ABSTRACT

We study the structure of the social graph of mobile phone users in the country of Mexico, with a focus on demographic attributes of the users (more specifically the users’ age). We examine assortativity patterns in the graph, and observe a strong age homophily in the communications preferences. We propose a graph based algorithm for the prediction of the age of mobile phone users. The algorithm exploits the topology of the mobile phone network, together with a subset of known users ages (seeds), to infer the age of remaining users. We provide the details of the methodology, and show experimental results on a network of more than 70 million users. By carefully examining the topological relations of the seeds to the rest of the nodes in the network, we find topological metrics which have a direct influence on the performance of the algorithm. In particular we characterize subsets of users for which the accuracy of the algorithm is 62% when predicting between 4 age categories (whereas a pure random guess would yield an accuracy of 25%). We also show that we can use the probabilistic information computed by the algorithm to further increase its inference power to 72% on a significant subset of users.

1. INTRODUCTION

Over the past decade, mobile phones have become prevalent in all parts of the world, across all demographic backgrounds. Mobile phones are used by men and women across a wide age range in both developed and developing countries. They have become one of the most important mechanisms for social interaction within a population, making them an increasingly important source of information to understand human behavior, human demographics, and the correlations between them. In particular, mobile phone usage information is being used to perform quantitative analysis on the demographics of users regarding their age group, gender, education, and socioeconomic status.

In this work we combine two sources of information: communication logs from a major mobile operator in Mexico, and information on the demographics of a subset of the users population. This allowed us to first perform an observational study of mobile phone usage, differentiated by categories such as gender and age groups. This study is interesting in its own right, since it provides knowledge on the structure and demographics of the mobile phone market in Mexico. Here we focus our study on one variable: the age of the users.

Furthermore, we tackle the problem of inferring demographic attributes, such as their age. The ability to predict such features has numerous applications for social organizations (for example, enabling them to perform specific health campaigns for women) and for businesses, allowing them to gain insights on their customers and on the open market, to segment (current and potential) customers by demographic attributes, and to perform directed marketing campaigns.

A semantic description of the network may consist in characterizing particular user attributes (e.g. gender, age or credit risk), or link attributes describing relationships between users (e.g. friendship, family or working relationship). One can also characterize individual users based on more abstract social attributes derived from their communication patterns or their geographic mobility, and study how these features relate to other social and behavioral properties. Methods such as logistic regression, readily used in this approach, emphasize finding similarity patterns between nodes given their known features and mostly rely on the assumption that the selected node features have a functional relationship with the particular attribute one wants to infer.

A complementary approach is to focus on the complex relationships between individuals given by their social network, and exploit both short and long range interactions to infer feature relationships among all individuals in the network. Methods such as Laplacian spectral decomposition and graph diffusion algorithms, which emphasize the topological relationship between nodes within the network, have
been widely used [4]. These methods rely on the existence of social or group properties such as homophily and other community structures prevalent in the network for a successful implementation.

Our main contribution in this work is to present an exclusively graph-based inference method, relying only on the topological structure of the mobile users network, together with a topological analysis of the performance of the algorithm. The equations of our algorithm can be described as a diffusion process with two added properties: (i) memory of its initial state, and (ii) the information is propagated as a probability vector for each node attribute (instead of the value of the attribute itself). We show that these two additions to the diffusion process significantly improve its inference power in the mobile network studied. Our algorithm can successfully infer different age groups within the network population given known values for a subset of nodes (the seed set). Most interestingly, we show that by carefully analyzing the topological relationships between correctly predicted nodes and the seed nodes, we can characterize particular subsets of nodes for which our inference method has significantly higher accuracy, going up to 62% (when predicting between 4 age categories). We then show that we can use the information in the probability vector of each node to again increase our predictive power (up to 72%).

The remainder of this paper is organized as follows. In section 5, we provide an overview of the datasets used in this study. We also describe the observations performed on the social graph, in particular concerning age homophily among users. In section 6, we describe our graph-based algorithm and the topological metrics used in the analysis. In section 7, we present the results of the inference power of our algorithm. We focus on understanding how the performance of the algorithm depends on age groups and the topological properties of correctly inferred nodes. We then reexamine our results on the subnetwork composed solely of the mobile operator’s clients. We further study how the model parameters and the added information contained in the probability vectors can enhance the inference power of the algorithm. Finally, in section 8, we conclude this work remarking the most significant findings and ideas for future work.

