee. These lifestyles are correlated with, and may reveal, certain chronic disease conditions of populations.

The connection between human mobility and diseases has been explored before. Several previous studies have investigated the outbreaks of infectious diseases via social ties and human mobility patterns [Meyers, 2007]. For chronic diseases, researchers have explored the association between human online activities and obesity [Mejova *et al.*, 2015]. However, little attention has been paid to lifestyles as a bridge between human mobility and chronic diseases nor exploring such correlation together with diseases-location similarity to predict residents' health.

In this work, we leverage the correlations between health and lifestyle reflected in human mobility to predict human health progression in populations. Using such data for these estimates gives unprecedented spatial and temporal granularity to the analysis but has also the potential to lower the costs of these studies and enable them to be applicable to regions for which other techniques would be deemed impractical or too expensive (e.g. developing regions). Here, we use human visitation patterns (check-ins) extracted from Foursquare (a location intelligence application) and the statistics of chronic disease morbidity in the London metropolitan area (presented on the government opening data website of the UK). We capture regional lifestyles as reflected in Foursquare mobility data, and apply a hybrid model to improve the prediction of public health conditions over simply using historic dataset. In summary, this paper offers the following contributions:

- We explore the correlations between human mobility pat-

terns and health conditions and apply a method which combines Gaussian mixture models (GMM) with collaborative topic modeling (CTM) to predict the health levels of a population, i.e., leveraging “where they go” to help predict “how healthy they are”.

- We get clues about human lifestyles from mobility patterns of residents, assuming that the groups of visited POIs are proxies for different lifestyles. We then exploit these inputs to identify fine-grained spatio-temporal associations between these lifestyles and chronic diseases for local populations.
- We collect real-world chronic disease and check-in data to evaluate our method and analyse the correlation between lifestyles and chronic diseases. Compared with methods using historic information solely, the proposed method shows a 45.7% reduction in mean square error (MSE) and a 1.67 times increase in R-squared value (R^2).

2 Related Work

Disease prediction. Many existing studies have tried to understand the spread of infectious diseases and forecast their outbreaks. Some works have analysed the social or contact networks formed by connections among individuals and human mobility patterns to model the outbreaks of infectious diseases [Meyers, 2007], while some others utilise large amounts of users’ status posts on social networks, such as Twitter, to analyse the public health on a large scale [Paul and Dredze, 2011]. Some other studies target on chronic diseases. Matic et al. [Matic and Oliver, 2016] seek help from smartphone-based health applications and wearable devices to continuously record human behaviour to analyse mental-health condition of individuals. The work [Mejova et al., 2015] employed both Foursquare and Instagram data to assess the relationship between fast food and obesity. Mason et al. [Mason et al., 2018] found that people who lived far away from fast-food restaurants were more likely to have small waist circumference, especially for women. However, there is a noticeable lack of research about the effects of human mobility patterns on the development of chronic diseases in small urban areas. To the best of our knowledge, no one has explored the similarities between chronic diseases and urban regions simultaneously. Our work aims to fill this gap.

Human mobility analysis. Patterns of human mobility are predictable and reflect how the residents of a certain area live in the physical world [Cho et al., 2011]. Many scholars have tried to learn such patterns in order to predict the movement of individuals [Gao et al., 2012]. Extensive research efforts have also been focused on, e.g., finding typical travel sequences by studying users’ check-in trajectories [Zheng et al., 2009], predicting users’ moving patterns by exploiting both the regularity of human mobility and influence of others [Wang et al., 2015], and making location recommendations with graphical models that integrates users’ preferences with their sequential movement patterns [Wang et al., 2016]. Different from above studies, in this paper we investigate the correlations between residents’ chronic disease development and their lifestyles indicated by their frequently visited venues and mobility habits.

Topic and Gaussian mixture models. Our analysis utilises topic modeling [Blei et al., 2003] and Gaussian mixture models [Friedman and Russell, 1997]. Topic modeling has been widely used to generate latent topics of documents [Wang and Blei, 2011] and learn human habits in daily lives [Yuan et al., 2013]. Gaussian mixture models are often used to describe the joint effect of multiple segments and factors [Bilmes and others, 1998]. It is widely utilised on cluster problems because of its unconstrained covariance structure and flexible application scenarios.

