#### 1 DIRICHLET RECORD LINKAGE

Recall that for fabl, we used the following prior for the linkage structure Z and the rate of matching parameter  $\pi$ :

$$p(Z_j = q | \pi) = \begin{cases} \frac{1}{n_A} \pi, & q > 0; \\ 1 - \pi, & q = 0; \end{cases}$$
 (1a)

$$\pi \sim \text{Beta}(\alpha_{\pi}, \beta_{\pi}).$$
 (1b)

This prior says that a record B has some match in A with probability  $\pi$ , and that each record in A is equally likely to be that match. We now generalize these prior distributions to allow the records in B to match with multiple records in A.

## 1.1 Prior Specification

We provide updated notation to allow us to describe one record in B having multiple matches in A. Let  $Z_j$  be a set containing the indices for all of the records in A that are a match with record  $B_j$ , and let  $Z = \{Z_j | j = 1, ..., n_B\}$  denote the collection of such sets for all records in B. Let  $|Z_j| = \sum_{k=1}^{\infty} I(Z_{j,k} > 0)$  denote the number of records in A that are linked to  $B_j$ . We use  $Z_j = \emptyset$  to denote when  $B_j$  has no match in A.

We modify the fabl model to allow each record in B to match to multiple records in A. Define a vector of probabilities  $\boldsymbol{\pi} = (\pi_0, \dots, \pi_{n_A})$  where  $\pi_k$  is the probability that some record in B has exactly k matches in A. Conditional on  $B_j$  having k matches, we construct a prior specification on Z such that each matching  $Z_j$  of length  $|Z_j|$  is equally likely. Since there are  $\binom{n_a}{|q|} = \frac{n_A!}{(n_A - |q|)!|q|!}$  ways to select |q| matching records out of all  $n_A$  possible records, we replace (1a) with

$$p(Z_j = q | \boldsymbol{\pi}) = \frac{(n_A - |q|)!|q|!}{n_A!} \pi_{|q|}.$$
 (2)

Since  $\pi$  is multivariate, we replace (1b) with

$$\pi \sim \text{Dirichlet}(\alpha_{\pi}).$$
 (3)

## 1.2 Full Conditional for $Z_i$

Following the observation of Wortman (2019) and elaborated by Kundinger et al. (2024), when  $B_j$  does not link to any record in A (such that  $|Z_j| = 0$ ) the contribution to the likelihood is simply a product of u parameters, which we will call  $c_i$ :

$$p(\Gamma_{.j}|\boldsymbol{m},\boldsymbol{u},\pi,Z_j=\emptyset) = \prod_{i=1}^{n_A} \prod_{f=1}^F \prod_{l=1}^{L_f} u_{fl}^{I(\gamma_{ij}^f=l)I_{obs}(\gamma_{ij}^f)} = c_j.$$

$$(4)$$

When  $Z_j = q = (q_1, \ldots, q_k)$  for some |q| > 0, we have

$$p(\Gamma_{.j}|\boldsymbol{m},\boldsymbol{u},\pi,Z_{j}=q) = \prod_{i \in q} \prod_{f=1}^{F} \prod_{l=1}^{L_{f}} m_{fl}^{I(\gamma_{ij}^{f}=l)I_{obs}(\gamma_{ij}^{f})} \prod_{i \notin q} \prod_{f=1}^{F} \prod_{l=1}^{L_{f}} u_{fl}^{I(\gamma_{ij}^{f}=l)I_{obs}(\gamma_{ij}^{f})}.$$
 (5)