2. DATA SOURCE AND OBSERVATIONS

2.1 Dataset Description

The dataset used in this work consists of cell phone calls and SMS (short message service) records (CDRs) for a three month period collected by a mobile phone operator for the whole country of Mexico. Each record contains detailed information about the anonymized phone numbers of both the caller and the callee, whether it is an incoming or outgoing call, the duration and timestamp of the call, and the location of the antenna servicing the client involved in the communication. The detailed information provided by this dataset allows for the construction of large-scale complex dynamic networks with added node (client) mobility based on the antenna location, which can be used for the study of human mobility [10].

A milestone of this work is to study how the “bare bones” structure of the mobile phone network contains useful information for the inference of the users age group. For our purposes, we summarize each call/SMS record as a tuple \((i, j)\) representing a call/SMS between users \(i\) and \(j\). We then aggregate these tuples for all calls/SMS for a three month period into an edge list \((n_i, n_j, w_{n,i,j})\) where nodes \(n_i\) and \(n_j\) represent users \(i\) and \(j\) respectively and \(w_{n,i,j}\) is a boolean value indicating whether these two users have communicated at least once within the three month period. This edge list will represent our mobile graph \(G = (N, E)\) where \(N\) denotes the set of nodes (users) and \(E\) the set of communication links. We note that only a subset \(N_C\) nodes in \(N\) are clients of the mobile operator, the remaining nodes \(N \backslash N_C\) are users that communicated with users in \(N_C\) but themselves are not clients of the mobile operator. This distinction implies that our network \(G\) is an incomplete network in the sense that communications among members in \(N \backslash N_C\) are not observed in \(G\).

Together with the CDR records, we had access to several demographic variables for over 500,000 clients. These demographic variables include their actual age. We therefore distinguish three sets of nodes in our graph \(G\): the whole set \(N\), the set of client nodes \(N_C\), and the set of client nodes with known age \(N_{GT}\) which we use as the ground truth nodes \((N_{GT} \subset N_C \subset N)\). Lastly we split our set \(N_{GT}\) into a set of seed nodes \(N_S\) consisting of approximately 75% of \(N_{GT}\) which we use as seeds to generate age predictions for the remaining nodes in \(G\), and the remaining nodes \(N_C\) which we use as a validation set to evaluate the performance of our predictions.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{population.png}
\caption{Population pyramid.}
\end{figure}

In Fig. 1 we plot the population pyramid of the seed nodes discriminating by gender. We can see a bimodal structure for both genders with peaks centered around the ages of 30 and 45 years. This double peak in the distribution is not observed in the Mexico age population census, thus it is possibly indicative of some particular feature of the population of mobile phone clients arising from special offers such as family plans.

The last step in our data preprocessing consisted in partitioning users ages into four age groups: below 25 years, from 25 to 34 years, from 35 to 49 years and 50 years and above. The motivation for the chosen partition is grounded in marketing purposes of the mobile phone operator. The predictive algorithm will therefore infer one of these age groups for each of the nodes in \(G\).
2.2 Age Homophily

Homophily is defined as “the principle that contacts between similar people occurs at a higher rate than between dissimilar people” [11]. This basic observation about human relations can be traced back to Aristotle, who wrote that people “love those who are like themselves” [1]. In particular, age homophily in social networks has been observed in the sociological literature, for instance in the context of friendship in urban setting [8] and other social structures [7]. These historical studies were limited to hundreds of individuals, due to the difficulty and cost of gathering data. A recent study on the structure of the Facebook social graph [17] shows that a strong age homophily can be observed in a social graph with hundreds of millions of users. Our dataset allowed us to verify that the age homophily principle is valid in the mobile phone communication network on a nationwide scale (the country of Mexico in our case).

As we mentioned in the introduction, graph based methods like the one we present in this work rely strongly on the ability of the graph topology to capture correlations between node attributes we are aiming to predict. A most fundamental structure is that of correlations between nodes and their neighborhoods. Figure 2(a) shows the correlation matrix $C$ where $C_{i,j}$ is the number of links between users of age $i$ and age $j$ for the nodes in the the ground truth $N_{GT}$. Though we can observe some smaller off diagonal peaks, we can see that it is most strongly peaked along the diagonal, showing that users are much more likely to communicate with users of their same age. Performing a linear regression between the ages of linked users yields a regression coefficient $r = 0.99$ and gradient $1.06$, confirming this observation.

