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Quality Indicators for Administrative Data¹ User Manual For R Package on GitHub

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1. Project aims and objectives

The Office for National Statistics (ONS) have strategic priorities on embedding and advancing the use of administrative data into their official statistics processes. Their immediate priority is to use administrative data in the quality assurance of the 2021 census and to develop the production of administrative-based population estimates (ABPEs). A more long-term priority is the Population Statistics Transformation Programme which will feed into a recommendation to Government due in 2023 on the future of census and population statistics. In particular, the objective is to create population characteristic estimates from administrative and integrated data sources.

Motivated by these initiatives, the University of Manchester was successfully awarded an ESRC grant from April 2021 to January 2023 titled: 'Methodological Advancements on the Use of Administrative Data in Official Statistics' (ES/V005456/1). The funded research enabled collaborations with the ONS on researching and developing methods to enhance the use of administrative data in official statistics.

This user manual concerns the sub-project related to developing a quality framework and quantitative measures to assess representativeness and coverage for a single administrative data source. We use univariate and bivariate distributions obtained from high- quality large random probability surveys (such as the UK Annual Population Survey) and compare them to distributions in the administrative data on a common set of variables based on distance metrics between these distributions. In addition, we developed a Representativity (R-) Indicator that is designed for quantifying the representativeness of population groups in the administrative data. Section 2 describes the methods underpinning the R-code hosted in GitHub and Section 3 describes the R-code with a running example of its outputs. We conclude in Section 4 with troubleshooting questions and answers.

2. Quality indicators

In the R-code on GitHub we focus on quality indicators to identify errors arising from coverage and representativeness of a single administrative data source. It is vital that statistical agencies have good quality indicators to ensure the fit of administrative data to the population and to identify those sub-groups that are missing or over-covered, especially when the administrative data contributes to an integrated dataset for multisource processing or to quality assure other data sources, such as surveys and censuses.

2.1 Distance Metrics

In this section we compare distributions obtained from the administrative data with external population auxiliary information, either obtained directly from a census or estimated from a large probability-based random survey.

Denote variable v having categories $h, h = 1, 2 \dots H$ in the administrative dataset having M individuals. Let $\Delta_{h,i}^{v}$ be the 0-1 indicator for individual i being a member of category h in variable v. We can then calculate the counts for category $h: m_{h}^{v} = \sum_{i=1}^{M} \Delta_{h,i}^{v}$ and note that $\sum_{h=1}^{H} m_{h}^{v} = M$. We can also obtain the probability distribution of variable v having category $h, h = 1, 2 \dots H$ and calculate: $p_{h}^{v} = \frac{m_{h}^{v}}{M}$. We note that the variable v can also represent a cross-tabulation of two or more variables, for example age group \times sex.

Now assume we have equivalent estimates of these distributions from a census, or alternatively from a large probability-based random sample of size n where every individual i in the sample has an associated survey weight w_i . The survey weights typically are calibrated to known population benchmarks and hence sum to the known population size N. In this case, $n_h^v =$

 $\sum_{i=1}^{n} w_i \Delta_{h,i}^{v}$ and $\sum_{h=1}^{H} n_h^{v} = N$. Moreover, the equivalent distribution is $q_h^{v} = \frac{n_h^{v}}{N}$.

The entropy measures the uniformity of the probability distributions and hence can be compared when calculated on the administrative data distribution versus the population distribution. The formula for the entropy on the probability distribution $\{p_h^v, h = 1, ..., H\}$ is: $\sum_h p_h^v \log(p_h^v)$ and similarly on the probability distribution $\{q_h^v, h = 1, ..., H\}$.

We can now use a variety of distance metrics to assess deviations between the distributions $\{p_h^v, h = 1, ..., H\}$ and $\{q_h^v, h = 1, ..., H\}$. In this research, we assess three distance metrics: the Indicator of Dissimilarity (Duncan and Duncan, 1955), Hellinger's Distance (HL) and the Kullback-Leibler divergence (KL). The formula for the three distance metrics is given in Table 1.

Table 1: Distance metrics between the distribution calculated from the administrative data $\{p_h^v, h = 1, ..., H\}$ and the distribution from the population $\{q_h^v, h = 1, ..., H\}$ on variable v

Distance Metrics	Formula	Standarize
Indicator of Dissimilarity (ID)	$\frac{1}{2}\sum\nolimits_{h} p_{h} - q_{h} $	1-ID
Hellinger's Distance (HL)	$\frac{1}{\sqrt{2}}\sqrt{\sum_{h} \left(\sqrt{p_{h}} - \sqrt{q_{h}}\right)^{2}}$	1-HL

Kullback-Leibler Divergence (KL)

There are subtle differences between these distance metrics. For example, Hellinger's Distance places more weight on the smaller proportions compared to the larger proportions whereas the Indicator of Dissimilarity treats all proportions equally. More work is needed on standardizing the distance metrics into meaningful quality measures. We have yet to determine which distance metric should be used and therefore propose to include all of them in the R-code. This will facilitate more empirical work for future recommendations.

2.2 R-indicators

The R-indicator and its related partial R-indicators were originally designed to assess the representativeness of responses from a survey and are particularly useful as an objective function in the optimization of adaptive survey designs (Schouten, et al. 2009, Schouten and Shlomo, 2017). The R-indicators measure the contrast between those who are missing and not missing in the data and identify those groups that are not represented in the data. Here, we develop the R-indicator and partial R-indicators to assess the representativeness and coverage of an administrative dataset compared to a target population. Recent research by Bianchi, et al. (2019) adapts the R-indicator to the case where only population-based auxiliary information are available instead of sample-based frame information. We draw upon this research and utilize population-based auxiliary information where the population auxiliary information is obtained by weighted survey counts from a large probability-based random sample.

To calculate the population-based R-indicator, denote the response indicator r_i equal to 1 for all units in the administrative dataset. We have information available on the values $x_i = (x_{1,i}, x_{2,i}, ..., x_{K,i})^T$ of a vector of K auxiliary variables X, for example, sex, age group, geographical region, ethnic minority group and employment status. Therefore, each $x_{k,i}$ is a binary indicator variable. We also assume that values of x_i are observed for all individuals in the administrative dataset so that $\{x_i; i \in r\}$ is observed.

Assume we know x_i at the aggregate level: the population total $\sum_U x_i$ and population crossproducts $\sum_U x_i x_i^T$. This information is known as the population–based auxiliary information. If this information is not available at the population level we can estimate the aggregates and crossproducts using a large probability-based random sample, denoted s, where each individual i in the sample has a survey weight w_i . The estimated population-based auxiliary information is then: $\sum_s w_i x_i$ and $\sum_s w_i x_i x_i^T$. We also know the overall total in the population, denoted by N.

Response propensities are defined as the conditional expectation of the response indicator variable r_i given the values of specified variables: $\rho_i \equiv \rho_X(x_i)$. In the population-based setting

we model the response propensities under an identity link function where the true response propensities satisfy: $\rho_i = x_i^T \beta$, $i \in U$. For the linear probability model, the estimate of ρ_i in the sample-based scenario is given by: $\hat{\rho}_i^{OLS} = x_i^T (\sum_{s'} d_i x_i x_i^T)^{-1} \sum_{s'} d_i x_i r_i$, $i \in s'$, where d_i is the design weight and s' denotes the sample under analysis.

