APPENDIX: Description of example code and Software Packages

A1. Walkthrough of RDD Application with R Code

In this appendix, we walk through a sample statistical R code for an RD Design analysis that can help to guide investigators interested in utilizing this method for their research. The R code is available at https://github.com/AdinaZekiAlHazzouri/RDD_Paper. Required R packages include *rdrobust*^{1,2} and *rddensity*³ (available at https://rdpackages.github.io/rdrobust/). Our code can be used to run a RD model on a teaching version of the Framingham Heart Study (FHS) dataset, which is available upon request at https://biolincc.nhlbi.nih.gov/teaching/. Please note that this dataset, which contains redacted data and is intended for teaching purposes only, should *not* be used for publishing or interpreting actual results.

The purpose of this walk-through is to provide a practical sequence of steps that can be followed to run a RD analysis. The FHS is a longitudinal cohort study which began in 1948 and has since helped to uncover some of the determinants of cardiovascular disease. In our application of RD to the teaching FHS dataset, we seek to estimate the causal effect of antihypertensive medications on cardiovascular disease incidence among individuals in this cohort. In our example, the cutoff (C) for treatment will be a systolic blood pressure of 140 mmHg, and the treatment will be the use of an anti-hypertensive medication. The following is a brief data dictionary of relevant variables and their purpose in our walk-through.

Variable	Description	Type of variable
CVD	Cardiovascular Disease in follow-up	Outcome variable (Y)
	period (0 = No, 1 = Yes)	
SYBP	Systolic Blood Pressure (mmHg)	Running variable (X)
BPMEDS	Use of Anti-hypertensive medication at	Treatment status variable (D)
	exam (0 = No, 1 = Yes)	
AGE	Age at exam (years)	Variable at baseline used to
		assess covariate balance

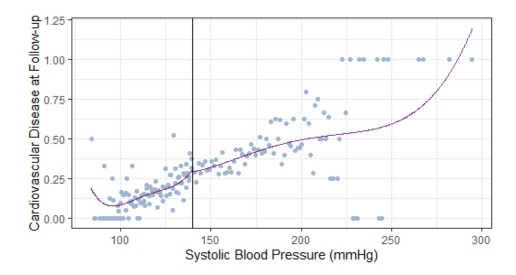
Next, we walk through the 7 steps of an RD analysis summarized in Section 3 of the commentary.

We first set up our R environment by attaching the necessary packages, and by loading and summarizing our dataset. In **Step 1**, we define the variables we would like to use for our RD analysis, comprised of the outcome value (Y), score/running variable (X), and the cutoff value (C).

```
rm(list=ls())
### Load R packages
library(rdrobust)
library(rddensity)
```

```
### Load dataset
setwd("Insert Directory Here") # Put location of your working directory here
frm <-read.csv("frmgham2.csv")</pre>
### For illustrative purposes, and to keep samples consistent between Sharp R
D and Fuzzy RD models, we will limit our sample to individuals with a non-mis
sing value of the outcome variable, running variable, and actual treatment ta
ke-up indicator
frm <-frm[complete.cases(frm[c("CVD", "SYSBP", "BPMEDS")]), ]</pre>
Y = frm$CVD # outcome variable
X = frm$SYSBP # running variable
D = frm$BPMEDS # actual treatment take-up indicator
C = 140 # cutoff value
T = 1*(X>=C) # treatment assignment indicator
summary(Y)
summary(X)
summary(D)
summary(T)
> summary(Y)
  Min. 1st Qu. Median Mean 3rd Qu.
                               Max.
 0.0000 0.0000 0.0000 0.2494 0.0000 1.0000
> summary(X)
  Min. 1st Qu. Median Mean 3rd Qu.
83.5 120.0 132.0 136.1 149.0
                                мах.
> summary(D)
  Min. 1st Qu. Median Mean 3rd Qu.
                                Max.
0.00000 0.00000 0.00000 0.08555 0.00000 1.00000
> summary(T)
  Min. 1st Qu. Median Mean 3rd Qu.
                                Max.
0.0000 0.0000 0.0000 0.3786 1.0000 1.0000
```

In **Step 2**, we generate an exploratory plot, which enables us to visualize the global relationship between the running variable and our outcome, separately at each side of the cutoff.