3 Disease Rate Evolution Prediction

3.1 Problem Definition

Human mobility records, e.g., the check-ins, reflect people’s movements in the physical world and to some extent reveal their lifestyles, which gradually affect their health conditions. Here we utilise the check-in dataset of Foursquare and the chronic disease dataset in London as an example for analysis. The Foursquare dataset contains check-in records from Dec. 2010 to Dec. 2013 created at 18,018 POI venues in 426 categories, e.g., fast food restaurant, gym, park, etc. There are over 4 million check-in transition between pairs of POIs, in each of which we have check-in timestamp and venue id (no user information). The chronic disease dataset contains the morbidity of 20 chronic diseases of 567 wards in London.

To exam the relationship between chronic diseases and human mobility patterns extracted from location-based services, we employ Pearson correlation analysis. More specifically, the correlation results between evolution rate of 7 most common chronic diseases and check-in amount of 17 categories of POIs from 2010 to 2013 in London are tested and presented in Figure 1(a). Here, for each chronic disease, we sort the disease evolution rates of 567 wards and split the ranking list into r segments averagely (we set $r = 19$, so there are 30 wards in each of the first 18 segments and 27 wards in the last one). We calculate the mean value of disease evolution rate and mean check-in volume of each category of POI in every segment to get two r -length sequences. Then, through calculating the correlation coefficient between the two types of sequences, we find that several categories, such as Malaysian restaurants, Chinese restaurants, and fast food restaurants, have high positive correlations with most of the 7 diseases except cancer. Some diseases have similar correlations with all the 17 categories of POIs, e.g. hypertension, heart failure, and obesity.

There are many confounding factors influencing population health across geographies. Our goal is not to draw causal conclusions regarding health. Instead we mine geo-referenced data emerging in urban environments so as to inform prediction models and attain better prediction results. Urban activities of users engaging with location intelligence systems can be indicative of lifestyle choices, some of which could be linked to health, yet describing a causal link is not the purpose of this paper.

People’s health conditions can be treated as dynamic factors. A healthy lifestyle may not prevent illness but may reduce the risk. Therefore, we focus on how people’s lifestyles may influence the development of these chronic diseases. Assume there are W regions in an area, represented as $\mathcal{R} =$

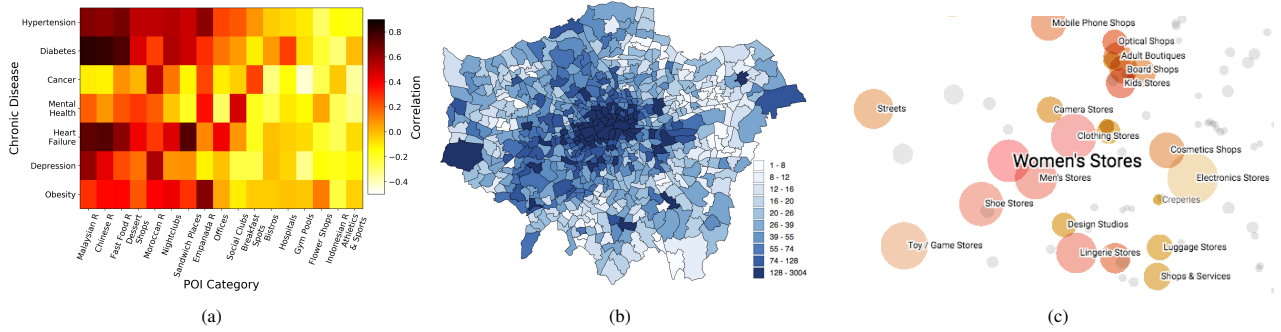


Figure 1: (a) Correlation between the evolution rate of 7 chronic diseases and check-in amount of 17 POI categories. (b) Distribution of the amount of POIs in the 630 wards of London. (c) 3-D projection of embedding results for POI categories.