We multiply and divide by the u parameters for the matching record pairs to obtain

$$p(\Gamma_{.j}|\boldsymbol{m},\boldsymbol{u},\pi,Z_{j}=q) = \prod_{i \in q} \prod_{f=1}^{F} \prod_{l=1}^{L_{f}} \left(\frac{m_{fl}}{u_{fl}}\right)^{I(\gamma_{ij}^{f}=l)I_{obs}(\gamma_{ij}^{f})} \prod_{i=1}^{n_{A}} \prod_{f=1}^{F} \prod_{l=1}^{L_{f}} u_{fl}^{I(\gamma_{ij}^{f}=l)I_{obs}(\gamma_{ij}^{f})}$$
(6)

$$=c_j\prod_{i\in q}w_{ij}. (7)$$

Lastly, we multiply the likelihood by the prior in (2) to obtain the posterior distribution. For  $Z_j = q$  where |q| = k, we have

$$p(Z_{j} = q | \gamma, \boldsymbol{m}, \boldsymbol{u}, \pi) = \frac{\frac{(n_{A} - k)!|k|!}{n_{A}!} \pi_{k} c_{j} \prod_{i \in q} w_{ij}}{\sum_{h \in \mathcal{Z}} \frac{(n_{A} - |h|)!|h|!}{n_{A}!} \pi_{|h|} c_{j} \prod_{i \in h} w_{ij}}$$
(8a)

$$= \frac{\frac{(n_A - k)!|k|!}{n_A!} \pi_k \prod_{i \in q} w_{ij}}{\sum_{h \in \mathcal{Z}} \frac{(n_A - |h|)!|h|!}{n_A!} \pi_{|h|} \prod_{i \in h} w_{ij}}$$
(8b)

$$\propto \frac{(n_A - k)!|k|!}{n_A!} \pi_k \prod_{i \in q} w_{ij}. \tag{8c}$$

# 1.3 Sequential Sampler

Sampling the full conditional in (8c) is infeasible for almost any record linkage task. In particular, there are  $2^{n_A}$  possible matchings for each  $B_j$ , meaning that computational complexity for the sampler would be  $O(n_B 2^{n_A})$ . One could reduce this computational burden by setting a maximum number of matches, K, per record in B, but this would still require  $\sum_{k=1}^{K} \frac{n_A!}{(n_A-k)!k!}$  possible options for the set  $Z_j$ , and would be prohibitive for

most values of  $n_A$  seen in record linkage problems. Through Gibbs sampling however, we can break this joint distribution into a sequence of more simple conditional univariate distributions. This allows for a more computationally efficient sampler, and allows us to learn K from the data, rather than setting it ahead of time.

We model each  $\pi_k$  as a product of conditional probabilities: let  $\eta_k$  be the probability that some record in B has at least k matches, given that it has at least k-1 matches. This gives us the stick breaking representation

$$\pi_k = (1 - \eta_{k+1}) \prod_{c=1}^k \eta_c, \tag{9}$$

where  $\eta_k$  are independent random variables from a Beta $(\alpha_{\eta}, \beta_{\eta})$  distribution.

We generalize the fast beta prior in (1a) to a sequence of priors that allows for multiple matchings. When  $B_j$  has been linked to k-1 records, we say that the probability that  $B_j$  has a  $k^{th}$  match is  $\eta_k$ , and that all remaining records in A are equally likely to be linked. Let  $Z_{j,-k} = (Z_{j,1}, \ldots, Z_{j,k-1})$  be the set of records linked to  $B_j$  before the  $k^{th}$  matching phase. We use

$$p(Z_{j,k} = q_k | \eta_k) = \begin{cases} \frac{\eta_k}{n_A - (k-1)}, & q_k \notin A_{j,k}, \\ 1 - \eta_k, & q_k = \emptyset; \end{cases}$$
(10)

where  $A_{j,k} = [n_A] \setminus Z_{j,-k}$  is the set of records in A that are available to be matched with  $B_j$ . This sequence of priors leads to sequence of posteriors that can be used to sample arbitrarily many links for record  $B_j$ . These posteriors are given by

$$p(Z_{j,k} = q_k | Z_{j,k-1}, \eta_k, \boldsymbol{m}, \boldsymbol{u}, \gamma) \propto \begin{cases} \frac{\eta_k}{n_A - (k-1)} w_{q_k, j}, & q_k \in A_{j,k}, \\ 1 - \eta_k, & q_k = \emptyset, \end{cases}$$
(11)

as derived in Appendix 5.1.

