

blocking: An R Package for Blocking of Records for Record Linkage and Deduplication

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Abstract Entity resolution (probabilistic record linkage, deduplication) is essential for bringing together data needed for estimation or other analyses based on multiple sources. It aims to link records that refer to the same entity (e.g., person, company) in absence of unique identifiers. Without identifiers, researchers must specify which records to compare to calculate matching probability and reduce computational complexity. Traditional deterministic blocking uses common variables like first letters of names, sex, or dates of birth, but assumes error-free, complete data. To address this limitation, we developed the R package **blocking**, which uses approximate nearest neighbor search and graph algorithms to reduce the number of comparisons. This paper presents the package design, functionalities, and two case studies.

1 Introduction

1.1 Blocking for record linkage

Entity resolution (probabilistic record linkage, deduplication) is essential for bringing together data needed for estimation or other analyses based on multiple sources (cf. [Fellegi and Sunter \(1969\)](#), [Binette and Steorts \(2022\)](#)). The goal is to link records that refer to the same entity (e.g., person, company, job position) in the absence of unique identifiers. This situation is frequently observed in administrative records, e.g., for foreign-national populations. For instance, the Social Insurance Institution register in Poland at the end of 2023 included 1.206 million records referring to approximately 1.105 million individuals, of which about 10% had missing information in the personal identifier (PESEL) and about 50% had missing address details. Note that the exact number of individuals is certainly lower than 1.105 million, as the 10% with missing identifiers may include duplicates (cf. [Beręsewicz \(2025\)](#)).

This drives the need to link records without unique identifiers, which often requires certain assumptions about how to reduce the large number of possible comparisons, as it is not feasible to compare all pairs of records in large datasets (e.g., the aforementioned example would require over 700 billion comparisons). Consequently, researchers aim to reduce the number of comparisons in various ways prior to the record linkage/deduplication stage. The rationale is twofold: computational resource constraints and clerical review workload.

Reducing the number of comparisons is accomplished through blocking, a method that limits possible comparisons by assuming that certain variables must match exactly or that combinations of variables should match above a specified threshold. For instance, a standard approach assumes that sex or birth year must match exactly, whilst other record characteristics may vary. Another method employs phonetic algorithms such as Soundex (cf. [Wright \(1960\)](#)) or its adaptations for non-English languages (cf. [Howard II \(2020\)](#)) to block records that sound similar but are spelt differently (e.g., Smith and Smyth, or Anna and Ania). Furthermore, with the growing popularity of large language models (both closed and open-source), one may consider using embeddings ([Mikolov et al., 2013](#)) to identify nearest neighbours and treat these as potential comparison pairs. For a comprehensive review of blocking methods, see [Steorts et al. \(2014\)](#) or [Papadakis et al. \(2020\)](#). Section 1.2 discusses existing R packages that implement blocking methods.

Reducing the number of pairs has inherent costs: missed comparisons lead to increased false positive rates (FPR) and false negative rates (FNR) in linkage studies. To validate

the blocking strategy, a subset of true pairs should be provided, or simulation studies of proposed methods should be conducted. Alternatively, one may consider approaches proposed by [Dasylva and Goussanou \(2021\)](#) and [Dasylva and Goussanou \(2022\)](#), who demonstrated how to estimate FPR and FNR without access to an audit sample.

1.2 Existing software and our contribution

The R ecosystem offers several packages that implement various blocking techniques, which we have grouped according to the following classification:

