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OPEN Estimating the epidemic risk using non-uniformly sampled contact data

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Many datasets describing contacts in a population suffer from incompleteness due to population sampling and underreporting of contacts. Data-driven simulations of spreading processes using such incomplete data lead to an underestimation of the epidemic risk, and it is therefore important to devise methods to correct this bias. We focus here on a non-uniform sampling of the contacts between individuals, aimed at mimicking the results of diaries or surveys, and consider as case studies two datasets collected in different contexts. We show that using surrogate data built using a method developed in the case of uniform population sampling yields an improvement with respect to the use of the sampled data but is strongly limited by the underestimation of the link density in the sampled network. We put forward a second method to build surrogate data that assumes knowledge of the density of links within one of the groups forming the population. We show that it gives very good results when the population is strongly structured, and discuss its limitations in the case of a population with a weaker group structure. These limitations highlight the interest of measurements using wearable sensors able to yield accurate information on the structure and durations of contacts.

An increasing number of studies on epidemic spreading processes use data-driven models. In particular, contact patterns between individuals are considered to play an important role in determining the possible outcome of the transmission of infectious diseases in a population¹⁻³. Many datasets describing contacts between individuals in various contexts have thus been gathered by different research groups, using techniques ranging from surveys or diaries to wearable sensors⁴⁻¹⁷. The resulting data are typically in the form of contact networks in which nodes represent individuals and edges represent the existence of at least one contact between the individuals linked.

Such network data can however be incomplete, for two main reasons. On the one hand, not all individuals agree to participate to the data collection (either not answering the surveys or not willing to wear a sensor), leading to node sampling. On the other hand, contacts between participating individuals might not all figure in the gathered data, so that links are missing from the data. In the case of diaries for instance, each individual only remembers a fraction of his/her contacts, the longest contacts being better reported 14, 18, 19. If instead the available data comes from a survey about friendship relations, it will typically miss many short encounters between people who are not friends¹⁴. Finally, even in the case of wearable sensors, some short contacts might not be detected, the actual detection of a contact might depend on its duration, depending on the sensitivity of the measuring infrastructure, and the temporal resolution may vary²⁰. It is thus of interest to understand how the resulting data incompleteness or limited resolution affects the properties of the measured contact network²⁰⁻²³, how it affects the outcome of data-driven models using incomplete data^{20, 23–25}, and most importantly if it is possible to infer the real network structure or statistical properties from incomplete information 26-28 and/or to devise methods to correctly estimate the epidemic risk even from incomplete data^{23,29} [Note that, for some types of wearable sensors, the opposite problem of false positives, i.e., of reported contacts that are not relevant for propagation events, can also arise. Here we focus on data incompleteness, but investigations of the impact of false positives would also be of clear interest]. To obtain such an estimation, a possibility is to try and construct surrogate datasets using only the information contained in the incomplete data such that these surrogate data, despite not being strictly equal to the original data, are "similar enough" with respect to the spreading process of interest. Here, "similar enough" means that the outcomes of simulations of spread using the surrogate data should be close to the ones using the real, complete data.

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This issue has been addressed in the case of uniform population sampling by Génois et al.²³. Uniform node sampling indeed maintains not only the network density, but also the whole contact matrix of densities that describes the structure of links in a population structured in distinct groups, such as classes in a school. We recall that the density of a network of N nodes and E edges is defined as the ratio of the number of edges to the maximal possible number of edges that could exist between the nodes, i.e., d = E/(N(N-1)/2). The contact matrix of densities gives for each pair of groups X and Y the ratio between the total number of links E_{XY} between the n_X individuals in X and the n_Y individuals in Y, and the maximum number of such possible links ($n_X n_Y$ if $X \neq Y$ or $n_X(n_X-1)/2$ if X=Y). Moreover, the sampling also maintains the temporal statistics of contacts. It is thus possible to measure this contact matrix and the temporal statistics in the incomplete data and to construct surrogate data having the same statistics as the original one. Spreading processes simulated using such surrogate data have been shown to reproduce well the outcome of simulations using the whole dataset²³. A similar method has been shown to work well also in a case study of contact diaries collected together with data from wearable sensors²⁹: although not all contacts were reported in the diaries, building surrogate data using the contact matrix measured in the diaries and publicly available statistics on contact durations made it possible to correctly estimate the outcome of simulations of spreading processes. In these two studies, the density of the sampled data was however either equal (for uniform population sampling) or close (for the diaries) to the one of the original data. In other cases, such as e.g. the friendship survey of ref. 14, the density of the data is much smaller than the one of the contact data, and the method has indeed been shown to fail in this case²⁹. Sampled data with smaller density than the original one occur as soon as the links between sampled nodes are not all present. This is expected to be the case in particular for data coming from diaries or surveys, and it is of interest to test the limits of the reconstruction method, as a function of the sampling properties, and possibly to understand how to overcome this obstacle.