This sequential sampler produces an output  $Z_j = q = (q_1, \ldots, q_k)$  when  $Z_{j,c} = q_c$  for steps  $c \in \{1, \ldots, k\}$ , and the k+1 step produces  $Z_{j,k+1} = \emptyset$ . While this vector is necessarily ordered, all reorderings of the same elements are equivalent for the purposes of record linkage. That is,  $Z_j = (i, i')$  and  $Z_j = (i', i)$  both communicate that record  $B_j$  is matched

to records  $A_i$  and  $A_{i'}$ . Let  $\sigma(q)$  denote all possible orderings of the elements of q, and note that there are |q|! such orderings. Observe that

$$p(Z_{j,k+1} = \emptyset | \Gamma_{.j}, \boldsymbol{m}, \boldsymbol{u}, \boldsymbol{\eta}) \sum_{q' \in \sigma(q)} \prod_{c=1}^{k} p(Z_{j,c} = q'_c | \Gamma_{.j}, \boldsymbol{m}, \boldsymbol{u}, \boldsymbol{\eta})$$
(12)

$$\propto (1 - \eta_{k+1}) \sum_{q' \in \sigma(q)} \prod_{c=1}^{k} \frac{\eta_c}{n_A - (c-1)} \prod_{c=1}^{k} w_{q'_c, j}$$
 (13)

$$= (1 - \eta_{k+1})k! \prod_{c=1}^{k} \frac{\eta_c}{n_A - (c-1)} \prod_{c=1}^{k} w_{q_c,j}$$
 (14)

$$= \frac{(n_A - k)!k!}{n_A!} (1 - \eta_{k+1}) \prod_{c=1}^k \eta_c \prod_{c=1}^k w_{q_c,j}$$
 (15)

$$= \frac{(n_A - k)!k!}{n_A!} \pi_k \prod_{c=1}^k w_{q_c,j}$$
 (16)

$$= p(Z_j = q | \gamma, \boldsymbol{m}, \boldsymbol{u}, \pi). \tag{17}$$

Thus, the probability that the sequential process samples each of the components of q (in any order) is equal to the joint probability of q as expressed in the joint distribution in (8c).

This sequential sampler amounts to an extension of fabl with an iterative matching phase. In each iteration of the Gibbs sampler, we sample an initial set of links using  $\eta_1$ . For each record in B that was found to have a link, we remove the linked record in A from consideration, and then sample another potential link with  $\eta_2$ . We continue, using  $\eta_k$  in the  $k^{th}$  matching step, until no new links are found, at which we point the matching phase terminates. The  $\eta$ , m, and u parameters are estimated based on all of the links identified, regardless of the order in which they are sampled. Crucially, there is no need to specify a maximum number of links per record, as this estimated through the model.

# 1.4 Efficient Sequential Sampling

Following Kundinger et al. (2024) we use hashing to reduce the computational complexity of the Gibbs sampler. Since each component  $\gamma_{ij}^f$  of the comparison vector is discrete, there are only finitely many possible realizations of the comparison vector  $\gamma_{ij}$ . Let P be the number of unique agreement patterns observed in  $\Gamma$ . This number is bounded above by

 $P^* = \prod_{f=1}^F (L_f + 1)$ , where the addition of 1 to  $L_f$  for each field accounts for the possibility of missing values. This upper bound does not scale with  $n_1$  or  $n_2$ , but rather is determined by F and  $L_f$ .

Let  $N_{p_j} = \sum_{i=1}^{n_B} I(\gamma_{ij} = h_p)$  denote the number of records in A with which record j in B has agreement pattern p. Collect these counts in  $\mathcal{N} = \{N_{p_j} \mid j \in [n_B], p \in [P]\}$ . Let  $N_p = \sum_{j=1}^{n_B} N_{p_j}$  denote the total number of record pairs with agreement pattern p. Finally, let  $r_{p_j} = \{i \in [n_A] \mid \gamma_{ij} = h_p\}$  be the set of records in A with which record j in B has agreement pattern p, and collect these sets as  $\mathcal{R} = \{r_{p_j} \mid p \in [P], j \in [n_2]\}$ .