- **Deterministic blocking:**
 - [reclin2](#) ([van der Laan \(2024\)](#), [van der Laan \(2022\)](#)) allows pairing records using `pair_blocking()` with a prespecified list of columns in a `data.frame`, and the `pair_minsim()` function, which allows specifying the minimal similarity score (e.g., 1 out of 3 variable values must match exactly).
 - [RecordLinkage](#) ([Sariyar and Borg \(2025\)](#), [Sariyar and Borg \(2010\)](#)) allows specifying blocking variables in the `blockf1d` parameter of either `compare.dedup()` or `compare.linkage()` functions as a vector (either character or numeric).
 - [fastLink](#) ([Enamorado et al. \(2023\)](#), [Enamorado et al. \(2019\)](#)) implements various blocking methods via the `blockData()` function, including exact matching, window matching (e.g., no more than 2-year difference between birth years), and k -means clustering. Notably, `fastLink` returns datasets split into separate lists, whilst `reclin2` and `RecordLinkage` packages create a single dataset.
- **Phonetic blocking:**
 - [RecordLinkage](#) allows direct specification of phonetic comparison via the `phonetic` argument in `compare.dedup()` or `compare.linkage()` functions using the `soundex()` function.
 - Additionally, [stringdist](#) ([van der Loo, 2014](#)) implements the Soundex algorithm, whilst [phonics](#) ([Howard II, 2021](#)) implements various phonetic algorithms that can be applied prior to the blocking procedure (e.g., to create a new column).
- **Probabilistic blocking:**
 - [klsh](#) ([Steorts, 2020](#)) is the only R package that implements probabilistic blocking using the k -means variant of locality sensitive hashing. The main `klsh()` function implements this approach, and the resulting object is a list containing row identifiers for the prespecified number of blocks (via the `num.blocks` argument).

In practice, the situation is more complicated, as missing data may be present in blocking/matching variables (such as birth dates) or typos may occur in names and surnames. Therefore, we developed [blocking](#), which leverages approximate nearest neighbour (ANN) algorithms and graphs to create numerous small blocks that can be used in subsequent analysis. This approach is similar to micro-clustering (cf. [Johndrow et al. \(2018\)](#)), but we do not aim at providing the final linkage of units between or within sources. The basic workflow of the [blocking](#) package consists of the following steps:

1. Create shingles of the input character vectors using [tokenizers](#) ([Mullen et al., 2018](#)) and [text2vec](#) ([Selivanov et al., 2023](#)) packages, or provide a matrix of vectors (e.g., embeddings via [ragnar](#), [Kalinowski and Falbel \(2025\)](#)) representing the input character vectors.
2. Search for nearest neighbours using ANN algorithms implemented in [rnndescent](#) ([Melville, 2024b](#)), [RcppHNSW](#) ([Melville, 2024a](#)), [mlpack](#) ([Curtin et al. \(2023\)](#), [Singh Parihar et al. \(2025\)](#)), and [RcppAnnoy](#) ([Eddelbuettel, 2024](#)).
3. Create final blocks using [igraph](#) ([Csárdi et al. \(2025\)](#), [Csárdi and Nepusz \(2006\)](#)).

This is the only package in the R ecosystem that readily applies modern ANN algorithms to reduce the number of comparisons and substantially accelerate record linkage and deduplication tasks. Additionally, we have developed the `pair_ann()` function for seamless integration with the `reclink2` package, as described in the package vignette entitled "[Integration with existing packages](#)". However, the `blocking` package is not limited to the `reclink2` and can be applied in various record linkage or deduplication contexts.

1.3 Outline of the article

This paper is structured as follows. Section 2 provides a description of the main functionalities of the `blocking` package and how results can be evaluated. Section 3 presents two case studies: probabilistic record linkage and deduplication. These examples demonstrate how our package can improve the entity resolution pipeline and integrate with existing R packages.

2 Blocking of records using `blocking()` function

2.1 The main function

The main functionality is available via the `blocking()` function, which contains the following key arguments:

- `x`, `y` – vectors of keys used for linkage, where `y` = `NULL` indicates a deduplication task;
- `representation` – whether `x` and `y` should be represented as shingles or vectors (e.g., provided by the user via the `model` argument);
- `ann` – which ANN algorithm should be applied (by default, we use the `rnndescent` package as it supports sparse matrices);
- `distance` – which distance metric should be applied (default is cosine distance);
- `graph` – whether a plot of the graph showing connected records should be returned (default `FALSE`);
- `true_blocks` – if a subset of true blocks is available, it can be provided here so that quality measures, presented in the next section, are returned;
- `n_threads` – number of threads used for computation;
- `control_txt` – controls provided via `controls_txt()` specifying how `x`, `y` are processed;
- `control_ann` – controls provided via `controls_ann()` allowing users to fine-tune the ANN algorithm (see documentation for the `controls_ann()` function and `control_*` functions with names referring to specific algorithms, e.g., `control_nnd()` for the NND algorithm).

