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Similar nodes in large graphs

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ABSTRACT

Given a large graph, in the range of millions or billions of nodes, it is difficult to efficiently find similar nodes. Examples of applications of finding similar nodes in data represented as graphs are all around the web nowadays: *people you may know* and *who to follow* suggestions on social networks, or topics that may interest you on different websites.

This project will explore ways to obtain meaningful information of this nature from large datasets represented as graphs.

The goals of this project are developing efficient ways to find similar nodes in large graphs for different use cases and implementing an open-source software library to provide everyone with this functionality.

Introduction

Plenty of datasets are or can be represented as graphs where vertices represent entities and edges represent relationships between entities. A problem of interest is to find entities that are similarly connected. Example instances of this problem are finding *people you may know* in a social network, people with common interests from research publications repositories or identifying possible duplicates in a dataset.

It is not computationally feasible to calculate the exact similarities between vertices of very large graphs. In this project, we investigate a method to compute an approximation of similarities between vertices and attempt to evaluate its performance (accuracy) on different datasets.

BRUTE-FORCE APPROACH

The similarity algorithm is described in regards to the undirected graph $G(v,\epsilon)$ with adjacency matrix M. The number of nodes is N = |v| and the number of edges is $|\epsilon|$. In order to derive an algorithm to find similarities between vertices in a graph, a measure of similarity between

two vertices must be defined. M is the adjacency matrix of the graph and $M_{ij} = [(i, j) \in \epsilon]$, that is $M_{ij} = 1$ if there is an edge between vertices i and j, otherwise $M_{ij} = 0$. Let D be a normalised adjacency matrix such that each column sums up to 1:

$$D_{ij} = \frac{1}{\sum_{i=1}^{N} M_{ij}} \tag{1}$$

Let $\delta_{\mathbf{i}}$ be the i^{th} column of the $N \times N$ identity matrix, and $\mathbf{C_i}$ a vector that represents how vertex i is connected to the graph. In a brute-force approach, $\mathbf{C_i}$ can be computed as:

$$\mathbf{C_i} = \sum_{t=0}^{\infty} \mu^t D^t \delta_i \tag{2}$$

Where $\mu \in (0,1)$ is a penalising factor. μ controls how much the far connections are considered into the similarities. Therefore a large μ enforces consideration of very far connections whereas a small μ considers far connections less. The similarity between two vertices i and j is the (squared) norm of the difference between $\mathbf{C_i}$ and $\mathbf{C_i}$:

$$\|\mathbf{C_i} - \mathbf{C_j}\|^2 \tag{3}$$

The brute-force approach was run on small randomly generated connected graphs (200 vertices) with 50 iterations and it gave meaningful results but it does not scale for large datasets. In the next section an algorithm that scales up to large datasets is described.

APPROXIMATIVE APPROACH

Geometrically, if vectors C_i and C_j start from the origin, then C_i , C_j and $C_i - C_j$ describe a triangle. Let θ be the angle between C_i and C_i . From the Law of Cosines:

$$\|\mathbf{C_i} - \mathbf{C_j}\|^2 = \|\mathbf{C_i}\|^2 + \|\mathbf{C_j}\|^2 - 2\|\mathbf{C_i}\|\|\mathbf{C_j}\|\cos\theta$$
 (4)

Take the dot product $C_i^T C_j = \|C_i\| \|C_j\| \cos \theta$. The angle between C_i and itself is 0 and $\cos 0 = 1$, thus $\|C_i\|^2 = C_i^T C_i$. Now *Equation 4* becomes:

$$\|\mathbf{C_i} - \mathbf{C_j}\|^2 = \mathbf{C_i}^{\mathrm{T}} \mathbf{C_i} + \mathbf{C_i}^{\mathrm{T}} \mathbf{C_j} - 2\mathbf{C_i}^{\mathrm{T}} \mathbf{C_j}$$
(5)

Therefore it is sufficient to be able to compute all dot products $\mathbf{C_i^T C_j}$ (for all $i, j \in v$) to compute the similarities between all vertices of the graph. Computing and storing all $\mathbf{C_i^T C_j}$; $i, j \in v$ takes too much time and memory for a very large dataset if we use the brute-force approach.

An approximative algorithm to efficiently compute the dot products $C_i^T C_j$ will be described. Let W be a diagonal matrix:

$$W_{ij} = \frac{[i=j]}{\sum_{i=1}^{N} M_{ij}}$$
 (6)

Now *Equation 1* can be written as D = WM, thus

$$D^t = (WM)^t \tag{7}$$

Define a new matrix *A*:

$$A = W^{1/2} M W^{1/2} \tag{8}$$

From Equation 7 and Equation 8:

$$D^{t} = W^{1/2} A^{t} W^{-1/2} (9)$$

The matrices $W^{1/2}$ and $W^{-1/2}$ are easy to compute because W is a diagonal matrix. Apply the matrix operation only on the diagonal elements and obtain $(W^{1/2})_{ii} = (W_{ii})^{1/2}$ and $(W^{-1/2})_{ii} = (W_{ii})^{-1/2}$.