In the case of population-based auxiliary information where we know both population totals and cross-products, we note that $\sum_{s'} d_i x_i$ and $\sum_{s'} d_i x_i x_i^T$ are unbiased estimates for $\sum_U x_i$ and $\sum_U x_i x_i^T$, respectively and that in large samples we may expect that $\sum_{s'} d_i x_i \approx \sum_U x_i$ and $\sum_{s'} d_i x_i x_i^T \approx \sum_U x_i x_i^T$. It follows that, in the population-based setting, we may approximate $\hat{\rho}_i^{OLS}$ by $\hat{\rho}_i^P = x_i^T (\sum_U x_i x_i^T)^{-1} \sum_r d_i x_i$, $i \in r$ and we refer to the propensities as 'participation' propensities. Note that $\hat{\rho}_i^P$ is computed only on the set of individuals in the administrative data. In addition, in our setting of assessing the representativeness in administrative data, the design weight d_i is the inverse of the coverage weight: $d_i = [M/N]^{-1}$ where M is the number of individuals in the administrative dataset.

In the population-based setting, an estimator for the R-indicator is given by $\hat{R}_{\hat{\rho}^P} = 1 - 2\hat{S}_{\hat{\rho}^P}$ where $\hat{S}_{\hat{\rho}^P}^2 = \frac{N}{N-1} \{\frac{1}{N} \sum_r d_i \hat{\rho}_i^P - \left[\frac{1}{N} \sum_r d_i\right]^2\}$ and $\hat{\rho}_i^P$ is estimated as above. This estimator of the R-indicator makes the estimator $\hat{S}_{\hat{\rho}}^2$ linear in $\hat{\rho}_i^P$ which provides an advantage for size bias adjustment computations (although given the large administrative datasets, a size bias adjustment is not needed). Furthermore, we use propensity weighting by $\hat{\rho}_i^{P^{-1}}$ to adjust for coverage bias. The R-indicator measures the variation of the sub-group participation propensities are all equal and there is no variation in sub-group participation, the R-indicator would obtain a value of 1.

The unconditional partial R-indicator measures the amount of variation of the participation propensities between the categories of a variable. The larger the between-category variation is, the stronger the relationship is and the stronger the impact of the variable on a lack of representativeness. As earlier, let x_k be one of the components of the vector X. The variable x_k is categorical and assume it has H categories. Let m_h denote the weighted respondent size in category h in the administrative data, for h = 1, 2, ..., H. That means $m_h = \sum_{i \in r} d_i \Delta_{h,i}$ where $\Delta_{h,i}$ is the 0-1 indicator for participating unit *i* being a member of category h and $\sum_{h=1}^{H} m_h = N$ given the definition of d_i as the inverse coverage weight. Define $\hat{\rho}_h$ the average of the participation propensities in category h of x_k for the units in the administrative dataset and $\hat{
ho}$ the overall average participation probability based on the estimated population-based participation propensities $\hat{\rho}_i^P$. The estimate for the unconditional partial R-indicator for variable x_k is: $R_U(x_k) = \sqrt{\frac{1}{N} \sum_{h=1}^{H} m_h(\hat{\rho}_h - \hat{\rho})^2}$. The upper bound of the unconditional partial Rindicator is 0.5. The larger the value of the partial R-indicator, the stronger the association of the variable with a lack of representativeness in the administrative dataset. By computing and comparing the unconditional partial indicators for a set of variables it can be established for which variables the relationships are strongest. The unconditional partial R-indicator at the

category level h for variable x_k is $R_U(x_k^h) = \sqrt{\frac{m_h}{N}} (\hat{\rho}_h - \hat{\rho})$ and can assume positive and negative values. Note that at the category-level, a negative sign represents under-representation and a plus sign represents over-representation.

Finally, we note that when producing estimates from the administrative dataset, one should weight each individual *i* by its inverse participation propensity: $\hat{\rho}_i^{P^{-1}}$ to adjust for coverage bias in the estimates.

3. User Guide on the R-package

This package assumes that there is a Census file to obtain auxiliary population estimates and the administrative datasets under analysis. However, in real settings we would not have a Census microdata to work with, rather we would have a large probability-based survey sample for estimating population distributions. Therefore, we draw a random sample from the Census microdata to support this scenario and obtain auxiliary population totals from weighted sample counts.

3.1 Download and inspect the contents

3.1.1 Download

Please visit the github site here: qualadmin link. Click on Code at the top-right corner. Then, click on Download ZIP to download to your local machine.

Now, the downloaded folder needs to be placed in the designated location. We recommend users decide the appropriate Drive (C, D, E, F, etc) to house the downloaded contents. Then, **create a new folder** called admindata in File Explorer of your PC. Users can customise the new folder name as appropriate. This is your **starting path**.

The screenshot showing starting path:



Under this Starting path, F:/admindata, place the downloaded folder from GitHub. Extract the zip folder as necessary.

As such, F:/admindata/qualadmin becomes the MASTER project folder. We'll set it as working directory¹ in RStudio later.

3.1.2 Downloaded contents explained

Example datasets

We provide two example data sources.

Data type	File name
Administrative	public_release_admin.csv
Census	pop_u_short_public_release_5vars.Rdata

Folders and R scripts

Under F:\admindata\qualadmin folder, you'll be presented with the following contents.

Functions
User_manual
0_Custom_Path.R
1_Create_Folders.R
1_Install_Packages.R
2_Prep_Wtsample_Freq_Table.R
3_A_Distance_Metrics.R
3_B_R-indicator.R
3_MASTER_Run_AB.R
pop_u_short_public_release_5vars.RData
public_release_admin.csv

The **"User_manual"** folder contains instructions on using the provided R code files.

The users do not need to do anything with the folder titled **"Functions"**. These pre-defined functions are used to either enclose complex procedures or perform repetitive tasks including cleaning and computing quality indicators. There are two files containing pre-defined functions. There is no need to run function files independently.

The functions will be automatically called in when the three main R script files are run: 2_Prep_Wtsample_Freq_Table.R 3_A_Distance_Metrics.R 3_B_R-indicator.R

The 2_Prep_Wtsample_Freq_Table.R file creates necessary data needed to compute distance metrics and R-indicators. The master file, 3_MASTER_Run_AB.R runs the above *two* main R script files, (3_A_Distance_Metrics.R 3_B_R-indicator.R) automatically in sequence.

¹ Notice that the terms, folder, directory, and path are used interchangeably in the user manual.

The first three files, O_Custom_Path.R, 1_Create_Folders.R and 1_Install_Packages.R can be run to get ready to run the above main analysis files, as discussed in the following section.

3.2 Launch RStudio and get ready

3.2.1 Open the entire master folder in RStudio

First, launch RStudio. Then, we need to **open the entire folder** F:/admindata/qualadmin where downloaded materials are located.

Unfortunately, RStudio has no feature in the menu, but you could do so by accessing **Files** tab. Click on ... as shown below.