In **Step 3**, we set up the local polynomial model by specifying a polynomial order (i.e., P = 1, corresponding to a linear fit), a weighting function (i.e., kernel), and a selection procedure for computing the bandwidth (i.e., MSE optimal).⁵

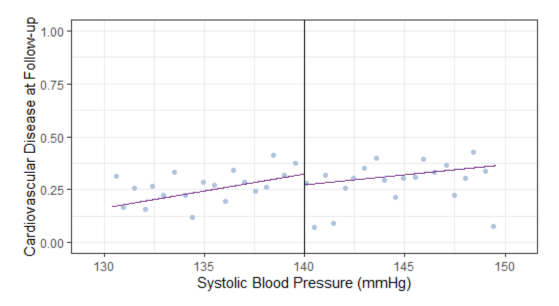
In **Step 4**, we estimate the RD effect by running *rdrobust* using our choices for the tuning parameters.¹ From the function output, we can recover the number of observations on each side of the cutoff, the bandwidth (h), and the number of observations within the bandwidth (i.e., Effective Number of Observations). In addition, the coefficient value on the row labeled "Conventional" corresponds to the RD causal estimate, the causal effect of the treatment on the outcome among subjects within the selected bandwidth. In this case, we obtain a reduction on the probability of a CVD event of 5.1 percentage points.

In addition, in **Step 5** we compute the 95% CI, corresponding to a plausible range of RD estimates consistent with our data, and the associated p-value. Here, the appropriate 95% CI can be recovered from the row labeled "Robust", [-xxx, xxx], which accounts for the nonparametric features of our estimation approach and should be preferred over the "Conventional" one. The p-value associated with this 95% CI is 0.072.⁶

```
call: rdrobust
Number of Obs.
                              11034
BW type
                              mserd
Kernel
                         Triangular
VCE method
                                NN
Number of Obs.
                               6857
                                            4177
Eff. Number of Obs.
Order est. (p)
                               1675
                                            1492
                               1 2
                                             1
Order bias (q)
                            9.571 9.571
17.892 17.892
0.535 0.535
BW est. (h)
BW bias (b)
                                         0.535
rho (h/b)
Unique Obs.
                               104
                                            155
```

Method	Coef. S	std. Err.	z	P> Z	[95% C.I.]
Conventional Robust	-0.051		-1.417 -1.610	0.157 0.107	[-0.122 , 0.020] [-0.147 , 0.014]

We can also plot our estimated RD model using the *rdplot* function, similar to the one in Figure 1, but now reflecting our choices for the estimation of the RD model instead of a global fit.



In the above figure, we can see a discontinuity at the cutoff. Individuals just above the cutoff (eligible for antihypertensive treatment) observed a decreased incidence in cardiovascular disease, as compared to individuals just below the cutoff. The distance between the lines at the cutoff gives you the ATE, which corresponds to the parameter estimate provided in the summary function (-0.051).

In **Step 6**, we can run a set of robustness checks and validation tests to assess the validity of our RD model. First, we explore the robustness of our RD effect estimates to the implementation choices of our local polynomial estimator. In particular, we look at alternative kernels, polynomial orders, and bandwidth choices.

```
# Change to uniform kernel
rd_2 \leftarrow rdrobust(y = Y, x = X, c = C,
            kernel = 'uniform', p = 1, bwselect = 'mserd')
summary(rd_2)
# Change to local quadratic regression
rd 3 <- rdrobust(y = Y, x = X, c = C,
            kernel = 'triangular', p = 2, bwselect = 'mserd')
summary(rd_3)
# Change to different bandwidths on each side:
rd_4 \leftarrow rdrobust(y = Y, x = X, c = C,
            kernel = 'triangular', p = 1, bwselect = 'msetwo')
summary(rd_4)
# Other bandwidth choices:
rd_bws <- rdbwselect(y = Y, x = X, c = C, all = TRUE)</pre>
```

summary(rd_bws)