$\{r_1, r_2, \dots, r_W\}$. Let $\mathcal{D} = \{d_1, d_2, \dots, d_S\}$ denote S chronic diseases and $\mathbf{H} \in \mathbb{R}^{W \times S}$ denote the evolution matrix of all the S diseases in W regions, where $h_{w,s}$ is the evolution rate of disease d_s from last year to this year in region r_w (a positive number denotes an increasing rate and a negative number denotes a decreasing rate). In practical terms, the disease data for some regions might be unavailable due to a tight budget. If we can only collect data in a subset \hat{R} regions of R , which fill in \hat{W} rows in the region-disease matrix \mathbf{H} and let others be zero (we denote \mathbf{H} matrix with zero rows by $\hat{\mathbf{H}}$), our goal is to predict the chronic disease development of the rest $\bar{W} = W - \hat{W}$ regions in set \bar{R} .

3.2 Overview

We adapt a prediction method which systematically integrates collaborative topic modeling and Gaussian mixture approach. Specifically, we firstly leverage an embedding method and a Gaussian mixture approach to aggregate categories of venues into several clusters according to their check-in patterns. Through that, we obtain a denser region-cluster-of-venues matrix. Then we apply a collaborative topic modeling method to extract lifestyle patterns of each region from human's check-in mobility. We hypothesise that lifestyle patterns in all regions and chronic disease evolution rates in some of the regions can be able to help exploring the chronic disease conditions in the missing regions.

3.3 Method

Venue Aggregation. Assuming there are N categories of venues, denoted as $\mathcal{V} = \{v_1, v_2, \dots, v_N\}$, we can collect the check-in amount of each category in \mathcal{V} and each region in \mathcal{R} (mentioned in Section 3.1), and build a region-category matrix $\mathbf{Y} \in \mathbb{R}^{W \times N}$. However, the spatial distribution of POIs is usually unbalanced in the city. In Figure 1(b), the amounts of Foursquare POIs in the 630 wards are presented. We can clearly see that central London and the region containing Heathrow Airport (the deep blue region on the left) have denser POI distribution. However, in some other regions the amount of POIs are considerably sparse, which leads to the sparsity of matrix \mathbf{Y} . To address this problem, we extract users' similar check-in preferences for some categories of POIs and aggregate these categories into a cluster. For example, if users often go shopping after having French or Italian food, we will cluster French and Italian food into a group.

To aggregate venues based on check-in patterns, we firstly represent each category of POI as a feature vector. Those categories with similar check-in patterns will have smaller vector distances from each other. This is similar to the embedding task in natural language processing, where each category of POI can be seen as a word, and users' transitions between categories of POIs is analogous to a sentence. Here we embed each category of POI into a P -length vector through word2vec method [Rehurek and Sojka, 2010]. Figure 1(c) shows a part of the 3-D projection¹ of embedding result for all the 426 categories of POIs when P is set to 100. Each circle denotes one category. The colored circles are the categories having shortest cosine distances to the category "women's stores". Redder colors represent closer relationships. The size of a circle indicates its "depth" from the surface of screen (foreshortening effects). We can observe that shoe stores, clothing stores, lingerie stores, and men's stores, have the most similar check-in patterns with women's stores.

We aggregate the N categories of POIs into C clusters by adopting Gaussian mixture method (GMM) [Friedman and Russell, 1997]. Compared with other cluster methods, e.g., k-means, GMM provides unconstrained covariance structure for each cluster, making the method more flexible. As illustrated in Figure 2, $\phi \in \mathbb{R}^{N \times P}$ is the venue-embedding matrix. The GMM can be described as follows:

$$\pi_n \sim \text{Dirichlet}(\gamma),$$

$$\sigma_c \sim \Gamma(\tau, \sigma_0), \quad \mu_c \sim \mathcal{N}(\mu_0, \nu \sigma_c), \quad c = 1, \dots, C,$$