Untitled1* ×	Console	Ter
	R R 4.	1.1 ·
2	>	
3		
4		
5		
3:1 (Top Level)		
Environment History Files Plots Packages Help F		
🔮 🗸 😂 📑 🌼 - 📿		7
□ > C:	$\langle \langle \rangle$	
▲ Name Size Modifie Got	to directory	
1		
🗍 🧰 \$Recycle.Bin		T
	Connectiv	nne

Then, locate the master folder. In our example, it is F:/admindata/qualadmin.

3.2.2 Set custom path

Click open the R script file, O_Custom_Path.R. Customise the starting path as needed, and set the path to indicate the master folder. The example code is:

```
# Starting path (CUSTOMISE PLEASE)
setwd("F:/admindata")
# Master project folder (USE AS IT IS)
setwd("./qualadmin")
# Check your current directory
getwd()
```

Please ensure to use a single forward slash / as above. R will print an error when backward slash $\$ is used in path. For instance,

```
setwd("F:\admindata)`
Error: '\a' is an unrecognized escape in character string starting ""F:\a
```

Please ensure your working directory is set at the master project path throughout the analytical steps.

3.2.3 Automatically create output folders

The three main R script files 2_Prep_Wtsample_Freq_Table.R, 3_A_Distance_Metrics.R, 3_B_R-indicator.R produce outputs. The outputs may be text, figure or in spreadsheet form. For the existing programmes to work, users need to create dedicated output folders.

To do so, please click on the 1_Create_Folders.R file to open. Then run line by line. The resulting folder structure is provided here:



3.2.4 Install packages

The final preparation step is installing packages. Open 1_Install_Packages.R file, and run line by line.

```
#-----
# Install packages (Run once)
#-----
install.packages("ggplot2")
install.packages("tidyverse")
install.packages("car")
```

Now, you're all set to proceed with quality measures indicators!

3.3 RUNNING 2_Prep_Wtsample_Freq_Table.R

This code file calculates the distributions of the weighted sample data. If the Census data or a weighted sample data are available, users consult 3.5.1 *Use the existing sample data*. For a scenario where these data are unavailable, users can generate the data as shown in 3.5.2 *Generate a weighted sample data*.

Open the 2_Prep_Wtsample_Freq_Table.R file.

The top of the code file concerns checking the current directory, reading in the pre-defined functions in the R environment and loading relevant R libraries.

```
getwd()
# Run the code file with functions.
source("Functions/1_Functions.R")
# Define output file folders, path
fn_output_folder_path()
# Disable scientific notation.
options(scipen = 999)
library("tidyverse") # data manipulation
library("ggplot2") # visualisation
library("janitor") # cross-tabulation
library("readxl") # read large csv file
library("writexl") # export to Excel
```

3.3.1 Use the existing weighted sample data

```
• Step 1: Read in the data.
```

```
df <- read_csv("custom_wtsample.csv")
dim(df)  # obs = 1163650 (example)
glimpse(df)  # Quick glance at the data</pre>
```

Users decide which variables are to be used for tabulation. For example, users may identify the following five variables.

```
names(df) # variable names
[1] "geog1" "sex" "agecode1" "eth_code5"
[5] "econg"
```

• Step 2: Declare variables to be tabulated. Here, we declare all five variables using var object². Users can customize the variable names here. By running the code below, R automatically saves the total number of variables, 5 in a macro called maxvar.

• Step 3: Obtain frequency table of categorical variables (count of categories).

This procedure is to assess and calculate *the distribution* of categories in the weighted sample. Users can run the pre-defined function, fn_maxvar5_freq_table() to perform the task. The function automatically obtains counts and structure the output in long form, organised by each variable, and by its discrete category. In case of using four variables, users can use fn_maxvar4_freq_table() instead³.

```
fn_maxvar5_freq_table()
```

• Step 4: Carry out checks to see if the calculated frequency tables are accurate. freq_table[1:8, 1:9]

##		seq	twdigits	n	р	oneway v	by1	by2
##	1	1	101	113250	0.09732308	3 11	geog1	01
##	2	2	102	148400	0.12752976	5 11	geog1	02
##	3	3	103	137450	0.11811971	. 11	geog1	03
##	4	4	104	175100	0.15047480) 11	geog1	04
##	5	5	105	92300	0.07931938	3 11	geog1	05
##	6	6	106	497150	0.42723327	11	geog1	06
##	7	7	201	565350	0.48584196	5 12	sex	01
##	8	8	202	598300	0.51415804	12	sex	02

When we printed the first 8 lines and 10 variables, we can see the count, n, and the corresponding proportion, p by each variable. The following code obtains the total observation size and confirms that the total proportion adds up to 1, for geog1 variable. Here, the total observation size can be viewed as the population size.

```
# Check whether the total adds up to 1
sum(freq_table[1:6, "n"])
```

² As the var object is treated as a global macro, the programme runs automatically using the information stored in global macro, and produces the results. Users can save var in a separate RData file and load it in each session, instead of repeating the procedure.

³ We do not provide functions for the maximum variable size beyond five. In such cases, users can create their own functions by consulting the provided functions.

[1] 1163650

```
sum(freq_table[1:6, "p"])
```

[1] 1

- Step 5: Rename and save.
- Step 6: Export the output frequency table in Excel with the file name, Weightedsample_freq_table.xlsx.
- Step 7: Save the R objects as RData. Done.

3.3.2 Generate a weighted sample data

• Step 1: Load a Census microdata. It is called **pop_u_short_public_release_5vars** in the provided example code.

```
#H-----
##> 1. Load Census data
#H-----
load("pop_u_short_public_release_5vars.RData")
df <- pop_u_short_public_release_5vars
dim(df)  # obs = 1163659
names(df)</pre>
```

In our example Census data, we have 1,163,659 observations with five categorical variables including geography, sex, age groups, ethnic groups, and economic activity status. One can declare which variable to tabulate. Here, we declare all five variables using var object⁴.

```
var <- c("geog1", "sex", "agecode1",
                              "eth_code5", "econg")
```

Variable Category Descr		Description	Ν	(%)
Total			1163659	
geog1	1	LA codes	116128	(10.0)
	2	LA codes	150139	(12.9)
	3	LA codes	137520	(11.8)
	4	LA codes	170624	(14.7)
	5	LA codes	90873	(7.8)
	6	LA codes	498375	(42.8)

The description of categories, and distribution is shown below.

⁴ As the var object is treated as global macro, the programme runs automatically using the information stored in global macro, and produces the results.

