

pared to this baseline, our approach is distinctive in that it handles multiple files, models distortion explicitly, offers a Bayesian treatment of uncertainty and error propagation, and employs a sophisticated graphical data structure for inference to latent individuals. Fellegi-Sunter methods based upon [5] can extend to $k > 2$ files [14], but they break down for even moderately large k or complex data sets. Moreover, they give little information about uncertainty in matches, or about the true values of noise-distorted records. The idea of modeling the distortion process originates with the “Hit-Miss Model” by [3], which anticipates part of our model in §2.1. The specific distortion model we use is however closer to that introduced in [7], as part of a nonparametric frequentist technique for matching $k = 2$ files. We differ from [7] by introducing latent individuals and distortion through a Bayesian model.

Within the Bayesian paradigm, most work has focused on specialized problems related to linking two files, which propagate uncertainty [1, 6, 12, 15]. These contributions, while valuable, do not easily generalize to multiple files and duplicate detection.

Two recent papers [4, 6] are most relevant to the novelty of our work, namely the linkage structure. To aid recovering information about the population from distorted records, [6] called for developing “more sophisticated network data structures.” Our linkage graphs are such a data structure with the added benefit of permitting de-duplication and handling multiple files. Moreover, due to exact error propagation, our methods are also easily integrated with other analytic procedures. Algorithmically, the closest approach to our linkage structure is the graphical representation in [4], for de-duplication within one file. Their representation is a unipartite graph, where records are linked to each other. Our use of a bipartite graph with latents individuals naturally fits in the Bayesian paradigm along with distortion. Our method is the first to handle record linkage and de-duplication, while also modeling distortion and running in linear time.

2 Notation, Assumptions, and Linkage Structure

We begin by defining some notation, where we have k files or lists. For simplicity, we assume that all files contain the same p fields, which are all categorical, field ℓ having M_ℓ levels. We also assume that every record is complete. (Handling missing-at-random fields within records is a minor extension within the Bayesian framework.) Let \mathbf{x}_{ij} be the data for the j th record in file i , where $i = 1, \dots, k$, $j = 1, \dots, n_i$, and n_i is the number of records in file i ; \mathbf{x}_{ij} is a categorical vector of length p . Let $\mathbf{y}_{j'}$ be the latent vector of true

field values for the j' th individual in the population (or rather aggregate sample), where $j' = 1, \dots, N$, N being the total number of *observed* individuals from the population. N could be as small as 1 if every record in every file refers to the same individual or as large as $N_{\max} \equiv \sum_{i=1}^k n_i$ if no datasets share any individuals.

Now define the linkage structure $\mathbf{\Lambda} = \{\lambda_{ij} ; i = 1, \dots, k ; j = 1, \dots, n_i\}$ where λ_{ij} is an integer from 1 to N_{\max} indicating which latent individual the j th record in file i refers to, i.e., \mathbf{x}_{ij} is a possibly-distorted measurement of $\mathbf{y}_{\lambda_{ij}}$. Finally, $z_{ij\ell}$ is 1 or 0 according to whether or not a particular field ℓ is distorted in \mathbf{x}_{ij} .

As usual, we use I for indicator functions (e.g., $I(x_{ij\ell} = m)$ is 1 when the ℓ th field in record j in file i has the value m), and δ_a for the distribution of a point mass at a (e.g., $\delta_{y_{\lambda_{ij\ell}}}$). The vector $\boldsymbol{\theta}_\ell$ of length M_ℓ denotes the multinomial probabilities. For clarity, we always index as follows: $i = 1, \dots, k$; $j = 1, \dots, n_i$; $j' = 1, \dots, N$; $\ell = 1, \dots, p$; $m = 1, \dots, M_\ell$.