This function returns an object of class `blocking` containing the following elements:

- `result` – a `data.table` with indices (rows) of `x`, `y`, `block`, and distance between points;
- `method` – name of the ANN algorithm used;
- `deduplication` – information about whether deduplication was applied;
- `representation` – information about whether shingles or vectors were used;
- `metrics` – quality assessment metrics, if `true_blocks` is provided;
- `confusion` – confusion matrix, if `true_blocks` is provided;
- `colnames` – variable names (`colnames`) used for search;
- `graph` – an `igraph` class object.

2.2 Assessment of results

The package implements several measures that can be used to assess results. The first is the *reduction ratio* (RR), which indicates the reduction in comparison pairs within the given

blocks. It has a value between $[0, 1]$, where 1 indicates perfect reduction whilst values close to 0 indicate poor reduction. The RR indicator for deduplication has the following form:

$$\text{RR}_{\text{dedup}} = 1 - \frac{\sum_{i=1}^k \binom{|B_i|}{2}}{\binom{n}{2}},$$

where k is the total number of blocks, n is the total number of records in the dataset, and $|B_i|$ is the number of records in the i -th block. $\sum_{i=1}^k \binom{|B_i|}{2}$ is the number of comparisons after blocking, whilst $\binom{n}{2}$ is the total number of possible comparisons without blocking. For record linkage, the reduction ratio is defined as follows:

$$\text{RR}_{\text{reclin}} = 1 - \frac{\sum_{i=1}^k |B_{i,x}| \cdot |B_{i,y}|}{m \cdot n},$$

where m and n are the sizes of datasets X and Y , and k is the total number of blocks. The term $|B_{i,x}|$ is the number of records from dataset X in the i -th block, whilst $|B_{i,y}|$ is the number of records from dataset Y in the i -th block. The expression $\sum_{i=1}^k |B_{i,x}| \cdot |B_{i,y}|$ represents the number of comparisons after blocking.

Another way to assess blocking is to examine the confusion matrix at the *block* level, i.e., blocking results are compared with ground-truth *blocks* in a pairwise manner (e.g., one true positive pair occurs when both records from the comparison pair belong to the same predicted *block* and to the same ground-truth *block* in the evaluation data.frame). The values in this table are defined as follows:

- True positive (TP): record pairs correctly matched in the same block.
- False positive (FP): record pairs identified as matches that are not true matches in the same block.
- True negative (TN): record pairs correctly identified as non-matches (different blocks).
- False negative (FN): record pairs identified as non-matches that are true matches in the same block.

Metrics calculated based on this confusion matrix are presented in Table 1.

Table 1: Evaluation Metrics

Metric	Formula	Metric	Formula
Recall	$\frac{TP}{TP+FN}$	Accuracy	$\frac{TP+TN}{TP+TN+FP+FN}$
Precision	$\frac{TP}{TP+FP}$	Specificity	$\frac{TN}{TN+FP}$
F1 Score	$2 \cdot \frac{\text{Precision} \times \text{Recall}}{\text{Precision} + \text{Recall}}$	False Positive Rate	$\frac{FP}{FP+TN}$
False Negative Rate	$\frac{FN}{FN+TP}$		

3 Case studies

3.1 An example of blocking for record linkage

Let us first load the required packages.

```
library("blocking")
library("data.table")
library("reclin2")
```

We demonstrate the use of the `blocking()` function for record linkage on the `foreigners` dataset included in the package. This fictional representation of the foreign population in Poland was generated based on publicly available information, preserving the distributions from administrative registers. It contains 110,000 rows with 100,000 entities (thus containing 10,000 duplicates). Each row represents one record, with the following columns: `fname` – first name, `sname` – second name, `surname` – surname, `date` – date of birth, `region` – region (county), `country` – country, and `true_id` – a person identifier.