Let λ_i be the i^{th} eigenvalue and $\bigvee^{(i)}$ be the i^{th} eigenvector ($\forall i \in \{1, 2, ..., N\}$). \bigvee is a matrix of all computed eigenvectors (such that $\bigvee^{(i)}$ is the i^{th} column of \bigvee), and \bigwedge is a diagonal matrix of all computed eigenvalues:

The eigendecomposition of the matrix A is $A = \bigvee \bigwedge \bigvee^{-1}$. The graph G is an undirected graph which makes A a real symmetric matrix, thus $\bigvee^{-1} = \bigvee^{T}$. The eigendecomposition of the

matrix *A* is therefore:

$$A = \bigvee \bigwedge \bigvee^{T} \tag{12}$$

$$\Rightarrow A^{t} = \left(\bigvee \bigwedge \bigvee^{T}\right)^{t} = \bigvee \bigwedge^{t} \bigvee^{T} \tag{13}$$

$$\Leftrightarrow A^t = \sum_{a=1}^N \lambda_a^t \bigvee^{(a)} \bigvee^{(a)^T}$$
 (14)

From Equation 9 and Equation 14:

$$D^{t} = W^{1/2} \sum_{a=1}^{N} \lambda_{a}^{t} \bigvee^{(a)} \bigvee^{(a)^{T}} W^{-1/2}$$
(15)

Plug D^t from Equation 15 into Equation 2:

$$\mathbf{C_{i}} = \sum_{t=0}^{\infty} \mu^{t} W^{1/2} \sum_{a=1}^{N} \lambda_{a}^{t} \bigvee^{(a)} \bigvee^{(a)^{T}} W^{-1/2} \delta_{i}$$
 (16)

Rearrange the equation and obtain:

$$\mathbf{C_{i}} = W^{1/2} \sum_{a=1}^{N} \sum_{t=0}^{\infty} \mu^{t} \lambda_{a}^{t} \bigvee^{(a)} \bigvee^{(a)^{T}} W^{-1/2} \delta_{i}$$
 (17)

Note that $\sum_{t=0}^{\infty} \mu^t \lambda_a^t$ is a sum of a geometric series with the first term 1 and ratio $\mu \lambda_a$, therefore:

$$\sum_{t=0}^{\infty} \mu^t \lambda_a^t = \frac{1}{1 - \mu \lambda_a} \tag{18}$$

Observe $\bigvee^{(a)^T} W^{^{-1/_2}} \delta_{\mathbf{i}}$ is a scalar:

$$\bigvee^{(a)^{T}} W^{-1/2} \delta_{\mathbf{i}} = \bigvee_{i}^{(a)} W_{ii}^{-1/2}$$
(19)

Substitute Equation 19 and Equation 18 back into Equation 17 and obtain:

$$\mathbf{C_i} = W^{1/2} \sum_{a=1}^{N} \frac{1}{1 - \mu \lambda_a} \bigvee_{i}^{(a)} \bigvee_{i}^{(a)} W_{ii}^{-1/2}$$
 (20)

An approximation of the vectors $C_i(\forall i \in v)$ can be obtained by only using the m leading eigenvectors and eigenvalues of A instead of all of them. Compute the largest m eigenvalues and eigenvectors of the matrix A using the Lanczos Method for real symmetric matrices[1]. For directed graphs the Arnoldi Iteration[2] will be used instead. These algorithms perform

well on large sparse matrices. In real datasets it is very often the case that the number of edges is orders of magnitude smaller than the number of vertices squared, resulting in sparse adjacency matrices and a sparse A. Define $\hat{\mathbf{C}}_{\mathbf{i}}$ to be an approximation of $\mathbf{C}_{\mathbf{i}}$:

$$\hat{\mathbf{C}}_{\mathbf{i}} = W^{1/2} \sum_{a=1}^{m} \frac{1}{1 - \mu \lambda_a} \bigvee^{(a)} \bigvee_{i}^{(a)} W_{ii}^{-1/2}$$
(21)

To simplify the equation, let Z be a $N \times m$ matrix such that:

$$Z_{ij} = \frac{\bigvee_{i}^{(j)} W_{ii}^{-1/2}}{1 - \mu \lambda_{j}}$$
 (22)

Substitute in *Equation 21* and get:

$$\hat{\mathbf{C}}_{\mathbf{i}} = W^{1/2} \sum_{a=1}^{N} Z_{ia} \bigvee^{(a)}$$
 (23)