Here, we tackle this issue by considering incomplete contact data stemming from a non-uniform sampling procedure intended to mimic a survey procedure in which (i) not all individuals participate and (ii) the contacts of each respondent are reported with a probability depending on their duration³⁰. We consider empirical contact datasets, resample them using this non-uniform sampling procedure and design surrogate data as described above. We show that, at low sampling, the use of such surrogate data in simulations of spreading processes is not enough to estimate the epidemic risk, even if it yields an improvement with respect to the use of the raw sampled data. We thus consider the case in which additional information is available for one of the groups forming the population: if one group is uniformly sampled (for instance if wearable sensors are available for this group), yielding a good estimate of its density, it is possible through a simple rescaling procedure to estimate a new contact matrix for the data and to use it to construct another surrogate dataset that yields better results. By using two datasets with different structures and varying the parameters of sampling and spreading processes, we explore the efficiency and limits of both procedures.

Data and Methodology

We first describe the datasets and the different steps of our methodology, which are in the same spirit as refs 23 and 29. Starting from a network of contacts between individuals in a population, we perform a specific resampling that leads to an incomplete dataset. We describe two methods to create surrogate datasets using the statistical information contained in the incomplete data. Spreading processes are then simulated on top of the incomplete and of the surrogate datasets, and their outcomes compared with the ones of simulations using the whole original contact network. The procedure is summarized in Fig. 1.

Data. We will use two datasets describing face-to-face contacts between individuals, collected and made publicly available by the SocioPatterns collaboration (see the SocioPatterns website http://www.sociopatterns.org/). The first dataset (Thiers13) has been collected in a French high school in December 2013. The resulting contact network has N = 327 nodes representing students (divided in 9 classes corresponding to different fields of study) and E = 5818 weighted edges (see ref. 14 for a detailed description and analysis of this dataset), for an average link density of $2E/(N(N-1)) \approx 0.11$. The participation rate reached 86.5% (there were overall 378 students in the 9 classes). The classes are of similar sizes (see Table 1), and most edges (69%, accounting for 93% of the weights, i.e., of the total contact time between students) are found within classes (Fig. 2). The second dataset (InVS) has been collected in the office buildings of the Institut de Veille Sanitaire (French Institute for Public Health Surveillance) in March 2015, and describes the contacts between 217 individuals divided in 12 departments. The contact network has 4274 edges, i.e., a density of ≈ 0.18 . The departments have very different sizes and participation rate, with an average participation rate of 60% (see Table 2) and, while most contact time occurs within departments (76%), the corresponding fraction of edges is only 42% (Fig. 2). In each dataset, an edge between two individuals corresponds to the fact that these individuals have been in contact at least once during the data collection, and the edge weight gives the total contact time between them. The contact matrices of edge densities are shown in Fig. 2 for both datasets. In the Thiers13 case, we have moreover access to a network describing friendship relations between students, obtained by a survey to which 135 of the 327 students answered. This friendship network has 413 unweighted edges.

Sampling method. We consider the sampling method put forward in ref. 30, called "EGOref", based on (i) a uniform sampling of nodes and (ii) a non-uniform sampling of edges, edges with larger weights being preferentially sampled. This procedure is designed to mimic a sampling of links obtained for instance through a survey on friendship relations in a population: it is inspired by the result of ref. 14 that the longest contacts measured in the Thiers13 dataset corresponded to reported friendships, while many short contacts did not. In particular, we have shown in ref. 30 that the outcome of simulations of spreading processes on the friendship network of the Thiers13 dataset can be reproduced if the EGOref sampling method is applied to the contact network of the same dataset, with correctly adjusted sampling parameters.

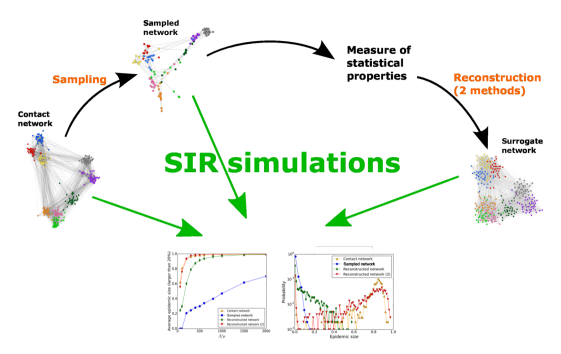


Figure 1. Sketch of the procedures considered in the article. We consider a dataset describing a contact network. We perform a resampling according to a certain (non-uniform) sampling method. We then measure a number of statistics of the resampled data and use these statistics to construct surrogate data. We simulate spreading processes on the original, resampled and surrogate data and compare the outcomes, as given by the fraction of large epidemics and by the whole distribution of epidemic sizes.

Classes	Number of individuals	% of the total population	Participation rate	Number of participants to the friendship survey
2BIO1	36	11%	92%	10
2BIO2	34	10.4%	81%	20
2BIO3	40	12.2%	98%	28
PC	44	13.5%	98%	21
PC*	39	12%	95%	10
PSI*	34	10.4%	77%	15
MP	33	10%	79%	21
MP*1	29	8.9%	69%	3
MP*2	38	11.6%	90%	7
Total	327	100%	86%	135

Table 1. Number of individuals in each class participating to the data collection in the highschool (Thiers13 dataset), percentage with respect to the population under study and participation rates.