We adapt the split sampler from Kundinger et al. (2024) for this multiple match setting. Let  $r_{p_j}(k) = r_{p_j} Z_{j,-k}$  be the set of records of agreement pattern p available for matching at the  $k^{th}$  matching step, and let  $N_{p_j}(k) = |r_{p_j}(k)|$  be the number of such records. We first sample among P+1 options for the agreement pattern between  $B_j$  and its potential link. Define r as an arbitrary set of records. We have

$$p\left(Z_{j,k} \in r \mid \tilde{\gamma}, \boldsymbol{m}, \boldsymbol{u}, \eta_{k}\right) \propto \begin{cases} \frac{\eta_{k} N_{p_{j}}(k)}{n_{A} - (k-1)} w_{p}, & r = r_{p_{j}}(k); \\ 1 - \eta_{k}, & r = \emptyset. \end{cases}$$

$$(18)$$

Since all remaining records in A sharing the same agreement pattern with  $B_j$  are equally likely, we then sample among candidate records uniformly using

$$p(Z_{j,k} = q \mid Z_{j,k} \in r, \boldsymbol{m}, \boldsymbol{u}, \eta_k) = \begin{cases} \frac{1}{N_{p_j}(k)}, & r = r_{p_j}(k) \text{ and } q \in r; \\ 1, & r = \emptyset \text{ and } q = \emptyset. \end{cases}$$
(19)

See Appendix 5.1 for a more detailed explanation of hashing.

We take a moment to emphasize the computational advantages to this approach. Sampling from the full conditional shown in (8c) would have complexity  $O(2^{n_A}n_B)$ , and would be nearly impossible for any reasonable record linkage task. Using the sequential sampler shown in (11) has complexity  $O\left(n_A(n_B + \sum_{k=1}^K n_k)\right)$ , which would still still grow quadratically as the size of the linkage task grows. However, sampling from (18) and (19) has complexity  $O\left(P(n_B + \sum_{k=1}^K n_k)\right)$ , which grows linearly in the size of the base dataset. Thus, we have produced a computationally efficient sampler with speed comparable to the  $O(Pn_B)$  complexity of base fabl.

#### 2 SIMULATIONS

We demonstrate the accuracy of the DRL approach at various levels of overlap between files and errors between matching records through an adaptation of the simulation study from Sadinle (2017) and Kundinger et al. (2024).

For each simulation, we construct two data files A and B such that there are in which there are 25, 125, or 225 records in B that have matching records in A. The matching record in A exhibits 1, 2, or 3 errors across the five fields used for linkage. Then, every record in A that has a matching record in B is duplicated, such that each simulation has 50, 250, or 450 matching record pairs in total. We use uniform priors for the m and u, with  $\alpha_{fl} = \beta_{fl} = 1$  for all f and l. We use uniform priors for  $\pi$  for standard fabl, and also uniform priors for each  $\eta_k$  in sequential sampler for DRL. We run the Gibbs sampler for 1000 iterations, and discard the first 100 as burn-in.

We run  $\mathtt{multilink}$  under two different parameter settings. We first use the default setting, where the prior distribution for the number of records in A that can match a single record in B is a Poisson distribution with mean 1. Second, we use a prior that is closer to the truth, and use a Poisson distribution with mean 2, and the maximum number of matching records also set to 2.

In these simulations, standard fabl drastically underperforms, so much so that results are omitted from the graph. For each record in B with matching records in A, the posterior match probability is split between the two matching records. Due to the randomness of the Gibbs sampler, one of the records occassionally has a posterior probability over 0.5 and is thus identified through the Bayes estimate, but often, both records have posterior probability below 0.5. Under vabl, the posterior probability of the two records is cut precisely in half, and thus vabl did not identify any matching record pairs in any of the simulations. Again, these results are omitted.