$$g_n \sim \text{Categorical}(\pi_n), \quad \phi_n \sim \mathcal{N}(\mu_{g_n}, \sigma_{g_n}), \quad n = 1, \dots, N,$$

where π_n and g_n are the parameter of categorical distribution and the component of the n^{th} observation respectively. μ_c, σ_c are the parameters of Gaussian distribution of component c . $\gamma, \tau, \sigma_0, \mu_0, \nu$ are the shared hyperparameters. Let $\mathbf{X} \in \mathbb{R}^{W \times C}$ represent the region-cluster matrix, which can be estimated through parameter π and check-in matrix \mathbf{Y} :

$$\mathbf{X} = \mathbf{Y} \cdot \pi, \quad (1)$$

where $x_{i,j}$ denotes the check-in amount of venue cluster j in region r_i . Figure 2 presents the association from \mathbf{Y} and π to matrix \mathbf{X} through dotted lines.

Collaborative Topic Model. Until now, we have obtained the region-cluster matrix \mathbf{X} . Our objective is to extract lifestyle information from \mathbf{X} helping to predict chronic disease conditions in missing regions. Traditional probabilistic matrix

¹<http://projector.tensorflow.org/>

factorization (PMF) is a perfect method for recommendation tasks, leveraging the similarity among different users and items to complement the user-item matrix [Salakhutdinov and Mnih, 2007]. Similarly, as presented in Figure 1(a), similarities exist among diseases when analysing the correlation between diseases and check-ins. If we assume that similarities could also be found among regions, we can in the same way factorize regions’ disease evolution rate matrix \mathbf{H} into two low dimensional latent matrices \mathbf{L} and \mathbf{A} , both with dimension K . We denote K as the number of latent lifestyles, vector \mathbf{L}_i as the weight of latent lifestyles in region r_i , and vector \mathbf{A}_j as the influence from latent lifestyles on chronic disease d_j . Thus we can generate \mathbf{H} through the distribution:

$$h_{i,j} \sim \mathcal{N}(\mathbf{L}_i \cdot \mathbf{A}_j^\top, \varsigma_{i,j} \lambda_H^2), \quad (2)$$

where $\varsigma_{i,j}$ is 0 if the data of $h_{i,j}$ is missing, and 1 otherwise. The distribution of region and disease vectors are:

$$\mathbf{L}_i \sim \mathcal{N}(0, \lambda_L^2 \mathbf{I}_K), \quad \mathbf{A}_j \sim \mathcal{N}(0, \lambda_A^2 \mathbf{I}_K),$$

where \mathbf{I}_K is a K -dimensional identity matrix. A common way to optimize parameters \mathbf{L} and \mathbf{A} is to minimize the squared-errors objective function with regularization terms:

$$\Omega = \mathbf{I} \odot \|\mathbf{H} - \mathbf{L}\mathbf{A}^\top\|_F^2 + \frac{\lambda_H^2}{\lambda_L^2} \|\mathbf{L}\|_F^2 + \frac{\lambda_H^2}{\lambda_A^2} \|\mathbf{A}\|_F^2, \quad (3)$$

where $\|\cdot\|_F$ denotes the Frobenius norm and \odot denotes the Hadamard product operator [Kolda and Bader, 2009].

However, as only the data of \widehat{R} regions are available, information for leveraging similarities among the missing rows and the existing rows in matrix $\widehat{\mathbf{H}}$ is lacking. Hence we use the region-cluster matrix \mathbf{X} obtained in the last section. We assume each region is characterized by a particular set of lifestyles, and each cluster of categories of POIs may reflect human’s various lifestyles in different probabilities. It is similar to a topic structure: if we regard all the regions as documents, the check-in patterns of various POI clusters in a region can be considered as the words in a document. In analogy with the assumption that each topic is described by several representative words, each lifestyle is reflected in different check-in patterns. Intuitively, a typical topic model, LDA [Blei *et al.*, 2003], can be applied here to model the lifestyles in different regions.