More precisely, the EGOref sampling depends indeed on two parameters. Starting from a weighted contact network with N_0 nodes, we select N of these nodes (called "egos") uniformly at random (a non-uniform selection could also be considered). For each ego i, each edge i-j is selected with a probability equal to $p\frac{W_{ij}}{S_i}$, with W_{ij} the weight of the edge between i and j, $S_i = \sum_{\ell} W_{i\ell}$ the strength of the ego node i and p the sampling parameter. We then keep only the egos and the selected edges linking them and we remove the other edges (between egos and non-egos and between non-egos) and nodes (non-egos). With this method, we end up with a tunable number of nodes N and a number of edges that depends on the parameter p. Figure 3 summarizes this process.

The parameter p clearly has an effect on the density of the sampled network. This is in contrast with the case of a uniform population sampling in which all the edges between sampled nodes are kept, and which thus conserves the network density²³. Figure 4 displays the ratio between the density of the EGOref sampled network and the original contact network for the two datasets used here: the density of the sampled network increases with the parameter p. On the other hand, the density of the sampled network does not depend on the number of sampled nodes (Fig. S1 in the Supplementary Information).

Despite the decrease in network density caused by the EGOref sampling, it is worth noting that the contact matrices of edge densities measured in the original contact network and in the EGOref sampled network remain very similar for both datasets (see Figs S2–S4 in the Supplementary Information). These matrices give, for each pair of groups in the population (here, classes or departments), the number of links between these groups

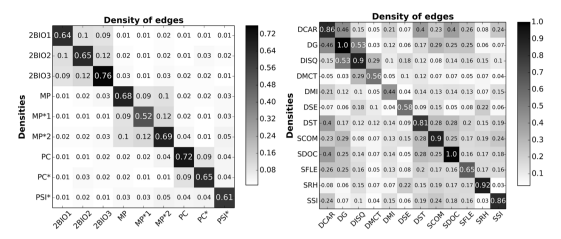


Figure 2. Density contact matrices. Left: contact matrix giving the density of edges between classes during the study (Thiers13), right: contact matrix giving the density of edges between departments during the study (InVS). For each matrix, the entry at row X and column Y is given by the total number of links between individuals in class or department X and individuals in class or department Y, normalized by the maximum number of observable links $(n_X n_Y \text{ or } n_X (n_X - 1)/2 \text{ if } X = Y$, with n_X the cardinality of X). These matrices give only structural information as they do not take into account edge weights.

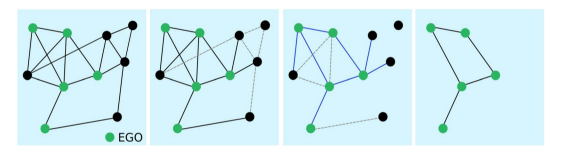


Figure 3. Sketch of the EGOref sampling process. We first select a certain number of nodes as egos. Each ego "chooses" to report some of its links, with probability depending on their weights. A link can be selected twice if it joins two *egos* (blue edges). We then finally keep only the egos and, among the chosen edges, only the ones joining *egos*.

Departments	Number of individuals	% of the total population	Participation rate
DCAR	13	6%	65%
DG	2	0.9%	22%
DISQ	18	8.3%	86%
DMCT	31	14.3%	63%
DMI	57	26.3%	79%
DSE	32	14.7%	56%
DST	23	10.6%	52%
SCOM	7	3.2%	70%
SDOC	4	1.8%	57%
SFLE	14	6.5%	37%
SRH	9	4.2%	64%
SSI	7	3.2%	32%
Total	217	100%	60%

Table 2. Number of individuals in each department participating to the data collection in the office buildings of InVS, percentage with respect to the population under study and participation rate.

normalized by the maximum possible number of such links (obtained if each member of one group is linked to all members of the other group). A large similarity between contact matrices indicates that the overall structure of the network is preserved by the sampling, even if the specific values of densities are changed.

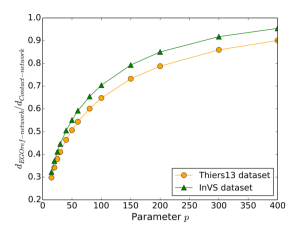


Figure 4. Impact of sampling on network density. Ratio between the density of the EGOref sampled network and the density of the whole contact network as a function of the parameter p for the two datasets. Here the number N of sampled individuals is 70% of the total population: changing this number does not change the ratio.

Surrogate data construction methods. As in refs 23 and 29, our initial goal is to use only the information contained in the sampled data to construct surrogate contact networks that are statistically similar to the full data. As already made clear in the introduction and in refs 23 and 29, we emphasize again that the point is not to infer the missing links but to build a "plausible" version of these links, such that the simulations of epidemic spread on the resulting network, as described below, yield an accurate estimation of the epidemic risk. We consider two distinct methods to construct such surrogate data.