2.1 Independent Fields Model

We assume that the files are conditionally independent, given the latent individuals, and that fields are independent within individuals. We formulate the following Bayesian parametric model, where the joint posterior is in closed form and we sample from the full conditionals using a hybrid MCMC algorithm:

$$\begin{aligned} \mathbf{x}_{ij\ell} \mid \lambda_{ij}, \mathbf{y}_{\lambda_{ij\ell}}, z_{ij\ell}, \boldsymbol{\theta}_\ell &\stackrel{\text{ind}}{\sim} \begin{cases} \delta_{\mathbf{y}_{\lambda_{ij\ell}}} & \text{if } z_{ij\ell} = 0 \\ \text{MN}(1, \boldsymbol{\theta}_\ell) & \text{if } z_{ij\ell} = 1 \end{cases} \\ z_{ij\ell} &\stackrel{\text{ind}}{\sim} \text{Bernoulli}(\beta_\ell) \\ \mathbf{y}_{j'\ell} \mid \boldsymbol{\theta}_{j'\ell} &\stackrel{\text{ind}}{\sim} \text{MN}(1, \boldsymbol{\theta}_\ell) \\ \boldsymbol{\theta}_\ell &\stackrel{\text{ind}}{\sim} \text{Dirichlet}(\boldsymbol{\mu}_\ell) \\ \beta_\ell &\stackrel{\text{ind}}{\sim} \text{Beta}(a_\ell, b_\ell) \\ \pi(\mathbf{\Lambda}) &\propto 1, \end{aligned}$$

where a_ℓ, b_ℓ , and $\boldsymbol{\mu}_\ell$ are all known, and MN denotes the Multinomial distribution.

Remark 2.1: We assume that every legal configuration of the λ_{ij} is equally likely a priori. This implies a non-uniform prior on related quantities, such as the number of individuals in the data. The uniform prior on $\mathbf{\Lambda}$ is convenient, since constructing either a subjective or an alternative objective prior is unclear. A uniform distribution on one quantity, i.e. Λ , implies a non-uniform distribution on other, related quantities (such as N). Making every entry in the matrix λ_{ij} uniformly distributed on $1, 2, \dots, N_{\max}$ implies that the distribution of N , a function of Λ , is not uniform

on $1, 2, \dots, N_{\max}$. This is a long-standing problem with “non-informative priors” [10].

Deriving the joint posterior and conditional distributions is now mostly straightforward. One subtlety, however, is that \mathbf{y} , \mathbf{z} and $\mathbf{\Lambda}$ are all related, since if $z_{ij\ell} = 0$, then it must be the case that $y_{\lambda_{ij\ell}} = x_{ij\ell}$. Taking this into account, the joint posterior is

$$\begin{aligned} \pi(\mathbf{\Lambda}, \mathbf{y}, \mathbf{z}, \boldsymbol{\theta}, \boldsymbol{\beta} \mid \mathbf{x}) & \propto \prod_{i,j,\ell,m} \left[(1 - z_{ij\ell}) \delta_{y_{\lambda_{ij\ell}}}(x_{ij\ell}) + z_{ij\ell} \theta_{\ell m}^{I(x_{ij\ell}=m)} \right] \\ & \times \prod_{\ell,m} \theta_{\ell m}^{\mu_{\ell m} + \sum_{j'=1}^N I(y_{j'\ell}=m)} \\ & \times \prod_{\ell} \beta_{\ell}^{a_{\ell} - 1 + \sum_{i=1}^k \sum_{j=1}^{n_i} z_{ij\ell}} \\ & \times (1 - \beta_{\ell})^{b_{\ell} - 1 + \sum_{i=1}^k \sum_{j=1}^{n_i} (1 - z_{ij\ell})}. \end{aligned}$$

We suppress derivation of the full conditionals, but note that the full conditionals of \mathbf{y} , \mathbf{z} and $\mathbf{\Lambda}$ always obey their logical dependence, and therefore never condition on impossible events. The full conditional of $\mathbf{\Lambda}$ must reflect whether or not there are duplicates within files. If we define $R_{ij'} = \{j : \lambda_{ij} = j'\}$, then not having within-file duplicates means that $R_{ij'}$ must be either \emptyset or a single record, for each i and j' . Graphically, this means allowing or forbidding links from a latent individual to multiple records within one file.

