

# Millennium Cohort Study

Data Handling Guide
with syntax in R, STATA and SPSS

August 2020





Contact

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## About the Millennium Cohort Study

The Millennium Cohort Study (MCS) is a longitudinal birth cohort study, following a nationally representative sample of approximately 19,000 people born in the UK at the turn of the century.

Through the study, we have captured rich information about the different aspects of cohort members' lives, from birth to childhood and adolescence, and we are continuing to keep up with them now they are adults.

As a multidisciplinary study, MCS is used by researchers working in a wide range of fields. Findings from MCS have influenced policy at the highest level, and today the study remains a vital source of evidence on the major issues affecting young people's lives.

Sweep	Fieldwork / data collection starting year	Cohort Members' average age
MCS 1	2001	9 months old
MCS 2	2004	3 years old
MCS 3	2006	5 years old
MCS 4	2008	7 years old
MCS 5	2012	11 years old
MCS 6	2015	14 years old
MCS 7	2018	17 years old

# About this user guide

The Data Handling guide aims at helping researchers use the data of the Millennium Cohort Study to its fullest potential. We focus on explaining the data structures of MCS and on providing solutions with different data handling strategies.

The structure of the guide per chapter is the following:

**Chapter 1 –** Overview of the data structure and the identifiers of the datasets of the MCS.

**Chapter 2 -** Explanation of how to use dataset of different structures separately but also how to merge between sweeps.

**Chapter 3 -** Detailed information on why restructuring may be needed and how to proceed with it, including walkthrough on merging datasets of different structures.

**Chapter 4 -** Finally, this guide provides the syntax that has been used for each chapter and sections. The syntax is in SPSS Syntax, R and STATA.

#### **Important notes** about this user guide:

- This user guide does not replace any of the User Guides of each sweep of MCS. Those guides contain specific information on survey design, questionnaire, survey outcomes and detailed information on the derived variables.
- The example codes are provided in the Appendices and they can be used for a hands-on experience with the data.

We hope you enjoy working with the Millennium Cohort Study!

#### **Abbreviations & shortcuts housekeeping**

There are some abbreviations used in the text to make text more condensed that are important to familiarise with:

- MCS Millennium Cohort Study
- CM Cohort Member
- CMs Cohort Members
- CAPI Computer Assisted Personal Interview
- CAPI name/code the 4-character code of a question as it appears in the questionnaire

In this guide we use the term **parent** and **carer** interchangeably.

### Data structures and identifiers of MCS

### 1.1 From the survey design to the data structures

This section explains how the data of the Millennium Cohort Study are structured. It also contains explanation of how the various data structures of MCS are generated by the distinctive design of the survey interview and the questionnaire.

### 1.2 Variations in the family structures

A typical interview setting of MCS includes the Main carer of the Cohort Member(s) (usually mentioned as Main), the Partner of the Main carer (usually mentioned as Partner) and the Cohort Member(s).

Family structures vary and MCS captures this variation:

- Some families had multiple births: twins and triplets of a family are all eligible to participate as Cohort Members in MCS.
- Sometimes the Partner respondent although s/he lives in the household s/he is not available for the interview and the Main respondent provides information about him/her (Proxy Partner module).
- Sometimes there is no person eligible for the Partner interview and there is only a Main respondent for that family.
- The largest number of individuals eligible for interview in one family in MCS is 5: 3 children (if triplets) and 2 carers/parents.

The key to using the data effectively is the questionnaire. The questionnaire provides information on whether:

- the question is addressed to Main only or Main and Partner or Partner only
- the question is about themselves and the family <u>or</u> it is about the Cohort Member(s), and,
- the questions get repeated (in a loop) for each Cohort Member of the household.

## 1.3 Distribution of variables between dataset types

The picture below shows how different questions collect information that varies in terms who the respondent is and whom the question is about (parents about themselves or about the cohort member).

The focus is on who answers the question and about whom.

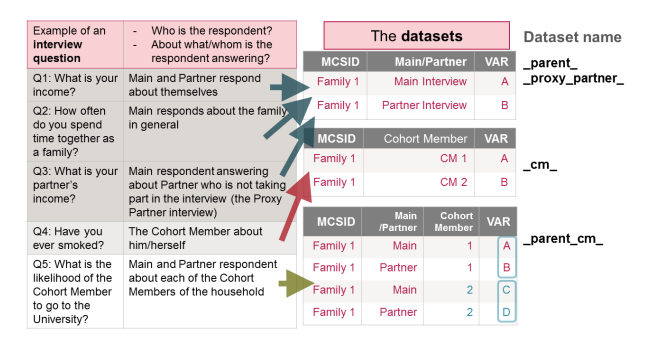


Figure 1: Interview setting -> Questionnaire -> Datasets

The datasets are available in 4 structures - or levels - depending on the information they include:

- Family level (one row per family)
- Parent level (one row for Main and one for Partner respondent)
- Cohort Member level (one row for each of the Cohort Members)
- Parent Cohort Member level (one row for each parent respondent and then one row for each of the Cohort Members)

This rule of distributing variables applies to the derived variable datasets too.

### 1.4 Dataset types

The structure of the data is part of the dataset title to make it easy for the data user to know the level(s) inside the dataset before accessing it.

Family level dataset \_family\_: These datasets contain information about the family as one entity. For example, in the mcs\*\_family\_derived dataset, one can find information about the family type, the total number of Cohort Members in the household, etc. The mcs\_longitudinal\_family\_file that contains the weights is also on the family level.

Parent level dataset \_parent\_: If the question is addressed towards the parent and it is about the family or about him/herself. This is regardless if the question is to be addressed to Main respondent only or to Main and Partner respondents), then the variable that corresponds to that question is in the \_parent\_ level dataset. For example, the variables of parental income questions can be found in the \_parent\_ level dataset. The Parent level dataset includes the Household Module which contains demographic information about the family.

The Proxy Partner interview occurs when there is a person eligible (ELIG, see identifiers) for the Partner interview but s/he does not participate as Partner (RESP, see identifiers). In these cases the Main is asked some questions about the Partner (hence, Proxy Partner). The \_proxy\_partner\_ dataset has similar structure to the \_parent\_ level dataset (one row per partner).

Cohort Member level dataset \_cm\_: If the respondent is the child answering questions or providing information about her/himself, then this is included in \_cm\_ level dataset. Physical measurements of the Cohort Members and cognitive assessments are in \_cm\_ level datasets. From sweep 4 onwards, the CMs get interviewed and this information can be found in the \_cm\_interview dataset.

Parent - Cohort Member level dataset \_parent\_cm\_: Many questions in the early sweeps of MCS ask for information about each Cohort Member from the parent(s).

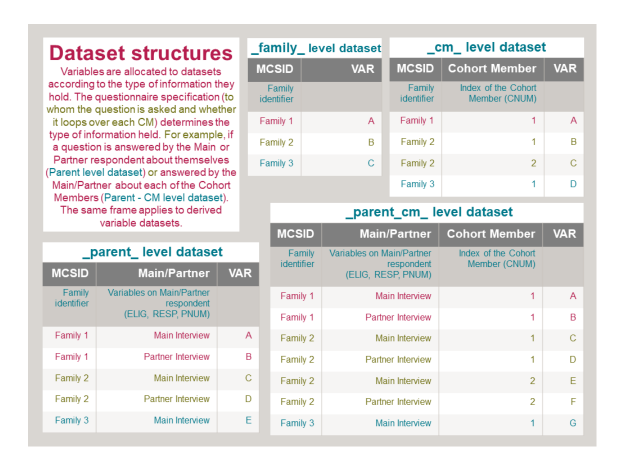


Figure 2: Overview of dataset structures and the identifiers they contain

The structure of this dataset type reflects the information collected by the survey questionnaire. For example, a question that is asking each of the parents **whether the CM cries at night** gets repeated for each Cohort Member.

This happens if the question about each CM has been addressed to both parent respondents (Main/Partner) or to the Main respondent only. The dataset includes identification of the respondent (Main/Partner) and the Cohort Member the question is asked about (Cohort Member number).

The **household grid** is on person level (PNUM / CNUM) since there is one row for each person of the household regardless if they were selected for an interview (for example, CM's siblings, grandparents, other relatives or non-relatives). The household grid is a source of information about the key respondents of the interview (Main/Partner/Cohort Members) as well as the rest of the members of the household that do not participate during the survey interview.

In certain sweeps, there is an **older sibling** dataset where there is a row per older sibling that has participated. Also, there have been teacher surveys of the Cohort Members which have a \_cm\_ structure since the teacher provides information about the CMs.

The \_derived datasets contain derived information about the demographic characteristics and socio-economic status of the household and its members. The derived datasets can be found in 3 different structures:

- cm derived with information about each Cohort Member (one row per CM)
- parent\_derived with information about each Main or Partner respondent (one row per parent)
- family\_derived with information about the family and the interview (one row per family)

Information about each variable of these datasets and how they have been derived can be found in the respective User Guide of the derived variables of each sweep.

### 1.5 Identifiers: MCSID, PNUM, CNUM, ELIG, RESP

The format of the different dataset types, require different identifiers that help the user manage the data for each family, parent/carer respondent (within the family), Cohort Member (within the family) and person (within the family). This section describes the identifiers and how they can be used.

The key identifiers of MCS are:

- MCSID is a family / household anonymised identifier and it is the same for all
  of its members, Cohort Members and parents per family.
- CNUM is Cohort Member number within a family. Namely, the CNUM in ascending order standing from 1 indicates the first Cohort Member within a family. The majority of the Cohort Members have CNUM=1, however, in families with twins and triplets the second and third Cohort Member have CNUM=2 and CNUM=3 respectively.
- PNUM is Person number for the individuals living in the household apart from the Cohort Member(s). This includes grandparents, siblings of the Cohort

Members, parents, etc. The PNUM is given in an ascending order starting from 1 randomly to each person that lives in the household that the Cohort Member lives or has lived (in a previous sweep). This means that a grandparent may be PNUM=1 and if s/he moves out in a later sweep s/he keeps the same PNUM. If another person moves in to the household or a sibling is born will receive the next available PNUM.

- ELIG provides the information on whether the individual has been eligible for
  the role of the Main or Partner respondent. For example, the mother of the
  Cohort Member may have PNUM=2 and the father PNUM=3, however, only
  the mother may have been eligible to be interviewed for the survey (ELIG=1)
  and not the father (ELIG=4).
- RESP marks whether the person selected at ELIG has participated in the interview or not (e.g. refusal). This variable is the outcome of the ELIG.

PNUM and CNUM are permanent cross-sweep identifiers. Namely, they do not change between sweeps and the person or CM holds this number for the rest of the survey. Moreover, the PNUM gets assigned at random. It is likely that the parents of the CM have a low PNUM as they have been present at Sweep 1 or 2 but this is not always the case.

ELIG and RESP are sweep-specific identifiers. Namely, they mark whether the person with a specific PNUM has been selected to participate in the interview and provide information about the Cohort Member(s) in this specific sweep (as Main or Partner respondent). The individuals eligible to respond to a survey can change from the one sweep to the other. For this reason PNUM is available in all datasets where there is a parent respondent like the \_parent\_ level dataset and the \_parent\_cm\_ level dataset.

Each dataset depending on its type and structure contains a different set of identifiers. For example:

- a dataset on the CM level \_cm\_, such as the self-completion questionnaire of the Cohort Member, contains only MCSID and CNUM and not any parent identifiers (PNUM, ELIG, RESP) since these are irrelevant,
- a dataset on the parent level \_parent\_, contains MSCID, PNUM, ELIG and RESP but does not contain CNUM.
- a dataset on the \_family\_ level contains only the MCSID which is the family identifier,
- figure 3 provides an example of the \_parent\_cm\_ level dataset that requires
  all the identifiers. This is because within a family (MCSID) a parent (PNUM,
  ELIG, RESP) provides information about each of the Cohort Members of the
  family (CNUM).

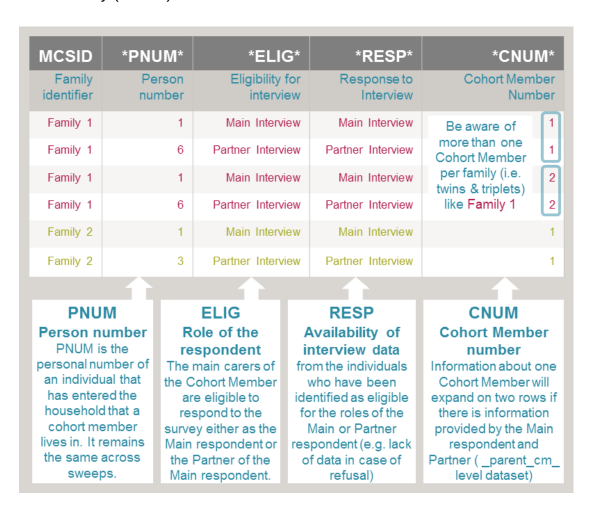


Figure 3: Example of how key identifiers look like in a parent cm dataset

# 1.6 Parent or Carer respondent and the distinction between PNUM and ELIG/RESP

The existence of PNUM, ELIG and RESP is a design characteristic of MCS that provides the data with an important advantage: there is a person providing information about the CM even if the natural parent does not live in the same household with the Cohort Member or is not available to respond.

When working with datasets that are on the \_parent\_ , \_parent\_cm\_ and \_proxy\_partner\_ levels it is important to consider that the parent respondent (Main/Partner) may change between sweeps.

The possible family structure changes vary. The most common one is for a natural parent not to be in the household (for example, divorced, moved out, passed away) and another carer (if available) of the Cohort Member, like a step parent or grand parent, becomes eligible to participate in the survey interview.

There is a different PNUM for the person who moved out and for the new person who moved in. For example, the natural parent may have a PNUM=3 and the step parent (or grandparent) may have a PNUM=5.

Although the PNUM will remain the same in the subsequent sweeps for both (in the household grid), ELIG contains their eligibility for interview for the role of the Main or Partner of the Main carer a different person in a specific sweep. Eligibility depends on who has been a resident in the same household as the Cohort Member at the time of the interview.

### 1.6.1 Merging between sweeps: focusing on data availability (ELIG)

For some research projects the focus may be on information provided by the Main/Partner respondent about the Cohort Member(s) regardless of the fact that the Main/Partner respondent may change between sweeps. For example, in the one sweep the Partner respondent may be a natural parent whereas in another a step

parent. In this case, a data user may prefer to merge the data between sweeps using MCSID and ELIG (or MCSID and RESP).

The variable **ELIG** contains information on what role the carer of the Cohort Member was selected for (eligibility). It is derived based on the information provided at the Household Grid and the main carers of the Cohort Member (usually mother and step/natural father) are prioritised for the role of the Main and Partner. The exact relationship of the Main/Partner to the Cohort Member can be found in the Household Grid in the variable **CREL**.

By merging datasets using ELIG, the corresponding parental interviews will be merged: Main with Main and Partner with Partner interview (where available). However, the actual person responding as Main (or Partner where available) may not be the same with the one of the other sweep. The mcs\*\_parent\_derived dataset provides information on whether there has been a change in the identity (PNUM) of the person responding as Main or Partner.

The RESP variable provides information on whether the individuals selected for the Main/Partner role (ELIG) proceeded with the interview. It can be used to remove missing information across the variables since some individuals did not participate in the interview even if they were eligible (ELIG).

Finally, if the project requires information about both parents (wherever there is a Partner respondent), a solution to increase the sample size of the two parent/carers families is to use the \_proxy\_partner\_ dataset in addition to the parent.

# 1.6.2 Merging between sweeps: specific person continuity across sweeps (PNUM)

If it is important for the research project that the information comes from the same respondent, then the use of MCSID and PNUM is recommended. The Person Number (PNUM) is the same for an individual across sweeps. The resulting dataset will contain the respondents who have remained the same across different sweeps.

However, respondents who have not been present in later sweeps will not appear in the dataset (\_parent\_ or \_parent\_cm\_ levels). This will result in a dataset that has lower sample size than the dataset merged using MSCID and ELIG (or MCSID and RESP). Even if the person has moved out, s/he still has a PNUM and is part of the \_hhgrid\_ dataset (across sweeps).

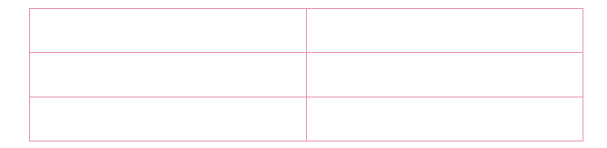
When merging focusing on specific person across sweeps (PNUM) the \_proxy\_partner\_ datasets will help increase the sample.

### 2 Datasets of MCS & how to merge them within sweep

This section applies to the long (stacked) format of the datasets of MCS as this is the format that MCS is and will be provided in future sweeps. The examples of code of this section can be applied on any sweep of MCS that is available in a long format. The description of the datasets and the data handling (e.g. merging) of this section have been conducted in R, SPSS Syntax and STATA. The syntax for each of these statistical packages is provided at the end of this guide.

Every **Example code** \_\_\_ of this section corresponds to respective syntax for R, SPSS Syntax and STATA.

It is important that the reader tries out and syntax provided while following the text.



### 2.1 Household grid dataset: mcs\*\_hhgrid

The household grid contains information about the individuals that live or have lived in the same household with the Cohort Member(s). There is one row per person (PNUM) and one row per Cohort Member (CNUM). Here, the data user can find gender and age of each person that lives in the household including for the Cohort Members.

This file contains all the key identifiers that are used in the rest of the datasets of MCS: MCSID, PNUM, CNUM, ELIG, RESP.