The first method is the equivalent for static networks of the one presented in refs 23 and 29. First, the contact matrix of edge densities is measured in the incomplete data. Assuming that the number of missing nodes in each group (class or department) is known, we add the missing nodes in each and we add links randomly in each group and between groups in such a way to keep the contact matrix fixed to its measured value. More precisely, we first measure the density d in the incomplete data as 2E/(N(N-1)) where N and E are the numbers of nodes and edges in these incomplete data. Knowing the number n of missing nodes, we can deduce the number e of additional links needed to keep the density constant when we add the *n* missing nodes, through d = 2(E + e)/((N + n))(N+n-1)). We moreover transform the contact matrix of edge densities ρ_{XY} into a row-normalized contact matrix C, in which the element $C_{XY} = \rho_{XY}/\sum_{Z}\rho_{XZ}$ gives the probability for a node of group X to have a link to a node of group Y. Then, for each missing edge, we proceed as follows: (i) we extract at random a node i (among the total population of N + n nodes); (ii) knowing the group X that i belongs to, we extract at random a target group Ywith probability given by C_{XY} ; (iii) we draw at random a node j in group Y such that i and j are not yet linked, and we add a link between i and j. The resulting network has the same number of nodes as the original network and the same contact matrix of densities and overall density as the sampled network. Weights are finally assigned to the edges: weights are taken at random from the empirical distribution of weights of the contact network, which is known to be a robust feature of human contact patterns and does not depend on the context¹⁵. This method has been shown to yield good results when the incomplete data results from a uniform population sampling of the original contact network²³, which preserves the overall network density. In ref. 29, it has also been shown to be able to create relevant surrogate data (i.e., yielding the same outcome as the original data when used in simulations of spreading processes) from contact diaries data. Also in this latter case, the density of the network deduced from diaries was similar to the one of the original contact data. In ref. 29 moreover, it has been shown that one can use a pool of publicly available contact duration statistics to assign weights to the edges of the surrogate data.

However, as the EGOref sampling method yields sampled network with densities smaller than the original data (Fig. 4) and as it is well known that density plays an important role in determining the outcome of a spreading process, we propose a second construction method, in which we assume in addition that the data includes the density of edges within one of the groups (chosen at random) in the original non-sampled network. We therefore first measure the contact matrix of edge densities ρ_{XY} in the sampled data, as in the first method. We then compute the ratio f between the real density $\widetilde{\rho}_{AA}$ in the original data for the group A that is assumed to be known and the measured density ρ_{AA} . We then rescale the average density of the graph by the factor f, $d' = f \times d$ and compute the new number e' of edges to be added through d' = 2(E + e')/((N+n)(N+n-1)). The procedure is then the same as the previous one: we add the missing nodes and e' links in order to preserve the normalized contact matrix C_{XY} , and weights are taken from the empirical distribution of aggregated contact durations and assigned at random to edges.

The rationale behind the second method is that the infrastructure of wearable sensors able to measure this density could have been available for only one group in the population (or even a random fraction of one group), for instance, while only partial information from a survey is available for the other groups. We then assume that the EGOref sampling method affects in a similar way all parts of the graph, so that a global rescaling of the

	N	E	d	Avg clustering	Avg shortest path
Contact network	327	5818	0.11	0.503	2.15
EGOref network	135	405	0.04	0.351	3.95
Reconstructed network 1	327	2386	0.04	0.263	3.12
Reconstructed network 2	327	4705	0.09	0.518	2.69

Table 3. Thiers13 dataset: Features of the original contact network, of the sampled one and of the surrogate data, for an EGOref sampling with p = 30 and N = 135.

density, which also rescales all the elements of the contact matrix of edge densities by the same factor, should yield values closer to the original data.

In the following, we will apply each method to contact networks sampled using the EGOref method for various values of the parameters p and N. As ref. 30 has shown the similarity between the friendship network of the Thiers13 dataset and the outcome of the EGOref sampling applied to the Thiers13 contact network, for a specific parameter value, we will also apply these methods to construct surrogate contact data using the friendship network instead of a sampled version of the original contact network.

Simulation of spreading processes. Our goal is to understand if it is possible to use incomplete datasets to estimate the epidemic risk in a population by constructing surrogate data and using them in the simulations of epidemic spread. As a paradigm of epidemic process, we consider the Susceptible-Infectious-Recovered (SIR) model: in this model, nodes are initially all susceptible (S), except one in the Infectious state, chosen at random and seed of the process. Each Susceptible (S) node i can become infectious when in contact with an Infectious one j. This occurs at a rate $\beta W_{ij}/T$ where W_{ij} is the weight of the link i-j and T the total measurement time^{30, 31} (i.e., the probability for i to become infectious during a time step dt is $\beta W_{ij}dt/T$). Infectious nodes become Recovered (R) at rate μ and cannot be infected anymore. The process ends when there are no Infectious nodes any more.

We perform numerical simulations of this model for each dataset on the original contact network, on the sampled networks at various values of the parameters p and N, and on the surrogate datasets built using the two methods described above (note that we will equivalently write "surrogate data" or "reconstructed networks" to describe the surrogate datasets). For the Thiers13 case, we also perform simulations on the friendship networks and the corresponding surrogate data. We also vary the ratio β/μ that modulates, in each given network, the impact of the modelled disease.

To quantify the epidemic risk, we measure in each simulation the epidemic size as given by the final fraction of recovered nodes. We compare the distributions of epidemic sizes, the fraction of epidemics with size larger than 20% and the average size of these epidemics (the cut-off of 20% is chosen arbitrarily to distinguish between small and large epidemics; changing the value of this threshold does not alter our results).

We finally note that we consider here static versions of the contact networks, while the original SocioPatterns data provides temporally resolved contacts. The EGOref sampling process indeed mimics a procedure yielding a static sample of the actual contact network. In the context of models of infectious diseases with realistic timescales of several days, this can represent enough information to obtain an estimate of the epidemic risk, as discussed in ref. 31. For faster spread, one could add to the surrogate construction method an additional step of building realistic contact timelines as in ref. 23.