Variable	Category	Description	Ν	(%)
sex	1	male	564905	(48.5)
	2	female	598754	(51.5)
agecode1	1	16-20	82426	(7.1)
	2	21-25	94643	(8.1)
	3	26-30	110296	(9.5)
	4	31-35	120398	(10.3)
	5	36-40	119393	(10.3)
	6	14-45	101711	(8.7)
	7	46-50	94209	(8.1)
	8	51-55	100159	(8.6)
	9	56-60	77799	(6.7)
	10	61-65	65833	(5.7)
	11	66-70	57305	(4.9)
	12	71-75	51263	(4.4)
	13	76-80	43678	(3.8)
	14	81+	44546	(3.8)
eth_code5	1	White	1081812	(93.0)
	2	Mixed/Multiple ethnic groups	10487	(0.9)
	3	Asian/Asian British	46446	(4.0)
	3	Black/African/Caribbean/Black British	16268	(1.4)
	4	Other ethnic group	8646	(0.7)
econg	1	In employment(FT, PT)	689140	(59.2)
	2	Unemployed	27744	(2.4)
	3	Out of workforce	446775	(38.4)

Using this prior information on the population distribution (based on the Census), we can mimic the distribution in a random sample. See the next step.

- Step 2: Then, we draw a random sample 1:50.
- Step 3: From the randomly selected sample (1163659/50 = 23273), we then obtain frequency table of categorical variables (count of categories). Users can run the predefined function, fn_maxvar5_freq_table() to perform the task. The function⁵ automatically obtains counts and structure the output in long form, organised by each variable, and by its discrete category.

⁵ We also provide fn_maxvar4_freq_table() for users who declare four categorical variables. We do not provide functions for the maximum variable size beyond five. In such cases, users can create their own functions by consulting the provided functions.

fn_maxvar5_freq_table()

This procedure is to assess and calculate *the distribution* of categories in a random sample (N = 23273). Based on the counts of the randomly selected sample, we multiply the counts by 50. One may wonder why we multiply. As we *reduced* the census sample by drawing a random sample by the 1:50 ratio, we need to *convert* the shrank sample back to the original size (with the priori distribution). This explains why we multiply by 50 (Weighted Sample N = 23273 * 50 = 1163650). This completes the process of generating weighted sample survey data.

 Step 4: Carry out checks to see if the calculated frequency tables are accurate. freq_table[1:8, 1:9]

##		seq	twdigits	raw_n	n	р	oneway	v	by1	by2
##	1	1	101	2265	113250	0.09732308	1	1	geog1	01
##	2	2	102	2968	148400	0.12752976	1	1	geog1	02
##	3	3	103	2749	137450	0.11811971	1	1	geog1	03
##	4	4	104	3502	175100	0.15047480	1	1	geog1	04
##	5	5	105	1846	92300	0.07931938	1	1	geog1	05
##	6	6	106	9943	497150	0.42723327	1	1	geog1	06
##	7	7	201	11307	565350	0.48584196	1	2	sex	01
##	8	8	202	11966	598300	0.51415804	1	2	sex	02

When we printed the first 8 lines and 10 variables, we can see the count, n, and the corresponding proportion, p by each variable. The following code obtains the total observation size and confirms that the total proportion adds up to 1, for geog1 variable. Here, the total observation size can be viewed as the population size.

```
# Check whether the total adds up to 1
sum(freq_table[1:6, "n"])
```

[1] 1163650

```
sum(freq_table[1:6, "p"])
```

[1] 1

- Step 5: Rename and save.
- Step 6: Export the output frequency table in Excel with the file name, Weightedsample_freq_table.xlsx.
- Step 7: Save the R objects as RData. Done.

3.4 RUNNING 3A_Distance_Metrics.R

To calculate distance metrics, we first obtain one-way, and two-way frequency tables of categorical variables, and calculates proportions of each sub-category by each data source.

Previously, we dealt with frequency tables for the weighted sample data. Here, we calculate distributions of administrative data. Next, we combine master (benchmark) and admin freq tables. Then, we create domains for quality indicators, and finally compute distance metrics as part of quality indicators. We offer three different types of distance metrics. To allow comparison across the metrics, we standardise the calculations.

Users can also produce a summary table of three types of distance metrics, and visualise the results.

The preliminary step is to ensure we have benchmark data. Simply *source* the previous file as below to update it.

source("2 Prep Wtsample Freq Table.R") Step 1: Read admin data df <- read_csv("public_release_admin.csv")</pre> ## Rows: 1033664 Columns: 6 ## -- Column specification ------_ _ _ _ _ _ ## Delimiter: "," ## dbl (6): person_id, geog1, sex, agecode1, eth_code5, econg ## ## i Use `spec()` to retrieve the full column specification for this data. ## i Specify the column types or set `show_col_types = FALSE` to quiet this m essage. tail(df) ## # A tibble: 6 x 6 ## person_id geog1 sex agecode1 eth_code5 econg ## <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> ## 1 1033659 1 1 6 1 1 1 1 ## 2 1033660 6 1 6 6 2 5 1 1 ## 3 1033661

The last six observations of the example admin data are shown above. As the person id goes up to 1033664, we can see there are 1033664 observations in admin data. As such, we declare all five variables for tabulations. If variables are already defined using var, users can skip this part.

1

1

1

3

1

1

12

9

9

```
var <- c("geog1", "sex", "agecode1",
                                 "eth_code5", "econg")
```

5

1

6

1

2

2

Step 2: Obtain admin freq tables.

4

5

6

1033662

1033663

1033664

As mentioned in the previous section, we obtain frequency table of categorical variables (count of categories). Users can run the pre-defined function, fn_maxvar5_freq_table() to perform

the task. The function⁶ automatically obtains counts and structure the output in long form, organised by each variable, and by its discrete category.

Once the frequency tables are obtained, we rename the object as Admin_f_table_one. Let's inspect Admin_f_table_one.

```
head(Admin_f_table_one)
```

##		seq	twdigits	admin_n	admin_perc	oneway	v	by1	by2	by3	by4	by5
##	1	1	101	137993	0.1334989	1	1	geog1	01	oneway	0	0
##	2	2	102	124051	0.1200110	1	1	geog1	02	oneway	0	0
##	3	3	103	131176	0.1269039	1	1	geog1	03	oneway	0	0
##	4	4	104	139867	0.1353119	1	1	geog1	04	oneway	0	0
##	5	5	105	142304	0.1376695	1	1	geog1	05	oneway	0	0
##	6	6	106	358273	0.3466049	1	1	geog1	06	onewav	0	0

tail(Admin_f_table_one)

##		seq	twdigits	admin_n	admin_perc	oneway	۷	by1	by2	by3	by4	by5
##	340	340	404501	8557	0.0082783187	2	4	eth_code5	04	econg	5	01
##	341	341	404502	791	0.0007652390	2	4	eth_code5	04	econg	5	02
##	342	342	404503	5007	0.0048439338	2	4	eth_code5	04	econg	5	03
##	343	343	405501	3867	0.0037410609	2	4	eth_code5	05	econg	5	01
##	344	344	405502	255	0.0002466953	2	4	eth_code5	05	econg	5	02
##	345	345	405503	3420	0.0033086187	2	4	eth_code5	05	econg	5	03

Step 3: Merge admin + Weighted sample freq tables

fn_merge_one_admin_wtsample_f_table_temp()

The code above merges two data sources.

Step 4: Create domains of quality indicators

fn_create_domain_temp()

To check the domains, we can use *Janitor* package's tabyl function⁷. The function creates 15 domains, including five single variables' domain, and ten bivariate domains.

	display_domain	%>%	% tabyl(fct_domain
##	fct_domain	n	percent
##	geog1	6	0.017391304
##	sex	2	0.005797101
##	agecode1	14	0.040579710

⁶ We also provide fn_maxvar4_freq_table() for users who wish to declare four categorical variables.