We see that fastLink generally has poorer performance than the more complex models at mid to high overlap. In particular, without a post-processing step tailors the scenario where one record in B can match to multiple records in A, fastLink tends to produce a large number of false positives, hindering precision.

The performance of multilink is more varied. When there is a moderate or high amount of matching records, multilink performs comparably with multiple match. However, when the number of matching records pairs, multilink underperforms. This may be because the model being fit is more complex, and so it is more difficult to learn the parameters from such few observations. Notably however, the performance of multilink is highly sensitive to prior specification. In FIGURE (to be added), we show accuracy suffers under various forms of mispecification.

We see that DRL is a strong alternative to multilink and fastLink in this setting. DRL maintains strong performance even in the low overlap scenario where multilink fails, and outperforms fastLink at all levels of error and overlap. multilink... In particular, the average computation time for 1000 iterations of the Gibbs sampler for multiple match was around 30 seconds, while it was around 1000 seconds for multilink.

### 3 CASE STUDY

We demonstrate our method on the North Carolina Voter Registration (NCVR) database taken two months apart (Christen 2014). The snapshots are filtered to include only those voters whose details changed over the two-month period, so there are matching records with full agreement on all fields. We use first name, middle name, last name, as string fields, and street address and age as categorical fields. Unique voter registration numbers are provided, however they are known to contain some errors. The NCVR dataset is not publicly available due to sensitive information. However, we have permission to utilize it for publication by its owner.

Using the voter registration numbers, we can see that each file has internal duplication rates of about 1%. In this analysis, we deduplicate file B, and left A with duplicates. In practice, such low amount of internal duplication may not warrant the use of Dirichlet Beta Linkage since vabl is considerably faster. However, we demonstrate here that even with such few internal duplications, DBL is effective at identifying this multiple matches and does not declare too many false matches.

I compare four approaches: standard fabl, our proposed DRL method, fastLink, and

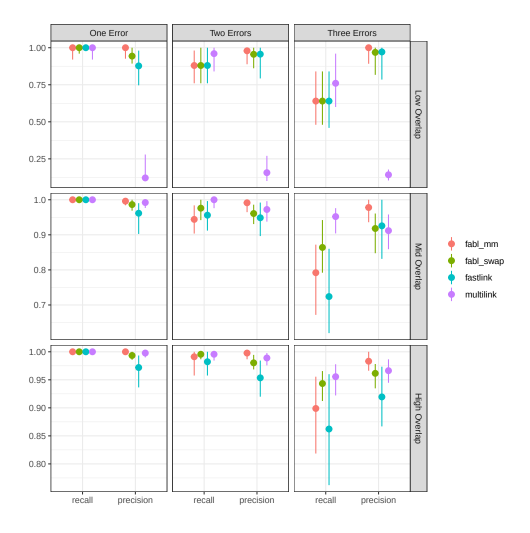


Figure 1: Sadinle Simulation

fastLink with the Jaro correction. This task is too large for running multilink in its current implementation in R, and is thus omitted. For fastLink, we use the threshold of 0.5 to declare matches, and for fabl and DRL, we use losses that are equivalent to a threshold of 0.5. Results are in Table 1.

We see that DRL has the highest f-measure, followed by fabl. When we use fastLink with no post-processing, we end up with an undesirable amount of false positives. Since the posterior match probability of each record pair is computed independently, we expect such behavior on large linkage tasks such as this. When we use the Jaro post-processing to achieve a one-to-one matching, we get considerably better results. However, we lose the possibility of identifying cases where one record in B matches to two records in A, and we

| method               | recall    | precision | $f_{-}$ measure |
|----------------------|-----------|-----------|-----------------|
| DRL                  | 0.9891623 | 0.9735309 | 0.9812843       |
| fabl                 | 0.9743739 | 0.9798799 | 0.9771191       |
| fastLink             | 0.9988472 | 0.8494865 | 0.9181321       |
| fastLink (with Jaro) | 0.9514490 | 0.9664229 | 0.9588775       |

Table 1: Accuracy results for NCVR, showing the DRL has the highest f-measure.

still have worse recall and precision than attained under standard fabl.