> summary(rd_2) Call: rdrobust

Number of Obs. BW type Kernel VCE method	11034 mserd Uniform NN	
Number of Obs. Eff. Number of Obs. Order est. (p) Order bias (q) BW est. (h) BW bias (b) rho (h/b) Unique Obs.	6857 1257 1 2 7.594 16.335 0.465 104	4177 1243 1 2 7.594 16.335 0.465

Method	Coef. S	td. Err.	z	P> z	[95% C.I.]
Conventional Robust	-0.042		-1.130 -1.348		[-0.116 , 0.031] [-0.137 , 0.025]

> summary(rd_3)
Call: rdrobust

Number of Obs. BW type Kernel VCE method	11034 mserd Triangular NN	
Number of Obs. Eff. Number of Obs. Order est. (p) Order bias (q) BW est. (h) BW bias (b) rho (h/b) Unique Obs.	6857 3050 2 3 15.413 23.186 0.665 104	4177 2188 2 3 15.413 23.186 0.665 155

Method	Coef. St	d. Err.	z	P> z	[95% C.I.]
Conventional Robust	-0.069		-1.632 -1.713		[-0.152 , 0.014] [-0.174 , 0.012]

> summary(rd_4) Call: rdrobust

Number of Obs. BW type Kernel VCE method	11034 msetwo Triangular NN	
Number of Obs. Eff. Number of Obs. Order est. (p) Order bias (q) BW est. (h) BW bias (b) rho (h/b) Unique Obs.	6857 1659 1 2 9.270 17.971 0.516 104	4177 2911 1 2 24.862 37.880 0.656 155

Method	Coef. S	td. Err.	Z	P> z	[95% C.I.]
Conventional Robust	-0.032		-0.968 -1.179	0.333 0.238	[-0.098 , 0.033] [-0.122 , 0.030]

> summary(rd_bws)

call: rdbwselect

BW type	All	
Kernel	Triangular	
VCE method	NN	
Number of Obs.	6857	4177
Order est. (p)	1	1
Order bias (q)	2	2
Unique Obs.	104	155

Number of Obs. 11034

BW est. (h) BW bias (b) Left of c Right of c Left of c Right of c mserd 9.571 9.571 17.892 17.892 msetwo 9.270 24.862 17.971 37.880 msesum 13.741 13.741 21.248 21.248 msecomb1 9.571 9.571 17.892 17.892					.=======
msetwo 9.270 24.862 17.971 37.880 msesum 13.741 13.741 21.248 21.248					Right of c
msecomb2 9.571 13.741 17.971 21.248 cerrd 6.009 6.009 17.892 17.892 certwo 5.820 15.610 17.971 37.880 cersum 8.627 8.627 21.248 21.248 cercomb1 6.009 6.009 17.892 17.892 cercomb2 6.009 8.627 17.971 21.248	msetwo msesum msecomb1 msecomb2 cerrd certwo cersum cercomb1	9.270 13.741 9.571 9.571 6.009 5.820 8.627 6.009	24.862 13.741 9.571 13.741 6.009 15.610 8.627 6.009	17.971 21.248 17.892 17.971 17.892 17.971 21.248 17.892	37.880 21.248 17.892 21.248 17.892 37.880 21.248 17.892

Overall, we see that the main results remain (i.e., a protective effect is observed between the indicated anti-hypertensive medication and cardiovascular disease). Next, we conduct a manipulation test, looking for potential violation of the assumed continuity of observations around the threshold. If this assumption holds, we should expect to see a relatively consistent frequency of observations at each side of the cutoff. This test can be easily computed using the *rddensity* package, which provides the results of a statistical test together with a density plot illustrating the findings.³

```
# Test for manipulation of the running variable
rdd <- rddensity(X = X, c = C)
summary(rdd)</pre>
```

Figure

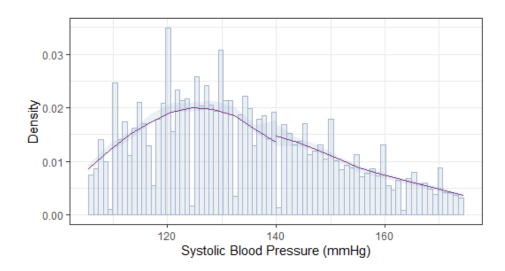
plot <- rdplotdensity(rdd, X, xlabel = "Systolic Blood Pressure (mmHg)", ylab
el = "Density", lcol = "mediumorchid4",pcol = "lightsteelblue3", histLineCol
= "lightsteelblue3", histFillCol = "lightsteelblue3", CIcol = "lightsteelblue
3")</pre>

> summary(rdd)

Manipulation testing using local polynomial density estimation.