⁷ This is essentially almost identical to table(display_domain\$fct_domain), but the approach by tabyl produces percent by default.

##	eth_code5	5	0.014492754
##	econg	3	0.008695652
##	geog1:sex	12	0.034782609
##	geog1:agecode1	84	0.243478261
##	<pre>geog1:eth_code5</pre>	30	0.086956522
##	geog1:econg	18	0.052173913
##	<pre>sex:agecode1</pre>	28	0.081159420
##	<pre>sex:eth_code5</pre>	10	0.028985507
##	<pre>sex:econg</pre>	6	0.017391304
##	agecode1:eth_code5	70	0.202898551
##	agecode1:econg	42	0.121739130
##	eth code5:econg	15	0.043478261

Step 5: Compute distance metrics

Run the functions to compute three types of distance metrics.

```
fn_unstd_distance_metrics_full()
fn_unstd_distance_metrics_tidy()
```

Step 6: Standardise distance metrics

Step 7: Reshape, then tidy

From wide form, the outputs have been reshaped to long form. Then, we keep standardised solutions. The results are as follows:

```
df <- distance_metrics_long</pre>
      # std_test(1-Duncan, 1-HD, 1-KL) only
      df <- df %>% filter(std test use == 1)
      df[1:9, c(1:2, 4:5, 9)]
## # A tibble: 9 x 5
##
     domain_id domain indicator index std_test_use
##
         <int> <chr>
                        <chr>
                                   <dbl>
                                                 <dbl>
## 1
             1 geog1
                        Std Duncan 0.897
                                                     1
## 2
             1 geog1
                       Std t HD
                                  0.911
                                                     1
## 3
                        Std_t_KL
                                   0.957
                                                     1
             1 geog1
                                                     1
## 4
             2 sex
                        Std Duncan 0.987
                        Std_t_HD
                                                     1
## 5
             2 sex
                                   0.991
## 6
                        Std_t_KL
                                                     1
             2 sex
                                   1.00
             3 agecode1 Std Duncan 0.946
                                                     1
## 7
                                                     1
## 8
             3 agecode1 Std_t_HD
                                   0.953
## 9
             3 agecode1 Std_t_KL
                                   0.987
                                                     1
```

Step 8: Visualise the distance metrics

plot(p)

△ Duncan ∘ HD □ KL



3.5 RUNNING 3B_R-indicator.R

The file computes the overall R-indicator. Users can also proceed with computing **partial** R-indicators by category level, and variable level. The procedure can be computationally extensive. This is noted in the relevant section, so that users can allow some time to execute the code.

On this example, we will prepare the sample and population distributions and calculate an R-indicator for 5 variables: geog1 (6), sex (2), agecode1 (14) and eth_cod5 (5) and econg (3).

As part of administrative data preparation, each of the variables should have their categories numbered 1,2,3.... We will use these numbers instead of the original names of the categories because it will enable us to do loops through the data. This is to facilitate building design matrix using dummy variables.

Along with administrative data, we also need benchmark data from Census. Assuming users have no access to Census data, we replace Census with weighted sample counts. The auxiliary data file contains these weighted sample counts.

As such, we will read in both administrative and auxiliary data files separately and compute Rindicators using matrix syntax in R. Due to the complexity of the procedure, we provide defined functions that users can execute with ease in practice. Users can consult Functions/2_Functions_R-indicators.R file for more details on the algorithm and operationalisation.

Step 1: Prepare an Auxiliary data file

From the frequency table generated by using the weighted sample, we will prepare an auxiliary data file to be used for R-indicator computation procedures. This auxiliary file is used as benchmark. From the previous steps, we identified the population size of 1,116,350 (popsize = 1163650) from the weighted sample.

For R-indicator calculations, we need to compute meanpop by variables which are geography, sex, age groups, ethnic groups, and economic activity status. Users can simply run the function, fn_meanpop_auxiliary() to achieve this goal.

```
load(file = "Output/04-RData/Weightedsample_freq_table.RData")
load(file="Output/04-RData/var.RData")
```

```
fn_meanpop_auxiliary()
```

Exerpts of the function is provided below to aid readers' understanding.

```
# Compute meanpop
auxiliary <- freq_table %>%
    filter(oneway == 1) %>%
    group_by(by1) %>%
    mutate(meanpop = n / popsize) %>%
    ungroup() %>%
    dplyr::select(seq, count = n, by1,
        v, by2, meanpop, raw_n)
```

We first remove two-way and keep the one-way frequency table only. Then, by variable-level (indicated by the variable, by1), we compute meanpop. As seen before, the count of each category is stored in n. The meanpop is obtained by dividing n by popsize. For example, the value of **meanpop** for first category of geog1 is calculated as 113250/1163650 = 0.0973, and the second category of geog1 is 148400/1163650 = 0.1275 and so on.

Let's look at the intermediate Auxiliary data. We can see the count of each variable (by1) and the sub-category (by2) for five variables. The population size is indicated as 1163560 (shown in the first row, under count column).

print(auxiliary)

##		seq	count	by1	v	by2	meanpop	raw_n	type
##	1	1	1163650	total	0	00	1.000000000	0	wtsample
##	2	2	113250	geog1	1	01	0.097323078	2265	wtsample
##	3	3	148400	geog1	1	02	0.127529756	2968	wtsample
##	4	4	137450	geog1	1	03	0.118119710	2749	wtsample
##	5	5	175100	geog1	1	04	0.150474799	3502	wtsample
##	6	6	92300	geog1	1	05	0.079319383	1846	wtsample
##	7	7	497150	geog1	1	06	0.427233275	9943	wtsample

##	8	8	565350	sex	2	01	0.485841963	11307	wtsample
##	9	9	598300	sex	2	02	0.514158037	11966	wtsample
##	10	10	84600	agecode1	3	01	0.072702273	1692	wtsample
##	11	11	93300	agecode1	3	02	0.080178748	1866	wtsample
##	12	12	111200	agecode1	3	03	0.095561380	2224	wtsample
##	13	13	124250	agecode1	3	04	0.106776092	2485	wtsample
##	14	14	118200	agecode1	3	05	0.101576935	2364	wtsample
##	15	15	99950	agecode1	3	06	0.085893525	1999	wtsample
##	16	16	95800	agecode1	3	07	0.082327160	1916	wtsample
##	17	17	95600	agecode1	3	08	0.082155287	1912	wtsample
##	18	18	78450	agecode1	3	09	0.067417179	1569	wtsample
##	19	19	67600	agecode1	3	10	0.058093069	1352	wtsample
##	20	20	57950	agecode1	3	11	0.049800198	1159	wtsample
##	21	21	50650	agecode1	3	12	0.043526834	1013	wtsample
##	22	22	42250	agecode1	3	13	0.036308168	845	wtsample
##	23	23	43850	agecode1	3	14	0.037683152	877	wtsample
##	24	24	1083250	eth_code5	4	01	0.930907060	21665	wtsample
##	25	25	10450	eth_code5	4	02	0.008980364	209	wtsample
##	26	26	45600	eth_code5	4	03	0.039187041	912	wtsample
##	27	27	16300	eth_code5	4	04	0.014007648	326	wtsample
##	28	28	8050	eth_code5	4	05	0.006917888	161	wtsample
##	29	29	691300	econg	5	01	0.594078976	13826	wtsample
##	30	30	28250	econg	5	02	0.024277059	565	wtsample
##	31	31	444100	econg	5	03	0.381643965	8882	wtsample