To account for the population density bias shown in Fig. 1, we compute a surrogate correlation matrix $R$ as the expected number of edges between ages $i$ and $j$ under the null hypothesis of independence, plotted in Fig. 2(b):

$$R_{i,j} = \frac{N_{GT}(i)}{|N_{GT}|} \times \frac{N_{GT}(j)}{|N_{GT}|} \times |E_{GT}|$$

(d) Number of links as function of age difference.

Figure 2: Age homophily plots showing the communications matrix $C$, the random links matrix $R$, the difference between $C$ and $R$, and the number of links according to the age difference.
where $N_{GT}(i) = \{x \in N_{GT} : age(x) = i\}$ and $E_{GT} = \{(x, y) \in E : x \in N_{GT} \land y \in N_{GT}\}$. This matrix represents a graph with the same nodes as the original, but with random edges (while maintaining the same number of edges as the original). Both $C$ and $R$ are represented with a logarithmic color scale. If we subtract the logarithm of $R$ to the logarithm of the original matrix $C$, we can isolate the “social effect” (i.e. homophily) from the pure random connections, as can be seen in Fig. 2(c).

Figure (2.d) summarizes the number of links as a function of the age difference $\delta$, showing a clear peak when the difference is $\delta = 0$. The number of links decreases with the age difference, except around the value $\delta = 25$, where an interesting inflection point can be observed; possibly relating to different generations (e.g. parents and children). This phenomenon can also be seen in Fig. 2(c), in the off diagonal bands at distance 25 years from the diagonal.

3. METHODS

The strong correlations shown in Fig. 2 indicate that the topological information of the network can be used to successfully infer the nodes age group (knowing the age group for a small subset of seed nodes). We describe below our algorithm that performs such task.

3.1 Reaction-Diffusion Algorithm

For each node $x$ in $G$ we define an initial state probability vector $g_{x,0} \in \mathbb{R}^C$ representing the initial probability of the nodes age belonging to one of the $C = 4$ age categories defined in section 2.1. More precisely, each component of $g_{x,0}$ is given by

$$g_{x,0} = \begin{cases} \delta_{i,a(x)} & \text{if } x \in N_S \\ 1/C & \text{if } x \notin N_S \end{cases}$$

(2)

where $\delta_{i,a(x)}$ is the Kronecker delta function and $a(x)$ the age category assigned each seed node $x$. For non seed nodes, equal probabilities are assigned to each category. These probability vectors are the set of initial conditions for the algorithm.

The evolution equations for the probability vectors $g$ are then as follows

$$g_{x,t} = (1 - \lambda) g_{x,0} + \lambda \sum_{y \sim x} w_{y,x} g_{y,t-1}$$

(3)

where $x \sim y$ is the set of $x$’s neighbours and $w_{y,x}$ is the weight of the link between nodes $x$ and $y$. For our purposes all link weights are set to one, thus the second term simplifies to a normalized average over the values of the neighborhood nodes for $x$:

$$g_{x,t} = (1 - \lambda) g_{x,0} + \lambda \sum_{y \sim x} g_{y,t-1}$$

(4)

The evolution equation is made up of two distinctive terms: the first term can be thought of as a memory term where in each iteration the node remembers its initial state $g_{x,0}$, while the second term is a mean field term of the average values of the nodes neighborhood for $x$. The model parameter $\lambda \in [0, 1]$ defines the relative importance of each of these terms. The influence of $\lambda$ on the performance of the algorithm will be analyzed in section 4.3.

We note that the evolution of each term in the probability vectors are totally decoupled from one another. This last assumption in the algorithm relies on the strong homophily suggested by the correlation plots of Fig. 2.

Next, we briefly outline our motivation for calling it a reaction-diffusion algorithm. A reaction–diffusion model explains how the concentration of one or more (typically chemical) substances changes under the influence of two processes: local reactions and diffusion across the medium.