MCSID	*PNUM	*CNUM	*ELIG	*RESP
Family identifier	Person number	CM number	Eligibility for interview	Response to Interview
Family 1	1	-1	Main Interview	Main Interview
Family 1	2	-1	Partner Interview	Partner Interview
Family 1	-1	1	-1	-1
Family 2	1	-1	Main Interview	Main Interview
Family 2	2	-1	Partner Interview	Partner Interview
Family 2	3	-1	-1	-1
Family 2	-1	1	-1	-1
Family 2	-1	2	-1	-1

Figure 4: Household Grid identifiers

When looking at the household grid (especially from Sweep 2 onwards) it is important to remember that not all the individuals that have a PNUM still live in the

household. Some people are not part of the household in a subsequent sweep and this is captured by PRES (presence of the individual during the interview).

An important variable is CREL that captures the relationship to the CM.

Many derived variables on relationships within the household are based on CREL. So, if the research project requires information about relationships in the household in general (mcs\*\_family\_derived) or of the main respondents (HTYP) it is worthwhile looking at the datasets that contain the derived variables and the derived variables user guide.

**Example code A** shows how the data of some families look like.

The Cohort Members do not have PNUM but only CNUM. We can see in PRES that the family structure has changed a lot for some families (where many people have left or passed away) whereas it has remained the same for some others.

### 2.1.1 Household relationships grid

The relationship of each person to any other person of the household is contained in the relationships grid. This information is very useful as one can draw varying family structures across sweeps, however, it may contain error. The information about the relationship of person X to person Y can contain measurement error firstly due to sensitivity of the question and secondly due to misunderstanding. Namely, a question about whether the relationship of person X to person Y is 'natural sibling' may be sensitive under certain circumstances. Moreover, the terms 'natural sibling' and 'half sibling' may not sound that different even if they imply one parent different between the siblings.

	Family	Person Number	Cohort Member Number	Info about person (age, gender, present in hh)	Relation- ship to cohort member	Relatio nship to Person 1	Relatio nship to Person 2	Relatio nship to Person 3	Relatio nship to Person 4
	Family 1	1			Parent				
$\stackrel{\circ}{\Box}$	Family 1	2			Parent				
	Family 1	3			Sibling				
	Family 1	4							
	Family 1		1						
	Family 2	1			Parent				
$\stackrel{\circ}{\Box}$	Family 2	2			Grandparent				
	Family 2		1						
	Family 2		2						
	Family 3	1			Partner				
	Family 3		1						

Figure 5: Relationships grid

#### 2.1.2 Generating person / cohort members identifiers

For various data manipulations the data user may need a person identifier instead of a family identifier that is the MCSID. It is possible to concatenate (join together) the MCSID with the PNUM or ELIG or CNUM to create a unique person identifier.

**Example code B** generates a person identifier for each person in the household and a person identifier for each individual of the household including Cohort Members. The syntax concatenates (joins) to create a PID (person ID) for people in the household only and for everyone in the household (including Cohort Members).

The person identifier can be constructed in any dataset, however, the household grid contains all the families regardless if they completed the entire interview of that sweep (full interview or partial completion).

### 2.2 Cohort member datasets: mcs\*\_cm\_

MCSID	CNU	JM
Family identifier	Cohort Mem Num	
Family 1	Be aware of more	1
Family 1	than one Cohort  Member per family	2
Family 2	(i.e. twins & triplets) like Family 1	1
Family 3		1

Figure 6: Cohort member structure dataset

The datasets that only contain \_cm\_ in the dataset name contain 1 row per cohort member. These can be interview information (mcs\*\_cm\_interview), results of the cognitive assessments (mcs\*\_cm\_cognitive\_assessment),

# 2.2.1 The number of Cohort Members within a sweep (CNUM versus NOCM)

MCSID	VAR
Family identifier	Variable
Family 1	1
Family 2	1
Family 3	2

Figure 7: Family structure dataset

The design of NOCM helps take a closer look to the long format of the MCS dataset. This variable tells us the total number of Cohort Members in each household. Some families have 1 Cohort Member but some have 2 and 3 (twins and triplets).

By running **Example code C**, we can get frequencies of NOCM from the mcs\*\_family\_derived and frequencies of CNUM from the mcs\*\_cm\_derived datasets.

NOCM contains the total numbers of Cohort Members in each household.

CNUM	NOCM = 1	NOCM = 2	NOCM = 3	Total in CNUM
CNUM = 1				
CNUM = 2				
CNUM = 3				

Figure 8: Correspondence of NOCM and CNUM (please use example code C to fill in)

The total number of Cohort Members per household (NOCM) indicates the variation of CNUM. For example, families that have 1 cohort member (NOCM = 1) cannot have CNUM higher than 1.

However, it is possible for a Cohort Member to decide not to participate and in this case the last column of the table would be slightly different, for instance, the CNUM = 1 in a NOCM = 2 family could be missing due to non-response.

### 2.2.2 Selecting families based on number of Cohort Members

This distinction between CNUM and NOCM can be helpful, depending on the need of the research project, to select households based on the one or the other. For example, the user can use NOCM and keep only families with 1 CM (instead of families with twins or triplets). In this case, the CNUM will be 1 across the file and there will be 11576 families in the dataset (that satisfy NOCM=1). Otherwise, the researcher may decide to keep only families that have 2 or 3 CMs (NOCM=2 or 3). The CNUM will range from 1 to 3 and it will contain only families with more than one CMs (no singleton families).

If the user selects Cohort Members using CNUM, and selects for example, only CNUM=1 (the first CM of each family), then the file will contain the first Cohort Member of families that have twins or triplets. If the user selects Cohort Members

that have CNUM=2 or CNUM=3 (the second and third CM of each family), then the file will not include any Cohort members that have CNUM=1 including those that belong to families with twins or triplets.

# 2.3 Parent and proxy partner datasets: mcs\*\_parent\_interview(1 row per parent/carer)

The datasets that contain \_parent\_ only in their title, refer to information coming from sections where the parent(s) provide information about themselves. The mcs\*\_parent\_interview dataset includes the household questionnaire with information about the household, e.g. language used at home.

MCSID	PNUM	ELIG	RESP
Family identifier	Person number	Eligibility for role of Main/Partner	Response to Interview
Family 1	1	Main Interview	Main Interview
Family 1	6	Partner Interview	Partner Interview
Family 2	1	Main Interview	Main Interview
Family 2	3	Partner Interview	Partner Interview
Family 3	2	Main Interview	Main Interview

Figure 9: Parent structure dataset

### 2.3.1 mcs\*\_parent\_derived dataset & PNUM vs ELIG/RESP

The mcs\*\_parent\_derived file contains information about the main and partner respondent. The output of **Example code D** focuses on specific families have been selected to illustrate the difference between PNUM, ELIG and RESP. Most of these families have Main and Partner respondents, hence, there are 2 rows per family. In the same output, a Partner respondent is eligible but not available for interview, therefore, the Main answers some questions about him/her (proxy\_partner\_interview dataset). In another family, Main and Partner respondents are eligible but only the

Main has participated in the interview (RESP=4 'No interview', for the Partner). In the last family, only a Main respondent is eligible for the interview.

# 2.3.2 Merging mcs\*\_parent\_interview between different sweeps with ELIG or PNUM

If the research project requires data on the \_parent\_ level from two plus sweeps, the merging of the datasets needs to take into account the MCSID and an identifier of the parent respondent. **Example code E** that demonstrates the merge of the \_parent\_ datasets of two sweeps uses firstly the MCSID and ELIG and secondly the MCSID and PNUM. This is a good exercise of merging MCS data from different sweeps focusing either on person continuity (PNUM) or information availability (ELIG/RESP).

In Example code E, MCSID and ELIG have been used to match between different sweeps of \_parent\_ level datasets of MCS. Assuming that the focus is on getting data available about carers of the CM regardless if they are different individuals compared to the previous sweep then MCSID and ELIG can be used. A certain number of Main and Partner respondents gets matched between MCS5 and MCS6 but not all cases. There are two reasons for unmatched cases: non-response on the household level (the family has not participated in the one sweep or the other) or non-response on the person level (for example, the Partner respondent may have moved out, thus there is no Partner respondent but only Main for that family).

Comparing the PNUM of Sweep 5 and Sweep 6, it is possible to see that PNUM has remained the same in more than \_\_\_\_ cases (please run example code for exact number). This means that in these rows the respondent (Main or Partner) is the same person in both sweeps.

In **Example code E**, **MCSID** and **PNUM** have been used assuming that the focus is on the same person answering questions about the family and the parental role.

In this case, more than \_\_\_\_ respondents (please run example code for exact number) have remained the same between MCS5 and MCS6 and they have been successfully matched. Both reasons for unmatched cases that occur when merging

with MCSID and ELIG apply here too: some families have not participated but even if they did a second carer of that family may have not participated.

A problem that may arise when merging by PNUM is the fact that a person participated in both sweeps as a parent respondent, however, the role may be different. The role that the person followed during the interview (ELIG) may change from the one sweep to the other. We see that the majority of the respondents out of the total number that participated in both sweeps have the same role (ELIG: Main or Partner). Most of the questions in the parent interview are addressed to both parents. However, some questions are addressed only to the Main respondent (for example, the Household Questionnaire, the Strengths and Difficulties Questionnaire) or only to the Partner. So, even though the same person (parent/carer identified by PNUM) participated in both sweeps, data may be missing (-1 Not applicable) if the individual has participated with a different interview role (ELIG) in each sweep when a particular variable has a Main/Partner only routing.

### 2.3.3 Composite score per family in a \_parent\_ level dataset

The research project may require one piece of information about the family regardless how many parents are in the household. At this point it is important to prioritise the derived variables datasets of any structure (\_cm\_, \_parent\_, \_family\_) because they include key information about the family, the survey and the individuals that participated in that sweep.

In **Example code F**, a variable is calculated that contains the mean of the parents'/carers' self-reported health from the variable GENA of MCS6. In the same example code, it is shown how to calculate a variable that contains the highest NVQ of the parents'/carers' using the DNVQ variable from the \_parent\_derived dataset.

### 2.3.4 Parent structure dataset: mcs\*\_proxy\_partner\_interview

The proxy partner interview can be used to increase the number of Partner respondents in the datasets. The proxy partner interview occurs when the second carer (Partner) of the Cohort Member may not be available to participate in the

interview. This means that the person lives in the household but s/he has not been available for the interview. In these cases, the Main respondent is asked whether s/he is willing to provide information about his/her Partner. The Proxy Partner part of the questionnaire focuses on the key questions that are asked in the parent interview and are important to collect about the person.

The proxy partner module is particularly useful for research focusing on 2-parent or 2-carers families or research that requires information about the same person (PNUM) across time. Through this module information (for example, health, income, employment) is available about the parent who is not able to participate.

Therefore, by merging the information of the \_proxy\_partner\_interview and the \_parent\_interview it is possible to increase the number of Partner respondents that we have information about as well as the families with 2-parents or 2-carers.

Using the syntax of **Example code G**, we take a look at CREL variable. We see that the majority of the parents that are not available for the interview (but still live in the household) are natural parents. The variables PXRE and PXIN are very helpful in understanding the reasons behind the need for a proxy interview (most common: the partner is working away) and whether the main respondent agreed to provide information in the proxy interview about the partner.

# 2.3.5 Using data of the mcs\*\_proxy\_partner\_interview to maximise the sample size of 2-carers families

Let us suppose that the focus is on increasing the parent sample size with information on general health. For this reason, the variable FXPXGE00 is used from the mcs\*\_proxy\_partner\_interview and the variable FPGENA00 from the mcs\*\_parent\_interview. When combining the mcs\*\_parent\_interview with the information coming from the mcs\*\_proxy\_partner\_interview, it is not possible to merge using parent identifiers. The two datasets do not contain the same respondents. The mcs\*\_parent\_interview includes the parents that have participated in the interview themselves, whereas, the mcs\*\_proxy\_partner\_interview contains

information about the parents that have not been available to participate in the interview. Because of this, we add rows to the dataset (append) rather than merge.

It is important to check what the variable looks like before attempting to combine the datasets. In this case the variables have the same values. The crosstabulation of the variable ELIG with the variable FPGENA00 shows that the dataset contains information from Main and Partner respondents as well as partners that were interviewed through Proxy (Main respondent).

The syntax of **Example code H** highlights the possibilities of handling mcs\*\_proxy\_partner\_interview. It is necessary for the same question to exist in both questionnaires: the parent one and the proxy partner.

The CAPI name is likely to be slightly different, so it is good to focus on the questionnaire to identify a pair of questions that tackle the same issue in the parent and the proxy partner interviews. Once the variables have been identified, then appending the one dataset (mcs\*\_proxy\_partner\_interview dataset) to the other (mcs\*\_parent\_interview) there will be approximately 200-400 additional respondents in the dataset (Proxy Partners).

### 2.4 CM structure datasets: mcs\*\_cm\_interview

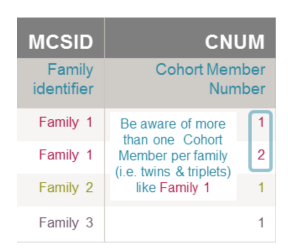


Figure 10: CM structure dataset

The datasets with \_cm\_ structure contain information that is collected directly from the Cohort Members like the Young Person interview, the Physical Measurements and the Cognitive Assessments.

As there is one row per child, the dataset is in a long (stacked) format. There is a family identifier (MCSID) and Cohort Member identifier (CNUM).

As in the mcs\*\_cm\_derived, CNUM is the identifier for the Cohort Member. The variables CSEX, CDBM, CDBY and CAGE come from the household grid (mcs\*\_hhgrid) where the data user can find information about the rest of the members of the household. CAGE has been calculated based on the interview date.

#### 2.4.1 Merging mcs\*\_cm\_interview between different sweeps

If the research project requires data on the \_cm\_ level from two sweeps, the merging needs to take into account the MCSID and the CNUM which is the key identifier for the Cohort Members. The syntax of **Example code I** merges \_cm\_ level datasets from two different sweeps in a similar way to the \_parent\_ datasets.

## 2.5 Parent / CM structure dataset: mcs\*\_parent\_cm\_interview

The mcs\*\_parent\_cm\_interview is an interesting dataset that contains information that the parent(s) provided about each Cohort Member.

MCSID	PNUM	ELIG	RESP	CNL	JM
Family identifier	Person number	Eligibility for role of Main/Partner	Response to Interview	Cohort Mem Num	
Family 1	1	Main Interview	Main Interview	Be aware of	1
Family 1	6	Partner Interview	Partner Interview	more than one Cohort Member per family (i.e. twins & triplets) like Family 1	1
Family 1	1	Main Interview	Main Interview		2
Family 1	6	Partner Interview	Partner Interview		2
Family 2	1	Main Interview	Main Interview		1
Family 2	3	Partner Interview	Partner Interview		1

Figure 11: parent\_cm structure dataset

The routing of the questions in the parent interview varies based on

- a) about whom the question is asked
- b) who is asked (both parents, just the Main or just the Partner).

The questionnaire is the most important source of information on whether a variable has \_parent\_cm\_ structure. As a means of illustration of point a), we can take a look at two questions: WALI and BFEV from the first sweep of MCS. The first question asks the parent how satisfied s/he is with life (WALI). The second one asks the parent whether s/he breastfed the Cohort Member (BFEV). The latter question is repeated for each Cohort Member of the family, namely, for the second and third child of families with twins and triplets. So, even if both questions appear in the parent questionnaire, the one will be located in the \_parent\_dataset (WALI), whereas information about the Cohort Member(s) will be in the \_parent\_cm\_ dataset (BFEV).

Examples of point b) are two questions where the one focuses on the income of both parents and the other one is addressed only towards the one parent. Both of these questions will produce data on the parent level.

Another example is the Strengths and Difficulties Questionnaire (SDQ) that the Main respondent (only) has filled in for each of the Cohort Members of the household. If there is a Partner respondent for a family there will be -1 'Not applicable' for the SDQ for his/her row.

The output of specific variables of a dataset of the \_parent\_cm\_ structure (Example code J) illustrates this structure. Firstly, we notice that there are some variables have '-1 Not applicable' in rows of Partner interview. This is because the question has been asked only from the Main respondent. Each family has provided different data depending on the number of parents participating and number of Cohort Members. For example, a family has Main and Partner respondent answering questions about Cohort Member 1. Another family has only Main respondent answer questions about Cohort Member 1. Also, there is a family that has Main and Partner respondents answering questions about 3 Cohort Members.

#### 2.5.1 Merging mcs\*\_parent\_cm\_interview between different sweeps

The mcs\*\_parent\_cm\_interview contains ELIG/RESP and PNUM as parent identifiers as the mcs\*\_parent\_interview dataset.

Therefore, it can get merged with either MCSID, ELIG, CNUM or MCSID, PNUM, CNUM. A selection of ELIG shows a focus on the data available, whereas selecting PNUM secures person continuity.

**Example code K** merges the data of mcs\*\_parent\_cm\_interview of two difference sweeps with the use of two keys: MCSID and a row ID that is either PNUM & CNUM or ELIG & CNUM. As it happens with the merge of mcs\*\_parent\_interview between sweeps, if we merge by ELIG, different people may have taken the role of the Main or Partner respondent of the CM compared to the previous sweep (specific person gets identified by PNUM). Whereas if we merge by PNUM, the same person may have participated with different roles: namely, the Main in the one sweep and the Partner in another.

### 2.5.2 Extracting information from a mcs\*\_parent\_cm\_interview dataset

There are many pathways for treating data of a mcs\*\_parent\_cm\_interview dataset and they depend on what the research aims are. In this section, we provide some ideas and the respective code.

In questions that have been asked only from the Main respondent (like the CSEN and the SDPF of the Strengths and Difficulties Questionnaire) the data user can select only the data of the Main respondent.