Results

We discuss separately the results obtained with the two datasets. The Thiers13 one is indeed much more strongly "structured" than the InVS one, in the sense that the fraction of interactions occurring within each class is very high. Moreover, all classes are of similar sizes and have similar link densities. Classes are also arranged in groups of 2 or 3 classes corresponding to the major topic of study of their students. In the InVS case, departments are of different sizes, their link densities vary more and the pattern of interactions between departments is less structured.

First case: highly structured network (Thiers13 dataset). Table 3 gives some basic features of the original, the EGOref sampled and the surrogate networks built with the two methods described above, for the EGOref parameter values yielding number of nodes and edges similar to the friendship network. This corresponds to N=135 and p=30. The sampled network has a much lower density than the original contact data, smaller clustering and larger average shortest path. The surrogate network built using the first method has by construction the same density, and has an even smaller clustering, while the second method yields values much closer to the original ones. Moreover, the similarities between the contact matrices of the original, sampled and surrogate data all exceed 98% (see also Fig. S3 of the Supplementary Information). However, the fraction of intra-class links is larger in the reconstructed network than in the original one (83% vs 69%, see Supplementary Information), while the fraction of contact durations these intra-class links carry is slightly smaller (83% vs 93%; note that since weights are put at random, the fraction of links and the fraction of weights they carry are the same in the reconstructed data).

Figures 5 and 6 compare the outcome of SIR simulations performed on the original contact network, on the EGOref sampled networks and on the surrogate data built using the two reconstruction methods, for various values of the sampling parameters. Figure 5 first displays the average size of large epidemics (i.e., the ones reaching at least 20% of the population) as a function of the spreading parameter β/μ . As expected and already explored³⁰, simulations performed on the EGOref sampled network yield a strong underestimation of the epidemic risk with

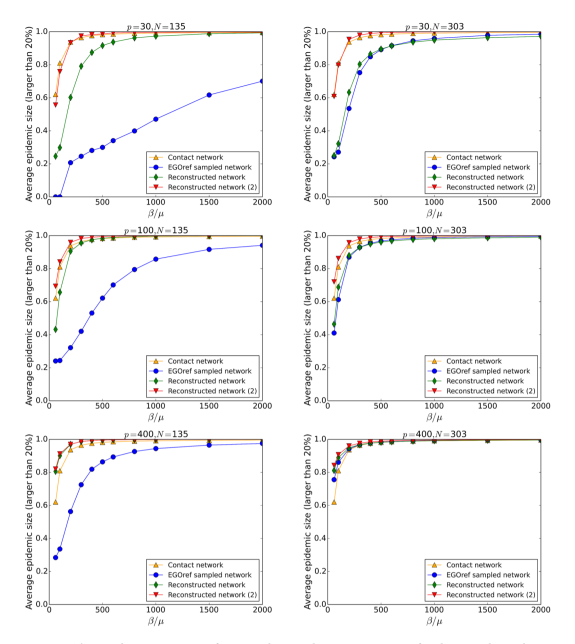


Figure 5. Thiers13 dataset: Outcomes of SIR spreading simulations. Average size of epidemics with size above 20% as a function of the spreading parameter β/μ for different values of p and N. The simulations are performed on contact network, EGOref sampled network and the reconstructed networks using the two methods of reconstruction described in the text.

respect to results obtained with the use of the contact network, except at large p and N (in this case, the sampled network is almost equal to the original one, and the random assignment of weights to the links leads in fact to a slight overestimation of the epidemic risk³⁰; this occurs only at unrealistically large values of p).

The use of surrogate data obtained with the first method improves the estimation of the epidemic risk with respect to the use of the sampled data but still leads to a clear underestimation for small and intermediate values of p. This is not unexpected given the reconstruction method maintains the density of the sampled network. The second method, which leads to surrogate data with densities closer to the original one, allows to obtain a much better estimation of the epidemic risk.

Figure 6 focuses on the case of N=135 and displays the whole distributions of epidemic sizes obtained from SIR simulations for different values of the spreading parameter β/μ and of the parameter of sampling p. In all cases, the distributions obtained with the sampled network remain narrow and do not develop a peak at large values of epidemic sizes. The distributions obtained with surrogate data are both broader but can differ strongly from each other depending on the value of p. At very large p, the sampling procedure almost does not affect the network density, so that both methods yield very close outcomes; as discussed above, the random assignment of weights leads then to a peak at large values of the epidemic size that is shifted to values larger than with the