Let $A \in \mathbb{R}^{[N] \times [N]}$ be the adjacency matrix of $G$, and $D \in \mathbb{R}^{[N] \times [N]}$ the diagonal matrix with the degree of each node in $G$, that is $A_{i,j} = w_{i,j}$ and $D_{i,i} = deg(i)$. Furthermore let $L \in \mathbb{R}^{[N] \times [N]}$ be the matrix defined as $L = D - A$. We can now rewrite equation (3) as

$$g_{x} = (1 - \lambda) g_{x,0} + \lambda L g_{x,t-1}$$

(5)

where $g^a \in \mathbb{R}^{[N]}$ now runs over all nodes, for each age probability category $a$. We are able to do this since from equation (1), the probabilities of each category evolve independently. We now rewrite the above equation using the graph Laplacian for $G$ defined as $L \equiv D^{-1/2}LD^{-1/2}[6]$

$$g_{x} = (1 - \lambda) g_{x,0} - \lambda L g_{x,t-1} + \lambda g_{x,t-1}$$

(6)

finally reordering terms we get

$$g_{x} - g_{x,t-1} = (1 - \lambda)(g_{x,0} - g_{x,t-1}) - \lambda L g_{x,t-1}$$

(7)

where on the left hand side we have the discrete derivative of $g^a$ with respect to time, the first term on the right, $(1 - \lambda)(g_{x,0} - g_{x,t-1})$, is the reactive term which drives $g^a$ towards $g_{x,0}$ and the second term $\lambda L g_{x,t-1}$ can be thought of as a diffusive term where the Laplacian operator in continuous space has been replaced by the Laplacian of the graph $G$ over which diffusion is taking place. From equation (7), we can see that $\lambda$ tunes the relative importance of the seed nodes information $g_{0}$ and the topological properties of $G$ given by the Laplacian $L$. We have therefore gone from a local description of the model in equation (3), to a global description in equation (7).

3.2 Topological Metrics

One of the main goals of this work is to uncover topological properties for a given node that increase its chances of being correctly predicted, and characterize (based on these properties) a subset of nodes in $G$ for which our algorithm works particularly well. For this, we describe three topological metrics used to characterize each node in $G$. The first metric is the number of seeds in a node’s neighborhood which we denote $SIN$ (seeds in neighborhood). We expect $SIN$ to be a relevant metric given the strong correlations shown in Fig. 2. The second metric is the topological distance of a node to the seed set which we denote $DTS$ (distance to seeds), where distance is defined as the shortest path from the node to its closest seed. The third metric is the degree $d$ of a node in the network, that is, the size of its neighborhood. Nodes with high degree are more likely to be strongly connected to the seed nodes, therefore we expect their response to the reaction diffusion algorithm, all else equal, to be significantly different than that of a node with low degree.

4. RESULTS

In this section we first present the results for the predictive power of the reaction-diffusion algorithm over the whole network $G$. To construct the mobile phone network
we start with a directed edge list consisting of 252,248,440 edges where an edge can represent outgoing communications from client \( x \) to user \( y \) and incoming communications from user \( y \) to client \( x \) distinctively. This set of edges involves 93,797,349 users. After pruning users with high degree (above 100), or not belonging to any connected component containing seed nodes, we end up with a resultant graph \( G_T \) with \(|E_T| = 125,433,585\) edges (where edges have been symmetrized) and \(|N_T| = 71,949,815\) nodes, of which \(|N_S| = 493,871\) are seed nodes and \(|N_V| = 164,768\) are validation nodes for which we know their actual age group. These are used to test the inference power of our algorithm.

As stated in equation 2, the initial conditions for each node’s probability vector \( g_{x,0} \) is that of total certainty for seed nodes and no information for the remaining nodes in \( G_T \). At each iteration, each \( g_{x,t} \) in (4) updates its state as a result of its initial state and the mean field resulting from the probability vector of its neighbors. To reduce diffusion noise, in every time step we consider the mean field generated only by the neighbors of \( x \) that have received some information from the seed nodes. After the last iteration \( t_{end} \) we assign the age group with highest probability in \( g_{x,t_{end}} \) to each node in \( G_T \). Unless stated otherwise, we set the value of \( \lambda = 0.5 \) and the number of iterations to 30 (which is 10 iterations more than the maximum distance from a node to the seed set: \( DTS_{\text{max}} = 20 \)).