2.2 Split and MERge REcord linkage and De-duplication (SMERED) Algorithm

Our main goal is estimating the posterior distribution of the linkage (i.e., the clustering of records into individuals). The simplest way of accomplishing this is via Gibbs sampling. We could iterate through the records, and for each record, sample a new assignment to an individual (from among the individuals represented in the remaining records, plus an individual comprising only that record). However, this requires the quadratic-time checking of proposed linkages for every record. Thus, instead of Gibbs sampling, we use a hybrid MCMC algorithm to explore the space of possible linkage structures, which allows our algorithm to run in linear time.

Our hybrid MCMC takes advantage of split-merge moves, as done in [9], which avoids the problems associated with Gibbs sampling, even though the number of parameters grows with the number of records. This is accomplished via proposals that can traverse the state space quickly and frequently visit high-probability modes, since the algorithm splits or merges records in each update, and hence, frequent updates of the Gibbs sampler are not necessary.

Furthermore, a common technique in record linkage is to require an exact match in certain fields (e.g., birth year) if records are to be linked. This technique of *blocking* can greatly reduce the number of possible links between records (see e.g., [17]). Since blocking gives up on finding truly co-referent records which disagree on those fields, it is best to block on fields that have little or no distortion. We block on the fairly reliable fields of sex and birth year in our application to the NLTCs below. A strength of our model is that it incorporates blocking organically. Setting $b_{\ell} = \infty$ for a particular field ℓ forces the distortion probability for that field to zero. This requires matching records to agree on the ℓ th field, just like blocking.

We now discuss how the split-merge process links records to records, which it does by assigning records to latent individuals. Instead of sampling assignments at the record level, we do so at the individual level. Initially, each record is assigned to a unique individual. On each iteration, we choose two records at random. If the pair belong to *distinct* latent individuals, then we propose merging those individuals to form a single new latent individual (i.e., we propose that those records are co-referent). On the other hand, if the two records belong to the *same* latent individual, then we propose splitting it into two new latent individuals, each seeded with one of the two chosen records, and the other records randomly divided between the two. Proposed splits and merges are accepted based on the Metropolis-Hastings ratio and rejected otherwise.

To choose the pair of records, one option is to sample uniformly from among all possible pairs. However, this is not ideal, for two reasons. First, most pairs of records are extremely unlikely to match since they agree on few, if any, fields. Frequent proposals to merge such records are wasteful. Therefore, we employ blocking, and only consider pairs of records within the same block. Second, sampling from all possible pairs of records will sometimes lead to proposals to merge records in the same list. If we permit duplication within lists, then this is not a problem. However, if we know (or assume) there are no duplicates within lists, we should avoid wasting time on such pairs. The no-duplication version of our algorithm does precisely this. (See Algorithm 1 for pseudocode.) When there are no duplicates within files, we call the SMERE (Split and MERge REcord linkage) algorithm, which enforces the restriction that $R_{ij'}$ must be either \emptyset or a single record. This is done through limiting the proposal of record pairs to those in distinct files; the algorithm otherwise matches SMERED.

Algorithm 1: Split and MErgE REcord linkage and De-duplication (SMERED)

Data: \mathbf{X} and hyperparameters

Initialize the unknown parameters $\theta, \beta, \mathbf{y}, \mathbf{z}$, and Λ .