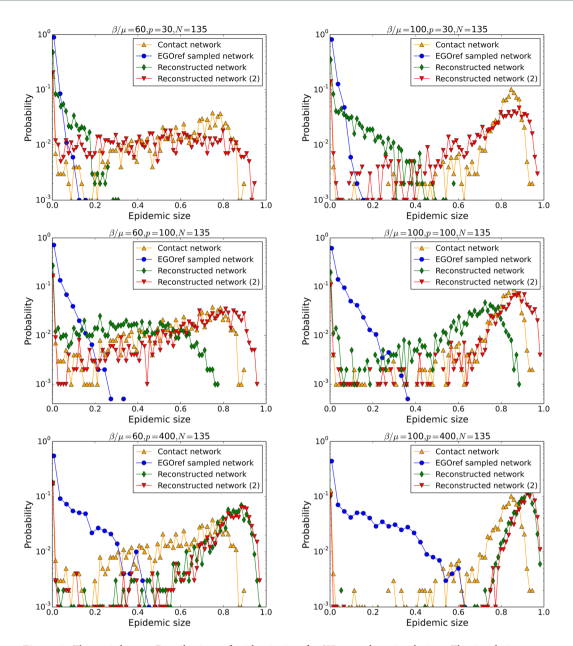


Figure 6. Thiers13 dataset: Distributions of epidemic sizes for SIR spreading simulations. The simulations are performed on the contact network on the EGOref sampled network and on the surrogate data (reconstructed networks) built using the two methods of reconstruction described in the text. The parameter of spreading β/μ , the parameter p and the number of sampled individuals N are given above each plot.

original network. For more realistic small and intermediate values of *p*, the first method leads to distributions that are much narrower than the outcome of simulations on the original network, while the second method yields a much better agreement, albeit with a systematic small overestimation of the largest epidemic sizes.

We now turn to the case of the friendship network. Table 4 compares the main features of the contact network, the friendship network and the surrogate data obtained by the second method of reconstruction applied to the friendship network. The reconstruction procedure allows to recover a density similar to the one of the contact data. Moreover, the contact matrix of these three networks are very similar (similarity values of more than 98%). We however note that the fraction of within-classes links is larger in the friendship network (75%) than in the contact network (69%), and that this characteristics holds also for the surrogate data.

Figures 7 and 8 show the outcomes of epidemic spreading simulations performed on the empirical networks and on the reconstructed network. While the simulations using the friendship network leads to a strong under-estimation of the epidemic risk, Fig. 7 shows that the use of the surrogate data yields a very good estimation of the fraction of epidemics with size above 20% and of the average epidemic size, across a large range of values of β/μ . Figure 8 displays the whole distributions of epidemic sizes for different values of the spreading parameter β/μ . It confirms that the surrogate data yields distributions with more similar shapes to the contact network case than the friendship network. However, the maximal sizes of epidemics are systematically overestimated, which

	N	E	d	Avg clustering	Avg shortest path
Contact network	327	5818	0.11	0.503	2.15
Friendship network	135	413	0.05	0.532	4.06
Reconstructed network	327	5376	0.10	0.464	2.46

Table 4. Thiers13 dataset: Basic features of the empirical networks and of the reconstructed network (obtained by applying the second method of construction of surrogate data to the friendship network).

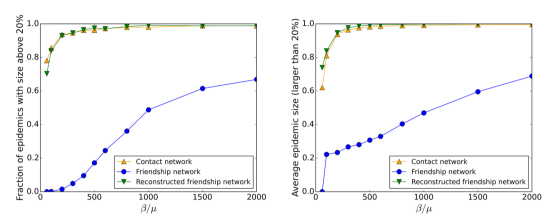


Figure 7. Thiers13 dataset: Outcome of SIR spreading simulations. Fraction of epidemics with size above 20% as a function of the spreading parameter β/μ (left) and average size of epidemic with size above 20% as a function of the spreading parameter β/μ (right). The simulations are performed on the contact network, on the friendship network and on surrogate data obtained by applying the second method of construction to the friendship network.

might be ascribed to the larger fraction of weights on inter-class links in the surrogate data with respect to the contact network.

Second case: a **less structured network (InVS dataset).** We now present the results obtained with the InVS dataset. As discussed above, the structuration in departments leads to a less structured contact matrix than in the highschool case. Table 5 and Fig. S4 of the Supplementary Information compare some characteristics of the sampled and surrogate data (with the second method of reconstruction) to the original network for p = 30 and N = 93 (i.e., a similar fraction of the population as used in the example of the Thiers13 dataset). Here, even the second method of reconstruction leads to a network density smaller than the original one, even if much closer than for the sampled data. The contact matrices of the sampled and of the surrogate data are very similar to the one of the original data (Fig. S4 of the Supplementary Information), but, as in the previous case, the fraction of intra-department edges is larger in the surrogate data than in the original one (58% versus 42%) while the fraction of the weights these links carry is much smaller (58% versus 76% in the original contact data).

Figures 9 and 10 compare the outcomes of SIR simulations performed on the contact network, the sampled and the reconstructed networks, for various values of the sampling and spreading parameters. As for the Thiers13 case, the simulations on the sampled networks strongly underestimate the epidemic risk, except obviously at large p and N. The use of surrogate data generally improves the estimation of the epidemic risk, except at large N for the first reconstruction method, as in this case almost no nodes have to be added so the network is almost unchanged by this method. For small and intermediate p, the second method gives better estimations of the average epidemic size while, at very large p, the effect of assigning randomly weights leads as before to a slight overestimation for reconstructed networks. At small β/μ and p, the second method of reconstruction can even overestimate this size substantially.