Next, we load the data and examine the first six records.

```
data("foreigners")
setDT(foreigners)
head(foreigners)

#>      fname  sname    surname      date region country true_id
#>      <char> <char>    <char>    <char> <char>   <num>
#> 1:  emin           imanov 1998/02/05       031      0
#> 2: nurlan         suleymanli 2000/08/01       031      1
#> 3: amio          maharrsmov 1939/03/08       031      2
#> 4: amik          maharramof 1939/03/08       031      2
#> 5: amil          maharramov 1993/03/08       031      2
#> 6: gadir         jahangirov 1991/08/29       031      3
```

In the next step, we split the dataset into two separate data.frames: one containing the first appearance of each entity in the `foreigners` dataset, and the other containing subsequent appearances. We then add row identifiers (`x` and `y`).

```
foreigners_1 <- foreigners[!duplicated(foreigners$true_id), ]
foreigners_1[, x := 1:N]
foreigners_2 <- foreigners[duplicated(foreigners$true_id), ]
foreigners_2[, y := 1:N]
```

Now, in both datasets we remove separators from the `date` column and create a new character column that concatenates information from all columns (excluding `true_id`) in each row. Information stored in the `txt` column will be used for blocking records in the `blocking()` function.

```
foreigners_1[, txt := paste0(fname, sname, surname, gsub("/", "", date),
                             region, country)]
foreigners_2[, txt := paste0(fname, sname, surname, gsub("/", "", date),
                             region, country)]
head(foreigners_1[, .(true_id, txt)])

#>      true_id          txt
#>      <num>        <char>
#> 1:      0  eminimanov19980205031
#> 2:      1  nurlansuleymanli20000801031
#> 3:      2  amiomaharrsmov19390308031
#> 4:      3  gadirjahangirov19910829031
#> 5:      4  zaurbayramova1996100601261031
#> 6:      5  asifmammadov19970726031
```

The default algorithm is the Nearest Neighbour Descent Method (Dong et al., 2011) implemented in the `rnnndescent` package. Note that a default parameter of the `blocking()` function is `seed = 2023`, which sets the random seed.

```
result_recln <- blocking(x = foreigners_1$txt,
                           y = foreigners_2$txt)
```

Now, we can examine the results by printing the `result_reclin` object. In this example, we have created 6,470 blocks utilizing 1,232 2-character shingles. Blocks are small, as we have 3,920 blocks of 2 elements, 1,599 blocks of 3 elements, ..., 2 blocks of 7 elements.

```
result_reclin

#> =====
#> Blocking based on the nnd method.
#> Number of blocks: 6470.
#> Number of shingles created for blocking: 1232.
#> Reduction ratio: 0.9999.
#> =====
#> Distribution of the size of the blocks:
#>   2   3   4   5   6   7
#> 3920 1599 928 19   2   2
```

To access the result, one should use `result_reclin$result`. The resulting `data.table` has four columns (as presented below):

- `x` – reference dataset (i.e., `foreigners_1`) – this may not contain all units of `foreigners_1`;
- `y` – query (each row of `foreigners_2`) – this will contain all units of `foreigners_2`;
- `block` – the block identifier;
- `dist` – distance between pairs.

```
head(result_reclin$result)

#>      x     y block     dist
#>    <int> <int> <num>     <num>
#> 1:    3     1    1 0.2216882
#> 2:    3     2    1 0.2122737
#> 3:   21     3    2 0.1172652
#> 4:   57     4    3 0.1863238
#> 5:   57     5    3 0.1379310
#> 6:   61     6    4 0.2307692
```