Number of obs = 11034 Model = unrestricted Kernel = triangular BW method = estimated VCE method = jackknife c = 140Left of c Right of c Number of obs 6857 4177 Eff. Number of obs 2182 1788 Order est. (p) 2 2 Order bias (q) 3 BW est. (h) 11.5 11.5 Method P > |T|Robust 1.0241 0.3058

P-values of binomial tests (HO: p=0.5).



From both the plot and the results of the test, we can conclude that there is no evidence of a jump in the density of the running variable at the cutoff, which is consistent with a lack of manipulation.

We also assess exchangeability around the cutoff by looking at covariate balance; this involves taking a given covariate (e.g., age) and treating it as an outcome in an RD model. One can subsequently view the RD effect and visualize a plot to see whether there is a significant difference in the level of the covariate of interest, comparing observations below and above the cutoff. If the exchangeability assumption holds, then we should expect the levels of this covariate to be comparable below and above the cutoff (i.e., the RD effect here should be close to 0).

```
## Covariate balance assessment (Age)

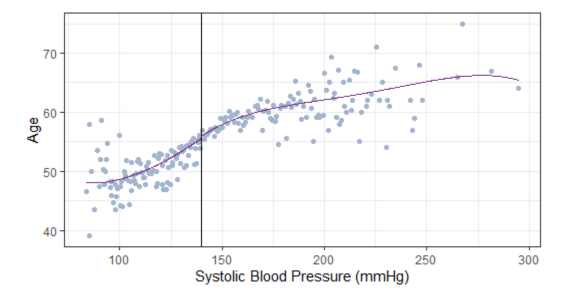
rd_covs = rdrobust(y = frm$AGE, x = X, c = C)
summary(rd_covs)

rdplot(y = frm$AGE, x = X, c = C,
    title = "Covariate Balance - RD Plot",
    x.label = "Systolic Blood Pressure (mmHg)",
    y.label = "Age", col.dots="lightsteelblue3",col.lines="mediumorchid4")
```

> summary(rd_covs) Call: rdrobust

Number of Obs. 11034 BW type mserd Kernel Triangular VCE method Number of Obs. 6857 4177 Eff. Number of Obs. 2812 2080 1 Order est. (p) 1 Order bias (q) 14.601 BW est. (h) 14.601 BW bias (b) 22.187 22.187 0.658 0.658 rho (h/b) Unique Obs. 104 155

Method	Coef. St	d. Err.	z	P> z	[95% C.I.]
Conventional Robust	0.596	0.565	1.055 0.927	0.292 0.354	[-0.512 , 1.704] [-0.708 , 1.978]



The RD model output and plot above show that there is no significant difference in age comparing individuals below and above the cutoff; this provides some evidence for the exchangeability assumption required for RD design.

Next, in **Step 7**, we account for imperfect compliance with treatment assignment by running a fuzzy RD model.

Eff. Number of Obs. 3713 24 Order est. (p) 1 Order bias (q) 2 Bw est. (h) 18.679 18.0 Bw bias (b) 30.694 30.0	> summary(RD_fuzzy) Call: rdrobust		
Eff. Number of Obs. 3713 24 Order est. (p) 1 Order bias (q) 2 BW est. (h) 18.679 18.0 BW bias (b) 30.694 30.0	BW type Kernel	mserd Triangular	
	Eff. Number of Obs. Order est. (p) Order bias (q) BW est. (h) BW bias (b) rho (h/b)	3713 1 2 18.679 30.694 0.609	4177 2448 1 2 18.679 30.694 0.609

Method	Coef.	Std. Err.	Z	P> z	[95% C.I.]
Conventional	-0.230	1.060	-0.217	0.828	[-2.307 , 1.847]
Robust	-		-0.581	0.561	[-3.184 , 1.727]

In this output, the coefficient value (-0.230) corresponds to the per-protocol treatment effect of anti-hypertensive medication on cardiovascular disease; that is, this is the complier-specific treatment effect, computed by dividing the sharp RD effect by the compliance rate (the percentage of participants who received the treatment that they were assigned to).