The dot product between \hat{C}_i and \hat{C}_j becomes:

$$\hat{\mathbf{C}}_{\mathbf{i}}^{\mathbf{T}}\hat{\mathbf{C}}_{\mathbf{j}} = \left(W^{1/2} \sum_{a=1}^{m} Z_{ia} \bigvee^{(a)}\right)^{T} \left(W^{1/2} \sum_{a'=1}^{m} Z_{ia'} \bigvee^{(a')}\right)$$
(24)

$$\Leftrightarrow \hat{\mathbf{C}}_{\mathbf{i}}^{\mathrm{T}} \hat{\mathbf{C}}_{\mathbf{j}} = \sum_{a=1}^{m} Z_{ia} \bigvee^{(a)^{\mathrm{T}}} W \sum_{a'=1}^{m} Z_{ia'} \bigvee^{(a')}$$
(25)

$$\Leftrightarrow \hat{\mathbf{C}}_{\mathbf{i}}^{\mathsf{T}} \hat{\mathbf{C}}_{\mathbf{j}} = \sum_{a=1}^{m} \sum_{a'=1}^{m} Z_{ia} Z_{ja'} \bigvee^{(a)^{\mathsf{T}}} W \bigvee^{(a')}$$
(26)

Define a new $m \times m$ matrix Q such that:

$$Q_{i,i} = \bigvee^{(i)^T} W \bigvee^{(j)} \tag{27}$$

 $\mathbf{Z_i}$ is the i^{th} row of the matrix Z. Substitute Q into $Equation\ 26$ and obtain:

$$\hat{\mathbf{C}}_{\mathbf{i}}^{\mathbf{T}}\hat{\mathbf{C}}_{\mathbf{j}} = \sum_{a=1}^{m} \sum_{a'=1}^{m} Z_{ia} Z_{ja'} Q_{aa'}$$
(28)

$$\Leftrightarrow \hat{\mathbf{C}}_{i}^{T} \hat{\mathbf{C}}_{j} = \mathbf{Z}_{i}^{T} Q \mathbf{Z}_{j}$$
 (29)

From *Equation 29* it is deduced that it is only required to store the matrixes Q and Z, which only takes $O(m^2 + Nm)$ memory, instead of $O(N^2)$ to store all $\hat{\mathbf{C}}_{\mathbf{i}}$ vectors.

To compute the similarity between two vertices, i and j, use the norm formula from Equa-

tion 29 and obtain:

$$\|\hat{\mathbf{C}}_{i}^{T} - \hat{\mathbf{C}}_{i}\|^{2} = \mathbf{Z}_{i}^{T} Q \mathbf{Z}_{i} + \mathbf{Z}_{i}^{T} Q \mathbf{Z}_{i} - 2 \mathbf{Z}_{i}^{T} Q \mathbf{Z}_{i}$$
(30)

After computing the matrices Z and Q, the time complexity of computing the similarity between two nodes is therefore the same as multiplying three small matrices (of sizes $1 \times m$, $m \times m$ and $m \times 1$).

Vectorisation

It is usually more efficient to have vectorised implementations (to use matrix operations in favour of iterative solutions) in programming languages like Matlab, Octave or R. In this subsection is presented a way of computing the matrices Z and Q using matrix operations. For this purpose, in this subsection, \bigvee only contains the computed eigenvectors (is of size $N \times m$), and \bigwedge only contains the computed eigenvalues (is of size $m \times m$).

To vectorise the computation of Z from Equation 22, define a new diagonal matrix R:

$$R = \begin{bmatrix} \frac{1}{1-\mu\lambda_1} & 0 & \dots & 0\\ 0 & \frac{1}{1-\mu\lambda_2} & \dots & 0\\ \vdots & \vdots & \ddots & \\ 0 & 0 & \dots & \frac{1}{1-\mu\lambda_m} \end{bmatrix}$$
(31)

And then define *Z* as:

$$Z = W^{-1/2} \bigvee R \tag{32}$$

A vectorised version of *Equation 27* (to compute *Q*):

$$Q = \bigvee^{T} W \bigvee \tag{33}$$

IMPLEMENTATION

A choice had to be made on what tools and languages to use for the implementation of this algorithm. It involves linear algebra so programming languages that support such operations or have good libraries for this purpose were candidates. Matlab was chosen for this task

because it has very strong linear algebra features built-in, it is easy and fast to try different experiments (can also be used interactively) and programs are usually smaller in size (lines of code) than their variants in C++ or other languages. At a later stage when the algorithm is closer to its final version, porting it to a language like C++ will be considered.