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for  $i \leftarrow 1$  to  $S_G$  do
  for  $j \leftarrow 1$  to  $S_M$  do
    for  $t \leftarrow 1$  to  $S_T$  do
      Draw records  $R_1$  and  $R_2$  uniformly and
      independently at random.
      if  $R_1$  and  $R_2$  refer to the same individual
      then
        propose splitting that individual,
        shifting  $\Lambda$  to  $\Lambda'$ 
      endif
      else
        propose merging the individuals  $R_1$ 
        and  $R_2$  refer to, shifting  $\Lambda$  to  $\Lambda'$ 
      endif
       $r \leftarrow \min \left\{ 1, \frac{\pi(\Lambda', \mathbf{y}, \mathbf{z}, \theta, \beta | \mathbf{x})}{\pi(\Lambda, \mathbf{y}, \mathbf{z}, \theta, \beta | \mathbf{x})} \right\}$ 
      Resample  $\Lambda$  by accepting proposal with
      Metropolis probability  $r$  or rejecting with
      probability  $1 - r$ .
    end
    Resample  $\mathbf{y}$  and  $\mathbf{z}$ .
  end
  Resample  $\theta, \beta$ .
end
return  $\theta | \mathbf{X}, \beta | \mathbf{X}, \mathbf{y} | \mathbf{X}, \mathbf{z} | \mathbf{X}$ , and  $\Lambda | \mathbf{X}$ .

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2.2.1 Time Complexity

Scalability is crucial to any record linkage algorithm. Current approaches typically run in polynomial (but super-linear) time in N_{\max} . (The method of [14] is $O(N_{\max}^k)$, while that of [4] finds the maximum flow in an N_{\max} -node graph, which is $O(N_{\max}^3)$, but independent of k .) In contrast, our algorithm is linear in both N_{\max} and MCMC iterations.

Our running time is proportional to the number of Gibbs iterations S_G , so we focus on the time taken by one Gibbs step. Recall the notation from §2, and define $M = \frac{1}{p} \sum_{\ell=1}^p M_{\ell}$ as the average number of possible values per field ($M \geq 1$). The time taken by a Gibbs step is dominated by sampling from the conditional distributions. Specifically, sampling β and \mathbf{y} are both $O(pN_{\max})$; sampling θ is $O(pMN) + O(pN_{\max}) = O(pMN)$, as is sampling \mathbf{z} . Sampling Λ is $O(pN_{\max}M)$ if done carefully. Thus, all these samples can be drawn in time linear in N_{\max} .

Since there are S_M Metropolis steps within each Gibbs step and each Metropolis step updates \mathbf{y} , \mathbf{z} , and Λ , the time needed for the Metropo-

lis part of one Gibbs step is $O(S_M p N_{\max}) + O(S_M p M N) + O(S_M p N_{\max} M)$. Since $N \leq N_{\max}$, the run time becomes $O(p S_M N_{\max}) + O(M p S_M N_{\max}) = O(M p S_M N_{\max})$. On the other hand, the updates for θ and β occur once each Gibbs step implying the run time is $O(p M N) + O(p N_{\max})$. Since $N \leq N_{\max}$, the run time becomes $O(p M N_{\max} + p N_{\max}) = O(p M N_{\max})$. The overall run time of a Gibbs step is $O(p M N_{\max} S_M) + O(p M N_{\max}) = O(p M N_{\max} S_M)$. Furthermore, for S_G iterations of the Gibbs sampler, the algorithm is order $O(p M N_{\max} S_G S_M)$. If p and M are all much less than N_{\max} , we find that the runtime is $O(N_{\max} S_G S_M)$.

Another important consideration is the number of MCMC steps needed to produce Gibbs samples that form an adequate approximation of the true posterior. This issue depends on the convergence properties (actual rate of convergence) of the hybrid Markov chain used by the algorithm, which are beyond the scope of the present work. Convergence diagnostics for our application to the NLTCs and hyperparameter sensitivity is discussed in Appendix B.

2.3 Posterior Matching Sets and Linkage Probabilities

In a Bayesian framework, the output of record linkage is not a deterministic set of matches between records, but a probabilistic description of how likely records are to be co-referent, based on the observed data. Since we are linking multiple files at once, we propose a range of posterior matching probabilities: the posterior probability of linkage between two arbitrary records and more generally among k records, the posterior probability given a set of records that they are linked, and the posterior probability that a given set of records is a maximal matching set (which will be defined later).