However, LDA does not tackle the main problem in this work: how to leverage the lifestyles extracted from check-in mobility to fill the missing parts in matrix $\widehat{\mathbf{H}}$. We adopt the collaborative topic model (CTM) proposed in [Wang and Blei, 2011] here to combine probabilistic matrix factorization and topic modeling. Different from the assumption in [Wang and Blei, 2011], that offsets exist between the document-topic factor factorized from document-user part and document-word part respectively, we propose that people’s general lifestyles, reflecting in check-in patterns and health conditions, are relatively consistent. POIs’ visiting and chronic disease condition are just two views of lifestyles, where the former one is the perspective in people’s daily life activities, and the latter one is how these lifestyles influence people’s health status. Therefore, we leverage the same factor \mathbf{L} , to represent region-lifestyle interactions in topic modeling.

Specifically, the generative process of the hybrid method is as follows (illustrated in the bottom part of Figure 2):

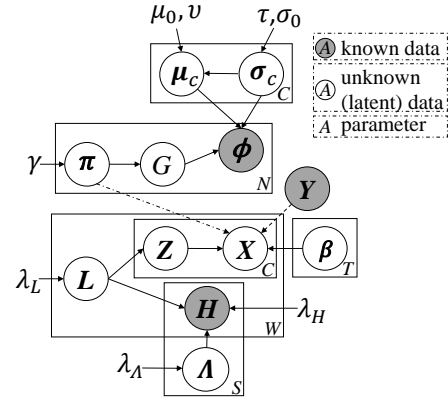


Figure 2: Illustration of CTM+GMM method.

asthma	chronic obstructive	heart failure
atrial fibrillation (AF)	pulmonary disease (COPD)	learning disabilities (LD)
cancer	diabetes	mental health
chronic kidney disease (CKD)	epilepsy	obesity
coronary heart disease (CHD)	hypertension	palliative care
dementia	hypothyroidism	smoking
depression	heart failure due to LVD (HFLVD)	stroke or transient ischaemic attacks (stroke or TIA)

Table 1: 20 chronic diseases and their abbreviations.

1. For each chronic disease j , draw disease latent factor $\mathbf{A}_j \sim \mathcal{N}(0, \lambda_A^2 \mathbf{I})$.
2. For each region i :
 - (a) Draw lifestyle factor $\mathbf{L}_i \sim \text{Dirichlet}(\lambda_L)$.
 - (b) For each category of venues $x_{i,c}$:
 - (i) Choose lifestyle assignment $z_{i,c} \sim \text{Mult}(\mathbf{L}_i)$.
 - (ii) Choose a category of venues $x_{i,c} \sim \text{Mult}(\beta_{z_{i,c}})$.
3. For each region-disease pair $h_{i,j}$, draw $h_{i,j} \sim \mathcal{N}(\mathbf{L}_i \cdot \mathbf{A}_j^\top, \varsigma_{i,j} \lambda_H^2)$,

where β_{z_t} is the distribution of POI clusters in lifestyle z_t .

Optimization. We apply EM method [Bilmes and others, 1998] to estimate the parameters $\{\mu_c, \sigma_c^2\} (c = 1, \dots, C)$ and π in GMM part. For the CTM part, we need to estimate the parameters $\{\mathbf{L}, \mathbf{A}\}$, where factor \mathbf{L} is employed both in PMF and topic modeling. We iteratively optimize parameters by two steps. Firstly, for topic modeling, we estimate \mathbf{L} through EM method, which is a typical optimization method for topic modeling. Then in the second step, we apply Gradient Descent method to estimate $\{\mathbf{L}, \mathbf{A}\}$ (we use the \mathbf{L} factor in the first step as the initialization here). We then go back to step 1 and set \mathbf{L} as the prior of region lifestyle distribution.