Matlab has the function eigs(A, m) to compute the leading m eigenvectors and eigenvalues. Internally, it uses the Lanczos Method for Hermitian matrices and the Arnoldi Iteration for non-Hermitian matrices.[3]

For the implementation of both the approximative approach and the brute-force algorithm, sparse matrices were used. A sparse matrix in Matlab only stores the non-zero values of a matrix, therefore a sparse matrix only uses memory for the positions where it holds values.[4] For instance, to store the adjacency matrix of a very large graph $G(v,\epsilon)$, using a dense matrix would take $O(|v|^2)$ memory whereas using a sparse matrix only takes $O(|\epsilon|)$ memory.

The approximative approach was implemented in Matlab as a couple of Matlab functions. The function signatures along with descriptions is presented in *Table 1*.

similarity(adj, mu, m)
Computes and returns the matrices Q and Z.

sim2(q,z,i,j)
Computes the similarity between vertices i and j.

Computes the similarity between vertex i and all other vertices.

top(q,z,i)
Uses simlist(q,z,i) and sorts the results.

randomGraph(nodes, vertices, directed)
readEprints()
Reads ePrints data from a pre-processed file.

Table 1: Implementation interface. Parameters: q represents the matrix Q, z represents the matrix Z, adj represents the adjacency matrix, and i and j represent vertex numbers.

The brute-force algorithm was implemented as a separate set of Matlab functions with a very similar interface.

EVALUATION

A simple evaluation metric of how well the approximative approach retains the most important information from the graph is to evaluate how different are the top similar vertices for the brute-force approach and the approximative approach with different number of eigenvalues and eigenvectors used.

A random connected graph of N=200 vertices and 350 edges was generated. The similarities between all pairs of nodes were computed using both the brute-force approach and the approximative approach, and nodes were sorted by their similarities to obtain 200 tops (the higher the position in the top, the more similar the nodes are). Let B_{ij}^m be the position of node j in the top similar nodes of node i for the brute-force approach with m iterations. Similarly let T_{ij}^m be the position of node j in the top similar nodes of node i for the approximative approach with m eigenvectors. Define the difference between two approaches to be:

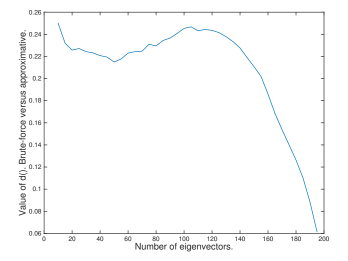
$$d(B^m, T^{m'}) = \frac{1}{N^3} \sum_{i=1}^{N} \sum_{j=1}^{N} abs(B_{ij}^m - T_{ij}^{m'})$$
(34)

The results can be observed in *Figure 1b*. The choice of the penalising factor $\mu=0.5$ was used for all the runs. The difference d was evaluated between the brute-force approach with m=20 iterations and the approximative approach with m'=5to195 eigenvectors. We can observe that if we use the majority of eigenvalues, the difference is insignificant. It is computationally unfeasible to do that on a large dataset thus we will aim for a number of eigenvectors m' in the range of 20 to 40, where the difference seems to be at a local minimum. The local minimum might differ from dataset to dataset depending mainly on the size of the dataset. For completeness, the experiment was run on a randomly generated graph with 500 vertices and 700 edges. The first local minimum of d() was at around 100 eigenvectors with a difference of approx. 0.2, and at 40 eigenvectors the difference is 0.2297, and at 20 eigenvectors the difference is 0.2557.

LIMITATIONS OF CURRENT IMPLEMENTATION

The current implementation of this algorithm has various limitations which are discussed along with possible improvements.

Only undirected graphs Directed graphs are not currently supported. In practice, datasets have meaningful unidirectional relationships (e.g. in a social network person A follows person B, but B does not follow A), and often datasets are represented as directed graphs rather than undirected graphs. The algorithm can be adapted to support both directed and undirected graphs but it will have the disadvantage of requiring to compute V^{-1} (for undirected graphs,



Parameters	Value of d()
$d(B^{20}, T^{10})$	0.2501
$d(B^{20}, T^{20})$	0.2257
$d(B^{20}, T^{30})$	0.2243
$d(B^{20}, T^{40})$	0.2208
$d(B^{20}, T^{50})$	0.2150
$d(B^{20}, T^{100})$	0.2453
$d(B^{20}, T^{199})$	0.0325

(a) A plot of the differences on a range of eigenvalues.

(b) Table of some of the values.

Figure 1: Difference of results between brute-force and approximative approaches.

$$V^{-1} = V^T$$
).

Ignored relationship types In the real world datasets might have different types of relationships between entities. Some relationship types might be more relevant than others in finding a specific result. For instance, in a social network two people being friends might be more relevant for recommending new friends than two people following the same topic. The current implementation of the algorithm can compute similarities if it is given a weighted (symmetric) adjacency matrix but it does not have a way to automatically obtain these weights from the dataset.

Not distributed What if the dataset is too large to fit into main memory? The algorithm is currently only designed to run on one machine. Methods of distributing the algorithm on more than one machine over a network will be investigated in the future.

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