We can also manually calculate this fuzzy RDD effect (-0.230) by running two separate steps: first, we can calculate the sharp RD effect (note that we are using the same sample size and optimal bandwidth from the fuzzy RD model). As shown below, this sharp RD effect is (-0.006).

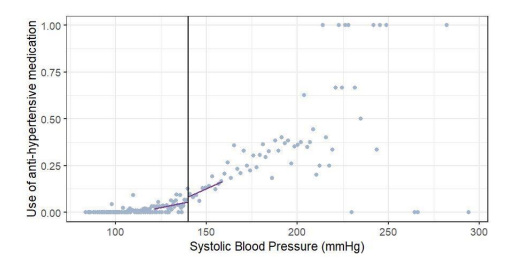
```
# First, assess sharp RDD effect within bandwidth from the above fuzzy model
RD_sharp = rdrobust(y = Y, x = X, c = C, h = RD_fuzzy$bws[1,1], b = RD_fuzzy$
bws[2,1], subset=!is.na(D))
summary(RD_sharp)
RD_sharp$Estimate[1] # Sharp RDD effect = -0.00558034
```

Next, we calculate the specific amount of non-compliance in our application. We do this by running an RDD where we use actual treatment take-up (BPMEDS) as the outcome, and the same running variable (here we use the same sample size and optimal bandwidth from the fuzzy RD model as well). Our estimated effect (the amount of compliance at the cutoff) is 0.024, which is the difference between treatment take-up in the treatment group (8.06%) and treatment take-up in the control group (5.64%).

> summary(RD_comp) Call: rdrobust

Number of Obs. BW type Kernel VCE method	11034 Manual Triangular NN	
Number of Obs. Eff. Number of Obs. Order est. (p) Order bias (q) BW est. (h) BW bias (b) rho (h/b) Unique Obs.	6857 3713 1 2 18.679 30.694 0.609 104	4177 2448 1 2 18.679 30.694 0.609

Method	Coef. S	td. Err.	Z	P> z	[95% C.I.]
Conventional Robust	0.024	0.014	1.677 1.599	0.094 0.110	[-0.004 , 0.053] [-0.006 , 0.061]



Finally, we can recover the fuzzy RDD estimate (-0.2301855) by dividing the sharp RDD estimate (-0.005580354) by the compliance rate (0.02424286).

```
# Final calculation
RD_sharp$Estimate[1] / RD_comp$Estimate[1] # -0.2301855
```

```
Box A2: R code
```

Sample statistical codes (in R) used in RDD analysis

Required R packages include *rdrobust*^{1,2} and *rddensity*³

(https://rdpackages.github.io/rdrobust/). The statistical codes presented below are based on the RDD application investigating the causal effect of blood pressure medication (BPMEDS) on CVD.