Now, we prepare the data to build design matrix using dummy variables. To do so, from the total number of categories for each variable, we need to remove the **last** category of each categorical variable (group_by(by1)). The last category is defined by max(by2) and removed accordingly. The first row (seq == 1) is not subject to this procedure and kept in the data. As shown in the code below, we keep rows if lastcat == 0, while filtering out cases which are the last category. We then drop the indicator for the last category (dplyr::select(-c(lastcat_, lastcat))). Here, we explicitly instruct R to use dplyr package to access select function⁸.

```
col_auxiliary <- auxiliary %>%
    group_by(by1) %>%
    mutate(
        lastcat_ = ifelse(by2 == max(by2), 1, 0),
        lastcat = ifelse(seq == 1, 0, lastcat_)
        ) %>%
        filter(lastcat == 0) %>%
        dplyr::select(-c(lastcat_, lastcat)) %>%
        ungroup()
```

⁸ This is to avoid warning messages from R when R searches for a particular function from two different packages.

Notice that from 5 variables, geog1 (6), sex (2), agecode1 (14) and eth_cod5 (5) and econg (3), we now have 1+(nvar-1) for each variable so here it is 1+5+1+13+4+3=26 rows. We create a macro, numcat to store this information on the total row.

```
nrow(col_auxiliary)
## [1] 26
numcat <- nrow(col_auxiliary)
numcat
## [1] 26</pre>
```

To inspect the setup on dummy variables (to be created later), let's produce a cross-tabulation by by1 and by2.

```
dummychk <- col_auxiliary</pre>
dummychk %>% tabyl(by1, by2)
          00 01 02 03 04 05 06 07 08 09 10 11 12 13
agecode1
          0 1
                1
                    1
                       1
                          1
                             1
                                1
                                   1
                                      1
                                         1
                                            1
                                               1
                                                  1
econg
           0 1 1
                    0
                       0
                          0
                             0
                                0
                                   0
                                      0
                                         0
                                            0
                                               0
                                                  0
eth_code5 0 1
                1
                   1
                       1
                         00
                               0
                                  0
                                     0
                                         0
                                            0
                                               0
                                                  0
geog1
           01
                1
                    1
                       1
                          1
                             0
                                0
                                   0
                                      0
                                         0
                                            0
                                               0
                                                  0
sex
           0
            1
                0
                    0
                       0
                          0
                            0
                               0
                                   0
                                     0
                                         0
                                            0
                                               0
                                                  0
           1
                0
                                            0
total
             0
                   0
                       0
                          0
                             0
                               0
                                   0
                                      0
                                         0
                                               0
                                                  0
```

All look good. At this stage, we keep meanpop only, dropping other columns, and print meanpop in a single column (column vector). This column will be transposed before we save it as row vectors. Before reshaping the data, we carefully inspect the values of meanpop.

```
# Print the column vector, meanpop
        print(col_auxiliary)
## # A tibble: 26 x 1
##
     meanpop
##
        <dbl>
## 11
## 2 0.0973
## 3 0.128
## 4 0.118
## 5 0.150
## 6 0.0793
## 7 0.486
## 8 0.0727
## 9 0.0802
## 10 0.0956
## 11 0.107
## 12 0.102
## 13 0.0859
```

14 0.0823
15 0.0822
16 0.0674
17 0.0581
18 0.0498
19 0.0435
20 0.0363
21 0.931
22 0.00898
23 0.0392
24 0.0140
25 0.594
26 0.0243

Step 2: Reshape and save benchmark data as row vectors

We use t(col_auxiliary) to transpose and tidy the reshaped data. We then generate ttt as an indicator of benchmark data. We use this indicator for merging with the administrative data later. Save the data as popmean_row_vector.

```
# Transpose to arrange in row vector format.
temp <- t(col_auxiliary)
temp <- as.data.frame(temp)
row.names(temp) <- 1:nrow(temp)
# generate merge id, ttt.
temp$ttt <- 0
# Rename variables "popmean1- popmean26"
names(temp) <- c(paste0("popmean", 1:numcat), "ttt")
# SAVE
```

popmean_row_vector <- temp</pre>

Let's see the row vector names, "popmean1- popmean26" and ttt.

```
names(popmean_temp)
## [1] "popmean1" "popmean2" "popmean3" "popmean4" "popmean5" "popmean6
"
## [7] "popmean7" "popmean8" "popmean9" "popmean10" "popmean11" "popmean1
2"
## [13] "popmean13" "popmean14" "popmean15" "popmean16" "popmean17" "popmean1
8"
## [19] "popmean19" "popmean20" "popmean21" "popmean22" "popmean23" "popmean2
4"
## [25] "popmean25" "popmean26" "ttt"
    popmean_temp[, 1:5]
```

popmean1 popmean2 popmean3 popmean4 popmean5
1 1 0.09732308 0.1275298 0.1181197 0.1504748

Now, our benchmark data preparation is completed.

Step 3: Declare variables in administrative data

The next step is to declare variables to be used for computing the R-indicator. These variable names should be from the administrative data. As mentioned earlier, users may skip this part if earlier defined var is unchanged and consistent throughout benchmark data and administrative data. In the provided example file, we save var object and load it, rather than defining at each stage.

Follow the instructions if users need to declare here. Users can define their own variables and save as a macro, var. For instance, users may use four variables and define as below.

```
var <- c("geog1", "sex", "agecode1", "econg")</pre>
```

For demonstration, we use five variables for R-indicator calculations and declared as such.

```
# Customise as needed.
var <- c("geog1", "sex", "agecode1",
               "eth_code5", "econg")
var
## [1] "geog1" "sex" "agecode1" "eth_code5" "econg"
# End of custom variables.
```

This concludes steps 1 through 3. Users can customise some setups up to this point.

Step 4: Define macro variables

Along with the defined variables, we need to ensure macro variables are generated correctly. These macro variables are not designed to customise and designed to run without altering the code.

var variablenum maxvar popsize

The total number of variables and the total number of categories of each categorical variables will be stored in a macro called variablenum and maxvar. We use var macro, as we defined earlier, to derive these two macros.

variablenum <- length(var) # No need to customise
maxvar <- length(var) # No need to customise</pre>

variablenum ; maxvar ## [1] 5 ## [1] 5

Earlier, we also generated popsize to feed the information on the (total observation) size of the the benchmark data to the programme. The remaining macros, such as respop, piinv and rrate, will be automatically generated by the pre-defined functions.

respop piinv rrate

respop is the (sample) size of the administrative data. piinv refers to the inverse pi. rrate is computed by resppop/popsize.