Let's examine the first block. Clearly, there are typos in the `fname` and `surname`. Nevertheless, all records refer to the same entity (as denoted by `true_id`).

```
rbind(foreigners_1[3, 1:7], foreigners_2[1:2, 1:7])

#>   fname  surname      date region country true_id
#>   <char> <char>      <char> <char> <char>   <num>
#> 1: amio    maharrsmov 1939/03/08        031      2
#> 2: amik    maharramof 1939/03/08        031      2
#> 3: amil    maharramov 1993/03/08        031      2
```

Now we use the `true_id` column to evaluate our approach.

```
matches <- merge(x = foreigners_1[, .(x, true_id)],
                  y = foreigners_2[, .(y, true_id)],
                  by = "true_id")
matches[, block := rleid(x)]
head(matches)

#> Key: <true_id>
#>   true_id     x     y block
```

```
#>      <num> <int> <int> <int>
#> 1:      2     3     1     1
#> 2:      2     3     2     1
#> 3:     20    21     3     2
#> 4:     56    57     4     3
#> 5:     56    57     5     3
#> 6:     60    61     6     4
```

We have 10,000 matched pairs, which can be used in the `true_blocks` argument of the `blocking()` function to specify the true block assignments. This `data.frame` must contain three columns: `x`, `y`, and `block`.

We obtain quality metrics for the assessment of record linkage.

```
res_reclin <- blocking(x = foreigners_1$txt,
                        y = foreigners_2$txt,
                        true_blocks = matches[, .(x, y, block)])
```

`res_reclin`

```
#> =====
#> Blocking based on the nnd method.
#> Number of blocks: 6470.
#> Number of shingles created for blocking: 1232.
#> Reduction ratio: 0.9999.
#> =====
#> Distribution of the size of the blocks:
#>   2   3   4   5   6   7
#> 3920 1599 928  19   2   2
#> =====
#> Evaluation metrics (standard):
#>   recall  precision      fpr      fnr  accuracy specificity
#>   96.7532    78.6700    0.0038    3.2468    99.9957    99.9962
#>   f1_score
#>   86.7795
```

For example, our approach results in a 3.25% FNR. To improve this, we can increase the `epsilon` parameter of the NND method from 0.1 to 0.5. To do so, we configure the `control_ann` parameter in the `blocking()` function using the `controls_ann()` and `control_nnd()` functions.

```
res_reclin2 <- blocking(x = foreigners_1$txt,
                        y = foreigners_2$txt,
                        true_blocks = matches[, .(x, y, block)],
                        control_ann = controls_ann(nnd = control_nnd(epsilon = 0.5)))
```

`res_reclin2`

```
#> =====
#> Blocking based on the nnd method.
#> Number of blocks: 6394.
#> Number of shingles created for blocking: 1232.
#> Reduction ratio: 0.9999.
#> =====
#> Distribution of the size of the blocks:
#>   2   3   4   5   7
#> 3800 1615 954  21   4
```

```
#> =====
#> Evaluation metrics (standard):
#>   recall    precision      fpr      fnr  accuracy specificity
#>   96.8776    80.0500    0.0036    3.1224    99.9960    99.9964
#>   f1_score
#>   87.6636
```

That decreases the FNR to 3.12%.