If the focus is on using the information provided by both respondents, Main and Partner, then it is possible to use the data as they are or create a composite score. **Example code L** gives an example where a mean of the parent-perceived likelihood to attend University for each CM is calculated. This can be used for example if we would like to examine whether there is difference in how each cohort member has been described by the carers (Main/Partner). In families where there are two parent respondents (Main and Partner) we can use the fact that the information is provided by both carers about the cohort member. The distance ((dis)agreement) between the responses of the two parents or the mean score of the two responses can be used to enhance analysis.

# 2.5.3 The mcs\_longitudinal\_family\_file: adding outcomes & weights for analysis

The mcs\_longitudinal\_family\_file holds information about every family that has been issued to participate in the MCS. It is therefore a point of reference about the total number of families.

Moreover, it includes families that have left the study in subsequent sweeps due to various reasons (refusal, untraced). This file includes important information about the outcome of the family in each sweep and the weights that can be used for analysis.

**Example code M** merges a dataset of each structure of MCS6 to the mcs\_longitudinal\_family\_file. This code can be used with any sweep.

## 3. Examples of data restructures

This chapter provides details and syntax on matching the different datasets of the same sweep while keeping the highest amount of information possible.

The research question dictates the data handling needed. The research scenarios of this chapter are hypothetical and they have been designed to help to illustrate how to restructure the data into one dataset. They are not example of best practice or recommendations for research.

#### The research scenarios are:

- 3.1 when two datasets with 1-level get matched (example: mcs\*\_cm\_interview & mcs\*\_parent\_interview)
- 3.2 when two datasets of 1-level and 2-level get merged into a 1-level dataset (example: mcs\*\_parent\_cm\_interview & mcs\*\_cm\_interview into a \_cm\_ level dataset)
- 3.3 when two datasets of 1-level and 2-level get merged into a 2-level dataset (example: mcs\*\_parent\_cm\_interview & mcs\*\_cm\_interview into a \_parent\_cm\_ level dataset)

In order to get the most out of this section, it is important to have read and tried the preceding chapters.

### 3.1 Merging two 1-level datasets with different identifiers

As we saw earlier, in MCS, the \_parent\_ level dataset and the \_cm\_ level dataset have 1-level. Namely, either 1 row per parent (identifiers PNUM / ELIG & RESP) or 1 row per cohort member (identifier CNUM). It may be required for a research project to merge these two datasets.

The most important decision is which one of the two datasets contains the main variable of interest or the outcome/dependent variable. This way it is possible to select the dataset that will remain the same (dependent / outcome variable) and which dataset will get restructured (independent / predictor variable).

#### Diagram of merging \_cm\_ and \_parent\_ to a \_cm\_ level dataset

The diagram shows the process of merging two datasets that have 1-level but contain different key identifiers. We need to manipulate the dataset that does not hold the main variable of interest (dependent / outcome variable). Namely, we focus

on restructuring the dataset that has the independent (or outcome) variables. We can either reshape it into wide or calculate a composite variable per family. After the manipulation of the one dataset we simply merge.

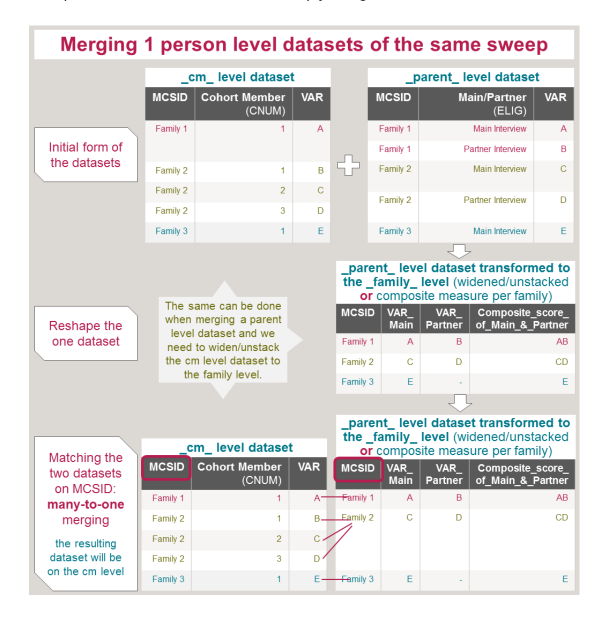


Figure 12: Merging 1 person level datasets of the same sweep (parent with CM)

Merging: \_cm\_ or \_parent <= \_cm\_ + \_parent\_

In this example, we want to examine to what extent the parents/carers' general health correlates with the general health reported by the CM.

The variables that we will be using for this example are:

Dataset	Variable name	Variable label

mcs6_cm_interview.sav	FCCGHE00	CM's general level of health
mcs6_parent_interview.sav	FPGENA00	Respondent's general health
mcs6_proxy_partner_interview.sav	FXPXGE00	Describe partner's health

We need the information on general health of both parents, so we will run **code** H that collapses the information coming from the \_parents\_ dataset as well as from the \_proxy\_partner\_ dataset for the partners that were not present during the interview. After this we follow the diagram. We can either restructure the new \_parent\_ dataset into wide format or create a composite score of the two parents/carers. **Example code** N does both. It generates a composite score (mean health of parents per household) and then restructures the data into a wide format.

It is important to remember that the variable you select to restructure from long to wide will split the variables in the wide format. In this case we restructure from long to wide using ELIG. The **Example code N** makes sure that ELIG has only 2 levels (Main and Partner&Proxy) because a larger number of levels in ELIG would mean additional variables in the wide format.

As we see in the example that uses the \_parent\_ dataset, turning into the wide format using the ELIG variable has created 2 variables per variable of the long format. If we turned into wide format the \_cm\_ dataset (one row per family) using the CNUM, that would create 3 variables per variable of the long format because the CNUM has 3 possible values occurring.

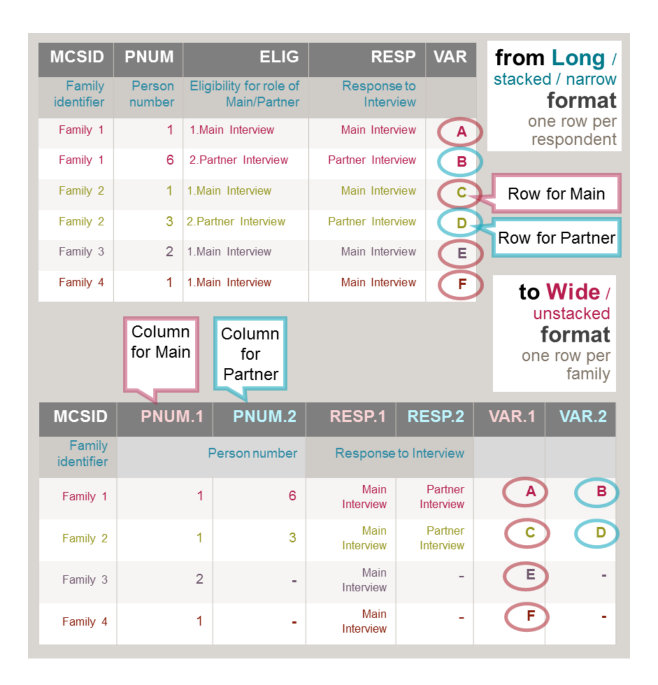


Figure 13: Restructuring long to wide

The **Example code N** concludes with merging the wide format dataset (one family per row) with the \_cm\_ level dataset. We have one row per CM and in each row there is information with the mean score of the parents of that family.

# 3.2 Merging 1-level dataset with a 2-level dataset resulting in a 1-level dataset

We may wish to connect information from a 2-level dataset (\_parent\_cm\_) to a 1-level dataset (\_cm\_ or \_parent). As in the previous example we need to select what is our main variable of interest (dependent variable) and restructure the other datasets (that have the independent variables) to match the dataset that our dependent variable is located.

In this section, we assume that the dependent variable is located in the 1-level dataset, so our focus is on ending up with a 1-level dataset (either \_parent\_ or \_cm\_ level).

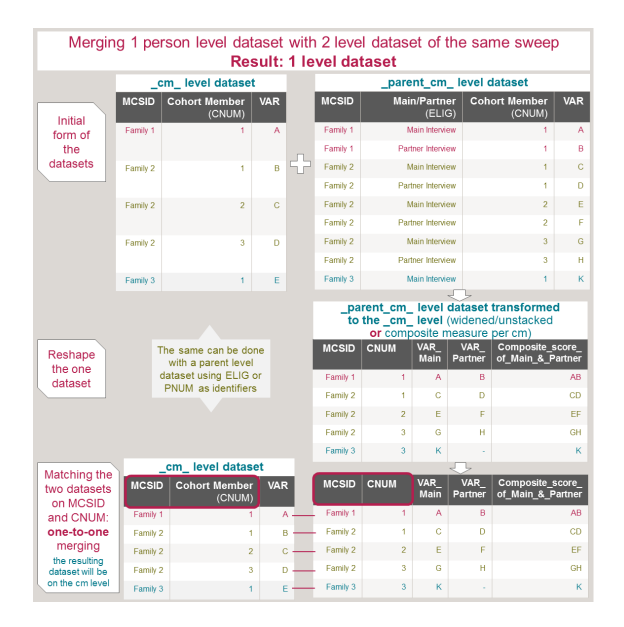


Figure 14: Merging 1 level with a 2 level dataset resulting in a 1 level dataset

The diagram shows a possible path for this. Assuming that our main variable of interest (dependent) is located on the \_cm\_ dataset, we manipulate the data of our independent variable that is located in the \_parent\_cm\_ dataset. We just need to either restructure the \_parent\_cm\_ or generate a composite score per child or per family, that will get merged to the \_cm\_ dataset.

In this example, we want to examine to what extent the parents/carers' aspiration that the CM will go to the University correlates with CM's perception on how likely it is that s/he will go to the University.

The variables that we will be using for this example are:

Dataset	Variable name	Variable label
mcs6_cm_interview.sav	FCSTYU00	How likely is it CM will go to university? (Scale 0-100%)
mcs6_parent_cm_interview.sav	FPASLU00	How (un)likely do you think it is that CM will attend university?

The first step is similar to **Example code L** that generates a composite score for variable ASLU for each child at the \_parent\_cm\_ dataset. This composite score is the mean score of the Main and the Partner for each CM. **Example code O** includes the code of this section.

We create a composite measure of ASLU (mean of parents' perception on how likely it is that the CM will go to the University). We turn the dataset into wide using ELIG, so the information of the parents will be split into two variables: ASLU.1 for the Main and ASLU.2 for the Partner whereas the composite score remains one variable as it is the same for Main and Partner.

You can recalculate the composite score now instead earlier. In the dataset there is one row per child. So, now that the \_parent\_cm\_ has been turned into a \_cm\_ level dataset, we just merge it with the \_cm\_interview.

# 3.3 Merging 1-level dataset with a 2-level dataset resulting in a 2-level dataset

This example is similar to the previous one, however, instead of trying to reduce the \_parent\_cm\_ dataset to a 1-level, we merge it with a 1-level dataset. In case the main variable of interest (dependent variable) is in a 2-level dataset (namely in the \_parent\_cm\_ dataset), then we need to keep the structure of the dataset the same

and merge to another datast, a 1-level dataset (a \_parent\_ or a \_cm\_ structure dataset) that has the independent variables.

#### Diagram of merging \_cm\_ and \_parent\_cm\_ to a \_parent\_cm\_ level dataset

As we see in the diagram, this merging is the easiest as no dataset requires restructuring.

Merging 1 person level dataset with 2 level dataset of the same sweep  Result: 2 level dataset								
Matching the	_cm_ level dataset				_parent_cm_ level dataset			
two datasets on MCSID and	MCSID	Cohort Member (CNUM)	VAR _1		MCSID	Main/Partner (ELIG)	Cohort Member (CNUM)	VAR _2
CNUM:	Family 1	1	L-		Family 1	Main Interview	1	Α
one-to-many merging			_		Family 1	Partner Interview	1	В
the resulting	Family 2	1	M –	53	Family 2	Main Interview	1	С
on the parent-					Family 2	Partner Interview	1	D
cm level	Family 2	2	N -		Family 2	Main Interview	2	E
The same can be					Family 2	Partner Interview	2	F
done with a parent level	Family 2	Family 2 3	0		Family 2	Main Interview	3	G
dataset using			_		Family 2	Partner Interview	3	Н
ELIG or PNUM as identifiers	Family 3	1	P -		Family 3	Main Interview	1	K

Figure 15: Merging 1 level with a 2 level dataset

#### Merging: \_parent\_cm\_ <= \_cm\_ + \_parent\_cm\_

In this example, we want to examine whether there is a correlation between how much the Cohort Member trusts others (\_cm\_interview) and how close the parents feel that they are to the CM (parent\_cm\_interview).

The variables that we will be using for this example are:

Dataset	Variable	Variable label
	name	

mcs6_cm_interview.sav	FCSTYU00	How likely is it CM will go to university? (Scale 0-100%)
mcs6_parent_cm_interview.sav	FPSCHC00	Overall, how close would you say you are to CM?

**Example code P** includes the syntax for this section.

### 4. Example code with R, SPSS Syntax and STATA

The syntax provided in this section is indicative of different working paths with the datasets of MCS and may contain errors. Users need to build syntax for their own project.

The code provided uses primarily MCS5 and MCS6, however, any long format dataset of MCS can be used. The MCSIDs selected to visualise the data structure were selected for demonstration purposes.

Minor differences between the code provided in SPSS, R and STATA exist due to the different commands and functions in each piece of software.

The calculation of the mean (**Example codes**) in each piece of software may require adjustment to the needs to a particular project as sometimes mean is calculated only for families that there are two carers respondents (Main and Partner) whereas other times for all families (including Main only).

Depending on the research project and type of analysis, clustering of standard errors may be needed if there are multiple rows per family (see User Guide S1-5, sections 2.9 and 6.4.3).

# Overview of the example codes

Table 1: Example code contents

Example code	Focus	Question
Example code A	Overview of the hhgrid	How does the household grid (hhgrid)dataset look like?
Example code B	Concatenating MCSID & person identifier to get a unique person identifier	How do I create a unique person identifier by concatenating MCSID & person identifier?
Example code C	Overview of the _family_derived	How does the _family_derived dataset look like?
Example code D	Overview of _parent_derived	How does the _parent_derived dataset look like?
Example code E	Merge _parent_ structure datasets from different sweeps	How do I merge _parent_ structure datasets from different sweeps?
Example code F	Create a composite variable per family in the _parent_ structure file	How do I create a composite variable per family in the _parent_ structure file?
Example code G	Overview of _proxy_partner_interview	How does the _proxy_partner_interview dataset look like?
Example code H	Combining proxy_partner_interview with parent_interview (append)	How do I combine the proxy_partner_interview dataset with the parent_interview dataset?
Example code I	Merge _cm_ structure datasets from different sweeps	How do I merge _cm_ structure datasets from different sweeps?
Example code J	Overview of parent_cm_interview	What does the parent_cm_interview dataset look like?
Example code K	Merge _parent_cm_ level datasets between sweeps	How do I merge the _parent_cm_ level datasets between sweeps?
Example code L	Create a composite variable per child in parent_cm dataset	How do I create a composite variable per child in parent_cm dataset?
Example code M	Merging datasets of different structures to the mcs_longitudinal_family_file	How do I merge datasets of different structures to the mcs_longitudinal_family_file?