4.1 Performance by Age Group

In Table 1 we present the results for the performance of the reaction diffusion algorithm discriminating by age group. The age group demographics predicted by the algorithm, both for the whole graph \( N_T \) and the validation set \( N_V \) are similar to that of the seed set \( N_S \), that is, the algorithm does a good job at preserving the age group demographics of the seed set for all nodes in \( G \). The last column shows the fraction of correctly validated nodes for each age group \( \text{hits} \) we refer to as the performance or inference power of the algorithm. The age group between 35 and 50 years is significantly better predicted than the other three age groups with a performance of 52.3%.

| Age group | \%\(N_T\) | \%\(N_S\) | \%\(N_V\) | \%\text{hits} |
|-----------|----------|----------|----------|-------------|
| <25       | 5.4%     | 8.6%     | 4.6%     | 26.6%       |
| 25-34     | 32.7%    | 27.7%    | 32.3%    | 44.6%       |
| 35-50     | 37.6%    | 35.9%    | 35.8%    | 53.6%       |
| 50+       | 24.4%    | 27.8%    | 27.3%    | 45.0%       |

Table 1: Performance by age group.

A possible explanation for the increased performance for the age group 35-50 can be found by looking at the correlation plot in Fig. 2, where we observe not only strong correlations and high population densities for this age group, but it is also less fooled by the offside peaks mostly concentrated within the age group 25-34. Again we see the importance of the correlation structure among neighboring nodes. Finally, the overall performance for the entire validation set was 46.6% and we note that a performance based on random guessing without prior information would result in an expected performance of 25%, or an expected performance of \( \sim 36\% \) if we set all nodes age group to the most probable category (35 – 50).

4.2 Performance Based on Topological Metrics

We next focus our attention on how our algorithm performs for different subsets of \( G_T \) selected according to the topological metrics described in section 3.2. The main purpose of this section is to show that, given a set of seed nodes, there are specific topological features characterizing each node in the network that allows us to, a priori, select a set of nodes with optimal expected performance. We label each node in our validation set with the three topological metrics described in section 3.2.

Figure 3 plots the performance \( \text{hits} \) of the algorithm over a set of nodes as a function of SIN (number of seeds in the nodes neighborhood). The algorithm performs worst for nodes with no seeds in the immediate neighborhood with \( \text{hits} = 41.5\% \), steadily rising as the amount of seeds increase with a performance of \( \text{hits} = 66\% \) for nodes with 4 seeds in their neighborhood. We also see that the amount of nodes decreases exponentially with the amount of seeds in their neighborhood. An interesting feature we observed is that it is the total amount of seed nodes and not the proportion of seed nodes that correlate most with the performance of the algorithm (this last observation is not shown in the figure).

In Fig. 4 we plot the population size of nodes as a function of their \( DTS \). The most frequent distance to the nearest seed is 2, and almost all nodes are at distance less than 4. This implies that after four iterations of the algorithm, the seeds information have spread to most of the nodes in \( G_T \). This figure also shows that the performance of the algorithm decreases as the distance of a node to \( N_S \) increases. Both SIN and DTS, although topological quantities themselves, are defined in terms of the distribution of the seeds set, and are not intrinsic properties the graph \( G_T \). We therefore look at a basic intrinsic topological property of a node \( x \) in \( G_T \), namely its degree \( d(x) \).

In Fig. 5 we see that the performance of the algorithm is lowest for nodes with small degree and gradually increases as the degree increases reaching a plateau for nodes with \( d(x) > 10 \).
it is here worth remarking that
g algorithm is something we will address in a future work, but
inflection region in the performance of the reaction-diffusion
Figure 5: Performance and population as function of DTS (distance to seeds set).

Figure 5: Performance and population as function of the nodes degree.

Understanding what makes degree values close to 10 an
inflection region in the performance of the reaction-diffusion
algorithm is something we will address in a future work, but
it is here worth remarking that $G_T$ is an essentially incomple-
graph, where we are missing all the communication
links among users that are not clients of the mobile operator.
An important topological consequence of this is that
whereas the average degree over the whole graph $G_T$ is 3.48,
the average degree for client nodes is 37.53.