Now, to use the result in the record linkage process, we add this information to both datasets and specify it in the appropriate argument of a given function. Below, we present an example using the `reclink2` package, where a simple score is calculated as the sum of Jaro-Winkler similarity measures (cf. [Winkler \(1990\)](#)) for the variables `fname`, `surname`, and `date`.

```
foreigners_1[res_reclink2$result, on = "x", block:= i.block]
foreigners_2[res_reclink2$result, on = "y", block:= i.block]

pair_blocking(x = foreigners_1,
              y = foreigners_2, on = "block") |>
  compare_pairs(on = c("fname", "surname", "date"),
                default_comparator = cmp_jarowinkler()) |>
  score_simple("score", on = c("fname", "surname", "date")) |>
  head(n= 4)

#> First data set: 100 000 records
#> Second data set: 10 000 records
#> Total number of pairs: 4 pairs
#> Blocking on: 'block'
#>
#>   .x     .y     fname   surname   date   score
#>   <int> <int>   <num>   <num>   <num>   <num>
#> 1:     3      1 0.8333333 0.8666667 1.0000000 2.700000
#> 2:     3      2 0.8333333 0.9333333 0.9666667 2.733333
#> 3:    21      3 0.8333333 0.9166667 1.0000000 2.750000
#> 4:    57      4 0.9259259 0.9259259 0.9666667 2.818519
```

3.2 An example of blocking for deduplication

In this section, we demonstrate a deduplication application using the `blocking()` function on the `RLdata500` dataset from the `RecordLinkage` package. Note that the dataset is included in the `blocking` package. It contains artificial personal data, and fifty records have been duplicated with randomly generated errors. Each row represents one record, with the following columns: `fname_c1` – first name, `fname_c2` – second name, `lname_c1` – last name, `lname_c2` – last name (second component), `by`, `bm`, `bd` – year, month, and day of birth, `rec_id` – record ID, and `ent_id` – entity ID.

```
data("RLdata500")
setDT(RLdata500)
head(RLdata500)

#>   fname_c1 fname_c2 lname_c1 lname_c2   by   bm   bd rec_id ent_id
#>   <char>   <char>   <char>   <char> <int> <int> <int> <int> <int>
#> 1: CARSTEN           MEIER       1949     7   22     1    34
#> 2: GERD             BAUER       1968     7   27     2    51
#> 3: ROBERT          HARTMANN    1930     4   30     3   115
#> 4: STEFAN          WOLFF       1957     9    2     4   189
```

```
#> 5:      RALF          KRUEGER        1966     1     13      5     72
#> 6: JUERGEN        FRANKE        1929     7      4      6    142
```

For the purpose of this example, we create a new column (`id_count`) that indicates how many times a given unit occurs, and then add leading zeros to the `bm` and `bd` columns. Finally, we create a new string column that concatenates information from all columns (excluding `rec_id`, `ent_id`, and `id_count`), as presented below.

```
RLdata500[, id_count := .N, ent_id]
RLdata500[, txt:=tolower(paste0(fname_c1,fname_c2, lname_c1, lname_c2, by,
                                sprintf("%02d", bm), sprintf("%02d", bd)))]
head(RLdata500[, .(rec_id, id_count, txt)])

#>   rec_id id_count           txt
#>   <int>   <int>           <char>
#> 1:     1       1 carstenmeier19490722
#> 2:     2       2 gerdbauer19680727
#> 3:     3       1 roberthartmann19300430
#> 4:     4       1 stefanwolff19570902
#> 5:     5       1 ralfkrueger19660113
#> 6:     6       1 juergenfranke19290704
```

As in the previous example, we use the `txt` column in the `blocking()` function. This time, we set `ann = "hnsw"` to use the Hierarchical Navigable Small World (HNSW; [Malkov and Yashunin \(2018\)](#)) algorithm from the [RcppHNSW](#) package.

```
res_dedup <- blocking(x = RLdata500$txt,
                        ann = "hnsw",
                        verbose = 1)

#> ===== creating tokens =====
#> ===== starting search (hnsw, x, y: 500, 500, t: 429) =====
#> ===== creating graph =====
```

The results are as follows. This time, the HNSW algorithm provided blocks varying from 2 to 17 units.

```
res_dedup

#> =====
#> Blocking based on the hnsw method.
#> Number of blocks: 133.
#> Number of shingles created for blocking: 429.
#> Reduction ratio: 0.9916.
#> =====
#> Distribution of the size of the blocks:
#>  2  3  4  5  6  7  8  9 10 11 12 17
#> 46 35 23  8  6  6  2  3  1  1  1  1
```