Two records (i_1, j_1) and (i_2, j_2) *match* if they point to the same latent individual, so $\lambda_{i_1 j_1} = \lambda_{i_2 j_2}$. The posterior probability of a match can be computed from the S_G MCMC samples:

$$P(\lambda_{i_1 j_1} = \lambda_{i_2 j_2} | \mathbf{X}) = \frac{1}{S_G} \sum_{h=1}^{S_G} I(\lambda_{i_1 j_1}^{(h)} = \lambda_{i_2 j_2}^{(h)}).$$

A one-way match is when an individual appears in only one of the k files, while a two-way match is when an individual appears in exactly two of the k files, and so on (up to k -way matches). We approximate the posterior probability of arbitrary one-way, two-way, \dots , k -way matches as the ratio of the number of times those matches happened in the posterior sample to S_G .

Although probabilistic results and interpretations provided by the Bayesian paradigm are useful both quantitatively and conceptually, we often report a point

estimate of the linkage structure. Thus, we face the question of how to condense the overall posterior distribution of Λ into a single estimated linkage structure.

Perhaps the most obvious approach is to set some threshold v , where $0 < v < 1$, and to declare (i.e., estimate) that two records match if and only if their posterior matching probability exceeds v . This strategy is useful if only a few specific pairs of records are of interest, but its flaws are exposed when we consider the coherence of the overall estimated linkage structure implied by such a thresholding strategy. Note that the true linkage structure is *transitive* in the following sense: if records A and B are the same individual, and records B and C are the same individual, then records A and C must be the same individual as well. However, this requirement of transitivity is in no way enforced by the simple thresholding strategy described above. Thus, a more sophisticated approach is required if the goal is to produce an estimated linkage structure that preserves transitivity.

To this end, it is useful to define a new concept. A set of records \mathcal{A} is a *maximal matching* set (MMS) if every record in the set has the same value of λ_{ij} and no record outside the set has that value of λ_{ij} . Define $\Omega(\mathcal{A}, \Lambda) := \Omega_{\mathcal{A}, \Lambda}$ to be 1 if \mathcal{A} is an MMS in Λ and 0 otherwise:

$$\Omega_{\mathcal{A}, \Lambda} = \sum_{j'} \left(\prod_{(i,j) \in \mathcal{A}} I(\lambda_{ij} = j') \prod_{(i,j) \notin \mathcal{A}} I(\lambda_{ij} \neq j') \right).$$

Essentially, the MMS contains all the records which match some particular latent individual, though which individual is irrelevant. Given a set of records \mathcal{A} , the posterior probability that it is an MMS in Λ is simply

$$P(\Omega_{\mathcal{A}, \Lambda} = 1) = \frac{1}{S_G} \sum_{h=1}^{S_G} \Omega(\mathcal{A}, \Lambda^{(h)}).$$

The MMSs allow a sophisticated method of preserving transitivity when estimating a single overall linkage structure. For any record (i, j) , its *most probable MMS* \mathcal{M}_{ij} is the set containing (i, j) with the highest posterior probability of being an MMS, i.e.,

$$\mathcal{M}_{ij} := \arg \max_{\mathcal{A}: (i,j) \in \mathcal{A}} P(\Omega_{\mathcal{A}, \Lambda} = 1).$$

Next, a *shared most probable MMS* is a set that is the most probable MMS of all records it contains, i.e., a set \mathcal{A}^* such that $\mathcal{M}_{ij} = \mathcal{A}^*$ for all $(i, j) \in \mathcal{A}^*$. We then estimate the overall linkage structure by linking records if and only if they are in the same shared most probable MMS. The resulting estimated linkage structure is guaranteed to have the transitivity property since (by construction) each record is an element of at most one shared most probable MMS.

2.4 Functions of Linkage Structure

The output of the Gibbs sampler also allows us to estimate the value of any function of the variables, parameters, and linkage structure by computing the average value of the function over the posterior samples. For example, estimated summary statistics about the population of latent individuals are straightforward to calculate. Indeed, the ease with which such estimates can be obtained is yet another benefit of the Bayesian paradigm, and of MCMC in particular.