Figure 10 sheds some more light by displaying the whole distributions of epidemic sizes for several values of the spreading parameter β/μ and of the sampling parameter p, for a rather small N of the order of 40% of the total population. While the distributions obtained with the second reconstructed networks are generally closer to the ones obtained with the original contact data than with the EGOref sampled network (which always leads to a very strong underestimation) or the first reconstruction method, a peak at very large epidemic sizes is observed for the reconstructed data, which is not present for the original contact data. The maximal sizes of epidemics is thus overestimated. As in the Thiers13 case, this effect is likely to be due to the fact that the amount of weights on inter-departments edges is larger in the surrogate data than in the original data, allowing for a more efficient spread across the whole population.

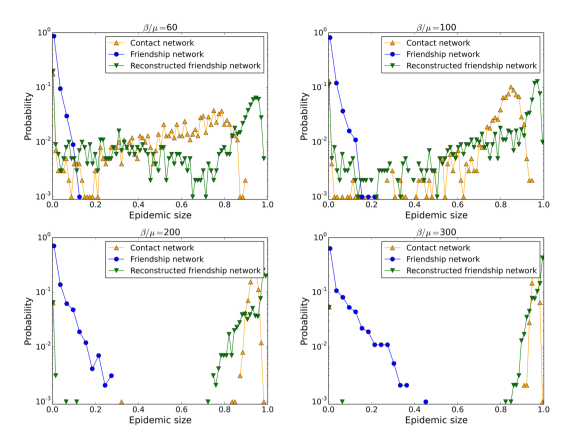


Figure 8. Thiers13 dataset: Distributions of epidemic sizes of SIR spreading simulations. The simulations are performed on the contact network, on the friendship network and on surrogate data obtained by applying the second method of construction to the friendship network. The value of β/μ used is given above each plot.

	N	E	d	Avg clustering	Avg shortest path
Contact network	217	4274	0.18	0.38	1.88
EGOref network	93	348	0.08	0.27	2.64
Reconstructed network	215	3048	0.13	0.32	2.07

Table 5. InVS dataset: Basic features of the contact network, of the EGOref sampled network (here with p = 30 and N = 40% of the total number of nodes) and of the surrogate data obtained using the second method of reconstruction.

Discussion

This paper positions itself in the context of the issue of data incompleteness in contact networks. More specifically, since many datasets are de facto incomplete, it is important to assess how data incompleteness affects the outcome of data-driven simulations, how the resulting biases can be compensated, and how much data is needed for the simulations^{23, 29–32}. We have here considered the case of non-uniformly sampled contact data and focused on a sampling procedure designed to mimic data resulting from surveys or diaries³⁰. This sampling procedure results in both population sampling, as not all individuals in the population are respondents, and in non-uniform link sampling, to mimic the fact that longer contacts have a larger probability to be remembered or to correspond also to friendship links.

We have applied this sampling procedure, called EGOref, on two datasets of contact networks in two different contexts and varied its two parameters, which determine the population participation rate and the fraction of sampled links. The datasets concern populations structured in groups with very different mixing patterns: in a high school, the class structure strongly determines contacts, with more than 90% of the duration of contacts occurring within classes; in office buildings on the other hand, the population is divided into departments but the impact on the contacts is less strong. As expected from previous investigations, using sampled data to run simulations of spreading processes leads to a strong underestimation of the epidemic risk, as quantified by the distribution of epidemic sizes. We have therefore considered the issue of building surrogate data from the sampled data, such that simulations using the surrogate data yield a better estimation of the epidemic risk. The first method we envisioned has been shown to yield good results in the context of uniform population sampling. It is based on the fact that the uniform sampling keeps invariant the contact matrix giving the densities of links between groups in the population. We have shown that the resulting surrogate data, when built from sampled data using the

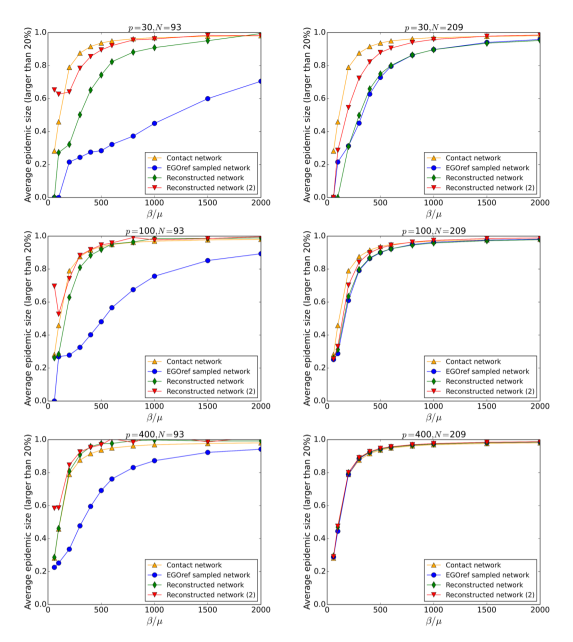


Figure 9. InVS dataset: Outcome of SIR spreading simulations. Average size of epidemics with size above 20% as a function of β/μ for several values of p and N. The simulations are performed on the contact network, on the EGOref sampled network and on the surrogate data obtained using two different methods of reconstruction.