Next, we create a long `data.table` with information on blocks and units from the original dataset. We add the block information to the final dataset. We can check in how many blocks the same entities (`ent_id`) are observed. In our example, all identical entities are in the same blocks.

```
# Reshape to long format
df_block_melted <- melt(res_dedup$result, id.vars = c("block", "dist"))
```

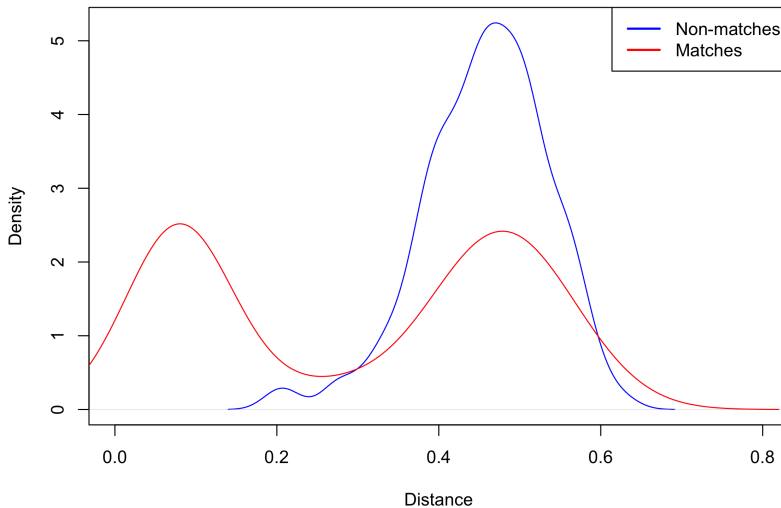


Figure 1: Distribution of distances between true matches and non-matches within blocks

```
# Unique record-block pairs
df_block_melted_rec_block <- unique(df_block_melted[, .(rec_id=value, block)])
# Join block IDs
RLdata500[df_block_melted_rec_block, on = "rec_id", block_id := i.block]
# Check block consistency per entity
RLdata500[, .(uniq_blocks = uniqueN(block_id)), .(ent_id)][, .N, uniq_blocks]

#>     uniq_blocks      N
#>           <int> <int>
#> 1:          1    450
```

Additionally, we visualise the result based on whether a block contains matches or not.

Finally, we compare the evaluation metrics across all ANN algorithms supported by the `blocking()` function, i.e., NND, HNSW, Annoy (from the [RcppAnnoy](#) package), Locality-Sensitive Hashing (LSH, from the [mlpack](#) package), and k -Nearest Neighbours (k NN – denoted as "kd", from the [mlpack](#) package). We use the `rec_id` and `ent_id` columns from the `RLdata500` dataset to specify the true blocks and then calculate evaluation metrics for all algorithms.

We compare our package with the `klsh()` function from the [klsh](#) package, configured to create 10 blocks (denoted as `klsh_10`) and 100 blocks (denoted as `klsh_100`), respectively. In both settings, we use 20 random projections and 2-character shingles. The results are presented in Table 2.

```
set.seed(2025)
true_blocks <- RLdata500[, c("rec_id", "ent_id"), with = FALSE]
setnames(true_blocks, old = c("rec_id", "ent_id"), c("x", "block"))
eval_metrics <- list()
ann <- c("nnd", "hnsw", "annoy", "lsh", "kd")
for (algorithm in ann) {
  eval_metrics[[algorithm]] <- blocking(
    x = RLdata500$txt,
    ann = algorithm,
    true_blocks = true_blocks)$metrics
}
blocks_klsh_10 <- klsh::klsh(
```