Step 5: Compute R-indicators

Once steps 1 through 4 are completed, we are set to carry out computing R-indicators. These procedures are automated via pre-defined functions using complex matrix and data management syntax. Please note that some procedures are computationally intensive.

We first open the corresponding administrative data, and notice the number of rows is 1,033,664.

```
aa <- read_csv("public_release_admin.csv")
## Rows: 1033664 Columns: 6
## -- Column specification ------
## Delimiter: ","
## dbl (6): person_id, geog1, sex, agecode1, eth_code5, econg
##
## i Use `spec()` to retrieve the full column specification for this data.
## i Specify the column types or set `show_col_types = FALSE` to quiet this m
essage.</pre>
```

nrow(aa) # 1033664

```
## [1] 1033664
```

We need several empty objects to hold data to get the functions to work as intended. Note that we have five functions that permit us to compute overall R-indicators, along with additional functions for obtaining partial R-indicators. Each function will generate objects including vvv, pop_respmean, des_pop_respmean, gh, and R_indicators as indicated in the function name. The partial R-indicators can be found in the data object, partial.

```
df <- NULL
between <- NULL
partial <- NULL
partialtemp <- NULL
fn_r_indicator_1_vvv()
fn_r_indicator_2_pop_respmean()
fn_r_indicator_3_des_pop_respmean()
fn_r_indicator_4_gh()
fn_R_indicators()
# Partial R-indicators
fn_r_indicator_partialtemp()
fn_r_indicator_domain_order_partial()</pre>
```

Let's go over one function at a time.

The utility of fn_overall_r_indicator_1_vvv() is to build design matrix. Starting with ensuring that the admin data only contains the declared variables and factorise them, the function defines macro variables, resppop and rrate using the data object, aa. To prepare for design matrix, the code creates dummy variables, using **fastDummies** R library package. Once all dummy variables generated, we need to remove the last category. As such, the function detects the categories of each variable and drops the last category.

Let's see the vvv object, which contains the design matrix with weights. The weights are calculated by the inverse of *rrate* (finalwgt = 1/rrate). The last five columns are printed below.

```
# fn_overall_r_indicator_1_vvv()
```

```
from <- ncol(vvv)-4
vvv[1:8, from: ncol(vvv)]</pre>
```

##	# A	A tibb.	le: 10	x 5		
##		des24	des25	des26	finalwgt	piinv
##		<int></int>	<int></int>	<int></int>	<dbl></dbl>	<dbl></dbl>
##	1	0	1	0	1.13	1
##	2	0	1	0	1.13	1
##	3	0	1	0	1.13	1
##	4	0	1	0	1.13	1
##	5	0	1	0	1.13	1
##	6	0	0	0	1.13	1
##	7	0	0	0	1.13	1
##	8	0	1	0	1.13	1

The fn_overall_r_indicator_2_pop_respmean() prepares the distributions of the administrative data. Using the data object, vvv, we obtain weighted sample counts and produces a row vector called *respmean_row_vector*. Merging the corresponding

popmean_row_vector from the benchmark data we prepared at the step 2 earlier, the function combines both mean vectors from the two data sources.

The combined mean vectors are stored at the pop_respmean data object. We will inspect the last five columns.

```
# fn_overall_r_indicator_2_pop_respmean()
from <- ncol(pop_respmean)-4
pop_respmean[, from: ncol(pop_respmean)]
## respmean25 respmean26 finalwgt piinv ttt
## 1 0.5896214 0.02252997 1.125753 1 0</pre>
```

The function, fn_overall_r_indicator_3_des_pop_respmean(), allows us to combine the design matrix with the pop_respmean. We store the data at des_pop_respmean.

fn_overall_r_indicator_3_des_pop_respmean()

names(des_pop_respmean)

##	[1]	"des1"	"des2"	"des3"	"des4"
##	[5]	"des5"	"des6"	"des7"	"des8"
##	[9]	"des9"	"des10"	"des11"	"des12"
##	[13]	"des13"	"des14"	"des15"	"des16"
##	[17]	"des17"	"des18"	"des19"	"des20"
##	[21]	"des21"	"des22"	"des23"	"des24"
##	[25]	"des25"	"des26"	"finalwgt"	"piinv"
##	[29]	"popmean1"	"popmean2"	"popmean3"	"popmean4"
##	[33]	"popmean5"	"popmean6"	"popmean7"	"popmean8"
##	[37]	"popmean9"	"popmean10"	"popmean11"	"popmean12"
##	[41]	"popmean13"	"popmean14"	"popmean15"	"popmean16"
##	[45]	"popmean17"	"popmean18"	"popmean19"	"popmean20"
##	[49]	"popmean21"	"popmean22"	"popmean23"	"popmean24"
##	[53]	"popmean25"	"popmean26"	"respmean1"	"respmean2"
##	[57]	"respmean3"	"respmean4"	"respmean5"	"respmean6"
##	[61]	"respmean7"	"respmean8"	"respmean9"	"respmean10'
##	[65]	"respmean11"	"respmean12"	"respmean13"	"respmean14'
##	[69]	"respmean15"	"respmean16"	"respmean17"	"respmean18'
##	[73]	"respmean19"	"respmean20"	"respmean21"	"respmean22'
##	[77]	"respmean23"	"respmean24"	"respmean25"	"respmean26'
##	[81]	"seq"	"responsesamp1"		
	from	n <- ncol(des_pop	respmean)-4		
	des	_pop_respmean[1:6	5, from: ncol(des	<pre>s_pop_respmean)]</pre>	
	_				
##	# A 1	tibble: 6 x 5			

##		respmean24	respmean25	respmean26	seq	responsesamp1
##		<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<int></int>	<dbl></dbl>
##	1	0.0139	0.590	0.0225	1	1
##	2	0.0139	0.590	0.0225	2	1

## 3	0.0139	0.590	0.0225	3	1	
## 4	0.0139	0.590	0.0225	4	1	
## 5	0.0139	0.590	0.0225	5	1	
## 6	0.0139	0.590	0.0225	6	1	

In the fn_overall_r_indicator_4_gh(), we compute the difference from the mean vectors and the weight variables. Exerpts of the code from the function below show that the differences are stored in rsam and psam. By adding *rsam* and *psam* to the intermediate data object, des_pop_respmean, we obtain the gh data. This concludes the pre-matrix preparation part.

```
# use df for programming.
df <<- data.frame(des_pop_respmean)</pre>
# Prep for loop.
des col
         <<- c(paste0("des" , 1:numcat))</pre>
respmean_col <<- c(paste0("respmean", 1:numcat))</pre>
popmean_col <<- c(paste0("popmean" , 1:numcat))</pre>
        <<- df[, des_col]</pre>
des
respmean <<- df[, respmean_col]</pre>
popmean <<- df[, popmean_col]</pre>
rsam <<- des - respmean
psam <<- des - popmean
temp <<- data.frame(rsam, psam)</pre>
# Rename variables
colnames(temp) <<- c(paste0("rsam", 1:numcat),</pre>
                      paste0("psam", 1:numcat))
# Combine
gh <<- cbind(des pop respmean, temp)</pre>
```

The fn_R_indicators() uses matrix syntax to calculate propensity scores, prior to computing R-indicators. We calculate two kinds of propensity scores – one that used only population information (roipop, or prop_pop) and the other which used a mixture of the response data and the population information (roimix, or prop_mix).