# 4 CONCLUSION

#### REFERENCES

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### 5 APPENDIX

## 5.1 Derivation of Sequential Sampler

We now provide the derivation of the sequential sampler, following the argument presented in Section 1.2. Suppose  $B_j$  has been linked to k records in A. Let  $Z_{j,-k} = (Z_{j,1}, \ldots, Z_{j,k-1})$  denote the vector of records already linked to  $B_j$ . When  $B_j$  has no additional link in A, the contribution to the likelihood is a product of the u parameters for all remaining records. That is,

$$p(\Gamma_{.j}|\boldsymbol{m},\boldsymbol{u},\pi,Z_{j,k}=\emptyset,Z_{j,-k}=q_{-k}) = \prod_{i \notin Z_{j,-k}} \prod_{f=1}^{F} \prod_{l=1}^{L_f} u_{fl}^{I(\gamma_{ij}^f=l)I_{obs}(\gamma_{ij}^f)} = c_{Z_{j,-k}}.$$
 (20)

When  $Z_{j,k} = q_k$  for some  $q_k > 0$ , we have

$$p(\Gamma_{.j}|\boldsymbol{m},\boldsymbol{u},\pi,Z_{j,k}=q_k,Z_{j,-k}=q_{-k}) = \prod_{f=1}^F \prod_{l=1}^{L_f} m_{fl}^{I(\gamma_{q_k,j}^f=l)I_{obs}(\gamma_{q_k,j}^f)} \prod_{i \notin (q_{-k},q_k)} \prod_{f=1}^F \prod_{l=1}^{L_f} u_{fl}^{I(\gamma_{ij}^f=l)I_{obs}(\gamma_{ij}^f)}$$

(21)

$$= \prod_{f=1}^{F} \prod_{l=1}^{L_f} \left( \frac{m_{fl}}{u_{fl}} \right)^{I(\gamma_{q_k,j}^f = l)I_{obs}(\gamma_{q_k,j}^f)} \prod_{i \notin (Z_{j,-k}} \prod_{f=1}^{F} \prod_{l=1}^{L_f} u_{fl}^{I(\gamma_{ij}^f = l)I_{obs}(\gamma_{ij}^f)}$$

(22)

$$=c_{Z_{j,-k}}w_{q_k,j} \tag{23}$$

To obtain the posterior, we multiply by the prior in (10). The posterior distribution this is given by

$$p(Z_{j,k} = q_k | Z_{j,k-1}, \eta_k, \boldsymbol{m}, \boldsymbol{u}, \gamma) = \frac{\left(\frac{\eta_k}{n_A - (k-1)} c_{Z_{j,-k}} w_{q_k,j}\right)^{I(q_k \in N_{j,k})} + \left(c_{Z_{j,-k}} (1 - \eta_k)\right)^{I(q_k = \emptyset)}}{\frac{\eta_k}{n_A - (k-1)} c_{Z_{j,-k}} \sum_{i \notin Z_{j,-k}} w_{ij} + c_{Z_{j,-k}} (1 - \eta_k)}$$

(24)

$$= \frac{\left(\frac{\eta_k}{n_A - (k-1)} w_{q_k, j}\right)^{I(q_k \in N_{j,k})} + (1 - \eta_k)^{I(q_k = \emptyset)}}{\frac{\eta_k}{n_A - (k-1)} \sum_{i \notin Z_{j-k}} w_{ij} + (1 - \eta_k)}$$
(25)

$$\propto \begin{cases}
\frac{\eta_k}{n_A - (k-1)} w_{q_k, j}, & q_k \in N_{j, k}, \\
1 - \eta_k, & q_k = \emptyset.
\end{cases}$$
(26)

Importantly, the constant  $c_{Z_{j,-k}}$  is not found in the final expression because the probability mass associated with every potential value for  $Z_j$  shares the same  $c_{Z_{j,-k}}$ . This does not occur due to proportionality.

# 5.2 Efficient Sequential Sampling Details