Having shown the strong correlations between the three
topological metrics and the algorithm’s performance – which
allows us to select optimal values for each metric independ-
ently – we next examine whether we can find an optimal
combination of these metrics. Table 2 shows the perfor-
ance of our algorithm for different values of their $DTS$

| Degree | Distance to Seeds |
|--------|------------------|
| 1      | 0.415057         |
| 2      | 0.465881         |
| 3      | 0.313735         |

Table 2: Performance as function of degree and dis-

tance to seeds.

nodes at distance 2 from the seed set. Most interestingly,
we find that selecting a subset based only on the nodes de-
gree and distance to seeds, the optimal set is at distance 1
from the seed set and with degree between 29 and 48 for
which the performance of the algorithm increases to $hits =$
61.8% with 20,050 nodes in $N_V$ satisfying these set of con-
ditions. This last result indicates that it is not sufficient
to look at each metric independently to select the subset of
nodes for optimal performance of the algorithm. We also
looked at the performance of the algorithms as a function of
$SIN$ and degree, and also found subsets of nodes for whom
performance increased to similar values (table not shown for
space reasons).

4.3 Preserving Age Demographics

A salient feature of our algorithm is that the demographic
information being spread is not the age group itself, but a
probability vector for each age group. In each iteration, the
algorithm does not collapse the information in each node
to a preferred value; instead it allows the system to evolve
as a probability state over the network, and only at the
end of the simulation, when we “observe” the system, are
we required to collapse the probability vectors to specific
age groups. Choosing how to perform this collapse is not
an obvious matter, in the previous sections we selected for
each node the category with highest probability. This is a
natural approach, and as we have shown, the algorithm’s
performance is satisfactory. A draw back of this approach
is that, when collapsing the probability vector at each node,
we ignore the resultant probabilities for all other nodes in
the network which could inform us in making a better se-
lection choice. We might want to collapse the probability
state of the system as a whole and not for each node in-
dependently. For instance, if we want to impose external
constraints on our solution, namely that the age group dis-
tribution for the whole network be that of the seed set, we
can do this optimally using the following algorithm, which
we call Population Pyramid Scaling (PPS).

Note that the population to predict has size $|N_T|$. For
each age group $a$ we compute the number of nodes $N_a$ that
should be allocated to group $a$ in order to satisfy the dis-
tribution constraint (the age distribution of $N_{GT}$), and such
that $\sum_{a=1}^{C} N_a = |N_T|$ (where $C$ is the number of age groups). The
PPS procedure is described in Algorithm 1

The performance of our algorithm constraining the de-

gographic population of $G_T$ to that of the seed nodes $N_S$
is 46.5%, almost identical to 46.6% obtained without PPS.
The reason for this is that the algorithm (without PPS) was
able to preserve a distribution over the whole graph very
similar to that of the seed nodes as we showed in Table 1.
Therefore in this case, the demographic constraint has not significantly changed our results.

4.4 Analysis of the Clients Subgraph

In section 4.2 we mentioned that the graph $G_T$ has two distinct types of nodes, client and non client nodes with considerably different average degree. Motivated by this difference, and also considering the distinctive importance from a business point of view of understanding the predictive power of our algorithm on the client nodes, we performed the same study as above for the graph $G_C$ consisting only of client nodes. This graph is made up of $|E_C| = 6,484,571$ edges and $|N_C| = 3,225,538$ nodes of which 349,542 nodes are seed nodes and 143,240 nodes are validation nodes. Note the average degree of the client nodes has dropped from 37.53 in $G_T$ to 4.02 in $G_C$.

| Age | $\%N_T$ | $\%N_S$ | $\%N_V$ | $\%hits$ |
|-----|---------|---------|---------|---------|
| <25 | 5.3%    | 7.1%    | 4.6%    | 27.6%   |
| 25-34| 28.7%   | 29.9%   | 28.4%   | 50.9%   |
| 35-50| 41.7%   | 34.6%   | 37.2%   | 53.4%   |
| 50+ | 24.3%   | 28.4%   | 29.8%   | 48.0%   |

Table 3: Performance by age group on the clients subgraph $G_C$.

In Table 3 we summarize the performance of the reaction-diffusion algorithm discriminated by age group. As for the full graph $G_T$, the algorithm did a good job in preserving the age group demographics of the seed nodes over all the nodes in $G_C$, in particular for the validation nodes. The last column indicates the performance of the algorithm for each age group, where two observations are in order: first we notice that the overall performance of 49.9% is significantly better than the performance of 46.6% for the full graph $G_T$. Second the relative performance of each age group is quite similar for both graphs, plausibly indicating that it is mainly governed by the correlation structure described in Fig. 2.