4 Evaluation

4.1 Set-Up

Data. We use three datasets in our experiment:

Foursquare dataset: The check-in records from Dec. 2010 to Dec. 2013 and the POI information in London. The details of Foursquare dataset are introduced in Section 3.1. In August 2015, Foursquare had more than 50 million active users and more than 10 billion check-ins already. The dataset used in

Cluster 1	Cluster 2
gardens, Russian r, malls, office supplies stores, banks, art galleries, music stores, bowling alleys, theme park, dessert shops, comedy clubs, Taiwanese r, college libraries, bookstores, souvenir shops, snack places, cosmetics shops	hotels, jazz clubs, skate parks, tattoo parlors, nightlife spots, Latin American r, social clubs, cupcake shops, smoke shops, beer stores, laundry services, dinners, modern European r, convenience stores, bridges

Table 2: Several categories of POIs in 2 clusters in venue aggregation part (r: restaurants).

this research is shared directly by Foursquare under a research contract agreement.

Boundary-line dataset of London: The dataset is collected from UK government websites², which contain the shapefiles of ward-level (electoral districts at sub-national level) boundary lines in London. In total, there are 630 wards. This spatial data contains the shape line, name, and id of each ward (shape lines are shown in Figure 1(b)).

Disease dataset: We collect the data from a government open data website of UK³. They publish the population, the annual morbidity (value of patients/population in an area) of 19 prevalence diseases (Table1), and the utilization rate of “palliative care” from year 2005 to 2015. Palliative care⁴ is a specialized medical care for people with life-threatening illness. Since we consider it as an indicator for the morbidity of malignant diseases, we call it “disease” here. The data for each year is from Apr. of one year to Mar. of the next year. For convenience, we use dataset “2013”, for example, to represent the annual dataset from Apr. 2013 to Mar. 2014 in the rest of the paper. We collected the data from 2009 to 2013, consistent with the period of check-in data. The dataset contains the popularity and morbidity data of 567 wards in London (no data for the rest 63 wards).

Metrics. We apply two metrics here to evaluate the prediction performance: (*MSE*) and R^2 score:

$$MSE(h, h')_{\tilde{\mathcal{R}}} = \frac{1}{\tilde{W} \cdot S} \sum_{r_i \in \tilde{\mathcal{R}}} \sum_{j=1}^S (h_{r_i, j} - h'_{r_i, j})^2,$$

$$R^2(h, h')_{\tilde{\mathcal{R}}} = \frac{1}{\tilde{W}} \sum_{r_i \in \tilde{\mathcal{R}}} \left(1 - \frac{\sum_{j=1}^S (h_{r_i, j} - h'_{r_i, j})^2}{\sum_{j=1}^S (h_{r_i, j} - \bar{h}_{r_i})^2} \right),$$

where $h'_{r_i, j}$ is the prediction result of item $h_{r_i, j}$ and $\bar{h}_{r_i} = \frac{1}{S} \sum_{j=1}^S h_{r_i, j}$. \tilde{W} is the amount of regions in testing set. R^2 score reflects how well the model performs in the prediction, considering the error and the mean of true values simultaneously. Here lower *MSE* and higher R^2 represent better result.

Baselines. We compare the hybrid method with 4 methods:

- *Regression:* We apply two regression methods using history data solely: boosting regression (BR) and support vector regression (SVR). They utilise the data of chronic dis-

²<https://www.ordnancesurvey.co.uk/opendatadownload/products.html#BDLINE>

³https://data.gov.uk/dataset/quality_and_outcomes_framework_achievement_prevalence_and_exceptions_data

⁴https://en.wikipedia.org/wiki/Palliative_care

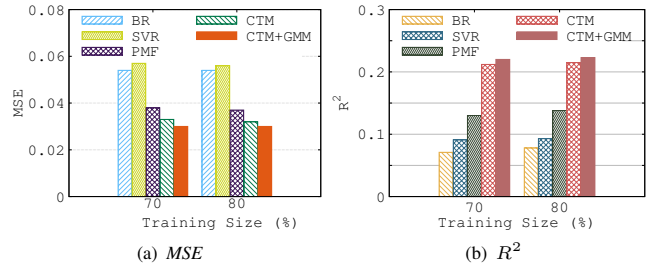


Figure 3: Prediction performance of all the evaluated methods on (a) *MSE* and (b) R^2 , when training size is 70% or 80%.