Example code N	Merging two 1-level datasets that have different identifiers	Merging two 1-level datasets that have different identifiers (_parent_interview dataset with _cm_interview dataset)
Example code O	Merging a 1-level dataset (_cm_) with a 2-level dataset (_parent_cm_) resulting into a 1-level structure (_cm_)	Merging a 1-level dataset (_cm_) with a 2-level dataset (_parent_cm_) resulting into a 1-level structure (_cm_)
Example code P	Merging a 2-level dataset (_parent_cm_) with a 1-level dataset (_cm_) resulting into a 2-level dataset (_cm_)	Merging a 2-level dataset (_parent_cm_) with a 1-level dataset (_cm_) resulting into a 2-level dataset (_cm_)

## R syntax

### # Setting up folders in R

```
# in case you need to clean the workspace
#rm(list=ls())

# the aim is to use *core* R functions in this syntax
# various packages exist for data management that users may
prefer

# download packages needed
install.packages("foreign")

# load packages needed
```

```
data folder_path = "//" # my folder path
# the data of mcs5 and mcs6 need to be in folders mcs6 and
mcs5
# Example code A
# -----.
# Overview of the hhgrid .
# -----
mcs6 hhgrid <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 hhgrid.sav", sep = ''), to.data.frame = TRUE,
use.missings=TRUE)
mcs6 hhgrid$MCSID <- trimws(mcs6 hhgrid$MCSID, which =</pre>
c('right')) # remove white space from MCSID
str(mcs6 hhgrid$MCSID)
# Overview of PRES / CREL / multiple Cohort Members per
family.
print(mcs6 hhgrid[which(mcs6 hhgrid$MCSID == 'M10002P' |
mcs6 hhgrid$MCSID == 'M10611J' | mcs6 hhgrid$MCSID ==
'M10611J' | mcs6 hhgrid$MCSID == 'M10106W' | mcs6 hhgrid$MCSID
== 'M10063C'), c('MCSID', 'FPNUM00', 'FELIG00', 'FRESP00',
'FCNUM00', 'FHCREL00', 'FHPRES00')])
# NA are the -1 not applicable of the original dataset
```

library(foreign)

```
# Example code B
# Concatenating MCSID & person identifier to get a unique
person identifier .
mcs6 hhgrid <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 hhgrid.sav", sep = ''), to.data.frame = TRUE)
mcs6 hhgrid$MCSID <- trimws(mcs6 hhgrid$MCSID, which =</pre>
c('right')) # remove white space from MCSID
str(mcs6 hhgrid$MCSID)
str(mcs6 hhgrid$FPNUM00)
str(mcs6 hhgrid$FCNUM00)
# a Person ID for each adult in the household (excluding
cohort members) .
mcs6_hhgrid$PnumID <- ifelse(is.na(mcs6_hhgrid$FCNUM00),</pre>
                          paste(mcs6_hhgrid$MCSID,
mcs6 hhgrid$FPNUM00, sep = ' P'),
                          mcs6 hhgrid$PnumID <- NA)</pre>
head(mcs6 hhgrid$PnumID)
```

```
\# a Person ID for each individual of the household (Cohort
Member or other person)
mcs6 hhgrid$PID <- ifelse(is.na(mcs6 hhgrid$FPNUM00),</pre>
                             paste(mcs6 hhgrid$MCSID,
mcs6 hhgrid$FCNUM00, sep = ' C'),
                             paste (mcs6 hhgrid$MCSID,
mcs6 hhgrid$FPNUM00, sep = ' P'))
head(mcs6 hhgrid$PID)
# Example code C
# -----.
# Overview of the family derived .
mcs6 family derived <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 family derived.sav", sep = ''), to.data.frame =
TRUE)
table(mcs6 family derived$FDNOCM00)
table(mcs6 family derived$FDRSP000)
# Overview of the cm derived .
mcs6 cm derived <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 cm derived.sav", sep = ''), to.data.frame = TRUE)
table(mcs6 cm derived$FCNUM00)
```

```
# Example code D
# -----.
# Overview of _parent_derived .
# -----.
mcs6 parent derived <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 parent derived.sav", sep = ''), to.data.frame =
TRUE)
mcs6 parent derived$MCSID <- trimws(mcs6 parent derived$MCSID,</pre>
which = c('right'))
mcs6 parent derived$FELIG00 <-</pre>
trimws(mcs6 parent derived$FELIG00, which = c('right'))
mcs6 parent derived$FRESP00 <-</pre>
trimws(mcs6 parent derived$FRESP00, which = c('right'))
print(mcs6 parent derived[which(mcs6 parent derived$MCSID ==
'M10002P' | mcs6 parent derived$MCSID == 'M10041W' |
mcs6 parent derived$MCSID == 'M23136V' |
mcs6 parent derived$MCSID == 'M10106W' |
mcs6 parent derived$MCSID == 'M10063C'),
                 c('MCSID', 'FPNUM00', 'FELIG00',
'FRESP00')])
```

```
# Example code E
```

# -----.

```
# merge parent structure datasets from different sweeps.
# \sim \sim \sim \sim merge the two datasets on MCSID & ELIG \sim \sim \sim
# load the mcs6 parent dataset.
mcs6 parent interview <- read.spss(file=</pre>
paste(data folder path, "/mcs6 /mcs6 parent interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent interview$MCSID <-</pre>
trimws(mcs6 parent interview$MCSID, which = c('right'))
mcs6 parent interview$FELIG00 <-</pre>
trimws(mcs6 parent interview$FELIG00, which = c('right'))
# prepare the mcs5 parent dataset.
mcs5 parent interview <- read.spss(file=</pre>
paste(data_folder_path, "/mcs5 /mcs5 parent interview.sav",
sep = ''), to.data.frame = TRUE)
mcs5 parent interview$MCSID <-</pre>
trimws(mcs5 parent interview$MCSID, which = c('right'))
mcs5 parent interview$EELIG00 <-</pre>
trimws(mcs5 parent interview$EELIG00, which = c('right'))
# create variables to be able to inspect the merge.
mcs6 parent interview$source mcs6 <- 'MCS6'</pre>
```

```
mcs5 parent interview$source mcs5 <- 'MCS5'</pre>
mcs6 parent interview$ELIG <- mcs6 parent interview$FELIG00</pre>
mcs5 parent interview$ELIG <- mcs5 parent interview$EELIG00</pre>
# merge .
str(mcs6 parent interview)
str(mcs5 parent interview)
mcs5 mcs6 parent interview <- merge(x=mcs6 parent interview,
y=mcs5 parent interview,
                                      by.x = c("MCSID", "ELIG"),
                                      by.y = c("MCSID", "ELIG"),
                                      all.x = TRUE, all.y =  
TRUE)
# inspect the merge
str(mcs5 mcs6 parent interview)
mcs5 mcs6 parent interview$rowsource <-</pre>
apply(mcs5 mcs6 parent interview[c('source mcs5',
'source mcs6')], # object to work with
                                                1, # 1 for rows
- 2 for columns
                                                function(x)
paste(na.omit(x), collapse = " ")) # function
table(mcs5 mcs6 parent interview$rowsource)
# Outcome perusal: Main and Partner respondents (ELIG) in both
sweeps.
```

```
str(mcs5 mcs6 parent interview$FELIG00)
str(mcs5 mcs6 parent interview$EELIG00)
mcs5_mcs6_parent interview$FELIG00 <-</pre>
as.factor(mcs5 mcs6 parent interview$FELIG00)
mcs5 mcs6 parent interview$EELIG00 <-</pre>
as.factor(mcs5 mcs6 parent interview$EELIG00)
table (mcs5 mcs6 parent interview$FELIG00,
mcs5 mcs6 parent interview$EELIG00)
# Let us see how many of the merged Main & Partner respondents
(ELIG) have
# the same PNUM, therefore they are they same person .
mcs5 mcs6 parent interview$same respondent <-</pre>
ifelse(mcs5 mcs6 parent interview$FPNUM00 ==
mcs5 mcs6 parent interview$EPNUM00, 1, 0)
# 1 the same respondent - 0 different respondent
table (mcs5 mcs6 parent interview$same respondent,
mcs5 mcs6 parent interview$ELIG)
\# ~ ~ ~ ~ ~ merge the two datasets on MCSID & PNUM ~ ~ ~ ~
# load the mcs6 parent dataset.
```

```
mcs6 parent interview <- read.spss(file=</pre>
paste(data folder path, "/mcs6 /mcs6 parent interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent interview$MCSID <-</pre>
trimws(mcs6 parent interview$MCSID, which = c('right'))
mcs6 parent interview$FELIG00 <-</pre>
trimws(mcs6 parent interview$FELIG00, which = c('right'))
# prepare the mcs5 parent dataset.
mcs5 parent interview <- read.spss(file=</pre>
paste(data folder path, "/mcs5 /mcs5 parent interview.sav",
sep = ''), to.data.frame = TRUE)
mcs5 parent interview$MCSID <-</pre>
trimws(mcs5 parent interview$MCSID, which = c('right'))
mcs5 parent interview$EELIG00 <-</pre>
trimws(mcs5 parent interview$EELIG00, which = c('right'))
# create variables to be able to inspect the merge.
mcs6 parent interview$source mcs6 <- 'MCS6'</pre>
mcs5 parent interview$source mcs5 <- 'MCS5'</pre>
mcs6 parent interview$PNUM <- mcs6 parent interview$FPNUM00</pre>
mcs5 parent interview$PNUM <- mcs5 parent interview$EPNUM00</pre>
# merge .
str(mcs6 parent interview)
str(mcs5 parent interview)
```

```
mcs5 mcs6 parent interview <- merge(x=mcs6 parent interview,</pre>
y=mcs5 parent interview,
                                     by x = c ("MCSID", "PNUM"),
                                     by.y = c("MCSID", "PNUM"),
                                     all.x = TRUE, all.y =
TRUE)
# inspect the merge
str(mcs5 mcs6 parent interview)
mcs5 mcs6 parent interview$rowsource <-</pre>
apply(mcs5 mcs6 parent interview[c('source mcs5',
'source mcs6')], # object to work with
                                               1, # 1 for rows
- 2 for columns
                                               function(x)
paste(na.omit(x), collapse = "&")) # function
table(mcs5 mcs6 parent interview$rowsource)
# Outcome perusal: parents/carers (PNUM) in both sweeps.
# Let us see how many of the merged parents/carers respondents
(PNUM) have
# the same role in the interview (ELIG: Main or Partner) .
table(mcs5 mcs6 parent interview$EELIG00,
mcs5 mcs6 parent interview$FELIG00, useNA = 'ifany')
# respondents that were not eligible in the one or the other
sweep have NA
```

```
# the crosstabulation shows the respondents whether
respondents have the
# same or different eligibility between sweeps
# Example code F
# -----.
# Create a composite variable per family in the parent
structure file .
-----.
mcs6 parent interview <- read.spss(file=</pre>
paste(data folder path, "/mcs6 /mcs6 parent interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent interview$MCSID <-</pre>
trimws(mcs6 parent interview$MCSID, which = c('both'));
str(mcs6 parent interview$MCSID)
# this syntax creates a mean of Main and Partner responses on
GENA variable .
# You can use other functions instead of MEAN, like SD, MIN,
MAX, etc.
str(mcs6 parent interview$FPGENA00)
table(mcs6 parent interview$FPGENA00, useNA = 'ifany')
mcs6 parent interview$GENA <-</pre>
as.numeric(mcs6 parent interview$FPGENA00)
```

```
table(mcs6 parent interview$GENA, useNA = 'ifany')
mcs6 parent interview small <-</pre>
mcs6 parent interview[c("MCSID", "FPNUM00", "FELIG00",
"FPGENA00")1
mcs6 parent interview small$GENA num <-</pre>
as.numeric(mcs6 parent interview small$FPGENA00)
str(mcs6 parent interview small)
mcs6 parent interview small composite <-</pre>
aggregate (mcs6 parent interview small$GENA num ~
mcs6 parent interview small$MCSID, FUN=mean, na.rm=TRUE,
na.action="na.omit")
str(mcs6 parent interview small composite)
colnames(mcs6 parent interview small composite) <- c('MCSID',</pre>
'GENA composite')
mcs6 parent interview small composite$MCSID <-</pre>
as.character(trimws(mcs6 parent interview small composite$MCSI
D, which = c('both'))
# connect to the rest of the data
mcs6 parent interview with GENA <- merge(x =
mcs6 parent interview small composite,
                                          y =
mcs6 parent interview small,
                                          by.x = 'MCSID', by.y =
'MCSID', all = TRUE)
str(mcs6 parent interview with GENA)
# let's take a look at the outcome
```

```
print(mcs6 parent interview_with_GENA[which(mcs6_parent_interv
iew with GENA$MCSID == 'M10002P' |
mcs6 parent interview with GENA$MCSID == 'M10611J' |
mcs6 parent interview with GENA$MCSID == 'M10106W' |
mcs6 parent interview with GENA$MCSID == 'M10063C'),
                           c('MCSID', 'FPNUM00', 'FELIG00',
'FPGENA00', 'GENA num', 'GENA composite')])
# this syntax selects the higher NVQ of Main and Partner
respondents.
# You can use other functions instead of MEAN, like SD, MIN,
MAX, etc.
mcs6 parent derived <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 parent derived.sav", sep = ''), to.data.frame =
TRUE)
mcs6 parent derived$MCSID <- trimws(mcs6 parent derived$MCSID,</pre>
which = c('both')); str(mcs6 parent derived$MCSID)
table(mcs6 parent derived$FDNVQ00, useNA='ifany')
mcs6 parent derived$NVQ num <-</pre>
as.numeric(mcs6 parent derived$FDNVQ00);
table(mcs6 parent derived$NVQ num, useNA='ifany')
mcs6 parent derived$NVQ num[mcs6 parent derived$NVQ num >= 6]
<- NA
table(mcs6 parent derived$NVQ num, useNA='ifany')
mcs6 parent derived composite <-
aggregate (mcs6 parent derived$NVQ num ~
mcs6 parent derived$MCSID, FUN=max, na.rm=TRUE,
na.action="na.omit")
```

```
str(mcs6 parent derived composite)
colnames(mcs6 parent derived composite) <- c('MCSID',</pre>
'NVQ composite')
mcs6 parent derived composite$MCSID <-</pre>
as.character(trimws(mcs6 parent derived composite$MCSID, which
= c('both')))
# connect to the rest of the data
mcs6 parent derived with NVQ <- merge(x =  
mcs6 parent derived composite,
                                          у =
mcs6 parent derived,
                                          by.x = 'MCSID', by.y
= 'MCSID', all = TRUE)
str(mcs6 parent derived with NVQ)
# let's take a look at the outcome
print (mcs6 parent derived with NVQ[which (mcs6 parent derived w
ith NVQ$MCSID == 'M10002P' |
mcs6 parent derived with NVQ$MCSID == 'M10611J' |
mcs6 parent derived with NVQ$MCSID == 'M10106W' |
mcs6 parent derived with NVQ$MCSID == 'M10063C'),
                                       c('MCSID', 'FPNUM00',
'FELIGOO', 'FDNVQOO', 'NVQ num', 'NVQ composite')])
```

#### # Example code G

# -----.

```
# Overview of proxy partner interview .
mcs6 proxy partner interview <- read.spss(file=</pre>
paste(data folder path,
"/mcs6 /mcs6 proxy partner interview.sav", sep = ''),
to.data.frame = TRUE)
mcs6 proxy partner interview$MCSID <-</pre>
trimws(mcs6 proxy partner interview$MCSID, which = c('both'));
str(mcs6 proxy partner interview$MCSID)
table(mcs6 proxy partner interview$FXCREL00)
table(mcs6_proxy_partner_interview$FXCREL00,
mcs6 proxy partner interview$FXPSEX00)
table(mcs6 proxy partner interview$FXPXRE00)
table(mcs6 proxy partner interview$FXPXIN00)
# Example code H
# -----.
# Combining proxy partner interview with parent interview .
-----.
```

```
# we keep only rows where the Main agreed to provide
information about the non-available Partner.
nrow(mcs6 proxy partner interview) # observations
mcs6_proxy_partner interview subset <-</pre>
subset (mcs6 proxy partner interview,
mcs6 proxy partner interview$FXPXIN00 == "Continue with PROXY
interview ")
nrow(mcs6 proxy partner interview subset)
table(mcs6 proxy partner interview subset$FXPXGE00)
# we rename the variable to the variable name that is used in
the parent interview dataset.
names (mcs6 proxy partner interview subset) [names (mcs6 proxy pa
rtner interview subset) == "FXPXGE00"] <- "FPGENA00"</pre>
mcs6_proxy_partner_interview_for_connection to parent <-</pre>
mcs6 proxy partner interview subset[c("MCSID", "FPNUM00",
"FELIG00", "FPGENA00")]
# open the parent interview dataset and keep only the
variables needed .
mcs6 parent interview <- read.spss(file=</pre>
paste(data folder path, "/mcs6 /mcs6 parent interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent interview$MCSID <-</pre>
trimws(mcs6 parent interview$MCSID, which = c('both'));
str(mcs6 parent interview$MCSID)
```

```
mcs6 parent interview small <-</pre>
mcs6 parent interview[c("MCSID", "FPNUM00", "FELIG00",
"FPGENA00")]
# we add cases/rows to the dataset - append.
str(mcs6 parent interview small)
str(mcs6 proxy partner interview for connection to parent)
table(mcs6 parent interview small$FPGENA00)
table (mcs6 proxy partner interview for connection to parent$FP
GENA00)
mcs6 parent plus proxy interview <-</pre>
rbind(mcs6 parent interview small,
mcs6 proxy partner interview for connection to parent)
str(mcs6 parent plus proxy interview)
table(mcs6 parent plus proxy interview$FPGENA00)
# there are 2 factor level 'excellent' because the value label
wording is slightly different
# they can get collapsed together
table(mcs6 parent plus proxy interview$FELIG00)
# Example code I
# merge cm structure datasets from different sweeps.
```

```
_________
# prepare the mcs6 cm dataset.
mcs6 cm interview <- read.spss(file= paste(data_folder_path,</pre>
"/mcs6 /mcs6 cm interview.sav", sep = ''), to.data.frame =
TRUE)
mcs6 cm interview$MCSID <- trimws(mcs6 cm interview$MCSID,</pre>
which = c('both')); str(mcs6 cm interview$MCSID)
mcs6 cm interview$CNUM <- mcs6 cm interview$FCNUM00;</pre>
table(mcs6 cm interview$CNUM)
# prepare the mcs5 cm dataset.
mcs5 cm interview <- read.spss(file= paste(data folder path,</pre>
"/mcs5 /mcs5 cm interview.sav", sep = ''), to.data.frame =
TRUE)
mcs5 cm interview$MCSID <- trimws(mcs5 cm interview$MCSID,</pre>
which = c('both')); str(mcs6 cm interview$MCSID)
mcs5 cm interview$CNUM <- mcs5 cm interview$ECNUM00;</pre>
table(mcs5 cm interview$CNUM)
\# merge the two datasets on MCSID & CNUM .
mcs6 cm interview$sweep 6 <- 'Sweep 6'</pre>
mcs5 cm interview$sweep 5 <- 'Sweep 5'</pre>
mcs5 mcs6 cm interview <- merge(x = mcs6 cm interview,</pre>
                                  y = mcs5 cm interview,
```

```
by x = c('MCSID', 'CNUM'),
                              by.y = c('MCSID', 'CNUM'), all
= TRUE)
# Outcome perusal: cohort members in both datasets.
table (mcs5 mcs6 cm interview$sweep 6,
mcs5 mcs6 cm interview$sweep 5, useNA='ifany')
table (mcs5 mcs6 cm interview$CNUM)
table (mcs5 mcs6 cm interview$FCNUM00,
mcs5 mcs6 cm interview$ECNUM00, useNA='ifany')
# Example code J
# -----.
# Overview of parent cm interview .
# -----.
# Parent's interview about the CM(s) of the household.
mcs6 parent cm interview <- read.spss(file=</pre>
paste(data folder path, "/mcs6 /mcs6 parent cm interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent cm interview$MCSID <-</pre>
trimws(mcs6 parent cm interview$MCSID, which = c('both'));
str(mcs6 parent cm interview$MCSID)
```