EGOref procedure, yields better estimations than the raw sampled data, but still yields a largely underestimated risk, since the link sampling of the EGOref procedure leads to a sampled network with a (possibly much) lower density than the original one, and the method of ref. 23 does not compensate for this bias. This implies that more information is needed on the data than just the contact matrix of densities measured in the sampled data. One of the simplest way to add information to the sampled data is to assume that the link density of one of the groups of the population is known: this can occur for instance if a measurement of contacts using wearable sensors is feasible only for a small subset of the population, for practical reasons, so that it is possible to correctly measure the link density for that group. We therefore considered a second method of construction of surrogate data, in which we first rescaled all the elements of the contact matrix measured in the sampled data by the ratio between the known and measured densities of the "known" group. The resulting surrogate contact network has thus a density closer to the original one.

In the case of a strongly structured population, such as the high school, we have obtained a strong improvement of the results and a good estimation of the epidemic risk, for a large range of realistic parameters, even if the maximum size of epidemics is slightly overestimated. We have also shown that this method gives good results when applied on the data obtained from a survey asking students about their friendship relations. In the case of the less-structured population (offices), we obtained a clear improvement of the epidemic risk estimation, but the probability of very large epidemics becomes strongly overestimated when using the

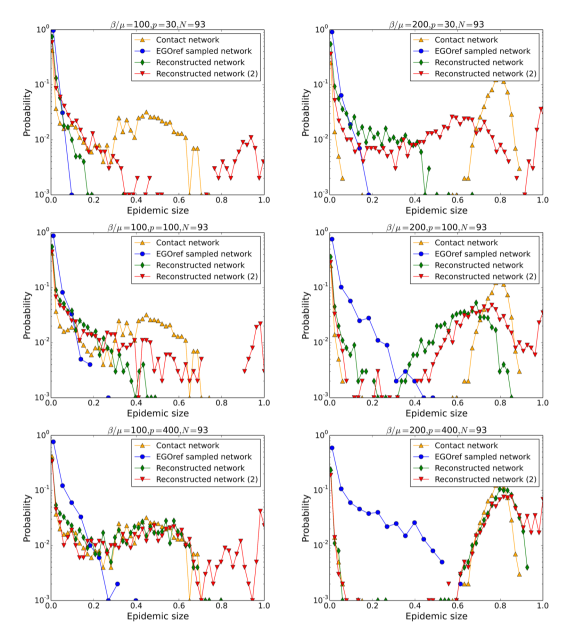


Figure 10. InVS dataset: Distributions of epidemic sizes of SIR spreading simulations. The parameter of spreading β/μ , the parameter p and the number of sampled individuals N are given above each figure. The simulations are performed on the contact network, on the EGOref sampled network and on the reconstructed networks using two different methods of reconstruction.

surrogate data. We have linked this overestimation with the fact that the total amount of weights carried by the inter-class or inter-department edges is larger in the surrogate than in the original data. Indeed, as shown in ref. 23, intra-class and intra-department links tend to carry larger weights than inter-class and inter-department ones. As weights are distributed randomly in the surrogate data, without taking this into account, the reconstruction tends to attribute more weight to the inter-groups edges with respect to the original data, which favors the spread.

Our results are overall two-sided. On the one hand, it is remarkable that, using very little information, namely the contact matrix of the sampled data and if possible the knowledge of the density of one single group, using surrogate data instead of the raw sampled data leads to a strong improvement of the epidemic risk estimation as quantified by the fraction of large epidemics and the average size of these epidemics. On the other hand, the lack of information about the precise values of the densities of links between pairs of groups, about the relative weights of intra- and inter-groups edges, as well as about the potential existence of small cohesive substructures, may lead, when the surrogate data is used, to distributions of epidemic sizes differing from the original ones, with for instance the over-estimation of the largest epidemic sizes and the presence of a peak at very large epidemic sizes. This shows both that survey data can be effectively used to construct surrogate data, but also that more detailed information coming from data collection with wearable sensors is of enormous value, even if such collection

concerns only a fraction of the population, as it allows (i) to have a correct estimation of the overall density, (ii) to obtain the distribution of contact durations, (iii) if enough sensors are available, to obtain a much better picture of the contact matrix and of the fraction of weights corresponding to intra- and inter-groups contacts, all elements having a role in the unfolding of spreading processes in a population.

Our study contributes to the discussion on the amount of details actually needed in contact data to be used in data-driven models. It is worth noting some of its limitations. We have focused here on one specific sampling model, while others might be of interest. We argue that this procedure is particularly relevant as it mimics surveys or diaries, as described in ref. 30. The procedure also assumes a uniform node sampling, while positive or negative correlations with actual contact activity might exist. Additional investigations concerning such non-uniform population sampling would certainly be of interest, as well as studies on other datasets or on synthetic populations with tuneable characteristics. Finally, the effect of using sampled or surrogate data for data-driven simulations of other types of processes ought to be investigated. Preliminary simulations of the Susceptible-Infectious-Susceptible model (SIS), in which individuals who recover become again susceptible and can catch again the disease, show that results similar to the SIR case are obtained (not shown): the sampling leads to a strong underestimation of the epidemic risk that is compensated only partially by the first method of reconstruction; the second method gives excellent results for the very structured dataset but still leads to an underestimation of the epidemic risk for the offices dataset. The issue of the performance of using surrogate data in more complex processes such as complex contagion remains open for future investigations.

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Author Contributions

A.B. and J.F. conceived and designed the study. J.F. performed the numerical simulations and the statistical analysis. A.B. and J.F. wrote the manuscript.

Additional Information

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