3 Assessing Accuracy of Matching and Application to NLTCS

We test our model on data from the NLTCS, a longitudinal study of the health of elderly (65+) individuals (<http://www.nltcs.aas.duke.edu/>). The NLTCS was conducted approximately every six years, with each wave containing roughly 20,000 individuals. Two aspects of the NLTCS make it suitable for our purposes: individuals were tracked from wave to wave with unique identifiers, but at each wave, many patients had died (or otherwise left the study) and were replaced by newly-eligible patients. We can test the ability of our model to link records across files by seeing how well it is able to track individuals across waves, and compare its estimates to the ground truth provided by the unique identifiers.

To show how little information our method needs to find links across files, we gave it access to only four variables, all known to be noisy: full date of birth, sex, state of residence, and the regional office at which the subject was interviewed. We treat all fields as categorical. We linked individuals across the 1982, 1989 and 1994 survey waves.² Our model had little information on which to link, and not *all* of its assumptions strictly hold (e.g., individuals can move between states across waves). We demonstrate our method's validity using error rates, confusion matrices, posterior matching sets and linkage probabilities, and estimation of the unknown number of observed individuals from the population.

Appendix A provides a simulation study of the NLTCS with varying levels of distortion at the field level. We conclude from this that SMERE is able to handle low to moderate levels of distortion (Figure 4). Furthermore, as distortion increases, so do the false negative rate (FNR) and false positive rate (FPR) (Figure 3).

²The other three waves used different questionnaires and are not strictly comparable.

3.1 Error Rates and Confusion Matrix

Since we have unique identifiers for the NLTCs, we can see how accurately our model matches records. A *true link* is a match between records which really do refer to the same latent individual; a *false link* is a match between records which refer to different latent individuals; and a *missing link* is a match which is not found by the model. Table 3 gives posterior means for the number of true, false, and missing links. For the NLTCs, the FNR is 0.11, while the FPR is 0.046, when we block by date of birth year (DOB) and sex.

More refined information about linkage errors comes from a confusion matrix, which compares records’ estimated and actual linkage patterns (Figure 1 and Appendix C, Table 4). Every row in the confusion matrix is diagonally dominated, indicating that correct classifications are overwhelmingly probable. The largest off-diagonal entry, indicating a mis-classification, is 0.07. For instance, if a record is estimated to be in both the 1982 and 1989 waves, it is 90% probable that this estimate is correct. If the estimate is wrong, the truth is most probably that the record is in all waves (4.4%), followed by the 1982 wave alone (1.4%) and waves 1982 and 1994 (0.15%), and then other patterns with still smaller probability.

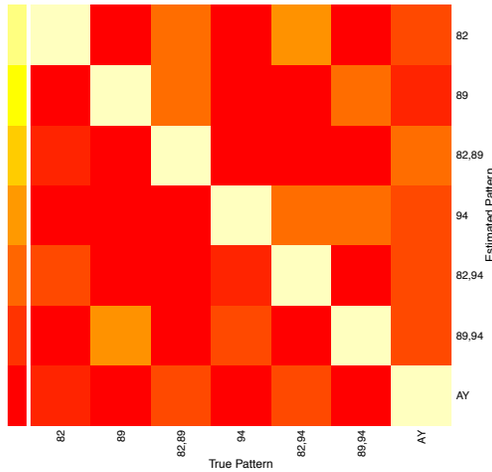


Figure 1: Heatmap of relative probabilities from the confusion matrix, running from yellow (most probable) to dark red (probability 0). The largest probabilities are on the diagonal, showing that the linkage patterns estimated for records are correct with high probability. Mis-classification rates are low and show a tendency to under-link rather than over-link.

3.2 Example of Posterior Matching Probabilities

We wish to search for sets of records that match record 10084 in 1982. In the posterior samples of Λ , this record is part of three maximal matching sets that occur with nonzero estimated posterior probability, one with high and two with low posterior matching probabilities (Table 1). This record has a posterior probability of 0.995 of simultaneously matching both record 6131 in 1989 and record 5583 in 1994. All three records denote a male, born 07/01/1910, visiting office 25 and residing in state 14. The unique identifiers show that these three records are in fact the same individual. If we threshold matching sets, ignoring ones of low posterior probability, we would simply return the set of records in last column of Table 1.