table(mcs6 parent cm interview\$FCNUM00)

```
# Overview of specific cases on key variables
print(mcs6 parent cm interview[which(mcs6 parent cm interview$
MCSID == 'M10002P' |
mcs6 parent cm interview$MCSID == 'M10611J' |
mcs6 parent cm interview$MCSID == 'M10106W' |
mcs6 parent cm interview$MCSID == 'M10063C'),
                               c('MCSID', 'FPNUM00',
'FELIGOO', 'FCNUMOO', 'FCCSEXOO', 'FCCAGEOO', 'FPSDPFOO',
'FPASLU00')])
# Example code K
# -----.
# Merge parent cm level datasets between sweeps.
______
# Create row identifier to connect for _parent_cm_ datasets.
\# ROWid = (ELIG or PNUM) + (CNUM Child 1/2/3).
\# ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ELIG + CNUM = ROWid ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~
~ ~ ~ ~
# prepare mcs6 parent cm interview .
```

```
mcs6 parent cm interview <- read.spss(file=</pre>
paste (data folder path, "/mcs6 /mcs6 parent cm interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent cm interview$MCSID <-</pre>
trimws(mcs6 parent cm interview$MCSID, which = c('both'));
str(mcs6 parent cm interview$MCSID)
table(mcs6 parent cm interview$FCNUM00)
table(mcs6 parent cm interview$FPNUM00)
mcs6 parent cm interview$ROWid <-</pre>
ifelse(mcs6 parent cm interview$FELIG00 == "Main Interview ",
                                             paste('M C',
mcs6 parent cm interview$FCNUM00, sep = ''),
                                             paste('P C',
mcs6 parent cm interview$FCNUM00, sep = '')
)
table(mcs6 parent cm interview$ROWid, useNA='ifany')
# Check that the ROWid matches the crosstabulation of ELIG &
CNUM .
table (mcs6 parent cm interview$FCNUM00,
mcs6 parent cm interview$FELIG00)
# prepare mcs5 parent cm interview .
mcs5 parent cm interview <- read.spss(file=</pre>
paste(data folder path, "/mcs5 /mcs5 parent cm interview.sav",
sep = ''), to.data.frame = TRUE)
```

```
mcs5 parent cm interview$MCSID <-</pre>
trimws(mcs5 parent cm interview$MCSID, which = c('both'));
str(mcs5 parent cm interview$MCSID)
table(mcs5 parent cm interview$ECNUM00)
table(mcs5_parent cm interview$EPNUM00)
mcs5 parent cm interview$ROWid <-</pre>
ifelse (mcs5 parent cm interview$EELIG00 == "Main Interview ",
                                           paste('M C',
mcs5 parent cm interview$ECNUM00, sep = ''),
                                           paste('P C',
mcs5 parent cm interview$ECNUM00, sep = '')
)
table(mcs5 parent cm interview$ROWid, useNA='ifany')
# Check that the ROWid matches the crosstabulation of ELIG &
CNUM .
table (mcs5 parent cm interview$ECNUM00,
mcs5_parent_cm interview$EELIG00)
# Merge .
mcs6 parent cm interview$sweep 6 <- 'Sweep 6'</pre>
mcs5 parent cm interview$sweep 5 <- 'Sweep 5'</pre>
mcs5 mcs6 parent cm interview <- merge(x =</pre>
mcs6 parent cm interview,
                                  y = mcs5 parent cm interview,
```

```
by x = c('MCSID', 'ROWid'),
                                 by.y = c('MCSID', 'ROWid'),
all = TRUE)
table(mcs5 mcs6 parent cm interview$sweep 6,
mcs5 mcs6 parent cm interview$sweep 5, useNA='ifany')
# Outcome perusal: parents (Main/Partner providing information
about
# each of the cohort members) are in both sweeps.
\# ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ PNUM + CNUM = ROWid ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~
~ ~ ~ ~ .
# prepare mcs6 parent cm interview .
mcs6 parent cm interview <- read.spss(file=</pre>
paste(data folder path, "/mcs6 /mcs6 parent cm interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent cm interview$MCSID <-</pre>
trimws(mcs6 parent cm interview$MCSID, which = c('both'));
str(mcs6 parent cm interview$MCSID)
table(mcs6 parent cm interview$FCNUM00)
table(mcs6 parent cm interview$FPNUM00)
mcs6 parent cm interview$ROWid <- paste('P',</pre>
mcs6 parent cm interview$FPNUM00, 'C',
mcs6 parent cm interview$FCNUM00, sep = '')
```

```
table(mcs6 parent cm interview$ROWid, useNA='ifany')
# Check that the ROWid matches the crosstabulation of PNUM &
CNUM .
table (mcs6 parent cm interview$FPNUM00,
mcs6 parent cm interview$FCNUM00)
# prepare mcs5 parent cm interview .
mcs5 parent cm interview <- read.spss(file=</pre>
paste(data folder path, "/mcs5 /mcs5 parent cm interview.sav",
sep = ''), to.data.frame = TRUE)
mcs5 parent cm interview$MCSID <-</pre>
trimws(mcs5 parent cm interview$MCSID, which = c('both'));
str(mcs5 parent cm interview$MCSID)
table(mcs5 parent cm interview$ECNUM00)
table(mcs5 parent cm interview$EPNUM00)
mcs5 parent cm interview$ROWid <- paste('P',</pre>
mcs5 parent cm interview$EPNUM00, 'C',
mcs5 parent cm interview$ECNUM00, sep = '')
table(mcs5 parent cm interview$ROWid, useNA='ifany')
# Check that the ROWid matches the crosstabulation of PNUM &
CNUM .
table (mcs5 parent cm interview$EPNUM00,
mcs5 parent cm interview$ECNUM00)
```

```
# Merge .
mcs6 parent cm interview$sweep 6 <- 'Sweep 6'</pre>
mcs5 parent cm interview$sweep 5 <- 'Sweep 5'</pre>
mcs5 mcs6 parent cm interview <- merge(x =</pre>
mcs6 parent cm interview,
                                        у =
mcs5 parent cm interview,
                                        by.x = c('MCSID',
'ROWid'),
                                        by.y = c('MCSID',
'ROWid'), all = TRUE)
table(mcs5 mcs6 parent cm interview$sweep 6,
mcs5_mcs6_parent_cm_interview$sweep 5, useNA='ifany')
# Outcome perusal: Individuals (PNUM) provide information
# about the cohort member(s) in both sweeps (either as Main or
Partner respondent).
# Example code L
# -----.
# Create a composite variable per child in parent cm dataset.
# this syntax creates a mean of Main and Partner responses on
ASLU variable .
```

```
# You can use other functions instead of MEAN, like SD, MIN,
MAX, etc.
mcs6 parent cm interview <- read.spss(file=</pre>
paste (data folder path, "/mcs6 /mcs6 parent cm interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent cm interview$MCSID <-</pre>
trimws(mcs6 parent cm interview$MCSID, which = c('both'));
str(mcs6 parent cm interview$MCSID)
# Mean score of ASLU assessment by Main/Partner respondent(s)
of the cohort member
table (mcs6 parent cm interview$FPASLU00, useNA='ifany')
mcs6 parent cm interview$ASLU num <-</pre>
as.numeric(mcs6 parent cm interview$FPASLU00)
table (mcs6 parent cm interview$ASLU num, useNA = 'ifany')
mcs6 parent cm interview small <-</pre>
mcs6 parent cm interview[c('ASLU num', 'MCSID', 'FCNUM00')]
mcs6 parent cm interview composite <-
aggregate (mcs6 parent cm interview small$ASLU num ~
mcs6_parent_cm interview small$MCSID +
mcs6 parent cm interview small$FCNUM00 , FUN=mean, na.rm=TRUE,
na.action="na.omit")
str(mcs6 parent cm interview composite)
colnames(mcs6 parent cm interview composite) <- c('MCSID',</pre>
'FCNUM00', 'ASLU mean')
mcs6 parent cm interview with ASLU <- merge(x =</pre>
mcs6 parent cm interview composite,
```

```
у =
mcs6 parent cm interview,
                                       by.x = c('MCSID',
'FCNUM00'),
                                       by.y = c('MCSID',
'FCNUM00'), all = TRUE)
# Overview of specific cases on composite score for each
Cohort Member
print(mcs6 parent cm interview with ASLU[which(mcs6 parent cm
interview with ASLU$MCSID == 'M10002P' |
mcs6 parent cm interview with ASLU$MCSID == 'M10611J' |
mcs6 parent cm interview with ASLU$MCSID == 'M10106W' |
mcs6_parent_cm_interview with ASLU$MCSID == 'M10063C'),
                                    c('MCSID', 'FPNUM00',
'FELIGOO', 'FCNUMOO', 'FPASLUOO', 'ASLU_num', 'ASLU_mean')])
# Example code M
# -----.
# Merging datasets of different structures to the
mcs longitudinal family file .
-----.
mcs longitudinal family file <- read.spss(file=</pre>
paste(data folder path,
"/mcs6 /mcs longitudinal family file.sav", sep = ''),
to.data.frame = TRUE)
```

```
mcs longitudinal family file$MCSID <-</pre>
trimws(mcs longitudinal family file$MCSID, which = c('both'))
mcs longitudinal family file$All sweeps <- 'longitudinal'
# merge with a parent level dataset .
mcs6 parent derived <- read.spss(file= paste(data_folder_path,</pre>
"/mcs6 /mcs6 parent derived.sav", sep = ''), to.data.frame =
TRUE)
mcs6 parent derived$MCSID <- trimws(mcs6 parent derived$MCSID,</pre>
which = c('both'))
mcs6 parent derived$Sweep 6 <- 'Sweep 6'</pre>
mcs6 parent derived plus longitudinal file <- merge(x =</pre>
mcs6 parent derived,
                                         y =
mcs longitudinal family file,
                                         by.x = c('MCSID'),
                                         by.y = c('MCSID'), all
= TRUE)
table (mcs6 parent derived plus longitudinal file$Sweep 6,
mcs6 parent derived plus longitudinal file$All sweeps,
useNA='ifany')
# merge with a cm level dataset .
mcs6 cm derived <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 cm derived.sav", sep = ''), to.data.frame = TRUE)
mcs6 cm derived$MCSID <- trimws(mcs6 cm derived$MCSID, which =</pre>
c('both'))
```

```
mcs6 cm derived$Sweep 6 <- 'Sweep 6'</pre>
mcs6 cm derived plus longitudinal file <- merge(x =</pre>
mcs6 cm derived,
                                                        у =
mcs longitudinal family file,
                                                        by.x =
c('MCSID'),
                                                        by.y =
c('MCSID'), all = TRUE)
table (mcs6 cm derived plus longitudinal file$Sweep 6,
mcs6 cm derived plus longitudinal file$All sweeps,
useNA='ifany')
# merge with a family level dataset .
mcs6 family derived <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 family derived.sav", sep = ''), to.data.frame =
TRUE)
mcs6 family derived$MCSID <- trimws(mcs6 family derived$MCSID,</pre>
which = c('both'))
mcs6 family derived$Sweep 6 <- 'Sweep 6'</pre>
mcs6 family derived plus longitudinal file \leftarrow merge(x =
mcs6 family derived,
                                                   y =
mcs longitudinal family file,
                                                   by.x =
c('MCSID'),
```

```
by.y =
c('MCSID'), all = TRUE)
table (mcs6 family derived plus longitudinal file$Sweep 6,
mcs6 family derived plus longitudinal file$All sweeps,
useNA='ifany')
# merge with a parent cm level dataset .
mcs6 parent cm interview <- read.spss(file=</pre>
paste (data folder path, "/mcs6 /mcs6 parent cm interview.sav",
sep = ''), to.data.frame = TRUE)
mcs6 parent cm interview$MCSID <-</pre>
trimws(mcs6 parent cm interview$MCSID, which = c('both'))
mcs6 parent cm interview$Sweep 6 <- 'Sweep 6'</pre>
mcs6 parent cm interview plus longitudinal file <- merge(x =</pre>
mcs6 parent cm interview,
                                                       у =
mcs longitudinal family file,
                                                      by.x =
c('MCSID'),
                                                      by.y =
c('MCSID'), all = TRUE)
table (mcs6 parent cm interview plus longitudinal file$Sweep 6,
mcs6 parent cm interview plus longitudinal file$All sweeps,
useNA='ifany')
```

```
# Example code N
```

```
# -----.
# Example I - merging two 1-level datasets that have different
identifiers.
# Please run example code H to get the parent level dataset
below.
str(mcs6 parent plus proxy interview)
mcs6 parent plus proxy interview$GENA num <-</pre>
as.numeric(mcs6 parent plus proxy interview$FPGENA00)
# create a composite score of the general health of the Main &
Partner.
# Mean self-assessed health of Main/Partner respondent(s) in
each family
mcs6 parent plus proxy interview small <-</pre>
mcs6 parent plus proxy interview[c('MCSID', 'GENA num')]
mcs6 parent plus proxy interview composite <-</pre>
aggregate(mcs6 parent plus proxy interview$GENA num ~
mcs6 parent plus proxy interview$MCSID, FUN=mean)
str(mcs6 parent plus proxy interview composite)
colnames(mcs6 parent plus proxy interview composite) <-</pre>
c('MCSID', 'GENA composite')
# connect to the rest of the data
mcs6 parent plus proxy interview with GENA <- merge(x =</pre>
mcs6 parent plus proxy interview,
mcs6 parent plus proxy interview composite,
```

```
by.x = 'MCSID', by.y
= 'MCSID', all = TRUE)
str(mcs6 parent plus proxy interview with GENA)
table (mcs6 parent plus proxy interview with GENA$FELIG00,
useNA='ifany')
mcs6 parent plus proxy interview with GENA$ELIG[mcs6 parent pl
us proxy interview with GENA$FELIG00 %in% c("Main Interview
")] <- 'Main'
mcs6 parent plus proxy interview with GENA$ELIG[mcs6 parent pl
us proxy interview with GENA$FELIG00 %in% c("Partner Interview
")] <- 'Partner'
mcs6 parent plus proxy interview with GENA$ELIG[mcs6 parent pl
us proxy interview with GENA$FELIG00 %in% c("Proxy
Interview")] <- 'Proxy'</pre>
mcs6 parent wide <-</pre>
reshape (mcs6 parent plus proxy interview with GENA,
        timevar = "ELIG",
        idvar = c("MCSID", "GENA_composite"),
        direction = "wide")
names(mcs6 parent wide)
head(mcs6 parent wide)
# merge cm level with the wide restructured parent dataset
(one row per family).
```

```
mcs6 cm interview <- read.spss(file= paste(data_folder_path,</pre>
"/mcs6 /mcs6 cm interview.sav", sep = ''), to.data.frame =
TRUE)
mcs6 cm interview$MCSID <- trimws(mcs6 cm interview$MCSID,</pre>
which = c('both')); str(mcs6 cm interview$MCSID)
\# merge the two datasets on MCSID & CNUM .
mcs6 cm interview$Source cm <- 'CM long one row per child'</pre>
mcs6 parent wide$Source parent <-</pre>
'Parent wide one row per family'
mcs6 cm interview with parent GENA <- merge(x =  
mcs6 cm interview,
                                 y = mcs6 parent wide,
                                 by.x = 'MCSID',
                                 by.y = 'MCSID', all = TRUE)
table (mcs6 cm interview with parent GENA$Source cm,
mcs6 cm interview with parent GENA$Source parent,
useNA='ifany')
# comparison between parents' general health and CM's.
table (mcs6 cm interview with parent GENA$GENA composite,
mcs6 cm interview with parent GENA$FCCGHE00)
```

## # Example code O

# -----.

```
# Example II - merging a 1-level dataset (cm ) with a 2-level
dataset ( parent cm ) resulting into a 1-level structure
( cm ).
# Please check example code L to get the parent cm level
dataset, it is similar.
# we generate a composite measure of ASLU like in the example
code L.
str(mcs6 parent cm interview with ASLU)
mcs6 parent cm interview with ASLU <-
mcs6 parent cm interview with ASLU[c('MCSID', 'FPNUM00',
'FELIG00', 'FCNUM00', 'FPASLU00', 'ASLU mean')]
# we turn the parent cm dataset into wide format (one row
per CM).
mcs6 parent cm interview with ASLU$ELIG[mcs6 parent cm intervi
ew with ASLU$FELIG00 %in% c("Main Interview ")] <- 'Main'
mcs6 parent cm interview with ASLU$ELIG[mcs6 parent cm intervi
ew with ASLU$FELIG00 %in% c("Partner Interview
                                                    ")] <-
'Partner'
mcs6 parent cm interview with ASLU wide <-
reshape (mcs6 parent cm interview with ASLU,
                                                    timevar =
'ELIG',
                                                    idvar =
c('MCSID', 'FCNUM00', 'ASLU mean'),
                                                    direction =
'wide')
names (mcs6 parent cm interview with ASLU wide)
```

```
# merge cm interview level with the wide restructured parent
dataset (one row per family).
mcs6 cm interview <- read.spss(file= paste(data folder path,</pre>
"/mcs6 /mcs6 cm interview.sav", sep = ''), to.data.frame =
TRUE)
mcs6 cm interview$MCSID <- trimws(mcs6 cm interview$MCSID,</pre>
which = c('both')); str(mcs6 cm interview$MCSID)
\# merge the two datasets on MCSID & CNUM .
mcs6 cm interview$Source cm <- 'CM long one row per child'</pre>
mcs6 parent cm interview with ASLU wide$Source parent cm <-
'Parent cm wide one row_per_child'
mcs6 parent cm interview with ASLU wide with cm <- merge(x =
mcs6 cm interview,
                                             у =
mcs6 parent cm interview with ASLU wide,
                                             by.x = c('MCSID',
'FCNUM00'),
                                             by.y = c('MCSID',
'FCNUM00'), all = TRUE)
table (mcs6 parent cm interview with ASLU wide with cm$Source c
m,
mcs6 parent cm interview with ASLU wide with cm$Source parent
```

cm, useNA='ifany')

head(mcs6 parent cm interview with ASLU wide)