Disease	ER	R^2 S	Disease	ER	R^2 S
asthma	-0.03	0.155	hypertension	0.07	0.156
AF	3.17	0.166	hypothyroidism	2.94	0.129
cancer	7.12	0.150	heart failure	0.59	0.160
CKD	-2.12	0.157	HFLVD	-44.27	0.150
CHD	-1.17	0.159	LD	4.36	0.175
COPD	2.94	0.166	mental health	2.37	0.170
dementia	11.13	0.163	obesity	-16.10	0.133
depression	12.15	0.144	palliative care	21.92	0.140
diabetes	3.10	0.141	smoking	1.18	0.181
epilepsy	2.02	0.168	stroke or TIA	1.11	0.142

Table 3: Average evolution rates of the 20 diseases and their R^2 scores in prediction (**ER**: evolution rate (%), **S**: score).

ease evolution rate (2009 - 2012) to predict the evolution rate from 2012 to 2013.

- *Probabilistic matrix factorization (PMF):* In PMF, the region-disease matrix \hat{H} (with missing rows) is decomposed through equation 3 (H is replaced with \hat{H}).
- *Collaborative topic modeling (CTM):* We compare our method with CTM to emphasize the improvement achieved through venue aggregation.

4.2 Results

We compare the prediction performance of our method with baseline models in this section. Here we embed each POI category into a 100-length vector, and produce matrix $\phi \in R^{426 \times 100}$. Each vector represents POI category’s features in check-in pattern space, where a shorter distance between two categories indicates more similar check-in patterns. We aggregate the 426 embedded vectors into C clusters through a GMM method. Table 2 shows POI categories in two typical clusters when $C=20$. We can see that most POIs in the first cluster are more about nature (parks, gardens, etc.), art (art galleries, music stores, comedy clubs, etc), and reading (libraries and bookstores), while the second cluster mainly includes nightlife-related venues. The clustering results provide us new perspectives on London citizens’ visitation to POIs, which are not totally consistent with our experience. For instance, it is unexpected that bridges have similar check-in patterns with some clubs and nightlife spots. This may be because that the fantastic lighting systems of some famous bridges in London, such as Albert Bridge and Tower Bridge, attract a large number of viewers during the night hours.

Next, we predict the missing rows in matrix H , in which item $h_{i, j}$ is the evolution rate of disease d_j in ward r_i from year 2012 to 2013. $h_{i, j}$ is generated from the equation

$h_{i,j} = (m_{i,j}^{2013} - m_{i,j}^{2012})/m_{i,j}^{2012}$, where m represent morbidity. Firstly, we obtain the region-cluster matrix \mathbf{X} through Equation 1. Then, the latent lifestyle factor \mathbf{L} and latent chronic disease factor \mathbf{A} are estimated through CTM in Section 3.3. Finally, we predict the missing regions $\tilde{\mathbf{R}}$ in matrix \mathbf{H} through Equation 2. We randomly select 70% and 80% rows in matrix \mathbf{H} as training data and evaluate the prediction performance in the rest rows, respectively. Due to the space limit, for all the methods, we tune the parameters through the training set and show the best prediction results on testing set. For each method, we run the analysis for 20 times and show the average results in Figure 3. The latent dimension length K and the amount of lifestyles T (in CTM part) are set to 50. As for C, the amount of component (in GMM part), is set to 20. It is obvious that CTM+GMM has the best performance on the two metrics. The two regression methods have the highest MSE results and lowest R^2 scores. We can infer that the historic evolutions of these chronic diseases cannot provide enough regular patterns (e.g. linear or periodic patterns) for future prediction. Compared with traditional PMF, CTM has an average improvement of 13% in MSE and 59% in R^2 score. Moreover, when we add the GMM part to CTM, it furtherly achieves a 7.6% improvement in MSE and a 3.7% improvement in R^2 score.