Step 1: Identify main components of the RD model

```
Y = frm$CVD
               # outcome variable
X = frm$SYSBP # running variable
D = frm$BPMEDS # actual treatment take-up indicator
          # cutoff value
T = 1*(X>=C) # treatment assignment indicator
Step 2: Plot the RD data (global fit)
rdplot1 = rdplot(y = Y, x = X, c = C,
                 x.label = "Systolic Blood Pressure (mmHg)",
                 y.label = "Cardiovascular Disease at Follow-up")
summary(rdplot1)
Step 3: Set up the local polynomial model
                  # local linear approximation
K = 'triangular' # triangular kernel
BW = 'mserd'
                  # MSE optimal bandwidth
Steps 4-5: RD estimation and inference
rd_{manual} = rdrobust(y = Y, x = X, c = C,
                     kernel = K, p = P, bwselect = BW)
summary(rd manual)
We can re-plot our RD data (as in Step 2) but now using our choices for the estima
tion of the RD model instead of a global fit.
bw = rd manual $bws[1]
RD_plot = rdplot(y = Y, x = X, c = C,
                 p = 1, kernel = 'triangular', h = bw,
                 x.label = "Systolic Blood Pressure (mmHg)",
                 y.label = "Cardiovascular Disease at Follow-up",
                 subset = (X >= (C - bw) & X <= (C + bw)), x.lim = c(C - bw - bw)
                 1 + C + hu + 1\ v 1im - c/a + 1\\
```

```
Box A2 (continued)
Sample statistical codes (in R) used in RDD analysis – continued
Step 6: Robustness checks and validation (a comprehensive list can be found in the
Appendix)
# Change to uniform kernel
rd_2 \leftarrow rdrobust(y = Y, x = X, c = C,
                 kernel = 'uniform', p = 1, bwselect = 'mserd')
summary(rd 2)
# Change to different bandwidths on each side:
rd_4 \leftarrow rdrobust(y = Y, x = X, c = C,
                 kernel = 'triangular', p = 1, bwselect = 'msetwo')
summary(rd 4)
# Test for manipulation of the running variable
rdd \leftarrow rddensity(X = X, c = C)
summary(rdd)
plot <- rdplotdensity(rdd, X)</pre>
## Covariate balance assessment (Age is the only covariate in this example)
rd\_covs = rdrobust(y = frm$AGE, x = X, c = C)
summary(rd covs)
rdplot(y = frm$AGE, x = X, c = C,
  title = "Covariate Balance - RD Plot",
  x.label = "Systolic Blood Pressure (mmHg)",
  y.label = "Age")
Step 7: Account for imperfect compliance
# Fuzzy RD model:
RD fuzzy = rdrobust(y = Y, x = X, c = C, fuzzy = D)
summary(RD fuzzy)
# Next, manually calculate this fuzzy RD effect by running the below 2 steps:
# First, assess sharp RDD effect within bandwidth from the above fuzzy model
RD_sharp = rdrobust(y = Y, x = X, c = C, h = RD_fuzzy$bws[1,1], b = RD_fuzzy$bws[2]
,1], subset=!is.na(D))
summary(RD sharp)
RD sharp$Estimate[1] # Sharp RDD effect = -0.005580354
# Next, assess compliance by computing RD effect using D as the outcome
RD_{comp} = rdrobust(y = D, x = X, c = C, h = RD_fuzzy$bws[1], b = RD_fuzzy$bws[2,1]
summary(RD_comp)
RD comp$Estimate[1] # Compliance RDD effect = 0.02424286
```

RD_comp\$beta_p_r[1] # Treatment Take-up for Treatment Group = 0.08062456 RD_comp\$beta_p_l[1] # Treatment Take-up for Control Group = 0.0563817 RD_comp_plot = rdplot(y = D, x = X, c = C, h = RD_fuzzy\$bws[1,1], p=1,

x.label = "Systolic Blood Pressure (mmHg)",

A3. Supplemental Stata Code for RDD Application

Box A3: Stata code

Sample statistical codes (in Stata) used in RDD analysis

Required Stata packages include *rdrobust*^{1,2} and *rddensity*³

xtitle(Systolic Blood Pressure (mmHg)) ///
graphregion(color(white)) legend(off))

(https://github.com/rdpackages). The statistical codes presented below are based on the RDD application investigating the causal effect of blood pressure medication (BPMEDS) on CVD.

```
Step 1: Identify main components of the RD model
                /* outcome variable */
global Y cvd
global X sysbp /* running variable */
global D bpmeds /* actual treatment take-up indicator */
                   /* cutoff value */
global C 140
gen T = 1*($X>=$C)
                          /* treatment assignment indicator */
Step 2: Plot the RD data (global fit)
rdplot $Y $X, c($C) ///
graph_options(title("") ///
      ytitle(Cardiovascular Disease at Follow-up) ///
      xtitle(Systolic Blood Pressure (mmHg)) ///
      graphregion(color(white)) legend(off))
Step 3: Set up the local polynomial model
local P = 1
                         /* local linear approximation */
local K = "triangular"
                         /* triangular kernel */
local BW = "mserd"
                         /* MSE optimal bandwidth*/
Steps 4-5: RD estimation and inference
rdrobust $Y $X, c($C) kernel(`K') p(`P') bwselect(`BW')
* You can also run RD using default choices, as shown here:
rdrobust $Y $X, c($C)
We can re-plot our RD data (as in Step 2) but now using our choices for the estima
tion of the RD model instead of a global fit.
rdplot Y X if -e(h_1) = X-C & X-C = e(h_r), c(C) kernel(K') p(P') bwsele
ct(`BW') ///
      h(`e(h_l)' `e(h_r)') ///
      graph options(title("") ///
      ytitle(Cardiovascular Disease at Follow-up) ///
```

```
Box A3 (continued)
```