# merge the two datasets on MCSID & CNUM .

## SPSS syntax

```
* Setting up folders in SPSS.
file handle mcs5\_folder /name = 'user_folder_path_\_of_mcs5'.
file handle mcs6_folder /name = 'user_folder_path_\_of_mcs6'.
file handle mcs working folder /name =
'user_folder_path_\_of_mcs_work_in_progress'.
* ======== .
SET TNUMBERS BOTH.
SET OVARS BOTH.
SET TVARS BOTH.
* Example code A
* -----.
* Overview of the hhgrid .
GET FILE = 'mcs6 folder/mcs6 hhgrid.sav'.
* Overview of PRES / CREL / multiple Cohort Members per
family.
TEMPORARY.
```

```
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10063C') OR
(MCSID EQ 'M10611J') OR (MCSID EQ 'M10106W') OR (MCSID EQ
'M10063C')).
SUMMARIZE
/TABLES = MCSID FPNUM00 FELIG00 FRESP00 FCNUM00 FHCREL00
FHPRES00
/FORMAT = VALIDLIST NOCASENUM TOTAL
/TITLE = 'Overview of specific cases on key variables'
/MISSING = VARIABLE
/CELLS = COUNT.
* Example code B
* -----.
* Concatenating MCSID & person identifier to get a unique
person identifier .
_____.
GET FILE = 'mcs6 folder/mcs6 hhgrid.sav'.
* a Person ID for each adult in the household (excluding
cohort members) .
STRING PnumID (A9).
```

```
COMPUTE PnumID = concat(rtrim(MCSID), ltrim(string(FPNUM00,
F2)) ).
EXECUTE.
SORT CASES BY PnumID.
* a Person ID for each individual of the household (Cohort
Member or other person) .
STRING CMrow (A3).
IF (FCNUM00 EQ 1) CMrow = '_C1'.
IF (FCNUM00 EQ 2) CMrow = '_C2'.
IF (FCNUM00 EQ 3) CMrow = 'C3'.
STRING PID (A11).
COMPUTE PID = concat(rtrim(MCSID), '_', ltrim(string(FPNUM00,
F2))).
EXECUTE.
IF (FCNUM00 EQ 1 OR 2 OR 3) PID = concat(rtrim(MCSID),
ltrim(CMrow)).
* Example code C
* -----.
* Overview of the _family_derived .
GET FILE = 'mcs6_folder/mcs6_family_derived.sav'.
```

```
FREQUENCIES FDNOCM00.
FREQUENCIES FDRSP000.
* Overview of the _cm_derived .
GET FILE = 'mcs6 folder/mcs6 cm derived.sav'.
FREQUENCIES FCNUM00 .
* Example code D
* -----.
* Overview of parent derived .
* -----
GET FILE = 'mcs6 folder/mcs6 parent derived.sav'.
TEMPORARY.
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10041W') OR
(MCSID EQ 'M23136V') OR (MCSID EQ 'M10106W') OR (MCSID EQ
'M10063C')).
SUMMARIZE
/TABLES = MCSID FPNUM00 FELIG00 FRESP00
/FORMAT = VALIDLIST NOCASENUM TOTAL
/TITLE = 'Overview of specific cases on key variables'
/MISSING = VARIABLE
```

```
/CELLS = COUNT.
```

```
* Example code E
* -----.
* merge parent structure datasets from different sweeps.
-----.
* ~ ~ ~ ~ ~ merge the two datasets on MCSID & ELIG ~ ~ ~ ~
* prepare the mcs6 parent dataset.
GET FILE = 'mcs6 folder/mcs6 parent interview.sav'.
FREQUENCIES FPNUM00 FELIG00.
COMPUTE ELIG = FELIG00.
EXECUTE.
SORT CASES BY MCSID ELIG (A).
SAVE OUTFILE =
'mcs working folder/mcs6 parent interview cross sweep merging.
sav'.
* prepare the mcs5 parent dataset.
GET FILE = 'mcs5 folder/mcs5 parent interview.sav'.
FREQUENCIES EPNUM00 EELIG00.
```

```
COMPUTE ELIG = EELIG00.
EXECUTE.
SORT CASES BY MCSID ELIG (A).
SAVE OUTFILE =
'mcs working folder/mcs5 parent interview cross sweep merging.
sav'.
* merge .
GET FILE =
'mcs working folder/mcs6 parent interview_cross_sweep_merging.
sav'.
MATCH FILES /FILE=*
  /IN source mcs6
/FILE='mcs working folder/mcs5 parent interview cross sweep me
rging.sav'
  /IN source mcs5
  /BY MCSID ELIG.
EXECUTE.
* Outcome perusal: Main and Partner respondents (ELIG) in both
sweeps.
CROSSTABS source mcs6 BY source mcs5 .
FREQUENCIES ELIG.
CROSSTABS FELIGOO BY EELIGOO .
* Let us see how many of the merged Main & Partner respondents
(ELIG) have
```

```
^{\star} the same PNUM, therefore they are they same person .
```

- \* Values in the diagonal of the crosstabulation are respondents who are the
- \* same in both sweeps.

TEMPORARY.

SELECT IF ELIG = 1.

CROSSTABS FPNUM00 BY EPNUM00 .

IF (FPNUM00 EQ EPNUM00) SAME RESPONDENT = 1.

IF SYSMIS (SAME RESPONDENT) SAME RESPONDENT = 0.

VARIABLE LABELS SAME\_RESPONDENT 'Is the Main/Partner respondent the same btw S5 & S6?'.

VALUE LABELS SAME\_RESPONDENT 1 'Same' 0 'Different or missing data'.

FREQUENCIES SAME RESPONDENT.

CROSSTABS ELIG BY SAME RESPONDENT.

SAVE OUTFILE =

'mcs working folder/mcs5 mcs6 parent interview by ELIG.sav'.

\* prepare the mcs6 parent dataset.

GET FILE = 'mcs6 folder/mcs6 parent\_interview.sav'.

COMPUTE PNUM = FPNUM00.

```
SORT CASES BY MCSID PNUM (A).
SAVE OUTFILE =
'mcs working folder/mcs6 parent interview cross sweep merging.
* prepare the mcs5 parent_dataset.
GET FILE = 'mcs5 folder/mcs5 parent interview.sav'.
COMPUTE PNUM = EPNUM00.
EXECUTE.
SORT CASES BY MCSID PNUM (A).
SAVE OUTFILE =
'mcs working folder/mcs5 parent interview cross sweep merging.
sav'.
* merge.
GET FILE =
'mcs working folder/mcs6 parent interview cross sweep merging.
sav'.
MATCH FILES /FILE=*
  /IN source mcs6
/FILE='mcs working folder/mcs5 parent interview cross sweep me
rging.sav'
  /IN source mcs5
  /BY MCSID PNUM.
EXECUTE.
* Outcome perusal: parents/carers (PNUM) in both sweeps.
```

CROSSTABS source mcs6 BY source mcs5 .

- \* Let us see how many of the merged parents/carers respondents (PNUM) have
- \* the same role in the interview (ELIG: Main or Partner) .

IF (FELIG00 EQ EELIG00) SAME ELIGIBILITY = 1.

IF SYSMIS (SAME ELIGIBILITY) SAME ELIGIBILITY = 0.

VARIABLE LABELS SAME\_ELIGIBILITY 'Is the role at the interview (ELIG) the same btw S5 & S6?'.

VALUE LABELS SAME\_ELIGIBILITY 1 'Same' 0 'Different or missing data'.

FREQUENCIES SAME ELIGIBILITY.

CROSSTABS PNUM BY SAME ELIGIBILITY.

SAVE OUTFILE =

'mcs working folder/mcs5 mcs6 parent interview by PNUM.sav'.

## \* Example code F

\* -----.

\* Create a composite variable per family in the \_parent\_ structure file .

\* -----

-----.

- $\star$  this syntax creates a mean of Main and Partner responses on GENA variable .
- \* You can use other functions instead of MEAN, like SD, MIN, MAX, etc.

```
GET FILE = 'mcs6 folder/mcs6 parent interview.sav'.
AGGREGATE
  outfile=*
  overwrite=yes
  mode=addvariables
  /break= MCSID
  /composite GENA = MEAN(FPGENA00)
  /groupsize = N.
VARIABLE LABELS composite GENA 'Mean score of GENA of
Main/Partner respondent(s) per family'.
VARIABLE LABELS groupsize 'Number of respondents (Main only,
Partner only, or Main&Partner) providing information in GENA'.
* Let's take a look at the result .
TEMPORARY.
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10611J') OR
(MCSID EQ 'M10106W') OR (MCSID EQ 'M10063C')).
SUMMARIZE
/TABLES = MCSID
FPNUM00
FELIG00
FRESP00
FPGENA00
composite GENA
```

```
groupsize
/FORMAT = VALIDLIST NOCASENUM TOTAL
/TITLE = 'Overview of specific cases on composite score for
each family (GENA = general health)'
/MISSING = VARIABLE
/CELLS = COUNT.
* this syntax selects the higher NVQ of Main and Partner
respondents.
* You can use other functions instead of MEAN, like SD, MIN,
MAX, etc.
GET FILE = 'mcs6 folder/mcs6 parent derived.sav'.
FREQUENCIES FDNVQ00 .
IF (FDNVQ00 GE 1 AND FDNVQ00 LE 5) NVQ = FDNVQ00.
EXECUTE.
CROSSTABS FDNVQ00 BY NVQ.
AGGREGATE
  outfile=*
  overwrite=yes
  mode=addvariables
  /break= MCSID
  /composite NVQ = MAX(NVQ)
  /groupsize = N.
```

VARIABLE LABELS composite\_NVQ 'Highest NVQ of Main/Partner respondent(s) in each family'.

VARIABLE LABELS groupsize 'Number of respondents (Main only, Partner only, or Main&Partner) with information on NVQ'.

\* Let's take a look at the result .

TEMPORARY.

SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10611J') OR (MCSID EQ 'M10451L') OR (MCSID EQ 'M10106W') OR (MCSID EQ 'M10063C')).

SUMMARIZE

/TABLES = MCSID

FPNUM00

FELIG00

FRESP00

FDNVQ00

composite NVQ

groupsize

/FORMAT = VALIDLIST NOCASENUM TOTAL

/TITLE = 'Overview of specific cases on composite score for each family (highest NVQ)'

/MISSING = VARIABLE

/CELLS = COUNT.

* Example code G
*
* Overview of _proxy_partner_interview .
*
<pre>GET FILE = 'mcs6_folder/mcs6_proxy_partner_interview.sav'.</pre>
FREQUENCIES FXCREL00 .
CROSSTABS FXCREL00 BY FXPSEX00 .
FREQUENCIES FXPXRE00 FXPXIN00 .
* Example code H  *
* Combining proxy_partner_interview with parent_interview .  *
^
* we keep only rows where the Main agreed to provide information about the non-available Partner.
SELECT IF FXPXIN00 EQ 1 .
FREQUENCIES FXPXGE00.
* we rename the variable to the variable name that is used in the parent interview dataset.

```
RENAME VARIABLES FXPXGE00 = FPGENA00 .
SORT CASES BY MCSID (A).
SAVE OUTFILE =
'mcs_working_folder/mcs6_proxy_partner_interview_for_connectio
n to parent.sav'
/KEEP
MCSID
FELIG00
FRESP00
FXCREL00
FPGENA00 .
* open the parent interview dataset and keep only the
variables needed .
GET FILE = 'mcs6 folder/mcs6 parent interview.sav'.
FREQUENCIES FPGENA00.
SAVE OUTFILE =
'mcs_working_folder/mcs6_parent_interview_for_connection_to_pr
oxy.sav'
/KEEP
MCSID
FELIG00
FRESP00
```

```
FPCREL00
FPGENA00 .
* open the reduced parent interview dataset.
GET FILE =
'mcs working folder/mcs6 parent interview for connection to pr
oxy.sav'.
* we add cases/rows to the dataset.
ADD FILES /FILE=*
/FILE='mcs working folder\mcs6 proxy partner interview for con
nection to parent.sav'.
EXECUTE.
FREQUENCIES FPGENA00.
CROSSTABS FPGENA00 BY FELIG00.
SAVE OUTFILE =
'mcs working folder/mcs6 parent plus proxy interview.sav'.
* Example code I
* -----.
* merge cm structure datasets from different sweeps.
-----.
* prepare the mcs6 cm dataset.
```

```
GET FILE = 'mcs6 folder/mcs6 cm interview.sav'.
FREQUENCIES FCNUM00 .
COMPUTE CNUM = FCNUM00.
EXECUTE.
SORT CASES BY MCSID CNUM (A).
SAVE OUTFILE =
'mcs working folder/mcs6 cm interview cross sweep merging.sav'
* prepare the mcs5 cm dataset.
GET FILE = 'mcs5 folder/mcs5 cm interview.sav'.
FREQUENCIES ECNUM00.
COMPUTE CNUM = ECNUM00.
EXECUTE.
SORT CASES BY MCSID CNUM (A).
SAVE OUTFILE =
'mcs working folder/mcs5 cm interview cross sweep merging.sav'
^{\star} merge the two datasets on MCSID & CNUM .
GET FILE =
'mcs working folder/mcs6 cm interview cross sweep merging.sav'
MATCH FILES /FILE=*
  /IN source mcs6
```

```
/FILE='mcs working folder/mcs5 cm interview cross sweep mergin
g.sav'
 /IN source mcs5
 /BY MCSID CNUM.
EXECUTE.
* Outcome perusal: cohort members in both datasets.
CROSSTABS source mcs6 BY source mcs5 .
FREQUENCIES CNUM.
CROSSTABS FCNUM00 BY ECNUM00 .
SAVE OUTFILE =
'mcs working folder/mcs5 mcs6 cm interview.sav'.
* Example code J
* -----.
* Overview of parent cm interview .
* -------
* Parent's interview about the CM(s) of the household.
GET FILE = 'mcs6 folder/mcs6 parent cm interview.sav'.
FREQUENCIES FCNUM00 .
```

TEMPORARY.

```
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10611J') OR
(MCSID EQ 'M10106W') OR (MCSID EQ 'M10063C')).
SUMMARIZE
/TABLES = MCSID
FCNUM00
FELIG00
FCNUM00
FCCSEX00
FCCAGE00
FPSDPF00
FPASLU00
/FORMAT = VALIDLIST NOCASENUM TOTAL
/TITLE = 'Overview of specific cases on key variables'
/MISSING = VARIABLE
/CELLS = COUNT.
* Example code K
* -----.
* Merge parent cm level datasets between sweeps.
* Create row identifier to connect for parent cm datasets.
```

```
* ROWid = (ELIG or PNUM) + (CNUM Child 1/2/3).
* ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ELIG + CNUM = ROWid ~ ~ ~ ~ ~ ~ ~ ~ ~ ~
~ ~ ~ ~ .
* prepare mcs6 parent cm interview .
GET FILE = 'mcs6 folder/mcs6 parent cm interview.sav'.
STRING ELIGROW (A1).
IF (FELIG00 EQ 1) ELIGrow = "M".
IF (FELIG00 EQ 2) ELIGrow = "P".
COMPUTE CNUM = FCNUM00.
EXECUTE.
CROSSTABS CNUM BY ELIGROW.
STRING ROWid (A2).
COMPUTE ROWid = concat (rtrim(ELIGrow), ltrim(string(CNUM,
F1))).
EXECUTE.
* Check that the ROWid matches the crosstabulation of ELIG &
CNUM .
FREQUENCIES ROWid.
SORT CASES BY MCSID ROWid (A).
SAVE OUTFILE =
'mcs working folder/mcs6 parent cm interview cross sweep mergi
ng.sav' .
```

```
* prepare mcs5 parent cm interview .
GET FILE = 'mcs5 folder/mcs5 parent cm interview.sav'.
STRING ELIGROW (A1).
IF (EELIG00 EQ 1) ELIGrow = "M".
IF (EELIG00 EQ 2) ELIGrow = "P".
COMPUTE CNUM = ECNUM00.
EXECUTE.
CROSSTABS CNUM BY ELIGROW.
STRING ROWid (A2).
COMPUTE ROWid = concat (rtrim(ELIGrow), ltrim(string(CNUM,
F1))).
EXECUTE.
* Check that the ROWid matches the crosstabulation of ELIG &
CNUM .
FREQUENCIES ROWid.
SORT CASES BY MCSID ROWid (A).
SAVE OUTFILE =
'mcs working folder/mcs5 parent cm interview cross sweep mergi
ng.sav' .
* Merge .
GET FILE =
'mcs working folder/mcs6 parent cm interview cross sweep mergi
ng.sav'.
MATCH FILES /FILE=*
```

```
/IN source mcs6
/FILE='mcs working folder/mcs5 parent cm interview cross sweep
merging.sav'
  /IN source mcs5
  /BY MCSID ROWid.
EXECUTE.
* Outcome perusal: Parents (Main/Partner providing information
about
* each of the cohort members) are in both sweeps.
CROSSTABS source mcs6 BY source mcs5 .
SAVE OUTFILE =
'mcs working folder/mcs5 mcs6 parent cm interview by ELIG.sav'
* ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ PNUM + CNUM = ROWid ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~
~ ~ ~ ~ .
* prepare mcs6 parent cm interview .
GET FILE = 'mcs6 folder/mcs6 parent cm interview.sav'.
STRING CMrow (A3).
IF (FCNUM00 EQ 1) CMrow = ' C1'.
IF (FCNUM00 EQ 2) CMrow = 'C2'.
IF (FCNUM00 EQ 3) CMrow = 'C3'.
```

```
EXECUTE.
STRING Prow (A3).
COMPUTE Prow = concat (rtrim('P'), ltrim(string(FPNUM00,
F2))).
EXECUTE.
STRING ROWid(A6).
COMPUTE ROWid = concat (rtrim(Prow), ltrim(CMrow)).
EXECUTE.
* Let's check that the totals match.
CROSSTABS FPNUM00 BY FCNUM00.
FREQUENCIES ROWid.
SORT CASES BY MCSID ROWid (A).
SAVE OUTFILE =
'mcs working folder/mcs6 parent cm interview cross sweep mergi
ng.sav'.
* prepare mcs5 parent cm interview .
GET FILE = 'mcs5 folder/mcs5 parent cm interview.sav'.
STRING CMrow (A3).
IF (ECNUM00 EQ 1) CMrow = 'C1'.
IF (ECNUM00 EQ 2) CMrow = 'C2'.
IF (ECNUM00 EQ 3) CMrow = 'C3'.
EXECUTE.
```

```
STRING Prow (A3).
COMPUTE Prow = concat (rtrim('P'), ltrim(string(EPNUM00,
F2))).
EXECUTE.
STRING ROWid (A6).
COMPUTE ROWid = concat (rtrim(Prow), ltrim(CMrow)).
EXECUTE.
* Let's check that the totals match.
CROSSTABS EPNUM00 BY ECNUM00.
FREQUENCIES ROWid.
SORT CASES BY MCSID ROWid (A).
SAVE OUTFILE =
'mcs working folder/mcs5 parent cm interview cross sweep mergi
ng.sav'.
* Merge .
GET FILE =
'mcs working folder/mcs6 parent cm interview cross sweep mergi
ng.sav'.
MATCH FILES /FILE=*
  /IN source mcs6
/FILE='mcs_working_folder/mcs5_parent_cm interview cross sweep
merging.sav'
  /IN source mcs5
```