Following the same analysis done for the full graph $G_T$, we looked at how the performance of the reaction-diffusion algorithm correlates to the three topological metrics. These results are shown in Figures 6 and 7. Comparing these plots with those for graph $G_T$ we can notice several topological differences between them. First we notice that while in $G_T$ only ~30% of the validation nodes have at least one seed in their neighborhood, over 65% of nodes in $G_C$ have at least one seed in their neighborhood. This is not surprising given that both validation and seed nodes are client nodes. Most interestingly, even though for both graphs the performance of the algorithm increases as their $SIN$ increases, for a fixed number of seeds the algorithm does better in the full graph than in the client graph. This is quite surprising given that the overall performance over the full graph is worse than in the client graph.

In Fig. 6 we see that $DTS$ for client nodes peaks at $DTS = 1$, falling exponentially as $DTS$ increases (whereas in the full graph the peak was observed at $DTS = 2$). Performance of the algorithm again correlates strongly with distance to the seed set.

Performance as function of the degree of the reaction-diffusion algorithm correlates to the three topological metrics. These results are shown in Figures 6 and 7. Comparing these plots with those for graph $G_T$ we can notice several topological differences between them. First we notice that while in $G_T$ only ~30% of the validation nodes have at least one seed in their neighborhood, over 65% of nodes in $G_C$ have at least one seed in their neighborhood. This is not surprising given that both validation and seed nodes are client nodes. Most interestingly, even though for both graphs the performance of the algorithm increases as their $SIN$ increases, for a fixed number of seeds the algorithm does better in the full graph than in the client graph. This is quite surprising given that the overall performance over the full graph is worse than in the client graph.

In Fig. 7 we see that $DTS$ for client nodes peaks at $DTS = 1$, falling exponentially as $DTS$ increases (whereas in the full graph the peak was observed at $DTS = 2$). Performance of the algorithm again correlates strongly with distance to the seed set.
4.5 Sensitivity to Model Parameters

So far we have analyzed the performance of the reaction-diffusion algorithm without addressing the issue of model parameters selection, namely the parameter $\lambda$ presented in section 4.1 which tunes the relative importance of the reaction and diffusion terms – together with the number of iterations of the algorithm, which determines to which extent the age information in the seed set spreads across the network. In this section we examine these two parameters for the client network $G_C$.

Parameter $\lambda$.

For $\lambda = 0.0$ (no diffusion) the performance is 28.1% but as soon as the diffusion term is slightly turned on, $\lambda = 10^{-7}$, the performance of the algorithm jumps to 48.3% and remains almost constant for $0 < \lambda < 1$ and drops suddenly to a performance of 42.5% for $\lambda = 1.0$ (total diffusion with no reactive term). We conclude from these results two things. First, the reactive term is important, increasing the performance from 42.5% (no reactive term) to 48.8% (optimally choosing the reactive term). Second, inside the boundary values for $\lambda$ the performance remains almost unchanged and thus the algorithm is very robust to parameter perturbations. In other words, our algorithm needs some diffusion and reactive terms but the relative weights of these are unimportant.

Convergence.

An important property of the model we address here is that of the model’s time scale, that is, how long it takes the system to reach a stationary state. Stationarity analysis in complex networks such as the one we study here is a hard problem, thus for the purpose of this work we limit ourselves to the notion of performance. We therefore say the algorithm reaches a stationary state after $t$ iterations if its performance remains unchanged for further iterations. We have seen that most nodes in the graph are not directly connected to a seed node, and that the diffusion term in Equation (1) spreads from neighbor to neighbor one iteration at a time. We therefore expect that as iterations increase, the seed nodes’ information will spread further across the network. Our experiments show that after a single iteration, the performance of the reaction-diffusion algorithm is already at 45.5% gradually increasing its performance to a stationary value of $\sim 50\%$ after 5 iterations. As conclusion, setting the number of iterations as $t_{\text{end}} = 30$ gives us a good margin to ensure convergence of the algorithm.