3.3 Estimation of Attributes of Observed Individuals from the Population

The number of observed unique individuals N is easily inferred from the posterior of $\Lambda|\mathbf{X}$, since N is simply the number of unique values in Λ . Defining $N|\mathbf{X}$ to be the posterior distribution of N , we can find this by applying a function to the posterior distribution on Λ , as discussed in §2.4. (Specifically, $N = |\#\Lambda|$, where $\#\Lambda$ maps Λ to its set of unique entries, and $|A|$ is the cardinality of the set A .) Doing so, the posterior distribution of $N|\mathbf{X}$ is given in Figure (2). Also, $\hat{N} := E(N|\mathbf{X}) = 35,992$ with a posterior standard error of 19.08. Since the true number of observed unique individuals is 34,945, we are overmatching, which leads to an overestimate of N . This phenomenon most likely occurs due to patients migrating between states across the three different waves. It is difficult to improve this estimate since we do not have additional information as just described above.

We can also estimate attributes of sub-groups. For example, we can estimate the number of individuals within each wave or combination of waves—that is, the number of individuals with any given linkage pattern. (We summarize these estimates here with posterior expectations alone, but the full posterior distributions are easily computed.) For example, the posterior expectation for the number of individuals appearing in lists i_1 and i_2 but not i_3 is approximately

$$\frac{1}{S_G} \sum_{h=1}^{S_G} \sum_{j'} I \left(\left| R_{i_1 j'}^{(h)} \right| = 1 \right) I \left(\left| R_{i_2 j'}^{(h)} \right| = 1 \right) I \left(\left| R_{i_3 j'}^{(h)} \right| = 0 \right).$$

(Note that the inner sum is a function of $\Lambda^{(h)}$, but a very complicated one to express without the R_{ij} .)

Table 2 reports the posterior means for the overlapping waves and each single wave of the NLTCs and

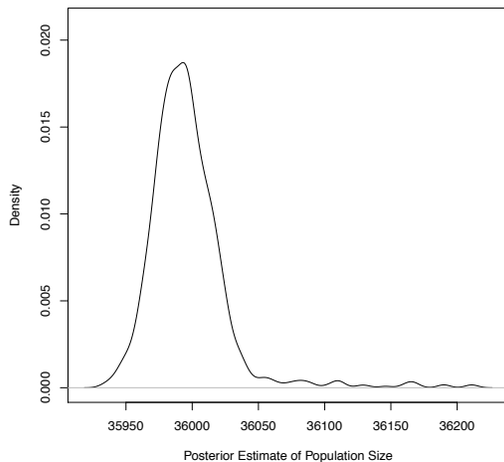


Figure 2: Posterior density of the number of observed unique individuals N .

compares this to the ground truth. In the first wave (1982), our estimates perform exceedingly well with relative error of 0.11%, however, as waves cross and we try to match people based on limited information, the relative errors range from 8% to 15%. This is not surprising, since as patients age, we expect their proxies to respond, making patient data more prone to errors. Also, older patients may move across states, creating further matching dilemmas. We are unaware of any alternative algorithm that does better on this data with only these fields available. Given these results, and considering how little field information we allowed it to use for matching, we find that our model performs overall very well.

4 De-duplication

Our application of SMERE to the NLTCS assumes that each list had no duplicates, however, many other applications will contain duplicates within lists. We showed in §2.1 that we can theoretically handle de-duplication across and within lists. We apply SMERE with de-duplication (SMERED) to the NLTCS by (i) running SMERED on the three waves to show that the algorithm does not falsely detect duplicates when there really are none, and (ii) combining all the lists into one file, hence creating many duplicates, to show that SMERED can find them.