/BY MCSID ROWid. EXECUTE. CROSSTABS source mcs6 BY source mcs5 . \* Outcome perusal: Only \_\_\_\_ Individuals (PNUM) provide information \* about the cohort member(s) in both sweeps (either as Main or Partner respondent). SAVE OUTFILE = 'mcs\_working\_folder/mcs5 mcs6 parent cm interview by PNUM.sav' \* Example code L \* -----. \* Create a composite variable per child in parent cm dataset. \* this syntax creates a mean of Main and Partner responses on ASLU variable . \* You can use other functions instead of MEAN, like SD, MIN, MAX, etc. GET FILE = 'mcs6 folder/mcs6\_parent\_cm\_interview.sav'. **AGGREGATE** outfile=\* overwrite=yes

```
mode=addvariables
  /break= MCSID FCNUM00
  /composite ASLU = MEAN(FPASLU00)
  /groupsize = N.
VARIABLE LABELS composite_ASLU 'Mean score of ASLU assessment
by Main/Partner respondent(s) of the cohort member'.
VARIABLE LABELS groupsize 'Number of respondents (Main only,
Partner only, or Main&Partner) providing information in ASLU'.
* Let's take a look at the result .
TEMPORARY.
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10611J') OR
(MCSID EQ 'M10106W') OR (MCSID EQ 'M10063C')).
SUMMARIZE
/TABLES = MCSID
FPNUM00
FELIG00
FRESP00
FCNUM00
FPASLU00
composite ASLU
groupsize
/FORMAT = VALIDLIST NOCASENUM TOTAL
```

```
/TITLE = 'Overview of specific cases on composite score for
each Cohort Member (groupsize = number of parents ELIG
providing information)'
/MISSING = VARIABLE
/CELLS = COUNT.
* Example code M
* -----.
* Merging datasets of different structures to the
mcs longitudinal family file .
* -----
-----.
* we do not need to sort files as they are all sorted by
MCSID,
* however, if you have worked on the file before make sure you
* sort it before this step.
* merge with a _parent_ level dataset .
GET FILE = "mcs6 folder/mcs6 parent derived.sav".
MATCH FILES /FILE = *
/IN source parent
/TABLE = "mcs6 folder/mcs longitudinal family file.sav"
/IN = source longitudinal file
```

```
/BY MCSID .
EXECUTE.
CROSSTABS source_parent by source_longitudinal_file.
* merge with a cm level dataset .
GET FILE = "mcs6 folder/mcs6 cm derived.sav".
MATCH FILES /FILE = *
/IN source cm
/TABLE = "mcs6 folder/mcs longitudinal family file.sav"
/IN = source longitudinal file
/BY MCSID .
EXECUTE.
CROSSTABS source cm by source longitudinal file.
* merge with a family level dataset .
GET FILE = "mcs6 folder/mcs6 family derived.sav".
MATCH FILES /FILE = *
/IN source familyDV
/TABLE = "mcs6 folder/mcs longitudinal family file.sav"
/IN = source longitudinal file
/BY MCSID .
EXECUTE.
```

```
* merge with a _{\rm parent\_cm\_} level dataset .
GET FILE = "mcs6 folder/mcs6 parent cm interview.sav".
MATCH FILES /FILE = *
/IN source parent cm
/TABLE = "mcs6 folder/mcs longitudinal family file.sav"
/IN = source longitudinal file
/BY MCSID .
EXECUTE.
CROSSTABS source parent cm by source longitudinal file.
* Example code N
* -----.
* Example I - merging two 1-level datasets that have different
identifiers.
* Please run example code H to get the parent level dataset
below.
GET FILE =
"mcs working folder/mcs6 parent plus proxy interview.sav".
SORT CASES BY MCSID (A).
```

CROSSTABS source familyDV by source longitudinal file.

```
* create a composite score of the general health of the Main \&
Partner.
AGGREGATE
  outfile=*
  overwrite=yes
  mode=addvariables
  /break= MCSID
  /composite HEALTH = MEAN(FPGENA00)
  /groupsize = N.
VARIABLE LABELS composite HEALTH 'Mean self-assessed health of
Main/Partner respondent(s) in each family'.
VARIABLE LABELS groupsize 'Number of respondents (Main only,
Partner only, or Main&Partner) with information on GENA'.
* Let's take a look at the result .
TEMPORARY.
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10611J') OR
(MCSID EQ 'M10451L') OR (MCSID EQ 'M10106W') OR (MCSID EQ
'M10063C')).
SUMMARIZE
/TABLES = MCSID
FELIG00
FRESP00
FPGENA00
```

```
composite HEALTH
groupsize
/FORMAT = VALIDLIST NOCASENUM TOTAL
/TITLE = 'Overview of specific cases on composite score for
each family (mean Health)'
/MISSING = VARIABLE
/CELLS = COUNT.
* at this stage we can just keep the one row for each family,
however, by
* restructuring in wide format (one row per family) we keep
all the information
* that we may need for crosschecking.
* restructure the datasets into wide format (one row per
family).
FREQUENCIES FELIG00 .
* We create a Main / Partner only ELIG.
IF (FELIG00 EQ 1) ELIG = 1.
IF (FELIG00 EQ 2 OR FELIG00 EQ 3) ELIG = 2.
EXECUTE.
VARIABLE LABELS ELIG 'Eligibility Partner collapsed'.
VALUE LABELS ELIG 1 'Main' 2 'Partner / Proxy'.
```

MISSING VALUES ALL ().

```
CROSSTABS FELIGOO BY ELIG .
* resctructure using ELIG.
CASESTOVARS
/ID = MCSID
/INDEX = ELIG.
* Let's look at the new dataset.
TEMPORARY.
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10611J') OR
(MCSID EQ 'M10451L') OR (MCSID EQ 'M10106W') OR (MCSID EQ
'M10063C')).
LIST.
* the variables containing the information of the Main have
been suffixed with 1 and of the Partner with 2.
SORT CASES BY MCSID (A).
SAVE OUTFILE =
"mcs working folder/mcs6 parent plus proxy interview wide.sav"
* merge cm level with the wide restructured parent dataset
(one row per family).
GET FILE = "mcs6 folder/mcs6 cm interview.sav".
MATCH FILES /FILE = *
/IN source cm
```

```
/TABLE =
"mcs_working_folder/mcs6_parent_plus_proxy interview wide.sav"
/IN = source parent wide
/BY MCSID .
EXECUTE.
CROSSTABS source cm by source parent wide.
* for some families there is no parent interview but the CM
has participated.
* comparison between parents' general health and CM's.
CROSSTABS composite HEALTH by FCCGHE00 .
* Example code O
* -----.
* Example II - merging a 1-level dataset (cm ) with a 2-level
dataset ( parent cm ) resulting into a 1-level structure
( cm ).
* Please check example code L to get the parent cm level
dataset, it is similar.
GET FILE = 'mcs6 folder/mcs6 parent cm interview.sav'.
* we generate a composite measure of ASLU like in the example
```

AGGREGATE

code L.

```
outfile=*
  overwrite=yes
  mode=addvariables
  /break= MCSID FCNUM00
  /composite ASLU = MEAN(FPASLU00)
  /groupsize = N.
VARIABLE LABELS composite ASLU 'Mean score of ASLU assessment
by Main/Partner respondent(s) of the cohort member'.
VARIABLE LABELS groupsize 'Number of respondents (Main only,
Partner only, or Main&Partner) providing information in ASLU'.
* Let's take a look at the result .
TEMPORARY.
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10611J') OR
(MCSID EQ 'M10106W') OR (MCSID EQ 'M10063C')).
SUMMARIZE
/TABLES = MCSID
FPNUM00
FELIG00
FRESP00
FCNUM00
FPASLU00
composite ASLU
groupsize
```

```
/FORMAT = VALIDLIST NOCASENUM TOTAL
/TITLE = 'Overview of specific cases on composite score for
each Cohort Member (groupsize = number of parents ELIG
providing information)'
/MISSING = VARIABLE
/CELLS = COUNT.
SORT CASES BY MCSID (A).
SAVE OUTFILE =
'mcs_working_folder/mcs6_parent_cm_interview_reduced.sav'
/KEEP
MCSID
FPNUM00
FELIG00
FRESP00
FCNUM00
FCCREL00
FPASLU00
composite ASLU
groupsize.
* we turn the parent cm dataset into wide format (one row
```

per CM).

```
GET FILE =
'mcs working folder/mcs6 parent cm interview reduced.sav'.
FREQUENCIES FELIG00 .
* We create a Main / Partner only ELIG.
IF (FELIG00 EQ 1) ELIG = 1.
IF (FELIG00 EQ 2 OR FELIG00 EQ 3) ELIG = 2.
EXECUTE.
VARIABLE LABELS ELIG 'Eligibility Partner collapsed'.
VALUE LABELS ELIG 1 'Main' 2 'Partner / Proxy'.
MISSING VALUES ALL ().
CROSSTABS FELIGOO BY ELIG .
* resctructure using ELIG.
CASESTOVARS
/ID = MCSID FCNUM00
/INDEX = ELIG.
* Let's look at the new dataset.
TEMPORARY.
SELECT IF ((MCSID EQ 'M10002P') OR (MCSID EQ 'M10611J') OR
(MCSID EQ 'M10451L') OR (MCSID EQ 'M10106W') OR (MCSID EQ
'M10063C')).
```

LIST.

\* the variables containing the information of the Main have been suffixed with 1 and of the Partner with 2.

\* there is one row per child, so the dataset is on the cm level. SORT CASES BY MCSID (A). SAVE OUTFILE = "mcs working folder/mcs6 parent cm interview reduced wide on c m level.sav". \* merge cm interview level with the wide restructured parent dataset (one row per family). GET FILE = "mcs6 folder/mcs6 cm interview.sav". MATCH FILES /FILE = \* /IN source cm /TABLE = "mcs working folder/mcs6 parent cm interview reduced wide on c m level.sav" /IN = source parent cm wide /BY MCSID FCNUM00 . EXECUTE. CROSSTABS source cm by source parent cm wide. \* for some families there is no parent interview but the CM has participated.

go to the University and CM's perception on the same topic.

CROSSTABS FCSTYU00 by composite\_ASLU .

\* comparison between parents' perception of CM's likelihood to

# \* Example code P \* -----. \* Example III - merging a 2-level dataset (\_parent\_cm\_) with a 1-level dataset ( cm ) resulting into a 2-level dataset ( cm ). GET FILE = 'mcs6 folder/mcs6 parent cm interview.sav'. MATCH FILES /FILE = \* /IN source cm /TABLE = "mcs6 folder/mcs6 cm interview.sav" /IN = source parent cm /BY MCSID FCNUM00 . EXECUTE. CROSSTABS source\_cm by source\_parent\_cm. CROSSTABS FCTRSTOA by FPSCHCOO.

# STATA syntax

### \* Setting up folders in STATA

```
global mcs5_folder "\user_folder_path\mcs5_"
global mcs6_folder "\user_folder_path\mcs6_"
global mcs_working_folder
"\user folder path\mcs working folder"
```

### \* Example code A

# \* Example code B

```
* ------
* Concatenating MCSID & person identifier to get a unique
person identifier .
* -------
use "$mcs6_folder/mcs6_hhgrid.dta", clear

* a Person ID for each adult in the household (excluding cohort members) .
gen PnumID = MCSID + string(FPNUM00) if (FPNUM00 > 0)

* a Person ID for each individual of the household (Cohort Member or other person) .
gen CMrow = "_C1" if FCNUM00 == 1
```

```
replace CMrow = "_C2" if FCNUM00 == 2
replace CMrow = " C3" if FCNUM00 == 3
gen PID = MCSID + " " + string(FPNUM00) if (FPNUM00 > 0)
replace PID = MCSID + CMrow if (FCNUM00 > 0)
* Example code C
* -----.
* Overview of the family derived .
use "$mcs6 folder/mcs6 family derived.dta", clear
tab FDNOCM00
tab FDRSP000
* Overview of the cm derived .
use "$mcs6 folder/mcs6 cm derived.dta", clear
tab FCNUM00
* Example code D
* -----
* Overview of _parent_derived .
* -----.
use "$mcs6 folder/mcs6 parent derived.dta", clear
gen example families = 1 if MCSID == "M10002P" | MCSID ==
"M10041W" | MCSID == "M23136V" | MCSID == "M10106W" | MCSID ==
"M10063C".
tab example families
ds MCSID FPNUM00 FELIG00 FRESP00
return list
local varlist = r(varlist)
list `varlist' if example families == 1, divider sep(4)
* Example code E
* -----.
* merge _parent_ structure datasets from different sweeps.
* -------
______.
* \sim \sim \sim \sim merge the two datasets on MCSID & ELIG \sim \sim \sim
* prepare the mcs6 parent dataset.
use "$mcs6 folder/mcs6 parent interview.dta", clear
tab FPNUM00
tab FELIG00
gen ELIG = FELIGOO
```

```
save
"$mcs working folder/mcs6 parent interview cross sweep merging
.dta", replace
* prepare the mcs5 parent dataset.
use "$mcs5 folder/mcs5 parent interview.dta", clear
tab EPNUM00
tab EELIG00
gen ELIG = EELIG00
"$mcs working folder/mcs5 parent interview cross sweep merging
.dta", replace
* merge .
"$mcs working folder/mcs6 parent interview cross sweep merging
.dta", clear
merge 1:1 MCSID ELIG using
"$mcs working folder/mcs5 parent interview cross sweep merging
* Outcome perusal: Main and Partner respondents (ELIG) in both
sweeps.
tab ELIG
tab FELIG00 EELIG00
* Let us see how many of the merged Main & Partner respondents
* the same PNUM, therefore they are they same person .
* Values in the diagonal of the crosstabulation are
respondents who are the
* same in both sweeps.
tab FPNUM0 EPNUM00 if (ELIG == 1)
gen same respondent = 1 if (FPNUM00 == EPNUM00)
replace same respondent = 0 if (same respondent != 1)
label variable same respondent "Is the Main/Partner respondent
the same btw S5 & S6?"
label define same respondent 1 "Same" 0 "Different or missing
data"
tab same respondent
tab ELIG same respondent
"$mcs working folder/mcs5 mcs6 parent interview by ELIG.dta",
replace
* \sim \sim \sim \sim merge the two datasets on MCSID & PNUM \sim \sim \sim
* prepare the mcs6 parent dataset.
use "$mcs6 folder/mcs6 parent interview.dta", clear
gen PNUM = FPNUM00
```

```
save
"$mcs working folder/mcs6 parent interview cross sweep merging
.dta", replace
* prepare the mcs5 parent dataset.
use "$mcs5 folder/mcs5 parent interview.dta", clear
gen PNUM = EPNUM00
save
"$mcs working folder/mcs5 parent interview cross sweep merging
.dta", replace
* merge.
use
"$mcs working folder/mcs6 parent interview cross sweep merging
.dta", clear
merge m:1 MCSID PNUM using
"$mcs working folder/mcs5 parent interview cross sweep merging
.dta"
* Outcome perusal: 17.211 parents/carers (PNUM) are in both
* Let us see how many of the merged parents/carers respondents
(PNUM) have
* the same role in the interview (ELIG: Main or Partner) .
gen same eligibility = 1 if (FELIG00 == EELIG00)
replace same eligibility = 0 if (same eligibility != 1)
label variable same eligibility "Is the role at the interview
(ELIG) the same btw S5 & S6?"
label define same eligibility 1 "Same" 0 "Different or missing
data"
tab same eligibility
tab PNUM same eligibility
"$mcs working folder/mcs5 mcs6 parent interview by PNUM.dta",
replace
* Example code F
* -----.
* Create a composite variable per family in the parent
structure file .
* ------
    -----
* this syntax creates a mean of Main and Partner responses on
GENA variable .
* You can use other functions instead of mean, like
use "$mcs6 folder/mcs6 parent interview.dta", clear
```

egen composite\_GENA = mean(FPGENA00), by (MCSID)
egen groupsize = count(FPGENA00), by (MCSID)