We show the average evolution rate of all the diseases and their R^2 scores in predictions through CTM+GMM method from 2012 to 2013 in Table 3 (different from the definition in Section 4.1, we here show the R^2 scores along each disease). The Pearson correlation coefficient of evolution rate list and R^2 score list in the table is 0.111, which means that there is no obvious correlation between these two factors: the predictability of a disease is neither positively nor negatively related to its own evolution rate. From this table, we can see a significant growth of palliative care during that year, indicating that the morbidity of malignant diseases, e.g., cancer, had increased. Also, the morbidity of all the psychiatric and mental diseases, e.g. depression, dementia, and mental health, increased dramatically in that period. Actually, as illustrated in [Mulyala and Varghese, 2010], depression and dementia have complex relationships: depression has been both a risk factor and a prodrome of dementia, which is also reflected in the close evolution rates between them. Good news is that the obesity cases in London decreased in the same period, which has been a serious health concern in UK for a long time. Moreover, we observe that obesity has a relatively low R^2 score in the experiment for both our method and baselines. This may be due to the fact that different from some incurable chronic diseases, e.g., diabetes [Etuk and others, 2010], of which the morbidity is more stable, people may lose weight through various approaches, making the evolution of obesity more difficult to predict.

5 Lifestyle and Chronic Diseases

We present the correlation between the 20 chronic diseases and various lifestyles. We leverage the check-in pattern as a projection of lifestyle. Specifically, we use the lifestyle-cluster factor (β in Section 3.3), and the disease-lifestyle factor \mathbf{A} , to uncover the hidden relationship between chronic diseases and POIs. Some of the observed correlations are

Check-in Lifestyles	Correlated Dis
government buildings, pubs, sushi r, plazas, candy stores, steakhouses, burger joints, Subways, Chinese r, churches, bookstores, fast food r, coffee shops, nightclubs	heart failure, attacks, asthma, COPD, coronary heart disease, diabetes, hypertension, obesity, HFLVD, stroke or TIA
Indian r, convention centers, fast food r, sandwich places, grocery stores, Thai r, dim sum r, hotel bars, bakeries, hostels, nightclubs, fried chicken joints	
dessert shops, organic groceries, gay bars, middle eastern r, sports bars, cocktail bars, whisky bars, nightclubs, offices, boutiques, Chinese r, hotel bars, hotels, Italian r	cancer, smoking chronic kidney, depression, disease, obesity,

Table 4: Typical check-in lifestyles and the highly-correlated diseases (Dis: Diseases, r: restaurants).

consistent with the research findings in clinical medicine and physiology, while others provide new insights into some open problems. We list 3 typical check-in lifestyles and their highly correlated diseases in Table 4.

The top two lifestyles are dominated by fast food venues: fast food restaurants, pizza places, fried chicken joints, and Asian food (Chinese and sushi restaurants). These two lifestyles show association with 10 chronic diseases listed on the right. This is coherent with the statement that some diseases, like heart-related diseases, hypertension, obesity, and diabetes, are correlated with diets high in sugar and fat. Additionally, asthma and chronic obstructive pulmonary disease are also highly correlated with these two lifestyles. This is consistent with previous studies in physiology [Rosenkranz *et al.*, 2010] claiming that a high-fat diet may contribute to chronic inflammatory diseases of airways and lungs.

The third lifestyle is alcohol-oriented, where 6 of the 14 categories of POIs are bars or clubs. The induction of alcohol on some diseases, like cancer, depression, have been proved in previous studies [Martínez, 2005]. However, as illustrated in [White *et al.*, 2009], evidence of an association between alcohol consumption and chronic kidney disease is conflicting. Here we draw inspiration from large-scale human mobility data and provide an intuitive perspective on the positive correlation between lifestyles related to alcohol and chronic kidney disease.

6 Conclusion

In this paper, we leverage human mobility data to study the evolution of human health conditions. We embed the POIs into vectors according to human transition patterns to capture their semantic meanings. Then we combine Gaussian mixture methods with collaborative topic modeling to predict health conditions, which is able to deal with data sparsity and extract human lifestyles from check-in patterns. Extensive experiments using real-world datasets indicated that CTM+GMM has a significant improvement on prediction tasks compared to other methods.

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