4.6 Optimal Node Subset using Probability Vector Information

The focus of our exploratory analysis to find an optimal subset of nodes (where our algorithm works best) has been centered on looking at the network topology. Here we present a conceptually different strategy which exploits the information in the probability vector for the age group on each node. Namely, we examine the performance of the reaction-diffusion algorithm as we restrict the performance to a subset of nodes whose selected category satisfies a minimal threshold value $\tau$ in its probability vector. In Fig. 8 we observe a monotonic increase in the performance as the threshold is increased and simultaneously a monotonic decrease of the validation set. We note that for $\tau = 0.5$ the performance increased to 72% with 3,492 out of the 143,240 (2.4%) of the validation nodes remaining. The performance of the algorithm increases to $\sim 81\%$ for $\tau = 0.55$ but the number of validation nodes remaining has decreased to 201 (0.1%) making this estimate very unreliable.

5. CONCLUSION AND FUTURE WORK

Large scale social networks such as those emerging from mobile phone communications are rapidly increasing in size and becoming more ubiquitous around the globe. Understanding how these community structures forming complex network topologies can inform mobile phone operators, as well as other organizations, on unknown attributes of their users is of vital importance if they are to better understand, not only their clients’ interests and behavior, but also their social environment including the users that are not clients of the mobile operator. A better description of this ecosystem can give the mobile operators a business edge in areas such as churn prediction, targeted marketing, and better client service among other benefits.

In this work we focused our attention on the bare bones topology of the mobile network, in other words, we aimed at uncovering the potential of the topology of the network itself to inform us on user attributes, in particular on user’s age groups. We proposed an algorithm where the only prior knowledge for inference is the age group of a subset of the network nodes (the seed nodes), and the network topology itself. Our algorithm achieved a performance of 46.6% for the entire network and of 53% for the age group 35–50 where pure random selections would have achieved a performance of 25%, or at most $\sim 36\%$ by labeling each node with the most frequent age group. To understand why the communication topology is allowing for this increase in performance we first studied the age correlations of users communicating between each other where we included the labeled set.

As shown in Fig. 2 we can observe a strong age homophily between callers which is being harnessed by our algorithm.

Next we searched for basic topological features to guide us in finding a subset of nodes where our algorithm performs best. Optimizing over all three metrics described in section 4.2 we found a subset of 20,050 nodes where the performance of our algorithm increased to 62%.
Given the particular nature of the mobile graph, where client nodes and non-clients nodes stand on a different footing, we tested the reaction-diffusion algorithm on the client only subgraph to examine differences in performance. Even though performance increased somewhat, relations of the topological metrics with performance behaved quite similarly and we were able to again find a sweet spot subset of nodes in $N_C$, characterized by the topological metrics, for which the performance of our algorithm increased to $\sim 67\%$.

Next we showed that the performance of the reaction-diffusion algorithm is robust to changes in the parameter $\lambda$ which governs the relative importance of the diffusion and reactive term, as long as both these terms are present. We also saw the performance reaches a stationary value after 5 iterations. We found that restricting the prediction over nodes with stronger inference given by their probability vector, we could achieve an increase of up to $72\%$ in performance for a significant subset of nodes.

In conclusion, in this work we have presented a novel algorithm that can harness the bare bones topology of mobile phone networks to infer with significant accuracy the age group of the network’s users. We have shown the importance of understanding nodes topological properties, in particular their relation to the seed nodes, in order to fine grain our expectation of correctly classifying the nodes. Though we have carried out this analysis for a specific network using a particular algorithm, we believe this approach can be useful to study graph based prediction algorithms in general.

As future work, one direction that we are investigating is to combine the graph based inference approach presented here with classical machine learning techniques based on node features [14]. We are also interested in applying our methodology to predict variables related to the users’ spending behavior. In [16] the authors show correlations between social features and spending behavior for a small population (52 individuals). We are currently tackling the problem of predicting spending behavior characteristics on the scale of millions of individuals.

Another research direction is to study the correlations between the user’s mobility and their demographic attributes. The geolocation information contained in the CDRs allows the analysis of users’ mobility patterns, e.g. their mobility motifs [15]. Recent studies have focused on the mobility patterns related to cultural events, such as sports [13] [15], where differences between genders and age groups may naturally be found. We expect that the interplay between mobility patterns and demographic attributes (such as gender and age) will provide relevant features to feed our prediction algorithms, and more generally will contribute to gain further insights on the behavior of different segments of the population.

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