Main/Partner respondent(s) per family"

label variable composite\_GENA "Mean score of GENA of

```
label variable groupsize "Number of respondents (Main only,
Partner only, or Main&Partner) providing information in GENA"
* Let's take a look at the result .
gen example families = 1 if MCSID == "M10002P" | MCSID ==
"M10611J" | MCSID == "M10106W" | MCSID == "M10063C".
tab example families
* run this code with/without nolabel to hide/see the value
labels
ds MCSID FPNUM00 FELIG00 FRESP00 FPGENA00 composite GENA
groupsize
return list
local varlist = r(varlist)
list `varlist' if example families == 1, divider sep(4)
nolabel
* this syntax selects the higher NVQ of Main and Partner
respondents.
* You can use other functions instead of MEAN, like sd, min,
max etc.
use "$mcs6 folder/mcs6 parent derived.dta", clear
tab FDNVQ00, nolabel
gen NVQ = FDNVQ00
replace NVQ = . if (NVQ < 1)
replace NVQ = . if (NVQ > 5)
egen composite NVQ = max(NVQ), by (MCSID)
egen groupsize = count(FPNUM00), by (MCSID)
label variable composite NVQ "Highest NVQ of Main/Partner
respondent(s) in each family"
label variable groupsize "Number of respondents (Main only,
Partner only, or Main&Partner) with information on NVQ"
* Let's take a look at the result .
gen example families = 1 if MCSID == "M10002P" | MCSID ==
"M10611J" | MCSID == "M10451L" | MCSID == "M10106W" | MCSID ==
"M10063C".
tab example families
* run this code with/without nolabel to hide/see the value
labels
ds MCSID FPNUM00 FELIG00 FRESP00 FDNVQ00 NVQ composite NVQ
groupsize
return list
local varlist = r(varlist)
list `varlist' if example families == 1, divider sep(4)
nolabel
```

# \* Example code G

```
use "$mcs6 folder/mcs6 proxy partner interview.dta", clear
tab FXCREL00
tab FXCREL00 FXPSEX00
tab FXPXRE00 FXPXIN00
* Example code H
* -----.
* Combining proxy partner interview with parent interview .
* -----
_______
* we keep only rows where the Main agreed to provide
information about the non-available Partner.
keep if FXPXIN00 == 1
tab FXPXGE00
* we rename the variable to the variable name that is used in
the parent interview dataset.
rename FXPXGE00 FPGENA00
keep MCSID FELIG00 FRESP00 FXCREL00 FPGENA00
save
"$mcs working folder/mcs6 proxy partner interview for connecti
on to parent.dta", replace
* open the parent interview dataset and keep only the
variables needed .
use "$mcs6 folder/mcs6 parent interview.dta", clear
tab FPGENA00
keep MCSID FELIG00 FRESP00 FPCREL00 FPGENA00
"$mcs working folder/mcs6 parent interview for connection to p
roxy.dta", replace
* open the reduced parent interview dataset.
"$mcs working folder/mcs6 parent interview for connection to p
roxy.dta", clear
* we add cases/rows to the dataset.
append using
"$mcs working folder\mcs6 proxy partner interview for connecti
on to parent.dta"
tab FPGENA00
tab FPGENA00 FELIG00
"$mcs working folder/mcs6 parent plus proxy interview.dta",
replace
* Example code I
```

\* -----

```
* merge cm structure datasets from different sweeps.
* -----
_____.
* prepare the mcs6 cm dataset.
use "$mcs6 folder/mcs6 cm interview.dta", clear
tab FCNUM00
gen CNUM = FCNUM00
"$mcs working folder/mcs6 cm interview cross sweep merging.dta
", replace
* prepare the mcs5 cm dataset.
use "$mcs5 folder/mcs5 cm interview.dta", clear
tab ECNUM00
gen CNUM = ECNUM00
"$mcs working folder/mcs5 cm interview cross sweep merging.dta
", replace
* merge the two datasets on MCSID & CNUM .
"$mcs working folder/mcs6 cm interview cross sweep merging.dta
", clear
merge 1:1 MCSID CNUM using
"$mcs working folder/mcs5 cm interview cross sweep merging.dta
* Outcome perusal: cohort members in both datasets.
tab CNUM
tab FCNUM00 ECNUM00
save "$mcs working folder/mcs5 mcs6 cm interview.dta", replace
* Example code J
* _____
* Overview of parent cm interview .
* ______
* Parent's interview about the CM(s) of the household.
use "$mcs6 folder/mcs6 parent cm interview.dta", clear
tab FCNUM00
gen example families = 1 if MCSID == "M10002P" | MCSID ==
"M10611J" | MCSID == "M10106W" | MCSID == "M10063C"
tab example families
* run this code with/without nolabel to hide/see the value
ds MCSID FPNUM00 FELIG00 FCNUM00 FCCSEX00 FCCAGE00 FPSDPF00
FPASLU00
return list
local varlist = r(varlist)
```

```
list `varlist' if example_families == 1, divider sep(4)
nolabel
```

## \* Example code K

```
* -----
* Merge parent cm level datasets between sweeps.
* -----
* Create row identifier to connect for parent cm datasets.
* ROWid = (ELIG or PNUM) + (CNUM Child 1/2/3).
* ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ELIG + CNUM = ROWid ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~
~ ~ ~ ~ .
* prepare mcs6 parent cm interview .
use "$mcs6 folder/mcs6 parent cm interview.dta", clear
gen ELIGrow = "M" if (FELIG00 == 1)
replace ELIGrow = "P" if (FELIG00 == 2)
gen CNUM = FCNUM00
tab CNUM ELIGrow
gen ROWid = ELIGrow + string(CNUM)
* Check that the ROWid matches the crosstabulation of ELIG &
CNUM .
tab ROWid
save
"$mcs working folder/mcs6 parent cm interview cross sweep merg
ing.dta", replace
* prepare mcs5 parent cm interview .
use "$mcs5 folder/mcs5 parent cm interview.dta", clear
gen ELIGrow = "M" if (EELIG00 == 1)
replace ELIGrow = "P" if (EELIG00 == 2)
gen CNUM = ECNUM00
tab CNUM ELIGrow
gen ROWid = ELIGrow + string(CNUM)
* Check that the ROWid matches the crosstabulation of ELIG &
CNUM .
tab ROWid
"$mcs working folder/mcs5 parent cm interview cross sweep merg
ing.dta", replace
* Merge .
use
"$mcs working folder/mcs6 parent cm interview cross sweep merg
ing.dta"
merge 1:1 MCSID ROWid using
"$mcs working folder/mcs5 parent cm interview cross sweep merg
ing.dta"
```

```
* Outcome perusal: parents (Main/Partner providing information
* each of the cohort members) are in both sweeps.
"$mcs working folder/mcs5 mcs6 parent cm interview by ELIG.dta
", replace
* ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ PNUM + CNUM = ROWid ~ ~ ~ ~ ~ ~ ~ ~ ~
* prepare mcs6 parent cm interview .
use "$mcs6 folder/mcs6 parent cm interview.dta", clear
gen CMrow = "C1" if (FCNUM00 == 1)
replace CMrow = " C2" if (FCNUM00 == 2)
replace CMrow = "C3" if (FCNUM00 == 3)
gen Prow = "P" + string(FPNUM00)
gen ROWid = Prow + CMrow
* Let's check that the totals match.
tab FPNUM00 FCNUM00
tab ROWid
save
"$mcs working folder/mcs6 parent cm interview cross sweep merg
ing.dta", replace
* prepare mcs5 parent cm interview .
use "$mcs5 folder/mcs5 parent cm interview.dta", clear
gen CMrow = "C1" if (ECNUM00 == 1)
replace CMrow = " C2" if (ECNUM00 == 2)
replace CMrow = "C3" if (ECNUM00 == 3)
gen Prow = "P" + string(EPNUM00)
gen ROWid = Prow + CMrow
* Let's check that the totals match.
tab EPNUM00 ECNUM00
tab ROWid
save
"$mcs working folder/mcs5 parent cm interview cross sweep merg
ing.dta", replace
* Merge .
"$mcs working folder/mcs6 parent cm interview cross sweep merg
ing.dta", clear
merge 1:1 MCSID ROWid using
"$mcs working folder/mcs5 parent cm interview cross sweep merg
ing.dta"
* Outcome perusal: Individuals (PNUM) provide information
* about the cohort member(s) in both sweeps (either as Main or
Partner respondent).
```

```
save
"$mcs working folder/mcs5 mcs6 parent cm interview by PNUM.dta
", replace
* Example code L
* -----.
* Create a composite variable per child in parent cm dataset.
* -----.
* this syntax creates a mean of Main and Partner responses on
ASLU variable .
* You can use other functions instead of MEAN, like sd, min,
max etc.
use "$mcs6 folder/mcs6 parent cm interview.dta", clear
tab FPASLU00
gen ASLU = FPASLU00
replace ASLU = . if (FPASLU00 <1)</pre>
egen composite ASLU = mean(ASLU), by (MCSID FCNUM00)
egen groupsize = count(FPNUM00), by (MCSID FCNUM00)
label variable composite ASLU "Mean score of ASLU assessment
by Main/Partner respondent(s) of the cohort member"
label variable groupsize "Number of respondents (Main only,
Partner only, or Main&Partner) providing information in ASLU"
* Let's take a look at the result .
gen example families = 1 if MCSID == "M10002P" | MCSID ==
"M10611J" | MCSID == "M10106W" | MCSID == "M10063C".
tab example families
* Overview of specific cases on composite score for each
Cohort Member (groupsize = number of parents ELIG providing
information)
* run this code with/without nolabel to hide/see the value
ds MCSID FPNUM00 FELIG00 FCNUM00 FPASLU00 ASLU composite ASLU
groupsize
return list
local varlist = r(varlist)
list `varlist' if example families == 1, divider sep(4)
nolabel
* Example code M
* -----.
```

\* Merging datasets of different structures to the mcs\_longitudinal\_family\_file .

\* -----

-----.

```
* we do not need to sort files as they are all sorted by
MCSID,
* however, if you have worked on the file before make sure you
* sort it before this step.
* merge with a parent level dataset .
use "$mcs6 folder/mcs6 parent derived.dta", clear
merge m:1 MCSID using
"$mcs6 folder/mcs longitudinal family file.dta"
* merge with a cm level dataset .
use "$mcs6 folder/mcs6 cm derived.dta", clear
merge m:1 MCSID using
"$mcs6 folder/mcs longitudinal family file.dta"
* merge with a _family_ level dataset .
use "$mcs6 folder/mcs6 family derived.dta", clear
merge m:1 MCSID using
"$mcs6 folder/mcs longitudinal family file.dta"
* merge with a parent cm level dataset .
use "$mcs6 folder/mcs6 parent cm interview.dta", clear
merge m:1 MCSID using
"$mcs6 folder/mcs longitudinal family file.dta"
* Example code N
* -----.
* Example I - merging two 1-level datasets that have different
identifiers.
* Please run example code H to get the parent level dataset
below.
use
"$mcs working folder/mcs6 parent plus proxy interview.dta",
* restructure the datasets into wide format (one row per
family).
tab FELIG00
* We create a Main / Partner only ELIG.
```

gen ELIG suffix = " M " if (FELIG00 == 1) // Main

Partner

Partner

\* resctructure using ELIG.

i(MCSID) j(ELIG suffix) string

replace ELIG suffix = " P " if (FELIG00 == 2) //(Proxy)

replace ELIG suffix = " P " if (FELIG00 == 3) //(Proxy)

reshape wide FELIG00 FRESP00 FPCREL00 FPGENA00 FXCREL00 ,

been suffixed with M and of the (Proxy) Partner with P.

\* the variables containing the information of the Main have

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```
save
"$mcs working folder/mcs6 parent plus proxy interview wide.dta
", replace
* merge cm level with the wide restructured parent dataset
(one row per family).
use "$mcs6 folder/mcs6 cm interview.dta", clear
merge m:1 MCSID using
"$mcs working folder/mcs6 parent plus proxy interview wide.dta
* for some families there is no parent interview but the CM
has participated.
* create a composite score of the general health of the Main &
Partner.
gen GENA M = FPGENA00 M
replace GENA M = . if (GENA M < 0)
gen GENA P = FPGENA00 P
replace GENA P = . if (GENA P < 0)
gen composite GENA = (GENA M + GENA P)/2
* Let's take a look at the result .
gen example families = 1 if MCSID == "M10002P" | MCSID ==
"M10611J" | MCSID == "M10106W" | MCSID == "M10063C".
tab example families
* Overview of specific cases on composite score for each
family (mean Health)
* run this code with/without nolabel to hide/see the value
ds MCSID GENA M GENA P composite GENA
return list
local varlist = r(varlist)
list `varlist' if example families == 1, divider sep(4)
nolabel
gen CGHE = FCCGHE00
replace CGHE = . if (CGHE < 0)</pre>
* comparison between parents' general health and CM's.
tabulate composite GENA CGHE
```

# \* Example code O

\* -----
\* Example II - merging a 1-level dataset (\_cm\_) with a 2-level dataset (\_parent\_cm\_) resulting into a 1-level structure (\_cm\_).

\* Please check example code L to get the \_parent\_cm\_ level dataset, it is similar.

use "\$mcs6\_folder/mcs6\_parent\_cm\_interview.dta", clear keep MCSID FPNUM00 FELIG00 FRESP00 FCNUM00 FCCREL00 FPASLU00

```
save
"$mcs working folder/mcs6 parent cm interview reduced.dta",
replace
* we turn the parent cm dataset into wide format (one row
per CM).
"$mcs working folder/mcs6 parent cm interview reduced.dta",
* restructure the datasets into wide format (one row per
family).
tab FELIG00
* We create a Main / Partner only ELIG.
gen ELIG suffix = " M " if (FELIG00 == 1) // Main
replace ELIG suffix = " P " if (FELIG00 == 2) //(Proxy)
Partner
replace ELIG suffix = " P " if (FELIG00 == 3) //(Proxy)
Partner
* resctructure using ELIG.
reshape wide FPNUM00 FELIG00 FRESP00 FCCREL00 FPASLU00 ,
i(MCSID FCNUM00) j(ELIG suffix) string
gen ASLU M = FPASLU00 M
replace ASLU M = . if (ASLU M < 0)
gen ASLU P = FPASLU00 P
replace ASLU P = . if (ASLU P < 0)
gen composite ASLU = (ASLU M + ASLU P) / 2
^{\star} impute the values of the one parent or the other if one is
missing and composite
* score has not been calculated - please adjust the
calculation of the composite score
* to the needs of your project
replace composite ASLU = ASLU M if missing(ASLU P)
replace composite ASLU = ASLU P if missing(ASLU M)
tab composite ASLU
* Let's take a look at the result .
gen example families = 1 if MCSID == "M10002P" | MCSID ==
"M10611J" | MCSID == "M10106W" | MCSID == "M10063C".
tab example families
* Mean score of ASLU assessment by Main/Partner respondent(s)
of the cohort member
* run this code with/without nolabel to hide/see the value
labels
ds MCSID FPNUM00 M FPNUM00 P FCNUM00 FPASLU00 M FPASLU00 P
composite ASLU
return list
local varlist = r(varlist)
list `varlist' if example families == 1, divider sep(4)
nolabel
```

```
save
```

- "\$mcs\_working\_folder/mcs6\_parent\_cm\_interview\_reduced\_wide\_on\_cm\_level.dta", replace
- \* merge  $\_{cm}$ \_interview level with the wide restructured parent dataset (one row per family).
- use "\$mcs6\_folder/mcs6\_cm\_interview.dta", clear
  merge 1:1 MCSID FCNUM00 using
- "\$mcs\_working\_folder/mcs6\_parent\_cm\_interview\_reduced\_wide\_on\_cm\_level.dta"
- $\mbox{\scriptsize \star}$  for some families there is no parent interview but the CM has participated.
- \* comparison between parents' perception of CM's likelihood to go to the University and CM's perception on the same topic. tab FCSTYU00 composite\_ASLU twoway lfitci composite\_ASLU FCSTYU00 || scatter composite\_ASLU FCSTYU00

#### \* Example code P

- \* -----.
- \* Example III merging a 2-level dataset (\_parent\_cm\_) with a 1-level dataset (\_cm\_) resulting into a 2-level dataset (\_cm\_).

use "\$mcs6\_folder/mcs6\_parent\_cm\_interview.dta", clear
merge m:1 MCSID FCNUM00 using

"\$mcs6\_folder/mcs6\_cm\_interview.dta" tab FCTRST0A FPSCHC00

use "\$mcs6\_folder/mcs6\_cm\_interview.dta", clear
merge 1:m MCSID FCNUM00 using
"\$mcs6\_folder/mcs6\_parent\_cm\_interview.dta"
tab FCTRST0A FPSCHC00