Table 2: Comparison of various approximate nearest neighbour algorithms implemented in the `blocking` and the `klsh` package for creation of blocks for deduplication (values in percentages)

	recall	precision	fpr	fnr	accuracy	specificity	f1_score
nnd	100	5.17	0.74	0	99.26	99.26	9.83
hnsw	100	4.76	0.80	0	99.20	99.20	9.08
annoy	100	4.80	0.79	0	99.21	99.21	9.17
lsh	98	1.04	3.74	2	96.26	96.26	2.06
kd	100	5.19	0.73	0	99.27	99.27	9.87
klsh_10	84	0.33	10.13	16	89.87	89.87	0.66
klsh_100	90	3.72	0.94	10	99.06	99.06	7.14

```
r.set = RLdata500[, c("fname_c1", "fname_c2", "lname_c1",
                      "lname_c2", "by", "bm", "bd")],
p = 20, num.blocks = 10, k = 2)

klsh_10_metrics <- klsh::confusion.from.blocking(
  blocking = blocks_klsh_10,
  true_ids = RLdata500$ent_id)[-1]

klsh_10_metrics$f1_score <- with(klsh_10_metrics,
                                    2*precision*recall/(precision + recall))

eval_metrics$klsh_10 <- unlist(klsh_10_metrics)

blocks_klsh_100 <- klsh::klsh(
  r.set = RLdata500[, c("fname_c1", "fname_c2", "lname_c1",
                        "lname_c2", "by", "bm", "bd")],
  p = 20, num.blocks = 100, k = 2)

klsh_100_metrics <- klsh::confusion.from.blocking(
  blocking = blocks_klsh_100,
  true_ids = RLdata500$ent_id)[-1]

klsh_100_metrics$f1_score <- with(klsh_100_metrics,
                                    2*precision*recall/(precision + recall))

eval_metrics$klsh_100 <- unlist(klsh_100_metrics)

round(do.call(rbind, eval_metrics) * 100, 2) |> kable(digits=2)
```

The results demonstrate a clear performance hierarchy among the ANN algorithms implemented in the `blocking` package, with traditional tree-based methods (NND, HNSW, Annoy, and kNN) achieving perfect recall (100%) whilst maintaining excellent precision and F1 scores around 5-10%. Notably, these methods exhibit minimal FPR (0.73-0.80%) and maintain high specificity (99.20-99.27%), indicating their effectiveness in creating tight, accurate blocks.

In contrast, the LSH-based methods show more variable performance: the `mlpack` LSH implementation achieves 98% recall but suffers from higher FPR (3.74%) and, importantly, FNR (2%), whilst the `klsh` package results reveal a trade-off between block granularity and performance – `klsh_10` with only 10 blocks shows poor recall (84%), high FPR (10.13%), and FNR (16%), whereas `klsh_100` with 100 blocks recovers much of the performance (90% recall, 0.94% FPR, but high FNR of 10%). This indicates that the `klsh`, particularly these implementations of the LSH approach, misses a large number of true matches.

These findings suggest that modern ANN algorithms like NND, HNSW, and Annoy provide superior blocking performance for entity resolution tasks, offering both computational

efficiency and high-quality results that minimise both missed matches and false linkages.

4 Summary

In this paper, we have demonstrated the basic use cases of the **blocking** package. We believe that the software will be useful for researchers working in various fields where integration of multiple sources is an important aspect. This is certainly of interest in the field of official statistics, where register-based statistics rely on high-quality linkage of administrative datasets, or medical studies, where assessment of health statistics relies on correct linkage of medical history with treatment outcomes or mortality records.

Furthermore, for users interested in integration with the **reclin2** package, we refer to the documentation of the `pair_ann()` function and the vignette entitled "[Integration with existing packages](#)", which provides case studies demonstrating how the **blocking** package can be included in existing record linkage/deduplication pipelines.

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We have also developed a Python version of the package, **BlockingPy**, which is available through PyPI. It has a similar structure but offers more ANN algorithms (e.g., FAISS) and enables the use of embeddings. For more details, see: Strojny, T., & Beresewicz, M. (2025). BlockingPy: Approximate nearest neighbours for blocking of records for entity resolution. arXiv preprint arXiv:2504.04266.

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