4.1 Application for NLTCS

We combine the three files of the NLTCS mentioned in §3 which contain 22,132 duplicate records out of

57,077 total records. We run SMERED on settings (i) and (ii), evaluating accuracy with the unique IDs.

In the the case of running SMERED on the three waves, we compare our results of SMERED and SMERE to that under ground truth (Table 2). In the case of the NLTCS, compiling all three files together and running the three waves separately under SMERED yields similar results, since we match on similar covariate information. There is no covariate information to add to from thorough investigation to improve our results, except under simulation study. Specifically, when running SMERED for the three files, the FNR is 0.11 and is 0.38 for FPR, while its FNR and FPR is 0.11 AND 0.37 for the one compiled file. We contrast this with the FNR of 0.11 and FPR of 0.046 under SMERE for the three waves (Table 3).

The dramatic increase in the FPR and number of false links shown in Table 2 is explained by how few field variables we match on. Their small number means that there are many records for different individuals that have identical or near-identical values. On examination, there are 2,558 possible matches among “twins,” records which agree exactly on all attributes but have different unique IDs. Moreover, there are 353,536 “near-twins,” pairs of records that have different unique IDs but match on all but one attribute. This illustrates why the matching problem is so hard for the NLTCS and other data sources like it, where survey-responder information like name and address are lacking. However, if it is known that each file contains no duplicates, there is no need to consider most of these twins and near-twins as possible matches.

5 Discussion

We have made two contributions in this paper. The first is to frame record linkage and de-duplication simultaneously, namely linking observed records to latent individuals and representing the linkage structure via Λ . The second contribution is our specific parametric Bayesian model, which, combined with the linkage structure, allows for efficient inference and exact error rate calculation. Moreover, this allows for easy integration with capture-recapture methods, where error propagation is *exact*. As with any parametric model, its assumptions only apply to certain problems, but it also serves as a starting point for more elaborate models, e.g., with missing fields, data fusion, complicated string fields, population heterogeneity, or dependence across fields, across time, or across individuals. Within the Bayesian paradigm, such model expansions will lead to larger parameter spaces, and therefore call for computational speed-ups, perhaps via online learning, variational inference, or approximate

Bayesian computation.

Our work serves as a first basis for solving record linkage problems using a noisy Bayesian model, a linkage structure that can handle large-scale databases, and a model that simultaneously combines record linkage and de-duplication for arbitrarily many files. We hope that our approach will encourage the emergence of new record linkage approaches, extensions of our method to non-categorical fields, and applications along with more state-of-the-art algorithms for this kind of high-dimensional data.

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sets of records	1.10084	3.5583; 1.10084	3.5583; 1.10084; 2.6131
posterior probability	0.001	0.004	0.995

Table 1: Example of posterior matching probabilities for record 10084 in 1982

	82	89	94	82, 89	89, 94	82, 94	82, 89, 94
NLTCS (ground truth)	7955	2959	7572	4464	3929	1511	6114
Bayes Estimates _{SMERE}	7964	3434.1	8937.8	4116.9	4502.1	1632.2	5413
Bayes Estimates _{SMERED}	7394.7	3009.9	6850.4	4247.5	3902.7	1478.7	5191.2
Relative Errors _{SMERE} (%)	0.11	16.06	18.04	-7.78	14.59	8.02	-11.47
Relative Errors _{SMERED} (%)	-7.04	1.72	-9.53	-4.85	-0.67	-2.14	-15.09

Table 2: Comparing NLTCS (ground truth) to the Bayes estimates of matches for SMERE and SMERED

	False links	True Links	Missing Links	FNR	FPR
NLTCS (ground truth)	0	28246	0	0	0
Bayes Estimates _{SMERE}	1298.9	25196	3050	0.11	0.05
Bayes Estimates _{SMERED}	10595	24900	3346	0.09	0.37

Table 3: False, True, and Missing Links for NLTCS under blocking sex and DOB year where the Bayes estimates are calculated in the absence of duplicates per file and when duplicates are present (when combining all three waves). Also, reported FNR and FPR for NLTCS, Bayes estimates.

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