In terms of partial R-indicators, we demonstrate using *prop_mix*. Running fn_r_indicator_partialtemp() users can yield partial R-indicators at the variable level and at the category level. The function, fn_r_indicator_domain_order_partial() helps us to organise domains.

```
fn_r_indicator_partialtemp()
fn_r_indicator_domain_order_partial()
# View(partial)
```

Step 6: Save in Excel and inspect

At this stage, users can inspect the output accordingly. Let's have a look. Here, we can see the overall R-indicator is estimated as 0.496 based the administrative data (N=1033664). Looking at the variable-level R-indicator (see rows 4-8), geog1 was seen to have the greatest R-indicator (0.04) compared to econg (0.0002).

```
partial[1:17, c(1:2, 4, 8:10)]
```

##		seq	domain	R_indicator	count	n_cat	domain_n
##	1	1	Overall	0.4960263148	NA	<na></na>	NA
##	2	2	mrphatall	0.9597768609	NA	<na></na>	NA
##	3	3	resppop	1033664.0000000000	NA	<na></na>	NA
##	4	4	geog1	0.0442418275	NA	<na></na>	NA
##	5	5	sex	0.0004983014	NA	<na></na>	NA
##	6	6	agecode1	0.0152841196	NA	<na></na>	NA
##	7	7	eth_code5	0.0000296805	NA	<na></na>	NA
##	8	8	econg	0.0002117090	NA	<na></na>	NA
##	9	9	des1	NA	0	1	0
##	10	10	geog1_1	0.0879501347	137993	1	1
##	11	11	geog1_2	-0.0192794816	124051	2	1
##	12	12	geog1_3	0.0219039079	131176	3	1
##	13	13	geog1_4	-0.0366161292	139867	4	1
##	14	14	geog1_5	0.1396946734	142304	5	1
##	15	15	geog1_6	-0.1216543425	358273	6	1
##	16	16	sex_1	-0.0162005503	489228	1	2
##	17	17	sex 2	0.0153571986	544436	2	2

Step 7: Visualising using scatterplots

The visualisation of R-indicator by the variable level is shown as an example:



And for R-indicator by the category-level:



This concludes the manual. Thank you for taking the time reading the material. Please get in touch with any query or errata at <u>fanfurcada@gmail.com</u>.

If you need technical support, please consult the following Troubleshooting Q & A section.

4. Troubleshooting Questions and Answers

4.1 Questions and Answers

How do I know where to customise the code to suit my needs?

Unless indicated as "Customise as needed", users can run the code as it is. Please consult each code file.

How to use Starting path in multiple machines?

If users plan to use different machines, simply by changing the "starting path", users can carry out the analysis with minimal disruption. To achieve this, please ensure to use the consistent master project folder name.

What are the commonly used commands?

Most commonly used commands in the tidyverse package are:

How to free up memory space and speed up RStudio?

You can remove objects that you no longer need.

```
# To remove objects except for certain objects
ls()
keepobjectslist <- c("a", "b", "c")
rm(list = ls()[!ls() %in% keepobjectslist])
ls()</pre>
```

I get error messages when a pre-defined function is used.

Users can inspect the codes used in the function, and identify the issues. It is recommended NOT edit the function file directly, as the functions are used repeatedly, and the interlinked sections may not run as expected. Where preferable, users may copy the codes in the function, and use locally with minor tweaks.

How do I modify pre-defined functions?

Users can modify 1_Functions and 2_Functions_R-indicators.R under *Functions* folder.

```
# 1_Functions.R
fn_output_folder_path <- function() {
    currentdate <<- Sys.Date()
    txtpath <<- "Output/01-Txt/"
    figpath <<- "Output/02-Figure/"
    xlsxpath <<- "./Output/03-ExcelOutput/"
    Rdatapath <<- "Output/04-RData/"
}</pre>
```

We can check how the output folder names are set as path to save the results during the analytical process.

```
fn_output_folder_path()
```

Let's run the function. We can see that xlsxpath is set as "./Output/03-ExcelOutput/".

xlsxpath

```
## [1] "./Output/03-ExcelOutput/"
```

Let's customise the xlsxpath, by renaming the folder name. If we customise 1_Functions.R file, we can edit the information enclosed in the brackets. Notice that we use <<- with functions so that the object created by a function will exist in the global R environment. This is very important.

Alternatively, we could ignore the pre-defined function and just write relevant lines of code and keep it in the main R script file. For instance, we could put output_folder_path at the top of the 2_Prep_Wtsample_Freq_Table.R. Here, we edited the xlsxpath. Notice that fn_output_folder_path <- function() { } is removed.

```
xlsxpath_2 <- "./Output/03-Excel/"
xlsxpath_2
## [1] "./Output/03-Excel/"
#H------
## > Step 1. Load Census data
#H------
# Load("pop u short before sim 5vars.RData")
```

Notice that we use <-. Using <<- is not necessary here. Users can remember the usage of <- and can modify the functions as appropriate, should the function incur errors.

What approaches are taken in programming?

When loop is used, base R functions were used (table, tapply, etc). For data manipulation, tidyverse package was used extensively. This strategy is partly to improve readability of the code.

To enhance users' workflow, output files are programmed to launch using the pre-defined functions.

Can I ignore Warning messages?

Some packages alert users with compatibility issues arising from old version. These can be ignored. For example,

```
library("fastDummies")
Warning message:
package 'fastDummies' was built under
R version 4.1.2
library(rlist)
Warning message:
package 'rlist' was built under R version 4.1.2
```

What version of R is used?

Tested with Windows PC. R version used: 4.1.1 RStudio version: RStudio 2022.12.0 Build 353.

4.2 Troubleshooting

Unused argument error

For example, sim %>% select(geog1) the select command can cause an error:

```
Error in select(., geog1) :
    unused arguments (geog1)
```

This maybe due to the conflict in packages.

The error can be fixed by adding the name of the package used, dplyr, explicitly. sim %>% dplyr::select(geog1)

I get errors when computing...

Please inspect zero cells, and ensure 0 (numeric value) is entered for n and perc, as well as admin_n and admin_perc. Errors may occur with NA coding and data attributes(character, factor, numeric).

I am experiencing slowness in computation.

R can be not responsive if memory is full. Please identify bottlenecks and remove them. It may be due to certain commands. For example, View(object) command could take a while if the object is huge in size. Unless one should inspect the data, suppress the View command to expedite the computation where possible.

It can also be the case that for loop functions can be slow as well. In some instances, removing objects may help as this procedure can free up memory space. See above commonly used commands for more information.

Error: cannot allocate vector of size xxxx.x Gb

If matrix symbols have entered mistakenly, R shows an error message like this. Please double check whether there are any mistakes. For instance, one may have typed a*b instead of a%*%b. Users can type memory.limit() to check the current memory limit and increase as necessary.

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