Sample statistical codes (in Stata) used in RDD analysis – continued

Step 6: Robustness checks and validation (a comprehensive list can be found in the Appendix)

```
* Change to uniform kernel
rdrobust $Y $X, c($C) kernel("uniform") p(1) bwselect("mserd")
* Change to different bandwidths on each side:
rdrobust $Y $X, c($C) kernel("triangular") p(1) bwselect("msetwo")
* Test for manipulation of the running variable: rddensity test
rddensity $X, c($C) plot ///
                     graph_opt(title("") ///
                     ytitle(Cardiovascular Disease at Follow-up) ///
                     xtitle(Systolic Blood Pressure (mmHg)) ///
                     graphregion(color(white)) legend(off))
** Covariate balance assessment (Age)
rdrobust age $X, c($C)
rdplot age $X, c($C) ///
                    graph options(title("Covariate Balance - RD Plot") ///
                     ytitle(Age) ///
                     xtitle(Systolic Blood Pressure (mmHg)) ///
                     graphregion(color(white)) legend(off))
```

Step 7: Account for imperfect compliance

A4. List of Statistical Software for RD Designs

- rdrobust: RD plots, estimation, robust inference, bandwidth selection and other related features
 for RD designs employing local polynomial and partitioning methods. It is described in more
 detail in Calonico et al. (2017). Stata and R packages: https://rdpackages.github.io/rdrobust/
- **frdboot**: iterative bootstrap from Bartalotti et al. (2017) and He and Bartalotti (2020). R package: github.com/yhe0802/FRD-bootstrap/
- **rdexo**: implements external validity tests of Bertanha and Imbens (2020). Stata and Matlab packages: https://sites.google.com/site/mbertanha/home/code-bertanha-imbens.
- **rdlocrand**: statistical inference and graphical procedures for RD designs employing local randomization methods. It provides point estimators, confidence intervals estimators, windows selectors, automatic plots, sensitivity analysis and other related features. Stata and R packages: https://rdpackages.github.io/rdlocrand/
- optrdd: optimized inference in RD designs, as proposed by Imbens and Wager (2019). R package: https://github.com/swager/optrdd
- **rdhonest**: honest confidence intervals in fuzzy and sharp RD designs using procedures from Armstrong and Kolesár (2020). R package: https://github.com/kolesarm/RDHonest
- **rdmulti**: RD plots, estimation, inference and extrapolation methods for RD designs with multiple cutoffs and multiple scores. Stata and R packages: https://rdpackages.github.io/rdmulti/
- rddensity: manipulation tests employing local polynomial density estimation methods. It
 provides hypothesis tests and bandwidth selectors for manipulation testing. Stata and R
 packages: https://rdpackages.github.io/rddensity/
- rdcont: continuity test for the density of the running variable based on g-order statistics developed in Bugni and Canay (2020). Stata package: https://bitbucket.org/iacanay/rdcontstata/overview

References

- 1. Calonico S, Cattaneo MD, Titiunik R. rdrobust: An R Package for Robust Nonparametric Inference in Regression-Discontinuity Designs. *R Journal*. 2015;7:38-51.
- 2. Calonico S, Cattaneo MD, Farrell MH, Titiunik R. Rdrobust: Software for Regression-discontinuity Designs. *The Stata Journal*. 2017;17(2):372-404.
- 3. Cattaneo MD, Jansson M, Ma X. Manipulation Testing Based on Density Discontinuity. *Stata Journal*. 2018;18:234-261.
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